

HHS Public Access

Author manuscript *Cell.* Author manuscript; available in PMC 2021 July 09.

Published in final edited form as:

Cell. 2020 July 09; 182(1): 38–49.e17. doi:10.1016/j.cell.2020.05.019.

CBASS Immunity Uses CARF-Related Effectors to Sense 3'-5'and 2'-5'-Linked Cyclic Oligonucleotide Signals and Protect Bacteria from Phage Infection

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SUMMARY

cGAS/DncV-like nucleotidyltransferase (CD-NTase) enzymes are immune sensors that synthesize nucleotide second messengers and initiate antiviral responses in bacterial and animal cells. Here, we discover *Enterobacter cloacae* CD-NTase-associated protein 4 (Cap4) as a founding member of

DECLARATION OF INTERESTS

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AUTHÔR CONTRIBÚTIONS

Experiments were designed by B.L., A.T.W., and P.J.K. Crystallography and biochemical experiments were conducted by B.L. with assistance from P.J.K. Bacterial cytotoxicity and phage challenge assays were conducted by A.T.W. with V.J.C. and J.J.M. Negativestain EM experiments were conducted by A.F.A.K. and S.S., and DNA fragment deep-sequencing experiments were conducted by S.P.A. and A.S.Y.L. CD-NTase nucleotide product purification was conducted by B.L. with assistance from B.R.M. and F.S. Synthetic nucleotide product synthesis and characterization experiments were performed by D.K., P.N., and F.S. LC-MS/MS experiments were performed by I.T.M. and M.J. The manuscript was written by B.L. and P.J.K., and all authors contributed to editing the manuscript and support the conclusions.

D.K., P.N., and F.S. are employed at Biolog Life Science Institute GmbH & Co. KG, which sells 3'3'3'-cAAG and 3'3'3'-cAAA and may sell 2'3'3'-cAAA as research tools.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.cell.2020.05.019.

a diverse family of >2,000 bacterial receptors that respond to CD-NTase signals. Structures of Cap4 reveal a promiscuous DNA endonuclease domain activated through ligand-induced oligomerization. Oligonucleotide recognition occurs through an appended SAVED domain that is an unexpected fusion of two CRISPR-associated Rossman fold (CARF) subunits co-opted from type III CRISPR immunity. Like a lock and key, SAVED effectors exquisitely discriminate 2'-5'- and 3'-5'-linked bacterial cyclic oligonucleotide signals and enable specific recognition of at least 180 potential nucleotide second messenger species. Our results reveal SAVED CARF family proteins as major nucleotide second messenger receptors in CBASS and CRISPR immune defense and extend the importance of linkage specificity beyond mammalian cGAS-STING signaling.

Graphical Abstract



In Brief

A family of bacterial immune effectors responds to 3'-5'- and 2'-5'-linked nucleotide signals in CBASS antiviral immunity.

INTRODUCTION

cGAS/DncV-like nucleotidyltransferases (CD-NTases) are enzymes that synthesize specialized oligonucleotide signals to amplify pathway activation and control downstream effector responses. CD-NTases are conserved in animal and bacterial signaling systems and play a key role in innate immunity and phage defense (Ablasser and Chen, 2019; Bernheim

and Sorek, 2020; Kranzusch, 2019). In human cells, the CD-NTase cyclic GMP-AMP synthase (cGAS) functions as a sensor for double-stranded DNA aberrantly localized in the cell cytosol during pathogen replication and cancer. When activated, cGAS produces the nucleotide second messenger 2'-5', 3'-5' cyclic GMP- AMP (2'3'-cGAMP) to induce antiviral immunity and interferon signaling (Ablasser et al., 2013; Diner et al., 2013; Gao et al., 2013a; Sun et al., 2013; Zhang et al., 2013). In an analogous system, the *Vibrio cholerae* CD-NTase dinucleotide cyclase in *Vibrio* (DncV) synthesizes the nucleotide second messenger 3'-5', 3'-5' cGAMP (3'3'-cGAMP) in response to an unknown stimulus during phage infection (Cohen et al., 2019; Davies et al., 2012; Kranzusch et al., 2014). Bacterial 3'3'-cGAMP activates a downstream effector response that results in cell death, limitation of phage replication through abortive infection, and protection of the remaining bacterial population (Cohen et al., 2019; Severin et al., 2018).

Bacterial CD-NTases include >5,600 unique enzymes (Whiteley et al., 2019) that control a diverse array of antiviral immune systems collectively called cyclic oligonucleotide-based antiphage signaling system (CBASS) immunity (Cohen et al., 2019; Ye et al., 2020). Synthesizing cyclic dinucleotide and cyclic trinucleotide products, bacterial CD-NTases are capable of using all four ribonucleotides as building blocks to dictate signal specificity and enable increased diversity of CBASS antiviral immune responses (Cohen et al., 2019; Whiteley et al., 2019). For example, in addition to the *V. cholerae* DncV product 3'3'-cGAMP, *Escherichia coli* CdnE synthesizes 3'3' cyclic UMP-AMP (3'3'-cUA), and *Enterobacter cloacae* CdnD (CD-NTase in clade D) synthesizes the cyclic trinucleotide second messenger 3'3'3' cyclic AMP-AMP-GMP (3'3'3'-cAAG) (Whiteley et al., 2019). In human cells, the non-canonical 2'-5' linkage in the cGAS product 2'3'-cGAMP is critical for immune specificity and potent activation of the downstream receptor stimulator of interferon genes (STING) (Ablasser et al., 2013; Diner et al., 2013; Gao et al., 2013b; Zhang et al., 2013), but it is unknown whether phosphodiester linkage specificity is an additional determinant of receptor activation in bacterial CBASS antiviral immunity.

The best-characterized family of receptors that respond to bacterial CD-NTase nucleotide second messengers are patatin-like phospholipases, which are activated to degrade membrane phospholipids upon nucleotide signal binding. The first known example is the *Vibrio cholerae* protein cGAMP-activated phospholipase in *Vibrio* (CapV), which responds to 3'3'-cGAMP and causes membrane rupture and bacterial cell death (Severin et al., 2018). Similarly, the *E. coli* CD-NTase CdnE signals through CapE, a CapV homolog that specifically recognizes 3'3'-cUA, indicating that CBASS operons function with high specificity for a single-nucleotide second messenger (Whiteley et al., 2019). However, a majority of bacterial CD-NTase enzymes are encoded in CBASS operons that do not contain proteins with homology to CapV-like receptors (Burroughs et al., 2015; Cohen et al., 2019; Whiteley et al., 2019), suggesting that still-uncharacterized proteins must be responsible for downstream effector functions.

Here we reconstitute *Enterobacter cloacae* CdnD signaling *in vitro* and *in vivo* to discover CD-NTase-associated protein 4 (Cap4) as the founding member of a major family of downstream receptors that specifically respond to nucleotide second messenger signals in CBASS immunity. High-resolution crystal structures of Cap4 combined with negative-stain

electron microscopy (EM) analysis reveals a DNA endonuclease effector domain activated through nucleotide second messenger-induced oligomerization. We demonstrate that a previously uncharacterized protein domain in Cap4 called SAVED is responsible for nucleotide second messenger recognition. Remarkably, SAVED is a fusion of two CRISPR-associated Rossman fold (CARF) domains derived from type III CRISPR immunity, revealing a common ancestry between these nucleotide second messenger-centric antiviral systems. Divergence in the SAVED nucleotide binding pocket enables recognition of an expanded range of CD-NTase products, including bacterial second messengers with alternative ring size, nucleobase, and 3'-5' or 2'-5' phosphodiester linkages. We further show that SAVED-containing effectors are essential for CBASS-mediated protection of bacteria from phage infection. Our results uncover a major family of nucleotide second messenger receptors and reveal a role of 2'-5'-linked nucleotide signals in bacterial resistance to phage infection.

RESULTS

Cap4 Proteins Are DNA Endonucleases that Respond to CD-NTase Nucleotide Second Messengers

The *E. cloacae* CD-NTase CdnD (*Ec*CdnD) is constitutively active *in vitro* and synthesizes the nucleotide second messenger 3'3'3'-cAAG (Whiteley et al., 2019). *Ec*CdnD is encoded in an operon containing three additional genes of unknown function, designated here CD-NTase-associated proteins 2, 3, and 4 (*Cap2, Cap3*, and *Cap4*) (Figure 1A). To define how bacterial CD-NTase enzymes control downstream signaling, we purified each protein from the *E. cloacae* CdnD02 operon and used a biochemical approach for receptor identification. Incubation of radiolabeled 3'3'3'-cAAG with purified *Ec*Cap4 (GenBank: WP_032676399.1) in an electrophoretic mobility shift assay resulted in complete shift and formation of a stable Cap4–3'3'3'-cAAG complex (Figures 1B and S1A). No interaction was observed between *Ec*Cap4 and the *V. cholerae* DncV product 3'3'-cGAMP, demonstrating that *Ec*Cap4 is a downstream receptor that specifically recognizes the cognate *Ec*CdnD nucleotide second messenger.

To understand the function of Cap4, we screened homologs for suitability in a structural analysis and determined a 2.4 Å crystal structure of Cap4 from the bacterium *Moraxella osloensis* (*Mo*Cap4; GenBank: WP_060996052.1) and a 2.6 Å crystal structure of Cap4 from *Acinetobacter baumannii* (*Ab*Cap4; GenBank: WP_008942236.1) (Figures 1C and S1B; Table S1). The structure of Cap4 reveals a two-domain architecture with an N-terminal domain containing a mixed β sheet braced on either side with α -helical bundles and a globular C-terminal domain that contains internal two-fold pseudosymmetry (Figures 1C and S1B). Sequence alignment based on the *Mo*Cap4 structure demonstrates that all Cap4 homologs contain the same domain architecture with *Mo*Cap4 versus *Ab*Cap4 sharing ~80% identity and *Mo*Cap4 versus *Ec*Cap4 sharing ~20% identity at the amino acid level (Figure S1B). The Cap4 N-terminal domain is a member of an uncharacterized protein domain classification, domain of unknown function 4297 (DUF4297), that is widespread in bacteria. Comparative analysis using the *Mo*Cap4 structure reveals that the DUF4297 domain shares structural homology with type II restriction endonucleases, including the enzymes AgeI and

HindIII (Figure 1D). Sequence alignment of Cap4 and type II restriction enzymes confirms conservation of all putative active-site residues required for metal coordination and suggests that Cap4 proteins are functional nuclease enzymes (Figure S1B; Burroughs et al., 2015; Watanabe et al., 2009). We therefore tested *Ec*Cap4 and *Ab*Cap4 proteins for the ability to cleave DNA substrates *in vitro*. In the presence of an activating nucleotide second messenger, Cap4 proteins catalyze complete degradation of plasmid DNA to small, <45 bp DNA fragments (Figure 1E). Cap4 alone exhibits no nuclease activity, indicating that recognition of a nucleotide second messenger is a strict requirement for nuclease activation. Cap4 activity is divalent cation metal dependent (Figure S1D), and mutation of a conserved lysine residue in the active site ablates nuclease activity (Figures 1F and S1E), further confirming that Cap4 proteins use a type II restriction enzyme-like reaction mechanism to degrade DNA.

Type II restriction endonucleases contain loops or helices that extend from the nuclease domain to form DNA major and minor groove contacts and control target sequence cleavage specificity (Tamulaitiene et al., 2017; Watanabe et al., 2009). The Cap4 N-terminal endonuclease domain is minimized, with nearly all extensions absent, suggesting relaxation of DNA targeting specificity (Figure 1D). Consistent with these structural observations, *Ec*-Cap4 degrades plasmid DNA, *E. coli* genomic DNA, and synthetic double-stranded DNA (dsDNA) *in vitro* with no apparent target sequence or DNA modification specificity (Figures 1G and S1F). In each case, Cap4 endonuclease activity remains strictly dependent on the presence of activating nucleotide second messenger. Together, these results demonstrate that Cap4 proteins are dsDNA endonucleases controlled through nucleotide second messenger-gated enzyme activation.

Cap4 Proteins Respond to Specific 3'-5' and 2'-5'-Linked Nucleotide Second Messengers

To determine the specificity of CD-NTase-Cap4 signaling, we next compared crossactivation between E. cloacae and A. baumannii CdnD operons. Although each CdnD activates the robust endonuclease activity of its cognate Cap4 protein in vitro, heterologous reactions with one CdnD or Cap4 component derived from each operon fail to reconstitute signaling and do not result in DNA degradation (Figures 2A and S2A). Selective Cap4 activation reveals that AbCdnD must synthesize a nucleotide second messenger distinct from the EcCdnD product 3'3'3'-cAAG. We tested nucleotide combinations and observed that ATP is necessary and sufficient to allow AbCdnD to synthesize the activating nucleotide second messenger signal (Figure S2B). However, all known canonically linked cyclic di-, tri-, and tetra-AMP RNA products failed to reconstitute activation of AbCap4 DNA degradation activity (Figure 2B). We therefore hypothesized that, like human cGAS-STING, AbCdnD-Cap4 signaling may require a noncanonical 2'-5'-linked second messenger. To assess whether the AbCdnD nucleotide product contains a non-canonical linkage, we analyzed the sensitivity of CD-NTase reaction products to digestion by nuclease P1, which specifically hydrolyzes 3'-5'-linked phosphodiester bonds and is unable to cleave noncanonical 2'-5' phosphodiester bonds (Ablasser et al., 2013; Diner et al., 2013; Whiteley et al., 2019). Unlike the V. cholerae DncV and EcCdnD products 3'3'-cGAMP and 3'3'3'cAAG, the mammalian cGAS 2'3'-cGAMP and the AbCdnD reaction products contain

phosphates resistant to nuclease P1 digestion, confirming the presence of a 2'-5'-linked bond (Figure 2C).

We next isolated the AbCdnD nucleotide second messenger directly from enzymatic reactions for further characterization. AbCdnD synthesizes two closely related products that co-elute with nearly all ion-exchange purification steps (Figure 2D) but can be separated on a C18 reverse-phase column (Figure 2E). Biochemical analysis demonstrates that the major (~65%) and minor (~35%) AbCdnD products are cyclic oligoadenylate species containing P1-resistant 2' -5' phosphodiester linkages (Figure S2C). Mass spectrometry analysis demonstrates that the major product is a cyclic trinucleotide and that the minor product peak is a mixture of cAAA and cA₄ species (Figure S2G). Interestingly, the minor product most potently activates AbCap4, with full enzyme activation and DNA degradation requiring low nanomolar concentrations of nucleotide second messenger (Figure S2D). Although the major product does not activate AbCap4 DNA degradation activity, this AbCdnD product is still capable of binding and stabilizing the AbCap4 enzyme, and we were able to determine a 2.2 Å co-crystal structure of the AbCap4-nucleotide second messenger complex (Figures 2F and S2E). Clear ligand density was observed for a cyclic trinucleotide bound within the AbCap4 globular C-terminal domain, allowing unambiguous assignment of this AbCdnD product as the cyclic trinucleotide 2'-5', 3'-5', 3'-5' c-AMP-AMP (2'3'3'-cAAA) and direct structural confirmation of the ability of a bacterial CD-NTase to synthesize noncanonical 2'-5'-linked RNAs (Figures 2G and S2F). To further confirm these findings, we verified, using mass spectrometry fragmentation analysis, that the AbCdnD major product matches a chemically synthesized 2'3'3'-cAAA standard (Figure S2G). 2'-5'phosphodiester linkages are rare in biology, and their role in nucleotide signaling has been suggested previously to be a unique adaptation evolved within eukaryotic innate immune signaling (Danilchanka and Mekalanos, 2013; Kranzusch et al., 2015; Margolis et al., 2017). These results demonstrate that 2'-5'-linked products are also involved in bacterial antiviral signaling and that Cap4 nucleases function as selective sensors that can use linkage specificity to adapt to distinct CD-NTase nucleotide second messenger signals.

Cap4 Contains a C-Terminal CARF Family Domain that Controls Ligand Specificity

The structure of the *Ab*Cap4–2'3'3'-cAAA complex reveals that the Cap4 C-terminal domain is responsible for nucleotide second messenger recognition. Previously, the Cap4 C-terminal SAVED domain was identified bioinformatically as enriched in CD-NTase-containing operons (Burroughs et al., 2015). Surprisingly, analysis of the *Ab*Cap4 SAVED domain structure reveals clear structural homology with CARF proteins, including Csm6 (DALI *Z* score, 4.2), with each half of the pseudo-symmetric SAVED domain containing homology to a single CARF domain (Figures 3A and 3B). The CARF domain of Csm6 from different CRISPR systems binds cA₄ or cA₆ cyclic oligoadenylate signals made by Cas10 following target recognition in type III CRISPR systems (Kazlauskiene et al., 2017; Niewoehner et al., 2017). Canonical CARF domain proteins like Csm6 homodimerize through CARF-CARF interactions to form a two-fold symmetric binding surface for nucleotide second messenger recognition (Jia et al., 2019; Niewoehner and Jinek, 2016). In contrast, the Cap4 SAVED domain is comprised of two individual CARF-like subunits joined with an ~25-amino acid internal linker between β strands 4 and 5. Using a thermal

shift assay to measure *Ec*Cap4 and *Ab*Cap4 nucleotide second messenger complex formation, we confirmed that the SAVED domain alone is sufficient to specifically recognize the activating signal (Figure S3A).

Comparison of the *Ab*Cap4 SAVED-2'3'3'-cAAA structure with previous CARF domain structures bound to RNA ligands reveals a mixture of shared and divergent features required for nucleotide second messenger recognition. The Csm6 homodimer has two-fold symmetry, with each monomer recognizing two nucleobases of the cA₄ signal (Jia et al., 2019; Molina et al., 2019). Unlike the larger cA₄ or cA₆ signals in type III CRISPR immunity, the majority of CD-NTase enzymes synthesize asymmetric nucleotide signals that lack internal two-fold symmetry (Whiteley et al., 2019). Pseudo-symmetry of the *Ab*Cap4 SAVED domain permits the existence of three unique pockets and specific recognition of each base of 2'3'3'-cAAA (Figures 3C, 3D, and S3B). The "A1 pocket" in *Ab*Cap4 is formed through a highly conserved aromatic W449 position that stacks against the nucleobase and Y454, which hydrogen-bonds with the adenine N7 nitrogen (Figures 3D and 3E). Nucleobase A2 is recognized through base-stacking interactions within a pocket formed by K299 and R301 (Figure S3B).

The A1 pocket is conserved in Cap4 and Csm6 homologs, demonstrating that nucleobase interactions at this site have been maintained throughout divergence of SAVED proteins from a CARF protein ancestor (Figure 3E). Mutagenesis confirms the importance of the A1 pocket in AbCap4 and EcCap4, with Cap4 proteins containing mutations within this pocket requiring >10-fold more ligand to induce similar levels of activation (Figure S3C). In contrast, the contacts to nucleobases A2 and A3 occur in pockets that do not exist in homodimeric CARF proteins. We determined an additional 2.4 Å structure of the AbCap4-3'3'3'-cAAA complex to further guide analysis of Cap4-ligand interactions and phosphodiester linkage specificity (Figure S4A). Comparison of the AbCap4-2'3'3'-cAAA and -3'3'3'-cAAA complexes demonstrates that a flat, parallel orientation of the nucleotide ribose at the A2 position along the SAVED domain surface allows the neighboring gap between the A1 and A2 pockets to accommodate a 3'-5' or 2'-5' linkage. In contrast, the perpendicular orientation of the A3 ribose constrains the nucleotide backbone to permit only a 3'-5' linkage and explains how the architecture within the SAVED domain binding pocket can dictate signal specificity. Diversification of the ligand binding interface through genetic fusion of two CARF-like subunits into a single-chain SAVED domain was likely a key evolutionary intermediate enabling specific recognition of diverse CD-NTase nucleotide second messengers that lack two-fold symmetry. Together, these data demonstrate that SAVED domains are divergent members of the CARF protein family and that the singlechain architecture of the Cap4 SAVED domain allows recognition of diverse asymmetric nucleotide recognition signals.

Cap4 Proteins Are Activated through Ligand-Dependent Oligomerization

To define the mechanism of Cap4 ligand-dependent activation, we next analyzed *Ec*Cap4 and *Ab*Cap4 proteins using negative-stain EM. In the presence of activating nucleotide ligand, Cap4 proteins oligomerize and form higher-order complexes (Figures 4A and S5A). The activated Cap4 complexes primarily adopt a dimeric state with two Cap4 proteins

stacked against each other in an SS-shaped configuration. We also observed Cap4 particles that correspond to higher-order oligomeric complexes (Figure S5B), and we confirmed, with size-exclusion chromatography-multiangle light scattering (SEC-MALS) analysis, that activated Cap4 in solution can oligomerize into multiple higher-order oligomeric states (Figures 4B and S5C). We classified the oligomerization status of ~20,000 particles from each condition to quantify the frequency of ligand-induced oligomerization. In agreement with Cap4-ligand interactions functioning as a requirement for high-order complex formation, <1% of the Cap4 particles occupied the dimeric or oligomeric state in the absence of an activating nucleotide signal (Figure 4A). The same higher-order oligomerization is observed in the presence of 45-bp target DNA, suggesting that nucleotide second messenger binding alone allows assembly of Cap4 into a fully active oligomeric state.

3D reconstruction at ~15 Å of the activated dimeric MoCap4 complex allowed docking of the high-resolution AbCap4-2'3'3'-cAAA crystal structure and further analysis of the mechanism of activation (Figure S5D). In the docked assembly, two SAVED domains stack against each other and result in alignment of the Cap4 endonuclease active sites. Docking of Cap4 within the 3D reconstructions suggests that no major domain rearrangement is required for activation (Figure S5B). In agreement, all four crystal structures of AbCap4 and MoCap4 share a fixed, rigid orientation of the Cap4 N-terminal endonuclease and Cterminal SAVED domains despite distinct packing and crystal forms (Figures S4B and S4C). These results support a model where recognition of the activating nucleotide second messenger likely induces a local conformational change in the SAVED domain that drives oligomerization and endonuclease activation. Oligomerization and positioning of two adjacent Cap4 endonuclease domains creates a singular surface to engage target DNA (Figure S5D). Using an electrophoretic mobility shift assay, we observed that Cap4 alone is unable to bind DNA and that ligand recognition is a pre-requisite for target DNA interaction (Figures 4C and S5E). An ~10- to 20-amino acid extension is conserved at the N terminus of each Cap4 effector but is not ordered in any of our AbCap4 or MoCap4 crystal structures (Figures S1B and S1C). Biochemical analysis of Cap4 proteins with N-terminal deletions shows that the N-terminal extension is dispensable for nucleotide signal recognition and oligomerization but required for DNA binding and endonuclease domain activation (Figure S6). Together, these results suggest a two-step model of Cap4 activation where nucleotide second messenger recognition in the SAVED domain induces Cap4 oligomerization and subsequent DNA binding and target degradation.

To further define the result of ligand-induced Cap4 activation, we next developed a deep sequencing approach to map the cleavage specificity and fragment distribution of Cap4 nuclease activity. Sequencing of DNA fragments remaining following degradation of genomic or plasmid DNA with *Ec*Cap4 reveals a consistent fragment length of ~17 bp and further suggests that DNA degradation occurs through defined oligomerization of multiple nuclease active sites (Figure 4D). In agreement with the promiscuous nuclease activity observed in biochemical assays (Figure 1), analysis of *Ec*Cap4 cut sites from mapped DNA fragments demonstrates a highly degenerate recognition sequence distinct from the strict sequence preference characteristic of type II restriction enzymes. *Ec*Cap4 exhibits preference for a minimal recognition sequence, 5' CNG, whereas *Ab*-Cap4 is less specific, targeting a 5' C or G and a final average fragment size of only ~6 bp (Figures 4D and S5F).

Mapping of the observed Cap4 cut sites in *E. coli* genomic DNA demonstrates complete degradation across nearly all regions of the genome (Figures 4E and S5G). Together, these data support a model of Cap4 regulation where ligand-induced oligomerization activates the endonuclease domain and results in promiscuous cleavage of DNA to minimal fragments.

SAVED Domain-Containing Proteins Are a Major Form of Viral Defense in Diverse Bacteria

SAVED domain-containing proteins occur in 29.8% of sequenced CD-NTase-containing operons and comprise one of the most prevalent effector modules in CBASS phage immunity (Figure 5A; Burroughs et al., 2015; Cohen et al., 2019; Whiteley et al., 2019). In addition to the endonuclease-SAVED architecture in Cap4 proteins, CBASS operons encode SAVED domains fused to additional protein partners, including putative HNH nucleases, caspase-like proteases, calcineurin-like phosphatases, Toll/interleukin-1 receptor (TIR) NAD ⁺ hydrolase enzymes, and transmembrane-containing segments (Figure 5B). Nucleotide second messenger-induced oligomerization of SAVED domains is likely a general strategy to activate diverse effector functions and coordinate abortive infection systems to limit phage replication.

Analysis of the bacterial CD-NTase phylogenetic tree reveals that SAVED domaincontaining proteins are encoded near vastly divergent CD-NTase enzymes from across clades B, C, D, G, and H (Figure 5A; Whiteley et al., 2019). The broad distribution of SAVED domain-containing effectors suggests that this domain can recognize diverse nucleotide second messengers. To test this hypothesis, we conducted a genetic screen to identify SAVED-containing proteins that specifically respond to the cyclic dinucleotide 3'3'-cGAMP. Co-expression analysis in *E. coli* with the promiscuously active CD-NTase V. cholerae DncV identified multiple SAVED domain-containing proteins in clades B and G that respond to 3'3'-cGAMP and induce cell death (Figure 5C). These effectors contain an HNH-SAVED architecture structurally distinct from the Cap4 DUF4297 endonuclease-SAVED architecture, and we therefore designated genes with this architecture CD-NTaseassociated protein 5 (Cap5). Purified Burkholderia pseudomallei Cap5 (GenBank: WP_004556385.1) is active in the presence of 3'3'-cGAMP and catalyzes robust dsDNA degradation in vitro (Figure 5D). Activation of each tested SAVED domain-containing protein requires recognition of a specific nucleotide second messenger, further demonstrating the specificity of CD-NTase signaling pathways (Figure 5D). Notably, the 3'3'-cGAMP-responsive SAVED domain effector *Bp*Cap5 failed to respond to the common bacterial second messengers 3'3' cyclic di-GMP and 3'3' cyclic di-AMP. Exquisite nucleotide ligand specificity therefore insulates CBASS immunity from other host signals and limits inappropriate activation.

CD-NTase operons protect bacterial populations during phage infection (Cohen et al., 2019; Ye et al., 2020). Interestingly, SAVED domain-containing proteins are also present within type III CRISPR operons, suggesting that the shared mechanism of SAVED- and CARF-dependent activation allows exchange of CBASS and CRISPR components and further diversification of antiviral immune defenses (Figure 6A). To verify that CD-NTase operons with SAVED domain-containing effectors also restrict phage replication, we used the *E. cloacae* operon containing *CdnD*, *Cap2*, *Cap3*, and *Cap4* and an *E. coli* operon containing

CdnG (GenBank: WP_000064266.1), *Cap2*, *Cap3*, and *Cap5* to analyze phage resistance (Figure 6B). Introduction of plasmids encoding one of these operons in *E. coli* mediated a >3-log decrease in efficiency of plaque formation for phage T2. Expression of the CdnG operon resulted in an ~1-log decrease in phage T5 plaque formation, whereas the CdnD operon had no effect. No effect was observed on replication of phage T7 (Figure 6B; Cohen et al., 2019; Ye et al., 2020). Restriction of phage replication is strictly dependent on the catalytic activity of the CD-NTase and the nuclease active site of Cap4 (Figure 6B). Together, these data demonstrate a broad strategy in bacteria that couples CD-NTase signaling to activation of downstream SAVED-containing proteins to protect against viral infection.

DISCUSSION

Here we identify Cap4 as the founding member of a widespread family of antiviral effector proteins in bacteria. We show that Cap4 proteins are nucleotide second messenger sensors that function as downstream DNA endonuclease effectors in CBASS immunity. Following phage infection, CBASS immunity begins with activation of a CD-NTase enzyme to catalyze nucleotide second messenger synthesis (Cohen et al., 2019; Whiteley et al., 2019). Although the mechanism of phage recognition remains unclear, many bacterial CD-NTase enzymes are constitutively active *in vitro* and may therefore function similar to guards in plant immunity and sense phage replication indirectly by detecting changes in metabolite concentrations or loss of homeostasis (Dodds and Rathjen, 2010; Kranzusch, 2019; Whiteley et al., 2019). After CD-NTase activation, Cap4 recognizes the resulting nucleotide signal and couples ligand binding with induction of a promiscuous DNA cleavage response that restricts phage replication. SAVED domain-containing proteins like Cap4 are found in ~30% of CBASS operons and provide a clear mechanism for how these systems mediate phage resistance.

Structures of Cap4 explain a molecular mechanism for ligand recognition and effector function activation. Following nucleotide second messenger recognition in the SAVED domain, Cap4 oligomerizes to closely stack two nuclease domains (Figures 3, 4, and S4D). The Cap4 nuclease effector domain is therefore maintained in an inactive monomeric state until detection of the correct CD-NTase signal enables higher-order complex assembly and target DNA degradation. The Cap4 SAVED domain is sufficient to recognize the activating nucleotide signal (Figure S3A), indicating that the effector domain does not participate in ligand specificity. Additionally, we characterize an HNH endonuclease-SAVED effector Cap5 from *B. pseudomallei* and demonstrate that SAVED domains can activate structurally divergent effector domains (Figure 5D). These results support a sensor-effector model where fusion of a SAVED domain allows nucleotide second messenger-gated control of diverse effectors. SAVED domains are fused to a wide variety of effector domains, including proteases, phosphodiesterases, and potentially pore-forming transmembrane proteins (Burroughs et al., 2015; Cohen et al., 2019), supporting that the generalizability of SAVEDdependent activation allows diverse domains to be co-opted as modules for viral defense.

We show that the SAVED domain formed through fusion of two CARF-like subunits into a single-chain nucleotide second messenger sensor (Figure 3). These results explain how

SAVED domains recognize diverse asymmetric ligands and reveal an unexpected shared evolutionary history of CBASS and CRISPR antiviral systems. Nearly all characterized type III CRISPR systems function through homodimeric CARF proteins that recognize a nucleotide second messenger with two-fold symmetry (Kazlauskiene et al., 2017; Niewoehner et al., 2017). Fusion of two CARF-like subunits into a SAVED domain breaks the requirement for two-fold symmetrical ligands and explains a key evolutionary event that enabled CD-NTase systems to signal with diverse asymmetric ligands. A similar evolutionary event was recently observed with fusion of two CARF domains in a Thermus thermophilus type III CRISPR effector (McMahon et al., 2020; Figure S4D), further highlighting how genetic fusion of multiple CARF domains can provide an evolutionary advantage in antiviral immunity. Interestingly, a structurally unrelated 3'3'3'-cAAAactivated homotrimeric DNA endonuclease, NucC, was identified in an E. coli CdnC CD-NTase system (Lau et al., 2020). NucC uses a three-fold symmetric ligand-binding pocket, suggesting that SAVED and NucC effectors are alternative evolutionary paths that enabled nucleotide second messenger diversification within CBASS immunity. CD-NTase enzymes synthesize nucleotide second messengers using all four nucleobases as building blocks (Whiteley et al., 2019), and we now show that CD-NTase product second messengers are further specified with incorporation of 2'-5' and 3'-5' phosphodiester linkages. These findings dramatically expand the known nucleotide second messenger signaling space and indicate that CD-NTase enzymes are theoretically capable of synthesizing at least 180 distinct oligonucleotide variants. Combined with cA4 and cA6 species in type III CRISPR immunity and linear 2'-5'-linked chains produced by mammalian oligoadenylate synthase enzymes (Hornung et al., 2014), these results reveal a vast diversity of natural nucleotide second messenger signals in viral defense.

The enormous diversity of CD-NTase nucleotide second messenger signals likely enables CBASS antiviral systems to specialize and adapt to phage resistance. Mammalian poxviruses encode a 2'3'-cGAMP-specific nuclease that degrades the cGAS product and prevents STING-dependent immune responses (Eaglesham et al., 2019). Likewise, a phage protein was recently identified that degrades cA₄ to evade type III CRISPR immunity (Athukoralage et al., 2020). Viral nucleases that specifically degrade second messengers likely create evolutionary pressure to diversify antiviral signaling systems. The ability of CBASS immunity to morph between cyclic dinucleotide and cyclic trinucleotide signals may represent larger "leaps" that allow escape and temporary relief from an evolutionary arms race (Daugherty and Malik, 2012). Our discovery of a role for 2'-5' linkages in A. baumannii CBASS immunity demonstrates that incorporation of noncanonical linkages is another mechanism that evolved in bacteria to subvert viral resistance. Along with the previous identification of diverse cGAS-like enzymes in bacteria and effector proteins with sequence homology to mammalian STING (Cohen et al., 2019; Whiteley et al., 2019), the discovery of bacterial signals with 2'-5' linkages demonstrates that all fundamental components of mammalian cGAS-STING signaling are functionally shared within ancestrally related bacterial CBASS immunity.

STAR * METHODS

RESOURCE AVAILABILITY

Lead Contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Philip J. Kranzusch (philip_kranzusch@dfci.harvard.edu).

Materials Availability—This study did not generate new unique reagents.

Data and Code Availability—Coordinates of the Cap4 structures have been deposited in the RCSB Protein Data Bank under the following accession numbers: *Mo*Cap4 (6VM5), *Ab*Cap4 (6WAM), *Ab*Cap4–2'3'3'-cAAA (6VM6), *Ab*Cap4–3'3'3'-cAAA (6WAN).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Escherichia coli strains—Recombinant CD-NTase and Cap4 proteins were expressed in *E. coli* BL21-RIL DE3 (Agilent) bacteria harboring a pRARE2 tRNA plasmid. Single transformants were inoculated into a 20 mL MDG media starter culture (0.5% glucose, 25 mM Na₂HPO₄, 25 mM KH₂PO₄, 50 mM NH₄Cl, 5 mM Na₂SO₄, 2 mM MgSO4, 0.25% aspartic acid, 100 mg ml⁻¹ ampicillin, 34 mg ml⁻¹ chloramphenicol, and trace metals), grown overnight at 37°C, and used to seed 1 L M9ZB media cultures (0.5% glycerol, 1% Cas-Amino Acids, 47.8 mM Na₂HPO₄, 22 mM KH₂PO₄, 18.7 mM NH₄Cl, 85.6 mM NaCl, 2 mM MgSO₄, 100 mg ml⁻¹ ampicillin, 34 mg ml⁻¹ chloramphenicol, and trace metals) grown at 37°C, 230 RPM until an OD₆₀₀ of ~2.5. Cultures were induced by chilling flasks on ice for 20 min, supplementing cultures with to 0.5 mM IPTG, and then incubating at 16°C, 230 RPM for ~15 h before harvest.

Phage challenge experiments were performed with electrocompetent *E. coli* MG1655 recovered and plated on selected LB media (carbenicillin 100 μ g ml⁻¹). Single colonies were inoculated into selective MMCG medium (1 × M9 Minimal Salts, 0.4% glucose, 0.02% MgSO₄, 0.001% CaCl₂, plus 100 μ g ml⁻¹ carbenicillin) and cultivated at 37°C shaking for ~20 h. Cultures were diluted 1:100 into selective MMCG medium and cultivated at 37°C shaking for 4 h to harvest mid-log cultures.

Bacteriophage strains—High-titer T2 phage (Coli Genetic Stock Center CGSC12141), T5 (CGSC12144), or T7 (CGSC12146) lysate stocks were purified from *E. coli* MG1655 cultures grown in LB.

METHOD DETAILS

Protein expression and purification—Recombinant CD-NTase and effector proteins were purified from *E. coli* as previously described (Whiteley et al., 2019; Zhou et al., 2018). Briefly, effector and CD-NTase enzymes were cloned from synthetic DNA (Integrated DNA Technologies) into a custom pET vector encoding a 6× His-SUMO2-tagged fusion protein and expressed in BL21-RIL *E. coli* (Agilent) containing the rare tRNA plasmid pRARE2. *E. coli* from large-scale M9ZB cultures (typically 2 L for each construct) were pelleted, washed with 1× PBS, and lysed by sonication in 1× Lysis Buffer (20 mM HEPES-KOH pH 7.5, 400

mM NaCl, 30 mM imidazole, 10% glycerol and 1 mM DTT). Recombinant protein was purified by gravity chromatography and binding to Ni-NTA resin (QIAGEN). Resin was washed with 1× Lysis Buffer supplemented to 1 M NaCl and then eluted with 1× Lysis Buffer supplemented to 300 mM imidazole. Purified protein was supplemented with recombinant human SENP2 protease (D364–L589, M497A) to remove the SUMO2 tag, and dialyzed overnight against 20 mM HEPES-KOH pH 7.5, 250 mM KCl, 1 mM DTT. Proteins were concentrated using a 30K-cutoff concentrator (Millipore) and further purified by size-exclusion chromatography on a 16/600 Superdex 200 column in 20 mM HEPES-KOH pH 7.5, 250 mM KCl, 1 mM TCEP. Final purified fractions were analyzed by SDS-PAGE and Coomassie staining, concentrated to > 30 mg ml⁻¹, flash-frozen in liquid nitrogen, and stored as 35 μ L aliquots at –80°C.

Electrophoretic mobility shift assay—Radiolabeled nucleotide products were enzymatically synthesized with purified *V. cholerae* DncV or *Ec*CdnD in reactions containing 25 mM ATP, 25 μ M GTP, trace α^{32} P-GTP in 13 Reaction Buffer (5 mM MgCl₂, 50 mM KCl, 1 mM TCEP and either 50 mM Tris-HCl pH 7.5 (DncV) or 50 mM CAPSO pH 9.4 (*Ec*CdnD)) in a final reaction volume of 20 μ l. Synthesis reactions were incubated ~15 h at 37°C and then inactivated by treating with 1 μ L of 5 units μ l⁻¹ Calf Intestinal Phosphatase (New England Biolabs) and incubating at 37°C for 1 h. Radiolabeled products were then diluted 1:20 into electrophoretic mobility shift reactions containing either no protein, ~1 μ M 6× His-SUMO2-*Ec*Cap2, 6× His-SUMO2-*Ec*Cap3, or 6× His-SUMO2-*Ec*Cap4 and 50 mM KCl, 50 mM Tris-HCl pH 7.5, 5 mM MgCl₂, and 1 mM TCEP. Reactions were incubated for 20 min at 25°C, then separated on an 8 cm 10% nondenaturing polyacrylamide gel run at 100 V for 45 min. The gel was dried and then exposed to a phosphor-screen and imaged with a Typhoon Trio Variable Mode Imager (GE Healthcare).

Crystallization and structure determination—Cap4 proteins were crystallized at 18°C using the hanging drop method. Concentrated protein stocks were thawed on ice and diluted in buffer (25 mM HEPES-KOH pH 7.5, 1 mM TCEP). Drops were set by mixing protein stock and reservoir solution in 2 µl drops over a 350 µl reservoir in Easy-Xtal 15well trays (QIAGEN). Each protein was crystallized as follows: 1) Apo MoCap4: Native or selenomethionine MoCap4 protein was diluted to 10 mg ml⁻¹, supplemented with 10.5 mM MgCl₂, mixed at a 1:1 ratio of protein:reservoir solution (110 mM Tris-HCl pH 8.5, 21% PEG-400), crystals were allowed to grow for 5 days and then were cryoprotected using reservoir solution supplemented with 10% ethylene glycol. 2) AbCap4-2'3'3'-cAAA: Native AbCap4 protein was diluted to 10 mg ml⁻¹ and pre-incubated with 10.5 mM MgCl₂ and ~200 µM purified AbCdnD product, mixed at a 1:1 ratio of protein:reservoir solution (10% PEG-3350, 200 mM lithium sulfate, 100 mM imidazole pH 8.0), crystals were allowed to grow for 25 days and then were cryoprotected using reservoir solution supplemented with 25% ethylene glycol; Selenomethionine AbCap4 was diluted to 10 mg ml⁻¹ and preincubated with 10.5 mM MgCl₂ and ~200 µM purified AbCdnD product, mixed at a 1:1 ratio of protein:reservoir solution (0.2 M lithium sulfate, 0.1 M Tris-HCl pH 8.5, 25% PEG-400), crystals were allowed to grow for 13 days and then were cryoprotected using reservoir solution supplemented with 10% PEG-400. 3) Apo AbCap4: Native AbCap4 protein was diluted to 10 mg ml⁻¹ and mixed at a 1:1 ratio of protein:reservoir solution (0.1

M Tris pH 8.5, 0.2 M LiSO₄, 16% PEG-3350), crystals were allowed to grow for 6 days and then were cryoprotected using reservoir solution supplemented with 30% ethylene glycol. 4) AbCap4–3'3'3'-cAAA: Native AbCap4 protein was diluted to 10 mg ml⁻¹ and preincubated with 500 μ M 3'3'3'-cAAA (Biolog), mixed at a 1:1 ratio of protein:reservoir solution (0.1 M Tris pH 8.5, 0.2 M LiSO₄, 16% PEG-3350), crystals were allowed to grow for 6 days and then were cryoprotected using reservoir solution supplemented with 25% ethylene glycol.

X-ray data were collected at the Lawrence Berkeley National Laboratory Advanced Light Source beamline 8.2.2 and at the Northeastern Collaborative Access Team beamlines 24-ID-C and 24-ID-E (P30 GM124165), and used a Pilatus detector (S10RR029205), an Eiger detector (S10OD021527) and the Argonne National Laboratory Advanced Photon Source (DE-AC02–06CH11357). X-ray crystallography data were processed with XDS and AIMLESS (Kabsch, 2010) using the SSRL autoxds script (A. Gonzalez, Stanford SSRL). Experimental phase information for *Mo*Cap4 and *Ab*Cap4 was determined using data collected from selenomethionine-substituted crystals. In total, 5 and 36 sites respectively were identified with HySS in PHENIX (Adams et al., 2010) and an initial map was produced using SOLVE/RESOLVE (Terwilliger, 1999). Model building was performed using Coot (Emsley and Cowtan, 2004), before refinement in PHENIX. Statistics were analyzed as described in Table S1 (Chen et al., 2010; Karplus and Diederichs, 2012; Weiss, 2001).

DNA degradation assays—DNA degradation assays were performed by incubating 50 nM Cap4 protein with 10 ng μ l⁻¹ pGEM9z plasmid (Promega), *E. coli* genomic DNA, or 100 bp synthetic DNA as specified in a 25 μ l reaction for 20 min at 37°C with 10 mM Tris-HCl pH 7.5, 25 mM NaCl, 5 mM MgCl₂, and 1 mM TCEP. Degradation reactions were supplemented with activating nucleotide second messenger product at 50 nM synthetic or purified product (*Ab*CdnD product), or unpurified cognate CD-NTase reaction (see below). Reactions were stopped by addition of 6× loading buffer (60 mM EDTA, 20 mM Tris-HCl pH 7.5, 30% glycerol, 0.1% SDS), and then 10 μ l was separated on a 1.5% or 2% TAE agarose gel. Gels were run at 100 V for 20 min, then stained by rocking at room-temperature in buffer with 10 μ g ml⁻¹ ethidium bromide for 30 min. Gels were de-stained in water for ~10 min and then imaged with a ChemiDoc MP Imaging System.

AbCdnD product purification—Large scale *Ab*CdnD product synthesis was carried out in 250 mL reactions with 500 nM purified *Ab*CdnD and 250 μ M ATP in 13 reaction buffer (50 mM KCl, 20 mM MgCl₂, 1 mM MnCl₂, 1 mM TCEP, 40 mM Tris pH 9.0), incubated overnight at 37°C with gentle shaking. The reaction was then treated with 20 μ l of 5 units μ l ⁻¹ Calf Intestinal Phosphatase (New England Biolabs) at 37°C for 3 h, then 0.2 mm filtered. The reaction was diluted to 12.5 mM salt with water, then loaded on to a HiTrap Q column (GE Healthcare) at 5 mL min⁻¹ with a peristaltic pump and eluted on an AKTA FPLC (GE Healthcare) with a linear gradient of ammonium acetate from 0 to 2 M over 50 min. The *Ab*CdnD product elutes as a single major peak at approximately 1.5 M ammonium acetate. Fractions containing the major product were identified by absorbance at 254 nm, pooled and dried by speedvac, and then re-suspended in approximately 1 mL of water. Initial ion

exchange-purified product was desalted by gel filtration on a Superdex 30 Increase 10/300 GL column (GE Healthcare) run at 0.5 mL min⁻¹ with water as running buffer.

Second-step purification was carried out using a C18 column (Agilent Zorbax Bonus-RP 4.6 3 150 mm, 3.5-micron). The column was heated to 50°C and run at 1 mL min⁻¹ with a mobile phase of 50 mM NaH₂PO₄ (pH 6.8 with NaOH) supplemented with 3% acetonitrile. The sample was injected with ~10 µg per run to ensure separation of major and minor products. Fractions from several runs were pooled and dried by speedvac overnight, then desalted using C18 cartridges (Thermo Scientific Hypersep C18 100 mg columns). Speedvac-dried fractions were re-suspended in nuclease-free water with 2% acetic acid. The C18 cartridge was pre-washed with 500 µl MeOH, then 2× with 500 µl water with 2% acetic acid before loading the sample over the cartridge 3×. The cartridge was washed 2× with 500 µl water with 2% acetic acid, then eluted with 500 µl MeOH. Sample was dried by speedvac for approximately 2 h, then re-suspended in nuclease-free water. Concentrations were estimated using an extinction coefficient of $\varepsilon = 40,500 \, \mathrm{l} \, \mathrm{mol}^{-1} \, \mathrm{cm}^{-1}$.

CD-NTase product thin layer chromatography—CD-NTase reactions were performed with 5 μ M recombinant enzyme in 20 μ l reactions with 25 μ M NTPs and trace α^{32} P-ATP (mcGAS: ATP, GTP; DncV: ATP, GTP; *Ec*CdnD: ATP, GTP; *Ab*CdnD: ATP) overnight at 37°C with 50 mM KCl, 10 mM MgCl₂, 1 mM TCEP, and 50 mM CAPSO pH 9.4 (CdnD) or 50 mM Tris-HCl pH 7.5 (cGAS, DncV). cGAS reactions were additionally supplemented with 2 μ M 45-bp DNA, and *Ab*CdnD reactions were additionally supplemented with 1 mM MnCl₂. Reactions were terminated with addition of 1 μ of 5 units μ l⁻¹ Calf Intestinal Phosphatase (New England Biolabs) and incubating at 37°C for 1 h. For P1 degradation assays, 10 μ l of each reaction was treated with 1 μ l P1 nuclease (Sigma Product N8630) for 1 h at 37°C. Reactions were analyzed by thin-layer chromatography by diluting each reaction 1:10 in 100 mM sodium acetate pH 5.2 and then spotting 1 μ l of each reaction on a PEI-cellulose plate (Millipore) developed in ~1 cm of 1.5 M KH₂PO₄ pH 3.8 until buffer was 1 cm from the top of plate. Plates were dried then expose to a phosphorscreen and imaged with a Typhoon Trio Variable Mode Imager (GE Healthcare).

AbCdnD product degradation analysis—Synthetic controls (3'3'3'-cAAA and 2'3'3'-cAAA) or purified *Ab*CdnD product were diluted to 25 µM in 100 µl reactions and supplemented with 10× P1 buffer (final 1× concentration: 30 mM NaOAc pH 5.3, 5 mM ZnSO₄, 50 mM NaCl) or snake venom phosphodiesterase buffer (final 1× concentration: 50 mM Tris pH 9, 10 mM MgCl2, 50 mM NaCl). Snake venom phosphodiesterase (0.8 mU) (Sigma, Phosphodiesterase I from *Crotalus adamanteus* venom, Product P3243) or Nuclease P1 (80 mU) (Sigma Product N8630) was added and reactions were incubated for 1 h or 5 h at 37°C, respectively. Reactions were diluted 1:2 in nuclease-free water, then filtered through a 10-kDa filter (Millipore) by centrifugation. The degradation products were analyzed using a C18 column (Agilent Zorbax Bonus-RP 4.6 × 150 mm, 3.5-micron) on HPLC held at 50°C and run at 1 mL min⁻¹ with 50 mM NaH₂PO₄ (pH 6.8 with NaOH) supplemented with 3% acetonitrile.

Chemical synthesis of 2'3'3'-cAAA—All reagents and solvents for chemical operations were of analytical grade or the best grade available from commercial suppliers.

Solvents for chromatographic operations were specified as analytical grade, HPLC grade, or gradient HPLC grade. YMC*Gel SIL (6 nm, S-75 μ m) was used for preparative flash chromatography and TLC was performed with Merck 60 F254 silica gel plates. All chromatographic operations were performed at ambient temperature. Evaporation of solvents was accomplished by rotary evaporation *in-vacuo* either with membrane pump vacuum or oil pump high vacuum with water bath temperatures not exceeding 30–33°C.

UV-spectra were recorded on a JASCO V-650 spectrometer in phosphate buffered aqueous solution (pH 7). Mass spectra were generated with a Bruker Esquire LC 6000 spectrometer in the electrospray ionization mass spectrometry (ESI-MS) mode with 50% water / 49.5% methanol / 0.5% NH₃ (pH 9–10) as matrix. Nuclear magnetic resonance (NMR) spectra were recorded with a 400 MHz Bruker Avance III HD and chemical shifts are expressed in parts per million (ppm). Chemical shifts were referenced to the DMSO solvent signal, 2.50 ppm for ¹H. 85% phosphoric acid was used as external standard for ³¹P NMR spectra with 0 ppm. All ³¹P NMR spectra were recorded with proton decoupling. VWR / Hitachi: LaChromElite L-2130 Pump; VWR / Hitachi: LaChromElite L-2420 UV/Vis detector; VWR / Hitachi: LaChromElite organizer, Agilent Technologies: OpenLAB Control Panel A.02.01.

3 mmol of cyanoethyl phosphoramidite 5'-DMTr-2'-TBDMS-3'-CEP-N⁶-Bz-adenosine (ChemGenes, Wilmington, MA, USA, Cat. No. ANP-5671) were used as starting material for the synthesis of the protected dimeric linear precursor 5'-OH-2'-TBDMS-N⁶-Bzadenosine-(3' /5')-cyanoethyl-phosphate-2'-TBDMS-3'-H-phosphonate-N⁶-Bz-adenosine with a standard oligonucleotide coupling protocol, originally developed for cyclic dinucleotides (Gaffney et al., 2010). After preparative flash chromatography purification on silica gel with chloroform / methanol (1:1), the linear dimer was evaporated to dryness. 7.5 mmol (2.5 eq.) 5-ethylthio-tetrazole as coupling reagent were added and the resulting mixture was evaporated 4 times from 40 mL absolute acetonitrile. The last evaporation was stopped at ~15 mL total volume, 0.5 g molecular sieves 3 Å were added and the mixture was stirred at room temperature for 30 min. 6 mmol (2 eq.) cyanoethyl phosphoramidite 5'-DMTr-2'-CEP-3'-TBDMS-N6-Bz-adenosine (ChemGenes, Wilmington, MA, USA, Cat. No. ANP-5681) were added and the protected trimeric linear precursor 5'-OH-3'-TBDMS-N⁶-Bz-adenosine-(2'/5')-cyanoethyl-phosphate-2'-TBDMS-N⁶-Bz-adenosine-(3'/5')cyanoethyl-phosphate-2'-TBDMS-3'-H-phosphonate-N6-Bz-adenosine was prepared as previously described (Gaffney et al., 2010). After preparative flash chromatography purification on silica gel with chloroform / methanol (1:1), the linear trimer was evaporated to dryness. The final cyclization step and the release of protection groups was performed according to the standard protocol in Gaffney et al. (2010), leading to the raw product 2'3'3'-cAAA after evaporation of solvents.

100 mL water was added and the resulting suspension was placed in an ultrasonic bath at room temperature for 15 min, followed by 3 extraction cycles with 50 mL chloroform each. The combined organic phases were extracted with 50 mL water and the combined product-containing aqueous phase was filtered with a 0.45 µm regenerated cellulose (RC) filter and partially concentrated under reduced pressure to remove traces of chloroform. The complex product solution was diluted with water to 1000 mL and applied to a Q Sepharose Fast Flow

anion exchange column (40–165 μ m; 380 \times 50 mm) Cl–-form (Sigma), previously regenerated with 2 M sodium chloride and washed with water. The column was washed with water (1000 ml), followed by a gradient of 0-600 mM triethylammonium bicarbonate buffer (TEAB, pH 7, 7500 ml) in water, 500 mL 600 mM TEAB and 1500 mL 1 M TEAB (detection wavelength 254 nm). The title compound eluted with ~400 mM TEAB. Productcontaining fractions were carefully concentrated to a final volume of approximately 20 mL with a rotary evaporator equipped with a drop catcher in-vacuo. Subsequent purification of 2'3'3'-cAAA were accomplished by repeated preparative reversed phase medium pressure liquid chromatography (MPLC). The product solution was applied to a LiChroprep RP-18 column (15–25 μ m; 450 \times 50 mm) (Merck) previously equilibrated with 100 mM triethylammonium formate (TEAF, pH 6.8) in water. Elution was performed with 100 mM TEAF, 20 mM TEAF, followed by a step-gradient of 1%, 2%, 3%, 4% and 10% 2-propanol, 20 mM TEAF (pH 6.8) in water. Final purification of product containing fractions was accomplished by a LiChroprep RP-18 column $(15-25 \text{ mm}; 430 \times 25 \text{ mm})$ (Merck) with a similar step-gradient of 2-propanol, 20 mM TEAF as organic modifier. For desalting, 2'3'3'-cAAA fractions of sufficient purity were applied to an in-line set-up of two LiChroprep RP-18 columns (15–25 μ m; 450 \times 50 mm each) (Merck), previously equilibrated with water. The columns were washed with water to remove excess TEAF buffer. Afterward, 2% 2-propanol in water was used to elute the desalted 2'3'3'-cAAA. To generate the sodium salt form of 2'3'3'-cAAA, pooled product-containing fractions were partially concentrated under reduced pressure and subsequently applied to a Toyopearl SP-650M cation exchange column (65 μ m; 90 \times 35 mm) Na⁺-form (Sigma), previously regenerated with 2 M sodium chloride and washed with water. For elution the column was washed with water until no UV-absorbance was detectable at 254 nm anymore. After filtration and careful evaporation under reduced pressure, 845.6 µmol 2'3'3'-cAAA, sodium salt was isolated with a purity of 99.02% by HPLC (theoretical yield: 28.2%).

Formula (free acid): C₃₀H₃₆N₁₅O₁₈P₃ (MW 987,63 g/mol)

UV-Vis (water pH 7.0): λ_{max} 259 nm; ϵ 40500.

ESI-MS pos. mode: m/z 988 (M+H)⁺, m/z 1010 (M+Na)⁺, m/z 1032 (M-H+2Na)⁺.

ESI-MS neg. mode: m/z 986 (M-H)-, m/z 1008 (M-2H+Na)-, m/z 1030 (M-3H+2Na)-.

¹H NMR (400 MHz, Deuterium Oxide) δ 8.51 (s, 1H), 8.47 (s, 1H), 8.24 (s, 1H), 8.22 (s, 1H), 8.21 (s, 1H), 8.18 (s, 1H), 6.34 (d, *J* = 7.8 Hz, 1H), 6.20 (d, *J* = 7.0 Hz, 1H), 6.17 (d, *J* = 4.2 Hz, 1H), 5.25 – 5.18 (m, 1H), 4.89 – 4.71 (m, 7H), 4.58 – 4.54 (m, 2H), 4.37 – 4.20 (m, 6H) ppm.

³¹P NMR (162 MHz, Deuterium Oxide) & 0.25 (s, 1P), 0.02 (s, 1P), -0.29(s, 1P) ppm.

Analytical HPLC: (Kromasil 100–10, RP-8 (10 μ m; 250 × 4 mm)) 3% acetonitrile, 50 mM sodium dihydrogen phosphate buffer, pH 6.8; 1.5 mL/min; UV 259 nm; t_{RET} 5.84 min.

Liquid chromatography-mass spectrometry—To characterize the enzymatic products of *Ab*CdnD, LC-MS/MS analysis was performed as previously described (Lau et

al., 2020; Ye et al., 2020). LC-MS/MS was performed on a Thermo Vanquish UHPLC coupled to a Thermo QExactive Hybrid Quadrupole-Orbitrap Mass Spectrometer (Thermo Scientific). Chromatography utilized a Sequant ZIC-pHILIC polymeric column (100 mm \times 2.1 mm, 5 μ m) (EMD Millipore) maintained at 45°C and a flow rate of 0.4 mL min⁻¹. AbCdnD products and cyclic oligonucleotide standards were separated by injecting $2 \mu l$ of sample and eluting on the following linear gradient: (A) 20 mM ammonium bicarbonate in water, pH 9.6, and (B) acetonitrile; 90% B for 0.25 min, a linear gradient to 55% B at 4 min, and sustained until 6 min. Column was re-equilibrated for 2.5 min at 90% B. Detection of cyclic oligonucleotides was performed in positive ionization mode using an heated electrospray ionization (HESI) source with the following parameters: spray voltage of 3.5 kV; sheath gas, auxiliary gas, and sweep gas flow rates of 40, 20, and 2 AU, respectively; capillary and auxiliary gas heater temperature of 275 and 350°C, respectively. Profile MS1 spectra were acquired under the following parameters: mass resolution of 35,000, AGC volume of 1×10^6 maximum IT of 75 ms and a scan range from 450 to 1400 m/z to account for z = 1 and z = 2 ions of cyclic tri- and tetraadenosine nucleotides. Data-dependent tandem mass spectra were acquired using CID of the following settings: mass resolution of 17,500, AGC volume of 1×105 , maximum IT of 50 ms; a loop count of 5, isolation window of 1.5 m/z; normalized collision energy of 25 eV; dynamic exclusion was not used. Data reported are for the most common ion for each indicated cyclic oligonucleotide.

Thermal shift assay—Proteins were mixed with $3 \times$ Sypro dye and 100 µM purified nucleotide ligand (3'3'3'-cAAG or IEX-purified *Ab*CnD product) in 20 mM HEPES-KOH pH 7.5 and 75 mM KCl, with a final protein concentration of 10 µM. Using a BioRad CFX96 thermocycler, samples were brought from 25 to 95°C, reading fluorescence in HEX channel every 0.5°C. The derivative of each curve over time was calculated using BioRad CFX Manager, then normalized as a percent maximum change in fluorescence for each sample.

Negative stain electron microscopy-0.05-0.1 mM purified Cap4 proteins were mixed with equimolar amounts of cyclic trinucleotide and DNA as indicated. For negative staining, all samples were adjusted to 0.0025-0.005 mg ml⁻¹ in 20 mM HEPES-KOH pH 7.5, 150 mM KCl, 1 µM TCEP without or with 1 mM activating ligand. Negative stain grids were prepared by applying the sample $(3 \ \mu)$ to a glow-discharged $(30 \ s, 30 \ mA)$ 400-mesh Cu grid (Electron Microscopy Sciences) coated with an ~15 nm layer of continuous carbon (Safematic CCU-010). After 30 s, the grid was blotted from the side, immediately stained with 1.5% uranyl formate, and blotted again. The staining procedure was repeated two times, with a 30 s incubation with uranyl formate before the final blotting step. The grid was allowed to dry for at least 15 min before imaging. Samples were imaged with a Tecnai T12 (FEI) transmission electron microscope operated at 120 keV and equipped with a Gatan 4K \times 4K CCD camera. Images were collected at a nominal magnification of 67,000 \times and pixel size of 1.68 Å with defocus values of \sim 1.0–2.0 µm. Between 80 and 333 micrograph images were collected for each of the 9 datasets. Image processing was done in RELION-3.0 (Zivanov et al., 2018). After CTF estimation with GCTF (Zhang, 2016), particle picking was carried out with gautomatch (Kai Zhang; https://www.mrc-lmb.cam.ac.uk/kzhang/) or LoGbased autopicking (Zivanov et al., 2018) and manually inspected. Particles were extracted

with a box size of 156 pixels and subjected to reference-free 2D classification. Particles in the best-resolved classes were selected and subjected to an additional one to two rounds of 2D classification. In addition, for each dataset, random subsets of 10,000 particles were subjected to 2D classification to compare the distributions of particles in different image groups. For datasets with more than 10,000 particles, the standard error of mean of the distributions between four independent subsets of 10,000 particles ranged from 0.1 to 1.1%. Particles retained after 2D classification were used to generate a *de novo* initial model using the stochastic gradient descent algorithm in RELION, followed by 3D refinement and classification. Overall resolutions of EM maps of dimer complexes were ~15 Å. Rigid body docking was performed using Chimera (Pettersen et al., 2004).

SEC-MALS—Cap4 and Cap4–nucleotide second messenger complex samples were prepared by diluting in SEC-MALS running buffer (150 mM KCl, 20 mM HEPES-KOH pH 7.5 and 1 mM TCEP) to 2 mg ml⁻¹ and incubating on ice for 5 min with 100 μ M nucleotide second messenger and then separated on an SRT SEC-300 column (Sepax). Protein concentration was calculated using refractive index on a Wyatt Optilab T-rex Refractive Index Detector assuming dn dc⁻¹ of 0.185 and a molar mass was calculated using a Wyatt Dawn Heleos II Multi-Angle Light Scattering detector and ASTRA software.

DNA EMSA assay—Catalytically inactive Cap4 proteins (*Ab*Cap4 K69A, *Ec*Cap4 K74A) were mixed on ice at a concentration of 10 μ M with 1 μ M 45 bp DNA and 50 μ M 3'3'3'-cAAG (*Ec*Cap4) or IEX-purified *Ab*CdnD product (*Ab*Cap4) in a final reaction volume of 20 μ L containing 20 mM HEPES-KOH pH 7.5, 50 mM KCl, 5 mM MgCl₂ and 1 mM TCEP. Reactions were incubated on ice for 5 min and then supplemented with to a final concentration of 5% glycerol (v/v). Samples were separated on a 2% native agarose TB gel containing 100 mM Tris and 45 mM boric acid by running at 250 V for 45 min at 4°C. The gel was stained by soaking in buffer with 10 μ g ml⁻¹ ethidium bromide solution for 20 min and then imaged with ChemiDoc MP Imaging System.

DNA fragment sequencing—Cap4 proteins were incubated with 400 ng plasmid or genomic DNA in 80 µl reactions for 2 h (500 nM protein, 100 nM activating ligand, 10 mM Tris-HCl pH 7.5, 25 mM NaCl, 1 mM TCEP, 5 mM MgCl₂), then DNA fragments were purified by phenol-chloroform extraction. Sequencing libraries were made from single stranded DNA fragments using an Accel-NGS 1S Plus DNA Library kit (Swift Biosciences) as previously described (Lau et al., 2020) and sequenced on a NextSeq500. For bioinformatics analyses, given that Cap4 digestion results in small fragments, we took advantage of that fact that reads containing the 3' adaptor sequence have the full fragment sequence. Reads containing the 3' adaptor were selected and adaptor trimmed using Cutadapt (Martin, 2011). The 8 nucleotide low complexity sequence was trimmed by Cutadapt, and reads were mapped to the pGEM9z(–) plasmid (Promega) or the *E. coli* K12 genomic sequence (GenBank U00096.3) using Bowtie2 (Langmead and Salzberg, 2012). The location of the 5' end of each read was used as the cut site and extracted from the output sam files. The 10 nucleotide sequence upstream and downstream of the cut sites were compiled and used to identify the consensus cut site using WebLogo3 (Crooks et al., 2004).

The lengths of the mapped reads were extracted using custom python scripts and histogram plots were made using ggplot2 (Wickham, 2016).

DncV co-expression with CBASS effectors—A plasmid expressing dncV (pAW1371pBAD33-dncV) and a second plasmid expressing a predicted CBASS effector were electroporated into competent Escherichia coli BL21-DE3 (Invitrogen). Bacteria were recovered for 1 h shaking at 37°C in Super Optimal Broth with Catabolite repression (SOC), then plated onto selective LB Agar with glucose to repress dncV expression (LB: 10 g L⁻¹ tryptone, 5 10 g L⁻¹ yeast extract, 5 10 g L⁻¹ NaCl, 20 μ g ml⁻¹ chloramphenicol, 100 μ g ml ⁻¹ carbenicillin, 0.2% w/v glucose). Single colonies were inoculated into selective LB Agar plus glucose medium and cultured at 37°C shaking for ~16 h. 5 µl spots of 10-fold serial dilutions in LB were pipetted onto selective high-salt LB (20 µg ml⁻¹ chloramphenicol, 100 µg ml⁻¹ carbenicillin, NaCl concentration adjusted to 2% w/v) under inducing conditions (0.2% arabinose, 5 µM IPTG) or repressive conditions (0.2% glucose). Data are measured as CFU per ml and the ratio of inducing to repressive conditions and are the mean of three independent experiments. The predicted effector plasmids are pETSUMO2 expressing either: green fluorescent protein (gfp, negative control); Vibrio dncV native effector capV (WP 001133548.1, positive control); Desulfotomaculum alkaliphilium CD-NTase005 effector Saf2TM-SAVED (WP_031517735.1); Escherichia coli CD-NTase010 effector Saf2TM-SAVED (WP 001057904.1); Acinetobacter baumannii CD-NTase011 effector AbCap5 (WP 031984940.1); Geobacillus sp. CD-NTase012 effector GsCap5 (WP 013400843.1); Myxococcus xanthus CD-NTase022 effector caspase-SAVED (WP_020479061.1); Acinetobacter baumannii CD-NTase037 effector a.k.a. AbCdnD effector AbCap4 (WP 008942236.1); Enterobacter cloacae CD-NTase038 effector a.k.a. EcCdnD effector EcCap4 (WP_032676399.1); Bradyrhizobium japonicum CD-NTase039 effector B_iCap5 (WP 011082906.1); Burkholderia pseudomallei CD-NTase041 a.k.a. bpCdnG effector bpCap5 (WP_004556385.1); Citrobacter freundii CD-NTase042 effector highly similar to EcCdnG effector CfCap5 (ETX65525.1); Acinetobacter baumannii CD-NTase043 effector AbCap5 (WP 000539314.1); Pseudomonas aeruginosa CD-NTase044 effector PaCap5 (WP 023082129.1); Bacillus coagulans CD-NTase046 effector BcCap5 (WP 013858316.1); Vibrio cholerae CD-NTase053 effector VcCap5 (WP 000259919.1).

Phage resistance assays—Electrocompetent *E. coli* MG1655 was electroporated with individual medium-copy plasmids encoding entire CBASS operons under their native promoters and analyzed similarly to previously described methods (Cohen et al., 2019; Doron et al., 2018). *E. coli* were recovered and plated on selective LB (carbenicillin 100 μ g ml⁻¹). Single colonies were inoculated into selective MMCG medium (1× M9 Minimal Salts, 0.4% glucose, 0.02% MgSO₄, 0.001% CaCl₂, plus 100 μ g ml⁻¹ carbenicillin) and cultivated at 37°C shaking for ~20 h. Cultures were diluted 1:100 into selective MMCG medium and cultivated at 37°C shaking for 4 h to harvest mid-log cultures. Phage resistance was measured by a modified double-agar overlay technique. Bacteria were immobilized in soft-agar overlays by thoroughly mixing 400 μ l of mid-log culture with 3.5 mL molten MMCG top agar (MMCG medium plus 0.35% agar, 5 mM MgCl₂, 0.1 mM CaCl₂, 1 mM MnCl₂). Bacteria combined with top agar was immediately poured onto a 100 × 15 mm Petri dish containing 20 mL solidified MMCG Agar (1.6%) and allowed to cool for 10 min at

room temperature. A high-titer T2 phage lysate (Coli Genetic Stock Center CGSC12141), T5 lysate (CGSC12144), or T7 lysate (CGSC12146) prepared from MG1655 *E. coli* was 10-fold serially diluted into SM Buffer (100 mM NaCl, 8 mM MgSO₄, 50 mM Tris-HCl pH 7.5). 3 μ l spots of dilutions were pipetted onto the solidified double-agar overlays. Spots were allowed to dry for 20 min at room temperature. Plates were incubated at 37°C for ~16 h and plaques were quantified to compare efficiency of plating. Data are the mean of at least three independent experiments. Plasmids expressing CBASS operons were pLOCO2-*dncV* operon (Genome AE003852.1, 178097–185003), pLOCO2-Ec-*cdnD02* operon (Locus JCKK01000002, 2261880–2268858), and pLOCO2-Ec-*cdnG* operon a.k.a. CD-NTase042 operon (Locus JSNY01000125, 10563–17743).

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical details for each experiment can be found in the figure legends and outlined in the corresponding methods details section. Data are plotted with error bars representing the standard deviation (SD) or standard error of the mean (SEM) as indicated.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

ACKNOWLEDGMENTS

The authors are grateful to Kevin Corbett (University of California, San Diego) and members of the Kranzusch lab for helpful comments and discussions, Kelly Arnett and Harvard University's Center for Macromolecular Interactions, and the Molecular Electron Microscopy Suite at Harvard Medical School. The work was funded by the Richard and Susan Smith Family Foundation (to P.J.K. and S.S.); a Cancer Research Institute CLIP grant (to P.J.K.); the Parker Institute for Cancer Immunotherapy (to P.J.K.); the Pew Biomedical Scholars Program (to P.J.K. and A.S.Y.L.); the Searle Scholars Program (to A.S.Y.L.); a Sloan research fellowship (to A.S.Y.L.); the Vallee Foundation (to S.S.); NIH S100D020025, R01ES027595, and P42ES010337 (to M.J.); and NIH R01AI018045 and R01AI026289 (to J.J.M.). B.L. is supported as a Herchel Smith Graduate Research Fellow. A.T.W. was supported as a Ruth L. Kirschstein NRSA postdoctoral fellow (NIH F32GM133063). I.T.M. was supported by NIH F31CA236405.

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Highlights

- Cap4 proteins are a major family of nucleotide second messenger receptors in bacteria
- Cap4 receptors degrade DNA and mediate phage resistance by CBASS operons
- The Cap4 ligand-binding SAVED domain evolved from CRISPR CARF proteins
- As in cGAS-STING signaling, bacteria use 2'-5'-linked signals for antiviral immunity

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Figure 1. Cap4 Proteins Are Endonucleases Activated by CD-NTase Nucleotide Second Messengers

(A) Architectures of CBASS operons used for biochemical analysis. *Enterobacter cloacae*, *Acinetobacter baumannii*, and *Moraxella osloensis* systems lack a CapV phospholipase effector homolog and instead encode the protein CD-NTase-associated protein 4 (Cap4).
(B) Electrophoretic mobility shift assay measurement of Cap4–3'3'3'-cAAG complex formation. *E. cloacae* Cap2, Cap3, and Cap4 were incubated with ³²P-labeled 3'3'-cGAMP or 3'3'3'-cAAG, and bound complexes were resolved by nondenaturing polyacrylamide gel electrophoresis.

(C) Cartoon schematic and crystal structure of apo *Mo*Cap4. *Mo*Cap4 contains an N-terminal DUF4297 domain (orange) with homology to restriction endonucleases and a C-terminal SAVED domain (blue) with homology to CARF domains found in type III CRISPR immunity. Red circles indicate locations of active-site residues.

(D) Structural comparison of the Cap4 DUF4297 domain and the restriction endonuclease HindIII, showing structural homology and shared catalytic residues. The core endonuclease domains in Cap4 and HindIII are highlighted in orange and green, respectively.

(E and F) Agarose gel analysis of plasmid DNA degradation by Cap4. (E) Cap4 proteins degrade target DNA only in the presence of activating nucleotide second messenger synthesized by the neighboring CD-NTase CdnD within the CBASS operon. (F) Activity is dependent on the conserved Cap4 catalytic active site, with no cleavage observed with the *Ec*Cap4 mutant K74A.

(G) Agarose gel analysis of Cap4 DNA cleavage promiscuity. In the presence of activating 3'3'3'-cAAG, *Ec*Cap4 is capable of degrading all sources of dsDNA, including plasmid DNA, *E. coli* genomic DNA, and synthetic 100-bp DNA. Biochemical data are representative of at least 3 independent experiments. See also Figure S1.

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Figure 2. Cap4 Proteins Respond to Specific $3^\prime-5^\prime$ - and $2^\prime-5^\prime$ -Linked Nucleotide Second Messengers

(A) Analysis of Cap4 activation specificity using nucleotide second messengers synthesized with CD-NTase enzymes (DncV reaction, 3'3'-cGAMP; *Ec*Cap4 reaction, 3'3'3'-cAAG). *Ec*Cap4 and *Ab*Cap4 activation requires the specific nucleotide second messenger synthesized by the cognate CD-NTase, with no cross-activation occurring between distantly related operons.

(B) Activation of *Ec*Cap4 and *Ab*Cap4 DNA degradation activity with synthetic nucleotide second messengers. Synthetic 3'3'3'-cAAA is able to activate the *Ec*Cap4 enzyme normally responsive to 3'3'3'-cAAG, but no tested canonically 3'-5'-linked cyclic oligonucleotide species is able to activate *Ab*Cap4.

(C) P1 nuclease digestion and thin-layer chromatography analysis of CD-NTase products. Nuclease P1 specifically degrades 3'-5' bonds, leaving non-canonical 2'-5' bonds intact and resistant to phosphatase treatment. The human enzyme cGAS product 2'3'-cGAMP and the bacterial *Ab*CdnD product contain nuclease P1-resistant 2'-5' linkages.

(D and E) Large-scale synthesis and purification of the *Ab*CdnD nucleotide second messenger products. *Ab*CdnD synthesizes two closely related cyclic oligoadenylate products that co-migrate as (D) a single peak on anion-exchange (IEX) chromatography but can be separated into (E) major and minor product species with C18 chromatography.

(F) Cartoon schematic and crystal structure of *Ab*Cap4 bound to the *Ab*CdnD major product 2'3'3'-cAAA. The cyclic trinucleotide 2'3'3'-cAAA binds within the Cap4 SAVED domain (blue), indicating that this domain is responsible for CD-NTase signal recognition.

(G) Simulated-annealing F_o - F_c omit map (contoured at 3 σ) of the ligand density, demonstrating unambiguous assignment of the major *Ab*CdnD nucleotide second messenger as 2'3'3'-cAAA. Biochemical data are representative of at least 3 independent experiments. See also Figure S2.

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Figure 3. Cap4 Ligand Specificity Is Controlled by the SAVED Domain, a Divergent CARF Family Domain

(A) Structure of the *Ab*Cap4 SAVED domain, colored in blue and gray to highlight internal two-fold pseudosymmetry. Each half of the SAVED domain monomer has homology to a CARF subunit. Within the SAVED domain, these two CARF subunits are fused into a single chain with an ~25-amino acid linker.

(B) Topology diagram of *Ab*Cap4 and comparison with the *T. onnurineus* Csm6 CARF domain from type III CRISPR immunity. Each half of the *Ab*Cap4 SAVED domain shares the common core CARF domain topology, including a β strand that leads into a bracing outer helix (β 1-to- α 2 and β 7-to- α 6) and a second β strand and central helix (β 2-to- α 3 and β 8-to- α 7) at the normal CARF dimerization interface. All CARF family proteins contain these conserved structural features, including Csm6 (PDB: 606V), Csx3 (PDB: 3WZI), and Csx1 (PDB: 2I71).

(C) Structural comparison of the *Ab*Cap4 SAVED-2'3'3'-cAAA and *To*Csm6 CARF-cA₄ complexes. SAVED and CARF domains recognize nucleotide ligands with a similar binding surface. The single-domain architecture of the SAVED domain breaks the restriction of two-fold symmetry (dashed line) and enables recognition of diverse CD-NTase nucleotide second

messengers. In contrast, the homo-dimeric architecture of canonical CARF proteins necessitates ligands with two-fold symmetry (solid line).

(D and E) Structure of *Ab*Cap4 adenosine 1 (A1) pocket interactions and conservation of key residues shared in the SAVED and CARF domains. (E) Strict conservation of the A1 pocket with an aromatic residue and a tyrosine residue further supports emergence of SAVED domains through duplication and fusion of an ancient CARF family protein. See also Figures S3 and S4.

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Figure 4. Cap4 Proteins Are Activated through Ligand-Dependent Oligomerization (A) Negative-stain EM class averages of *Ec*Cap4 (K74A) with or without activating 3'3'3'cAAG and 45 bp DNA. Particle classification and quantification for each condition demonstrates that Cap4 oligomerization only occurs in the presence of activating nucleotide second messenger.

(B) Size-exclusion chromatography-multiangle light scattering (SEC-MALS) analysis of EcCap4 in the presence or absence of activating ligand 3'3'3'-cAAG. Absolute molecular masses of apo and ligand-bound complexes confirms EcCap4 oligomerization in the presence of activating nucleotide second messenger (EcCap4 expected molecular weight [MW], 56.1 kDa).

(C) Electrophoretic mobility shift assay assessing DNA binding of EcCap4 (K74A). Cap4 was incubated with 45-bp DNA and activating ligand 3'3'3'-cAAG as indicated. Stable EcCap4-DNA complex formation only occurs in the presence of activating nucleotide second messenger.

(D) Deep sequencing of *Ec*Cap4 DNA fragments and analysis of *Ec*Cap4 target specificity and distribution of DNA fragment sizes following *E. coli* genomic DNA or plasmid DNA degradation by *Ec*Cap4 (left). Cut site mapping demonstrates a minimal 5' CNG cut site preference (right).

(E) Mapping of *Ec*Cap4 cleaved DNA fragments across the *E. coli* genomic DNA confirms the relaxed targeting specificity of *Ec*Cap4 endonuclease activity. y axis positive and negative values indicate strand-specific cutting. Data are representative of at least 3 independent experiments.

See also Figures S5 and S6.

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Figure 5. SAVED Domain-Containing Proteins Are a Major Form of Viral Defense in Diverse Bacteria

(A) Phylogenetic tree showing distribution of SAVED-containing effectors in bacterial CD-NTase operons. Bacterial CD-NTase clades are displayed as an unrooted tree (Whiteley et al., 2019), with clades containing SAVED effectors (~30% of all bacterial CD-NTase operons) outlined in bold (clades B, C, D, G, and H). Dots indicate locations of CD-NTases of interest.

(B) Domain organization of diverse SAVED-containing effectors (Burroughs et al., 2015). The prevalence of each domain architecture in sequenced bacterial genomes is listed as a percentage of all SAVED-containing effectors (REase, restriction endonuclease).

(C) A genetic screen to identify SAVED effectors responsive to 3'3'-cGAMP. SAVED effectors were co-expressed with the 3'3'-cGAMP-producing CD-NTase DncV, and a spotdilution assay with quantification of recovered colony-forming units (CFUs) was used to assess SAVED activation and cytotoxicity. SAVED domain effectors from CD-NTase clades B and G respond to 3'3'-cGAMP, demonstrating that SAVED domains are capable of responding to cyclic dinucleotide and cyclic trinucleotide second messengers. Data represent the mean \pm SD of 3 independent experiments. See STAR Methods for CBASS effector species designations.

(D) *In vitro* reconstitution of nucleotide second messenger specificity for divergent Cap4 proteins and the HNH endonuclease-SAVED fusion *B. pseudomallei* CD-NTase-associated protein 5 (Cap5). DNA cleavage by *Bp*Cap5 demonstrates that SAVED domain-dependent nucleotide second messenger sensing is capable of activating structurally distinct enzymatic domains. Data are representative of at least 3 independent experiments.

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Figure 6. SAVED Effectors Are Essential for CBASS Antiviral Defense

(A) Example CBASS and CRISPR immunity operons encoding SAVED domain-containing effector proteins: Vibrio cholerae (WP_001901330), Enterobacter cloacae (WP_032676400), Citrobacter freundii (WP_032942206), Methylibium petroleiphilum (WP_011829962), Escherichia coli (WP_000058223), Myxococcus xanthus (YP_635404), and Fervidobacterium nodosum (WP_011994539).

(B) CBASS operons encoding SAVED domain effectors protect E. coli from phage replication. E. coli containing either an empty vector or a vector encoding E. cloacae CdnD or *E. coli* CdnG CBASS operons, as indicated, were infected with phage, and efficiency of plating was quantified by plaque-forming units. Data represent the mean \pm SEM of 3 independent experiments.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
E. coli BL21-DE3 RIL	Agilent	230245
E. coli MG1655	Coli Genetic Stock Center	CGSC6300
E. coli BL21-DE3	Invitrogen	C600003
T2 phage	Coli Genetic Stock Center	CGSC12141
T5 phage	Coli Genetic Stock Center	CGSC12144
T7 phage	Coli Genetic Stock Center	CGSC12146
Chemicals, Peptides, and Recombinant Proteins		
Ni-NTA Agarose	QIAGEN	30250
HiTrap Q HP Column	GE Healthcare	17115401
HiLoad 16/600 Superdex 75 PG	GE Healthcare	28989333
Zorbax Bonus-RP	Agilent	863668–901
Superdex 30 Increase 10/300 GL	GE Healthcare	29219757
SRT SEC-300	Sepax	215300-7830
SeQuant ZIC-pHILIC	EMD-Millipore	1504620001
[α- ³² P] ATP	Perkin Elmer	BLU003H250UC
[a- ³² P] GTP	Perkin Elmer	BLU006H250UC
PEI-Cellulose F TLC plate	EMD Biosciences	EM1.05579.0001
Nuclease P1 from Penicillium citrinum	Sigma-Aldrich	N8630
Snake Venom Phosphodiesterase	Sigma-Aldrich	P3243
Alkaline Phosphatase, Calf Intestinal (CIP)	New England Biolabs	M0290S
DNA Extraction Kit	QIAGEN	69504
ATP, GTP	New England Biolabs	N0450S
SYPRO Orange	ThermoFisher	S6650
HEPES	VWR	97061-824
Tris base	VWR	97062-416
PEG-400	Sigma-Aldrich	202398
PEG-3350	Sigma-Aldrich	202444
Lithium sulfate	Sigma-Aldrich	62609
Imidazole	VWR	97065-016
Tris[-2carboxyethyl] phosphine hydrochloride (TCEP)	GoldBio	TCEP50
Ethylene glycol	VWR	97061–964
3'3'3'-cAAA	Biolog Life Science Institute GmbH & Co. KG	C 362
3'3'3'-cAAG	Biolog Life Science Institute GmbH & Co. KG	C 361
Deposited Data		
MoCap4	This paper	PDB: 6VM5
AbCap4–2'3'3'-cAAA	This paper	PDB: 6VM6

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
AbCap4-3'3'3'-cAAA	This paper	PDB: 6WAN
AbCap4	This paper	PDB: 6WAM
Oligonucleotides		
5′ -TACAGATCTACTAGTGATCTATGACTG ATCTGTACATGAT CTACA-3′	Integrated DNA Technologies	DNA45 sense
5′ -TGTAGAT CATGTACAGAT CAGTCATAG ATCACTAGTAGATCTGTA-3′	Integrated DNA Technologies	DNA45 antisense
Software and Algorithms		
Phenix 1.13–2998	Adams et al., 2010	https://www.phenix-online.org/
Coot 0.8.9	Emsley and Cowtan, 2004	https://www2.mrc-lmb.cam.ac.uk/ personal/pemsley/coot/
Pymol vl.7.4.4	Schrödinger	https://pymol.org/2/
Prism 7.0d	GraphPad software	https://www.graphpad.com/ scientific-software/prism/