

RESEARCH ARTICLE

The structure of a Type III-A CRISPR-Cas effector complex reveals conserved and idiosyncratic contacts to target RNA and crRNA among Type III-A systems

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Abstract

Type III CRISPR-Cas systems employ multiprotein effector complexes bound to small CRISPR RNAs (crRNAs) to detect foreign RNA transcripts and elicit a complex immune response that leads to the destruction of invading RNA and DNA. Type III systems are among the most widespread in nature, and emerging interest in harnessing these systems for biotechnology applications highlights the need for detailed structural analyses of representatives from diverse organisms. We performed cryo-EM reconstructions of the Type III-A Cas10-Csm effector complex from *S. epidermidis* bound to an intact, cognate target RNA and identified two oligomeric states, a 276 kDa complex and a 318 kDa complex. 3.1 Å density for the well-ordered 276 kDa complex allowed construction of atomic models for the Csm2, Csm3, Csm4 and Csm5 subunits within the complex along with the crRNA and target RNA. We also collected small-angle X-ray scattering data which was consistent with the 276 kDa Cas10-Csm architecture we identified. Detailed comparisons between the *S. epidermidis* Cas10-Csm structure and the well-resolved bacterial (*S. thermophilus*) and archaeal (*T. onnurineus*) Cas10-Csm structures reveal differences in how the complexes interact with target RNA and crRNA which are likely to have functional ramifications. These structural comparisons shed light on the unique features of Type III-A systems from diverse organisms and will assist in improving biotechnologies derived from Type III-A effector complexes.

Introduction

CRISPR-Cas systems provide adaptive immunity to prokaryotes by capturing fragments of genetic information from mobile genetic elements, such as plasmids and bacteriophage, and

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storing the information content in the spacers of a CRISPR array. The spacers are transcribed and processed into crRNAs, which when bound to effector Cas proteins, facilitate an interference response upon the detection of a complementary foreign nucleic acid [1,2]. CRISPR-Cas systems have been organized into two classes, class 1 multi-Cas protein effectors versus class 2 single Cas protein effectors, and six types by bioinformatics analyses [3]. The *S. epidermidis* effector complex, known as Cas10-Csm, consists of a crRNA and five Cas proteins: Cas10, Csm2, Csm3, Csm4 and Csm5 (Fig 1A). As a multi-protein effector complex, it is a member of class 1 and further categorized as Type III-A due to the presence of Cas10 and the small signature subunit Csm2.

Type III CRISPR-Cas systems are among the most abundant in nature and considered the most complex [3,4]. Type III systems comprise approximately 25% of all CRISPR systems, a prevalence which suggests their physiological importance in prokaryotes [5]. There are six subtypes currently described (A-F), and among these, the III-A and III-B system are the best characterized [3]. These systems possess a distinct interference activity: upon sensing foreign RNA transcripts, the synthesis of second-messenger molecules, cyclic oligoadenylates (cOAs), is activated by the Palm-2 domain of Cas10 [6,7]. The second-messenger binds to Csm6, a nuclease that is not a part of the Cas10-Csm complex, and activates its latent, indiscriminate RNase activity, which in turn drives the cell to dormancy to block viral replication (Fig 1B) [6,7]. Type III CRISPR systems also possess the ability to degrade foreign RNA complementary to the crRNA (target RNA) via the Csm3/Cmr4 protein, and many complexes also have the ability to degrade single-stranded (ss) DNA via the HD nuclease domain of Cas10, comprising a complex multi-faceted interference response [8–10].

Recently, multiple investigators noted the intrinsic ability of Type III systems to specifically detect RNA and amplify this detection event by Cas10-mediated cOA synthesis makes them well suited to serve as a diagnostic tool for RNA viruses [11–14]. A Type III CRISPR complex was incubated with SARS-CoV-2 RNA initiating cOA synthesis. The cOA molecules stimulated the RNase activity of Csm6 or a Csm6 homolog which then cleaved a fluorophore-quencher reporter RNA unleashing a fluorescent signal indicating the presence of the viral RNA [11–13]. Alternatively, NucC, a cOA stimulated DNase, was used in the place of Csm6 in a reaction that uses a fluorophore-quencher double-stranded DNA to report on the presence of viral RNA [14]. The Type III virus detection schemes achieved specificity similar to RT-qPCR assays but required coupling of an isothermal amplification step to achieve similar sensitivity, attomolar level [11–13]. The synthesis of cOA produces H^+ and PP_i products during the course of the reaction and Santiago-Frangos and co-workers showed these molecules can also be sensed to report on the presence of viral RNA [11]. In all cases the Type III CRISPR based assays proceeded as rapid and isothermal reactions indicating a path to achieve sensitive and specific molecular diagnostics without expensive equipment—critical features for deployment in point-of-care settings. Realizing the full potential of this biotechnology will require a detailed understanding of the structure and mechanism of Type III CRISPR systems.

Structures of Type III CRISPR systems from several organisms of varying resolution are currently available. Low resolution Type III structures appeared in 2014, providing a good description of the location of the Cas proteins in the complex, and a high resolution structure of a Type III-B appeared shortly after [15–18]. Type III-A and Type III-B CRISPR systems differ structurally in two important ways: the presence of the small subunit protein Csm2 in Type III-A versus the presence of the small subunit protein Cmr5 in Type III-B and in the presence of the Cmr1 protein which produces a slightly different architecture adjacent to the 3' end of the crRNA [5,13,16,19]. Since the small subunit proteins, Csm2 and Cmr5, directly interact with target RNA upon its binding to the complex, it is likely these proteins play a critical role in sensing and activating interference [20,21]. Therefore the first high resolution structures of

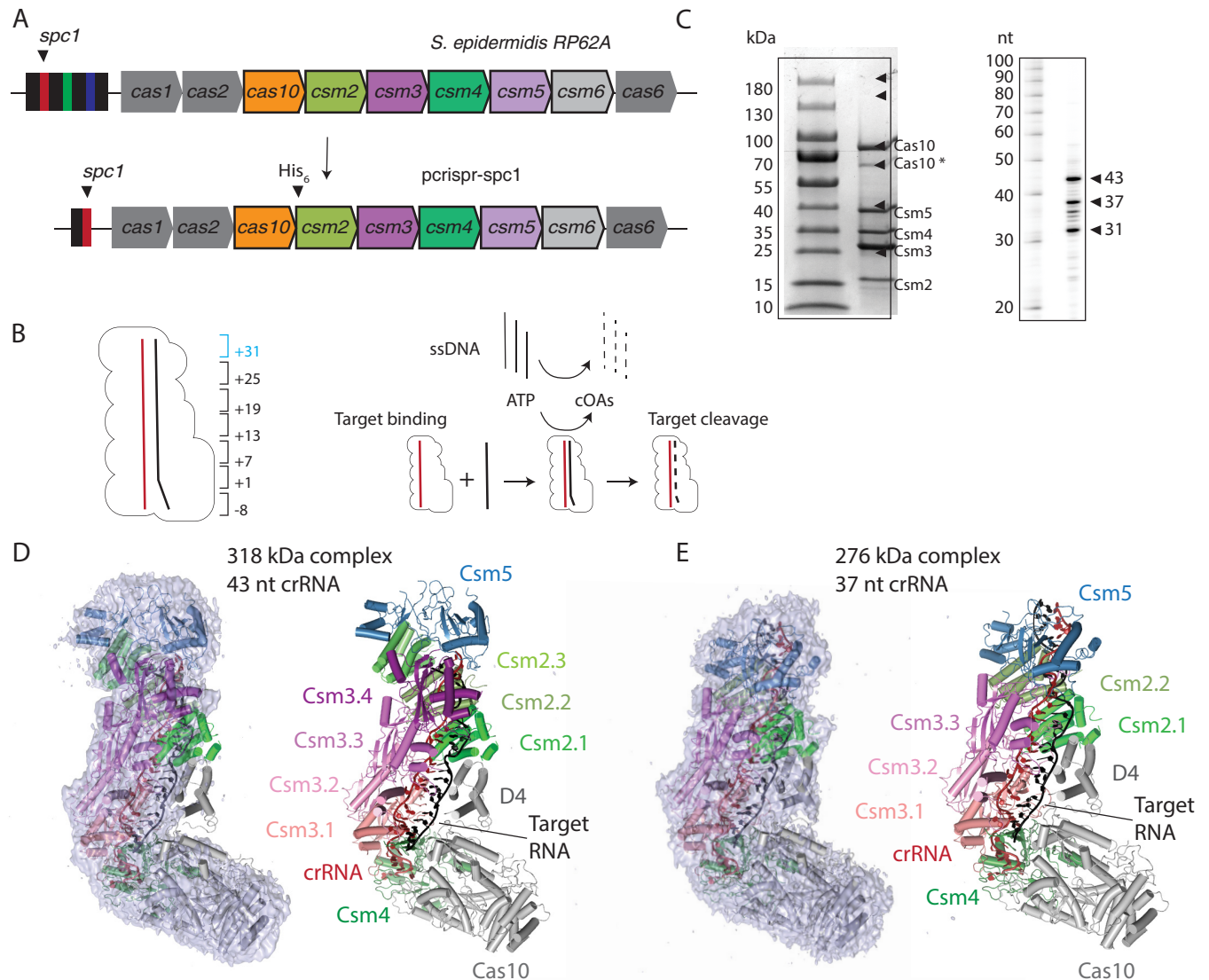


Fig 1. Overall architecture of two oligomeric states of *S. epidermidis* Cas10-Csm revealed by cryo-EM. (A) A schematic of the *S. epidermidis* Type III-A crisper locus. Cas10-Csm purification was facilitated by the construction of the plasmid *pcrispr-spc1*. (B) The numbering scheme describing positions within the crRNA-target duplex is shown. The six-nucleotide region beginning at +31 (blue) is present in the 43 nt crRNA complex but not the 37 nt crRNA complex. A schematic of steps in the interference reaction is shown. Target RNA binding activates single-stranded DNA (ssDNA) cleavage and cyclic oligoadenylates (cOAs) synthesis activities of Cas10-Csm. (C) SDS-PAGE and urea-PAGE analysis of purified Cas10-Csm. Cas10* denotes a band that has been identified by mass spectrometry as a truncated Cas10. CrRNA extracted from purified SeCas10-Csm possesses lengths of 31, 37 and 43 nt. (D) The larger complex consists of a 43 nt crRNA, three copies of Csm2 and four copies of Csm3. The cryo-EM density map is shown at left revealing that very little density is present for Cas10 domain 4 (D4) and Csm2.1. The map is contoured at level 3.0σ . (E) The smaller complex consists of a 37 nt crRNA, two copies of Csm2 and three copies of Csm3. Density contoured at 3.0σ is shown. In contrast to the larger complex, density is observed for Cas10 domain 4 and Csm2.1. A molecular model of the 276 kDa complex has been deposited with the PDB, code 8d06.

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Type III-A CRISPR systems in early 2019, were a welcome development [20,22,23]. These structures of Cas10-Csm from the archaeobacterium *T. onnurineus* and the eubacterium *S. thermophilus* provided critical insights into the contacts of Cas10-Csm with crRNA and target RNA, but also differed in a key aspect: the *T. onnurineus* structure indicated little conformational change upon target binding, while the *S. thermophilus* structure underwent substantial conformational change upon target binding [20,22]. In 2022, new high resolution structures of Cas10-Csm appeared. An *L. lactis* structure was reported further enabling structural biologists

ability to compare and contrast across the available structures to formulate hypotheses regarding the mechanisms of target RNA sensing and activation [21]. While this manuscript was in preparation, high-resolution structures of *S. epidermidis* Cas10-Csm also were reported; however, in these structures, the cognate target RNA is not intact [24]. Thus, the conformation of the complex in the active pre-cleavage state and corresponding contacts with the target RNA remain unclear.

To enrich the mechanistic understanding of how Type III systems function, we investigated the structure of the *S. epidermidis* Cas10-Csm complex bound to cognate target RNA by cryo-electron microscopy. A 3.1 Å map facilitated building atomic models for four of the five protein components, crRNA and target RNA. *S. epidermidis* Cas10-Csm is a leading model system in the study of Type III CRISPR systems dating to the earliest days of the field and therefore our structural model can assist in the design of detailed structure-function experiments utilizing the wealth of tools developed to study *S. epidermidis* Cas10-Csm [25,26]. Additionally, we conducted a detailed comparison into the similarities and differences in how the *S. epidermidis* proteins interact with crRNA and target RNA compared to two well-resolved cognate, target RNA bound structures from *S. thermophilus*, an exemplar of bacterial Cas10-Csm systems, and *T. onnurineus*, an exemplar of archaeal systems. Our observations shed light on the unique features of Type III-A systems from diverse organisms and are likely to guide efforts in optimizing the biotechnologies derived from Type III-A effector complexes.

Materials and methods

Modification of pcrispr to contain a single spacer

The pcrispr-*spc1* plasmid which contains a single repeat and *spc1* was constructed using a two-piece Gibson assembly with primers noted in [S1 Table](#). Briefly, the repeat-*spc1* region was amplified using primers F063/A010 from a plasmid containing the wild-type repeat-spacer array, and primers F062/L162 were used to amplify the backbone from a pcrispr-cas plasmid lacking all repeats and spacers [27]. PCR products were then purified with EZNA Cycle Pure Kit (Omega Bio-tek) and Gibson assembled. The assembled construct was introduced into *S. aureus* RN4220 via electroporation, and transformants were subjected to PCR and DNA sequencing using primers A200/F111 to confirm the presence of a single repeat and spacer. The confirmed construct was then purified with EZNA Plasmid DNA Mini Kit (Omega Bio-tek) and introduced into *S. epidermidis* LM1680 strains via electroporation.

Purification of the Cas10-Csm complex with bound crRNA

S. epidermidis LM1680 was transformed with pcrispr-*spc1* for expression of SeCas10-Csm complex. Cell growth, lysis and immobilized metal affinity chromatography (IMAC) were performed as described previously [28]. IMAC fractions were analyzed by SDS-PAGE and peak fractions were pooled, concentrated and loaded to a 5–20% w/v sucrose gradient for further purification by ultracentrifugation, a procedure previously reported [29]. Ultracentrifugation was performed for 41 hours in a SW32 rotor at 31,000 RPM and fractions were collected and analyzed by A_{280} and SDS-PAGE. Fractions with pure Cas10-Csm were pooled, concentrated to an A_{280} of 5.44, aliquoted and flash frozen in liquid nitrogen. Aliquots were stored at -80°C until needed.

Structure determination of SeCas10-Csm by cryo-EM

Target RNA (ssRNA-01, [S1 Table](#)) aliquots were thawed, heated at 70° for two minutes, then snap cooled on ice. SeCas10-Csm was thawed on ice, mixed with target RNA at a 1:1.6 molar

ratio in the presence of 2 mM EDTA and incubated at 37°C for 5 minutes to form the target RNA bound SeCas10-Csm complex. The complex was then diluted 10-fold with 50 mM Tris-HCl pH 7.5, 150 mM NaCl buffer and applied to UltrAuFoil grids with 1.2 μm diameter holes and 1.3 μm hole separation for flash freezing into liquid ethane using a Vitrobot Mark IV and a blotting time of 3–4 seconds. Movie frames were collected at the New York Structural Biology Center on a Titan Krios with a 300 keV FEG equipped with a Cs corrector, K3 camera, and a BioQuantum energy filter using Legikon at a pixel size of 0.846 \AA with a total dose of 44.70 $\text{e}/\text{\AA}^2$ fractionated over 40 frames. About 5000 movie frames were collected over a period of 24 hours.

All image processing was performed in cryoSPARC 2 [30]. After patch motion correction and CTF estimation, the first set of particles were picked with the blob picker. A 2D classification was done on this set of particles and the resulting good 2D classes were used as templates for template matching. In order to remove false positives and bad particles, 2D classification followed by multiple rounds of ab initio and 3D heterogenous refinement were carried out. The strategy was to curate a set of good particles to be used for neural network training in Topaz [31]. After pooling together particles from different 3D classes, manual curation was done to create a set of 5,000 particles. This set of particles were used for Topaz training. Topaz cross validation was executed to optimize for the expected number of particles. The initial value for this parameter was set to 500 with 4 increments of the size 100. Training radius was set to 2. ResNet8 model architecture was trained for 15 epochs. After Topaz training, Topaz extract was carried out. With a particle threshold of -1, about 1.4 million particles were extracted and then binned by a factor of two. The rest of the processing pipeline is illustrated in S1 Fig. The dataset was heterogenous both in terms of stoichiometry and conformation. The smallest to the largest stoichiometry are labelled (complex I to III in S1 Fig). Complex II was used for the molecular models presented herein. Conformational heterogeneity was seen in Cas10 in 2D classifications. When refined to high resolution, complex II lost almost all the density for Cas10, but complex III retained the secondary structures. Variability analysis was used to tease out complex II and complex III [32]. This was confirmed by a heterogenous refinement. Final refinements were done using nonuniform refinement [33]. The local resolution map was generated using CryoSPARC's local resolution tool.

An atomic model consisting of the Csm2-5 proteins, crRNA and target RNA was constructed using the Complex II density map. Iterative manual modeling and real-space refinement in Coot was performed followed by cycles of real-space refinement in Phenix [34,35]. Docking of the coordinates for Csm3 and Csm2, described by PDB codes 6nbt and 6nbu, respectively, into the Complex II volume described in emd-27593 was performed in Coot. AlphaFold2 models of Csm4 and Csm5 (sequence references in S2 Table) were remodeled within Coot to fit the volume [36,37]. RNA molecules were built in Coot from the sequences described in S2 Table. Molprobit was used to monitor coordinate geometry and minimize clashes [38]. An AlphaFold2 model of *S. epidermidis* Cas10 was docked into the corresponding density and real-space refinement was conducted with two rigid-groups: the HD domain, Palm1 and Palm 2 domains as a group and domain 4 as a second group. Density in the Cas10 region was of sufficient quality to identify domain locations but was not sufficient to generate an atomic model. The coordinates generated from the Complex II density could be docked into the Complex III density to confirm the stoichiometry and overall architecture of Complex III as a 318 kDa complex. However, Complex III had poor density for the Csm2 proteins and therefore an all-atom atomic model was not generated from the Complex III density. Local resolution plots are given in S4 Fig and S1 and S2 Movies.

Collection of SEC-SAXS data

SEC-SAXS was performed at the SIBYLS beamline at the Advanced Light Source at Lawrence Berkeley National Laboratory. Three concentrations of SeCas10-Csm complex bound to target RNA (ssRNA-01) were formed by the brief incubation of SeCas10-Csm with a 1:1 mol:mol ratio of target RNA followed by application of the complex to a Shodex 804 SEC column flowing at 0.5 mL/min with running buffer of 50 mM Tris-HCl pH 7.5, 150 mM NH₄Cl, 2% v/v glycerol. The concentrations of complex were 20 μM, 10 μM and 5 μM which corresponds to ~6 mg/mL, ~3 mg/mL and ~1.5 mg/mL. Frames collected at the SEC peak were averaged and buffer-subtracted using ScÅtter (<https://bl1231.als.lbl.gov/scatter/>). Subtracted curves were further analyzed using PRIMUS [39], and pairwise distance distribution functions were calculated using GNOM in PRIMUS [40]. Ab initio models were calculated using DAMMIN [41], and molecular models were calculated using SASREF [42]. To generate molecular models, coordinate files for individual subunits bound to RNA were generated by partitioning the *S. thermophilus* Cas10-Csm structure (PDB ID 6ifu) [20] into individual subunits bound to RNA segments and, in the case of the Csm1 subunit, partitioning the chain into two components at a domain boundary at position Lys-654 (S3 Table). During rigid body model calculation, constraints were applied to limit distances between these components at covalent bonds linking nucleotides or the peptide backbone. CRY SOL [43] and OLIGOMER were used to generate theoretical scattering profiles from molecular models (those reported here and previously) and compare them to the experimental scattering profile. Because SAXS solutions are inherently degenerate and cannot distinguish enantiomorphs, mirror images of ab initio and molecular models are displayed wherever alignments with molecular models indicate that this is the biochemically relevant model. Enantiomorph transformation was performed using ALPRAXIN [44].

Sequence logos and structural superpositions

All sequence logos were constructed using WebLogo [45]. For Csm2, input was a multiple sequence alignment given by PFAM of all Uniprot entries in the protein family identified by PFAM, PF03750 [46]. The sequence logos for Csm3 or Csm5 were constructed from a multiple sequence alignment of 250 homologs identified by Protein BLAST and aligned with Clustal Omega. The structural superpositions shown in Fig 3 were made in Pymol using the align command with Csm4 as the basis for the alignment. Structural superpositions in Figs 4 through 10 were made in Pymol using the fit command with P atoms of the target or crRNA backbone selected.

Results

The overall architecture of *S. epidermidis* Cas10-Csm by cryo-EM and SAXS

The *S. epidermidis* Type III-A CRISPR array contains three spacer segments (*spc1-sp3*). To facilitate purification of an in vivo assembled Cas10-Csm complex with a single crRNA, the segments encoding *spc2* and *spc3* were removed from *pcrispr* (Fig 1A). The *pcrispr* plasmid is designed for expression of *S. epidermidis* Cas10-Csm (SeCas10-Csm) in *S. epidermidis* cells and we have previously demonstrated this expression and purification approach leads to a ribonucleoprotein complex with robust activity in cyclic oligoadenylate synthesis and target RNA cleavage (Figs 1B and S2) [29]. Cas10-Csm bound to a crRNA derived from *spc1* was purified and SDS-PAGE analysis demonstrated the presence of the five expected proteins. Urea-PAGE analysis of RNA extracted from the complex produced bands of the typically observed sizes of 31, 37 and 43 nucleotide-length crRNAs (Fig 1C).

To obtain a cryo-EM structure of SeCas10-Csm bound to target RNA, purified complex was incubated with a slight excess of synthetic target RNA containing sequence for the nickase mRNA, which is complementary to *spc1* derived crRNA and is its natural target. SeCas10-Csm bound to the target was plunge frozen. Csm3 mediated target RNA cleavage was inhibited during the incubation by the presence of EDTA, which chelates divalent metal ions required for the cleavage reaction. From 5,000 micrographs, 1.4 million particles were extracted and through the classification and refinement steps detailed in [S1 Fig](#), two distinct volumes at 3.1 Å resolution were reconstructed. Docking of the *S. epidermidis* Csm2 and Csm3 crystal structures, given by PDB codes 6nbu and 6nbt respectively, and manual building of crRNA, target RNA, Csm4 and Csm5 into the two volumes revealed that each corresponded to a different oligomeric state of SeCas10-Csm, a 318 kDa complex and a 276 kDa complex ([Fig 1D and 1E](#)). The larger complex is composed of a 43 nt crRNA and an extra copy of Csm2 and Csm3 compared to the 276 kDa complex which possesses a 37 nt crRNA ([Fig 1D and 1E](#)). An additional difference between the complexes is present: Csm2 is well ordered in the 276 kDa complex but, while present, poorly ordered in the 318 kDa complex ([Fig 1D and 1E](#)). Cas10 is present in both complexes, but is highly flexible with a hinge motion observed in 2D classes. The flexibility of Cas10 led to density that is not of sufficient quality to support the construction of an atomic model for the protein. Since the 276 kDa complex possesses high quality density for crRNA, target RNA and proteins Csm2-5, an atomic model for these components was constructed ([S3 and S4 Figs and S4 Table](#)). The complex was completed by docking of an AlphaFold2 model of the SeCas10 component for which there is agreement on the underlying secondary structures and domain positions ([S5 Fig](#)). A C_{α} model of the 318 kDa complex was constructed by docking the molecular models of the Csm2-5 coordinates and the Cas10 AlphaFold2 model into density. The molecular model of the 276 kDa complex facilitates identifying the detailed interactions of the Csm2, Csm3, Csm4 and Csm5 proteins with crRNA and target.

We performed a complementary analysis of the SeCas10-Csm architecture utilizing size-exclusion chromatography coupled with small-angle x-ray scattering (SEC-SAXS). Purified SeCas10-Csm bearing a single *spc1*-derived crRNA was purified and incubated with target RNA in a manner nearly identical to our cryo-EM experiments, then passed over a SEC column and subjected to in-line SAXS data collection. Data frames from the center of the SEC peak were used to generate a scaled, averaged scattering curve. This scattering curve was then used to generate sets of independently calculated *ab initio* envelopes and rigid body models of the SeCas10-Csm complex based on the previously published *S. thermophilus* 6ifu structure [[20](#)]. Replicate rigid body models showed good agreement with each other and with the set of *ab initio* envelopes ([Figs 2A, 2B, S6 and S7; S5 Table](#)). We further compared our SAXS data and models to both the 276 kDa and 318 kDa EM models reported here. We found that our SAXS-derived *ab initio* and rigid body models both showed better agreement with the 276 kDa complex than the 318 kDa complex ([Fig 2C, S6 Table](#)). To further validate the agreement between the solution scattering data and the 276 kDa complex, we calculated theoretical SAXS profiles (via CRY SOL) from the 276 kDa and 318 kDa EM structures and compared them to our experimental SAXS data. The profile calculated from the 276 kDa model showed significantly better agreement with our experimental SAXS data than did the 318 kDa model profile ([Fig 2D, S7 Table](#)). Finally, attempts to deconvolute the experimental scattering curve (via OLIGOMER) with multiple theoretical scattering profiles generated from plausible molecular models consistently revealed the presence of only a single component with the same stoichiometry as the 276 kDa complex ([S8 Fig](#)). The low resolution afforded by SAXS reconstructions (as well as the inherent degeneracy of three-dimensional structural solutions derived from two-dimensional scattering data) do not allow us to unambiguously confirm the stoichiometry

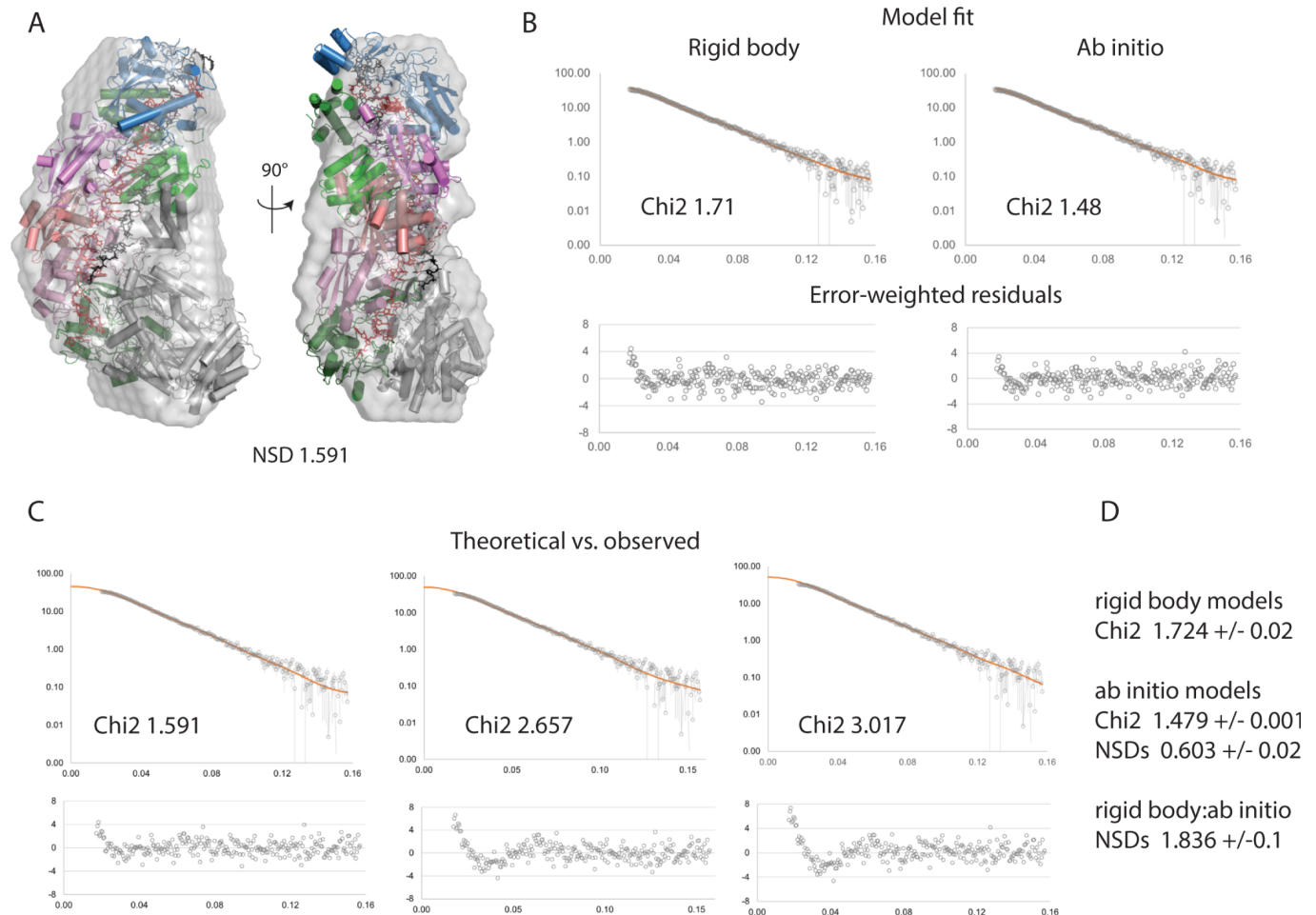


Fig 2. Support for Cas10-Csm complex stoichiometry from SEC-SAXS. (A) Multiple *ab initio* and rigid body models were calculated from the averaged peak frames of the SEC-SAXS run. Representative *ab initio* (gray envelope) and rigid body (cartoon) models are shown superposed (via SASREF). (B) The fits of the models from panel A (orange) to the experimental scattering curve (gray) are shown along with error-weighted residuals (lower panels). The rigid body model fit is shown at left, the *ab initio* model, at right. (C) EM structures reported here, overlaid with a representative *ab initio* model from our SAXS analysis. (D) CRYSOLE was used to generate theoretical scattering curves from the EM models reported in this study, and these theoretical curves (orange) were scaled to the experimental SAXS data (gray). From left to right, the models shown are the 276 kDa complex, the 318 kDa complex, and the 318 kDa complex minus Csm2 subunits. As in B, error-weighted residuals are shown below each plot. Throughout this figure normalized spatial discrepancy is abbreviated NSD.

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of the complex using SAXS data alone, but the agreement between the 276 kDa complex and our SAXS-derived models suggests that the 276 kDa model stoichiometry exists and may predominate in solution.

Superposition of the coordinates of the 276 kDa complex and the 318 kDa derived from cryo-EM reveal nearly identical positioning of Csm4 and the Csm3 oligomer. The Csm2 oligomer shifts slightly toward Cas10 in the 276 kDa complex a movement that could indicate sampling of a different conformational state, the influence of the differing stoichiometry or a combination of both (S9A Fig). A superposition of the SAXS-derived rigid-body model that best fit the *ab initio* envelope to the 276 kDa complex also reveals that the 276 kDa complex has a slight shift of the Csm2 oligomer towards Cas10 (S9B Fig). The conformational differences observed are not surprising given the documented dynamic nature of the Cas10-Csm complex [20,21,47]. However, a further detailed analysis of the differences cannot be conducted at this time due to the entanglement of stoichiometric differences and biases arising from the differing techniques used, SAXS versus cryo-EM.

Comparison of the 276 kDa complex to previously published Cas10-Csm complexes of the same stoichiometry determined by cryo-EM indicates that we have captured a state of Cas10-Csm poised for catalysis (Fig 3A). The 276 kDa complex (PDB code 8do6) superimposes well with the *S. thermophilus* structure reported in PDB code 6ig0 which is bound to intact, cognate target RNA with two ATP and two Mg²⁺ ions in the Palm2 domain, the active

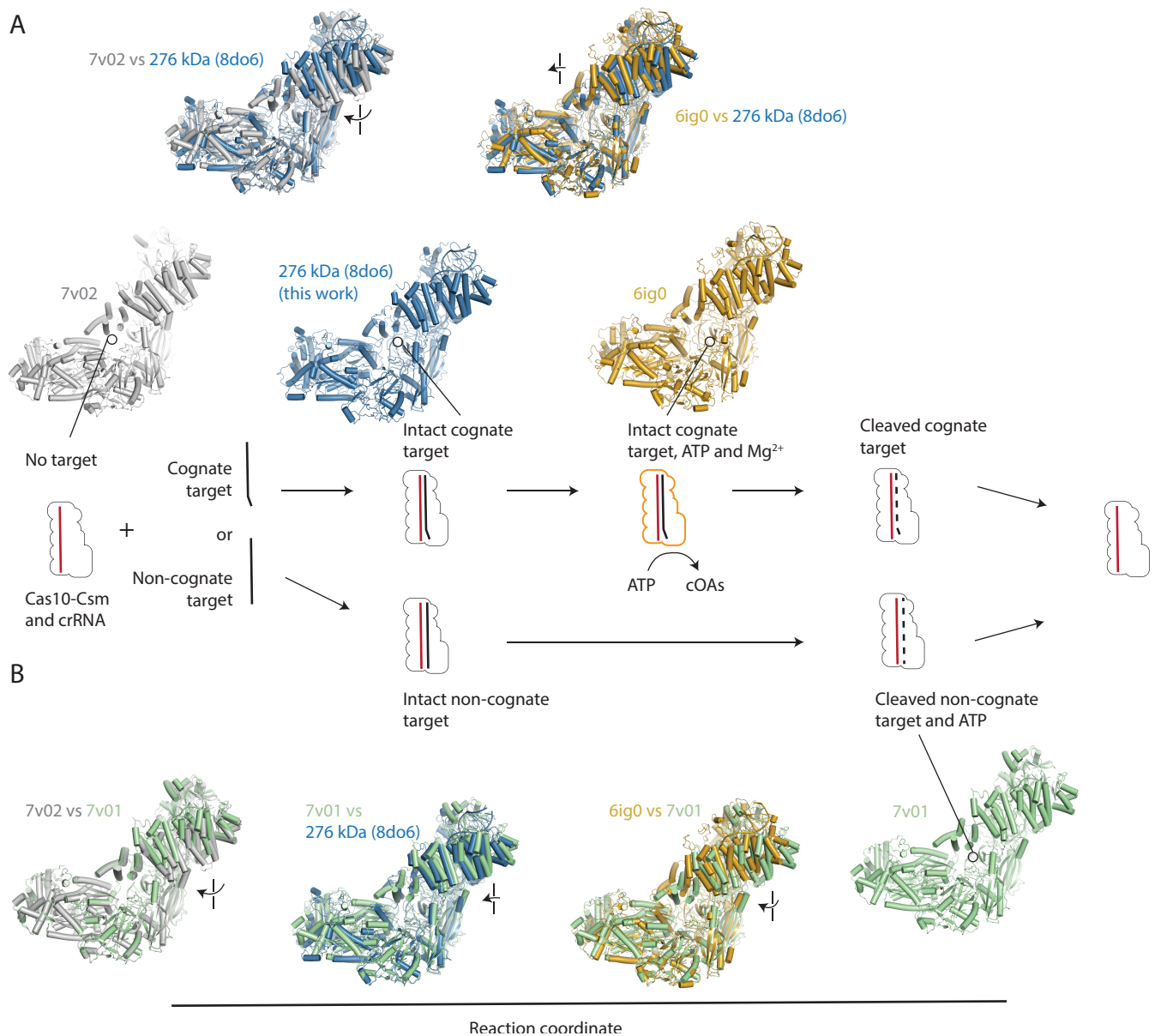


Fig 3. Placing the structure of the Cas10-Csm bound to intact, cognate target RNA on the reaction coordinate. (A) A diagram of the states of Cas10-Csm shows that it can exist unbound to target RNA, bound to cognate target RNA leading to the activation of cOA synthesis and bound to cleaved, cognate target RNA after the Csm3 mediated RNase reaction. Alternatively, Cas10-Csm can bind to non-cognate target RNA, which possesses base-pairing with crRNA in the 5' tag-3' flank region which leads to Csm3-mediated RNase activity but not cOA synthesis. Superpositions of PDB code 8do6, reported in this work, with an unbound *S. epidermidis* Cas10-Csm (7v02) via the Csm4 protein reveals target binding causes a rotation of Csm2 clockwise and toward Cas10. Superposition of 8do6 with *S. thermophilus* Cas10-Csm bound to cognate target RNA, two ATP and two Mg²⁺ reveals its Csm2 proteins are located slightly closer to Cas10 than 8do6. (B) A comparison of *S. epidermidis* Cas10-Csm unbound to target (7v02) and bound to a non-cognate target (7v01) displays clockwise and downward rotation of Csm2. Comparison of 7v01 with 8do6 or 6ig0 shows all three structures are similar, however, the Csm2 proteins of 8do6 are positioned slightly closer to 6ig0.

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site for cOA synthesis [20]. Comparisons to previously published *S. epidermidis* Cas10-Csm structures bound to non-cognate, target RNA reveals the two structures are similar but in the 276 kDa complex reported here, the Csm2 oligomer is rotated slightly more toward Cas10 (Fig 3B) [24]. We believe the structure reported in this work better models *S. epidermidis* Cas10-Csm poised for catalysis because it is bound to a cognate and intact target RNA rather than a partially cleaved non-cognate target RNA as in 7v01.

The detailed interactions of the Csm2-5 proteins with crRNA and target RNA

We used the atomic models of the Csm2, Csm3, Csm4 and Csm5 proteins to answer two questions. First, in *S. epidermidis* Cas10-Csm what are the specific interactions each of these proteins make with target and crRNA? Second, to what degree are the interactions between the Csm proteins and target and crRNA conserved versus idiosyncratic among Cas10-Csm complexes? To address this question we used structures available for another bacterial Cas10-Csm (*S. thermophilus*, PDB code 6ifu) and archaeal Cas10-Csm (*T. onnurineus*, PDB code 6mus). The second question is important because it directly bears on the larger question: to what degree is there a conserved mechanism for sensing the binding of target to crRNA and activating interference in Type III-A CRISPR-Cas? Recent biochemical data have identified site-directed mutants of Csm2, Csm3 and Csm4 with specific defects in interference and this literature will be described below as necessary [20–22].

Csm2 is present in two copies in the 276 kDa complex, contacting target RNA in the vicinity of positions +13 to +24 (Figs 1B and 4A). Helix- α 2 is positioned in the major groove of the target-crRNA duplex and conserved residues within the N-terminal region of α 2 participate in hydrogen bonding and electrostatic interactions with the sugar-phosphate backbone of target RNA (Fig 4B and 4C). Two conserved residues in α 3, Y87 and R91, interact with phosphates at the +23 and +22 positions of target RNA, respectively (Csm2.2 protomer, Fig 4C). Superposition of the bacterial Cas10-Csm complex from *S. thermophilus* with the *S. epidermidis* complex shows nearly identical interactions between Csm2 and target (Fig 4C). A superposition of *T. onnurineus* Cas10-Csm with the *S. epidermidis* complex, however, reveals substantial differences in how Csm2 contacts target RNA (Fig 4D). ToCsm2 residues Y129 and K133 in α 3 are equivalent to SeCsm2 residues Y87 and R91, yet the ToCsm2 residues are located greater than 8 Å away from target RNA (Fig 4D). This is due to a substantial difference in how Csm2 is positioned within the ToCas10-Csm complex that also affects ToCsm2 α 2: the N-terminal residues of α 2 contact target RNA but contact the +18 position rather than the +22 and +23 positions and make fewer overall contacts to target RNA (Fig 4D). The Arg residue equivalent to Csm2 R49 (ToCsm2 R96) has been analyzed by site-directed mutagenesis in *S. thermophilus* Cas10-Csm and *L. lactis* Cas10-Csm and shown to be critical for target RNA cleavage activity confirming that the Csm2 contacts to target RNA have functional importance [20,21].

Csm3, like other Cas7 family proteins, forms a filament that cradles crRNA presenting it to solvent (Fig 1E). Csm3 is composed of an RNA recognition motif (RRM) core elaborated with an RNase loop following β 1, the α 2 region and the thumb (or hook) region (Fig 5A and 5B). The α 2 loop contacts the minor groove of the crRNA-target duplex and the RNase loop contacts the major groove. The RNase loop performs divalent metal dependent target RNA cleavage (Fig 5A and 5B). The thumb region inserts into the duplex disrupting nucleotide stacking (Fig 5A and 5B). The bacterial Csm3 proteins, SeCsm3 and StCsm3, are very similar (S8 Table). However, the archaeal, ToCsm3, possesses a more extensive α 2 region (Fig 5C).

The SeCas10-Csm 276 kDa complex possesses three copies of Csm3 which have similar interactions with crRNA. Our analysis below focuses on Csm3.3 contacts to crRNA. The

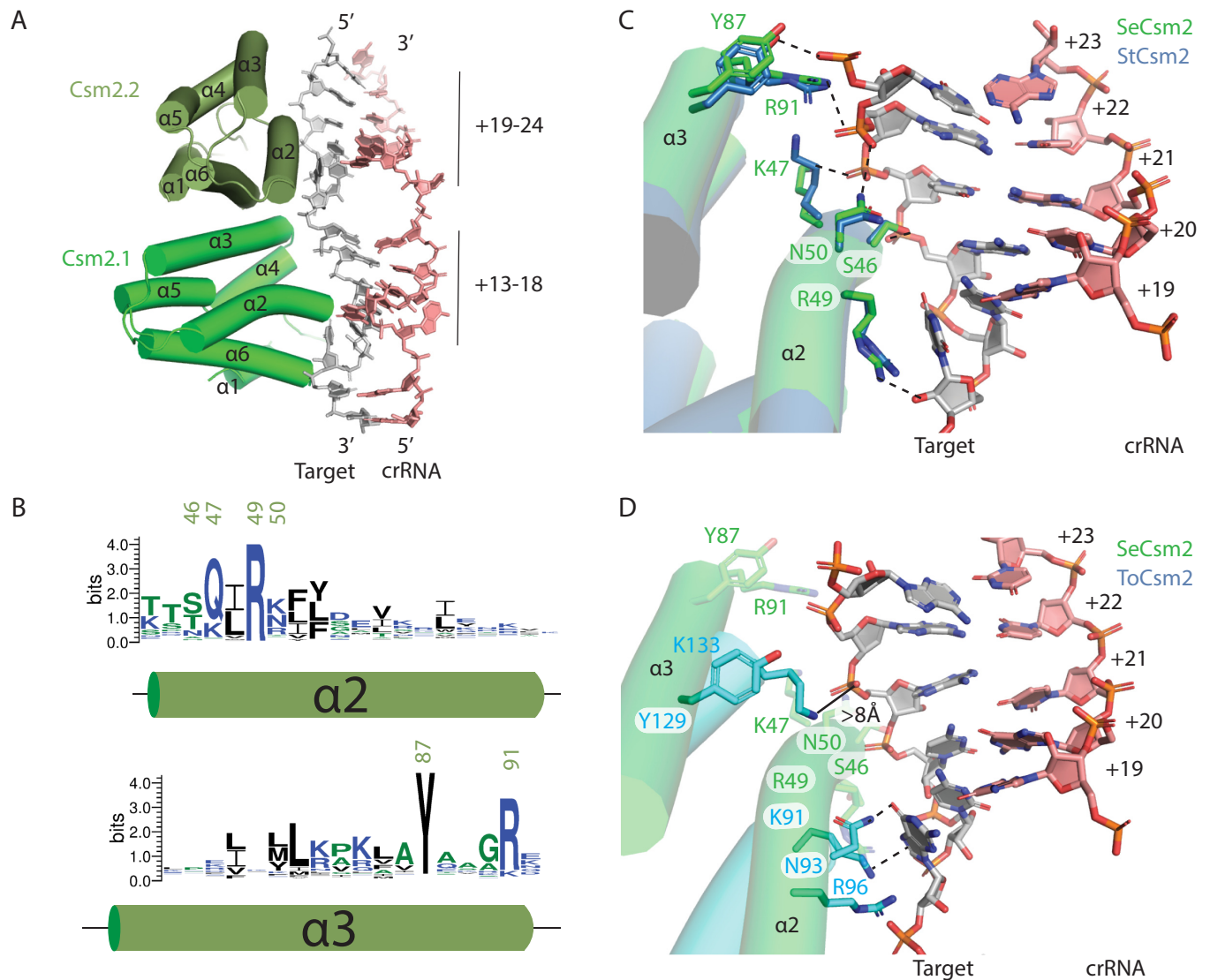


Fig 4. Conserved and idiosyncratic interactions of Csm2 with target RNA. (A) Helices $\alpha 2$ and $\alpha 3$ of *S. epidermidis* Csm2 (SeCsm2) form extensive interactions with target RNA in the major groove of the target-crRNA duplex. (B) A sequence logo depicting the conservation of $\alpha 2$ and $\alpha 3$ of Csm2. Residues of interest are numbered according to the SeCsm2 sequence. (C) A superposition of *S. thermophilus* Csm2 (StCsm2), given by PDB code 6ifu, with SeCsm2 was performed using the phosphate atoms of six nucleotides of target RNA (+19 to +24 positions). The superposition reveals that $\alpha 2$ and $\alpha 3$ in SeCsm2 and StCsm2 interact with target RNA in a similar manner. (D) A superposition of *T. onnurineus* Csm2 (ToCsm2), given by PDB code 6MUS, with SeCsm2 was performed as in (C) which reveals that $\alpha 2$ and $\alpha 3$ in ToCsm2 are translated in space relative to SeCsm2, form fewer interactions with target RNA and these differ from the pattern seen in StCsm2 and SeCsm2. K91 of ToCsm2 forms a cation-pi interaction with the target-RNA nucleotide at position +18, analogous to the interaction of SeCsm2 R49 with the +18 position.

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majority of contacts to crRNA occur via $\alpha 1$ residues. K52 and R56 make electrostatic interactions with the backbone of crRNA while S49, K54 and N57 contact the flipped nucleotide (+12 for Csm3.3, Fig 6A). Thumb residues, N125 and I127 make additional contacts to the +12 crRNA position (Fig 6A). The $\alpha 2$ region can make a single hydrogen bond to the minor groove via S86. The D32 residue of the RNase loop, which has been shown to be critical for target cleavage, is visible in the structure but is not coordinating a divalent metal ion, as it is expected to, because of the EDTA treatment of our sample prior to structural analysis (Fig 6A and 6B).

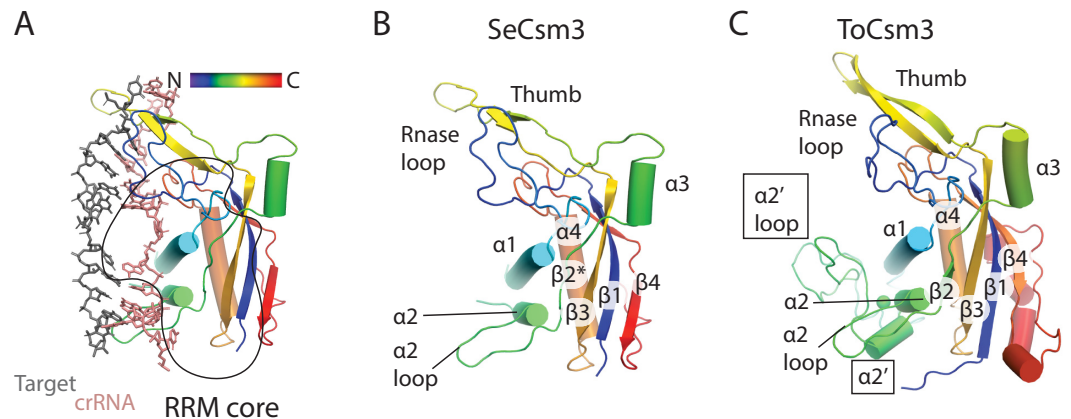


Fig 5. Variation in the secondary structure of Csm3. (A) The hydrophobic core of Csm3 is composed of an RNA recognition motif (RRM core). The secondary structure of Csm3 is colored by the position of each element on the chain from the N-terminus (blue) to the C-terminus (red). (B) The secondary structure of *S. epidermidis* Csm3 (SeCsm3) is shown with key functional components highlighted: The RNase loop contains Asp 32 which coordinates a metal to catalyze target RNA cleavage, $\alpha 1$ forms key interactions with crRNA, the $\alpha 2$ loop interacts with crRNA and lastly the thumb, composed of a β -hairpin protrudes through the crRNA-target RNA duplex interrupting it. $\beta 2^*$ indicates that SeCsm3 adopts only β -strand like geometry at the position of $\beta 2$ in the canonical RRM fold. (C) The secondary structure of *T. onnurineus* Csm3 (ToCsm3) differs from SeCsm3 because it contains additional elements of interest including the $\alpha 2'$ helix and the $\alpha 2'$ loop which make contacts to the target RNA that do not occur in SeCsm3. The structure depicted is taken from PDB code 6mus.

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A comparison of bacterial to archaeal Csm3, reveals that ToCsm3 has similar interactions to crRNA along $\alpha 1$, however, ToCsm3 uses an electrostatic interaction from thumb residue R173 to stabilize the +12 position flipped nucleotide (Fig 6C). A substantial difference is seen in the addition of the $\alpha 2'$ region in ToCsm3 (Figs 5C and 6C). The $\alpha 2'$ loop spans the minor groove of crRNA-target duplex and contacts target RNA via residues I105 and R107 (Fig 6B). Since the bacterial Csm3 structures do not possess the $\alpha 2'$ region and contact only crRNA, not target, the ToCsm3 $\alpha 2'$ region could be the source of a substantial mechanistic difference between archaeal and bacterial Cas10-Csm [20,22].

Csm4 is a Cas5 family protein, which in both Type I and Type III CRISPR systems, interacts with the 5' end of crRNA [3,48]. Like Csm3, Csm4 consists of an RRM core with a thumb region. Helices $\alpha 3$ and $\alpha 4$ are present in both SeCsm4 and ToCsm4, but interact differently with Cas10 in each case (Fig 7A and 7B). The 5' nt of crRNA, position -8, is gripped by SeCsm4 via pi-stacking with residues F40 and H291 (Figs 1B and 8A). The -7 nt is splayed across $\alpha 1$ with a hydrogen-bonding interaction between Q251 and R191 contributing to nt positioning by steric occlusion (Fig 8A). F249 stabilizes an A-form helix segment that spans the -5 to -2 positions of crRNA by pi-stacking to the -5 nt while Y148 pi-stacks to the -2 nt (Fig 8A). Van der Waal's interactions of residues H17, L23 and R190 also appear to contribute to formation of the A-form geometry of the -5 to -2 nts, a geometry that is pre-formed to inspect the 3' flank region of target RNA for complementarity [20,22,49]. Residues Y148, V132 and L134 appear to promote the flipped conformation of the -1 nt by steric occlusion (Fig 8A).

The pattern of interaction with crRNA is conserved between SeCsm4 and ToCsm4 and several critical residues are conserved, including the His that pi-stacks to the -8 nt, the Lys residue that sits beneath the -3 nt and the Tyr that pi-stacks with the -2 nt. An additional Lys residue, K241 in ToCsm4, is conserved between the archaeal and bacterial Csm4 however appears to interact with the -6 nt in SeCsm4 versus the -7 nt in ToCsm4 (Fig 8A and 8B). A difference between the two Csm4 proteins is in the thumb region where ToCsm4 appears to use R136 to stabilize the flipped -1 nt. In contrast to Csm2 and Csm3 where comparison of archaeal and

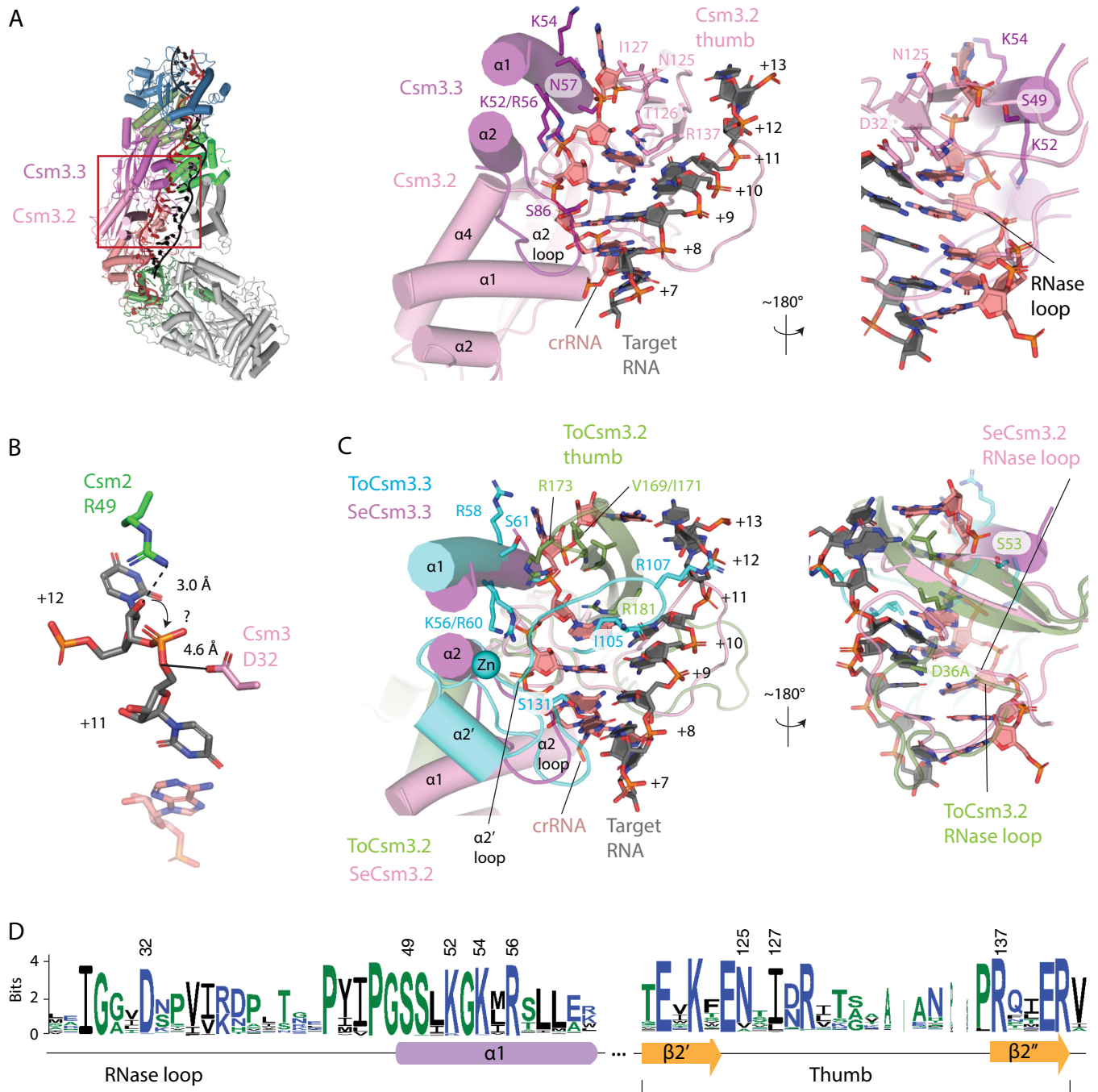


Fig 6. *S. epidermidis* Csm3 only contacts crRNA, however, *T. onnurineus* Csm3 contacts both crRNA and target RNA, utilizing the $\alpha 2'$ loop. (A) A red rectangle marks the area of Cas10-Csm which is magnified to the right. The detailed interactions of *S. epidermidis* Csm3 with crRNA are shown. A basic patch in α -helix 1 interacts with the terminal paired and the unpaired crRNA nucleotides of the repeating hexa-nucleotide structure. The S86 residue in the $\alpha 2$ loop hydrogen bonds to the minor groove of the duplex and residues N125-I127 of the Csm3.2 thumb form van der Waal's interactions and hydrogen bonds with crRNA. An extensive loop structure that follows β -strand 1 of Csm3 and sits on the major groove of the crRNA-target duplex, is termed the RNase loop because it contains the catalytically essential residue D32. (B) Csm2 R49 and Csm3 D32 are positioned to catalyze target RNA cleavage but the mechanism of cleavage, in-line attack (arrow) or hydrolysis is unclear. (C) A superposition of *T. onnurineus* Csm3 (ToCsm3), given by PDB code 6mus, is shown revealing similar interactions of α -helix 1 and the $\alpha 2$ loop with crRNA in the two structures. However, ToCsm3 uses the $\alpha 2'$ loop, a structure not present in SeCsm3, to form hydrogen bonds (I105) and electrostatic interactions (R107) to target RNA. Additionally, the ToCsm3 thumb interacts differently with crRNA utilizing an electrostatic interaction via R173 that is absent in SeCsm3. The crRNA-target duplex shown is from PDB code 6mus. (D) A sequence logo depicting the conservation of Csm3 residues of interest is given. Numbering of the logo is according to the SeCsm3 sequence.

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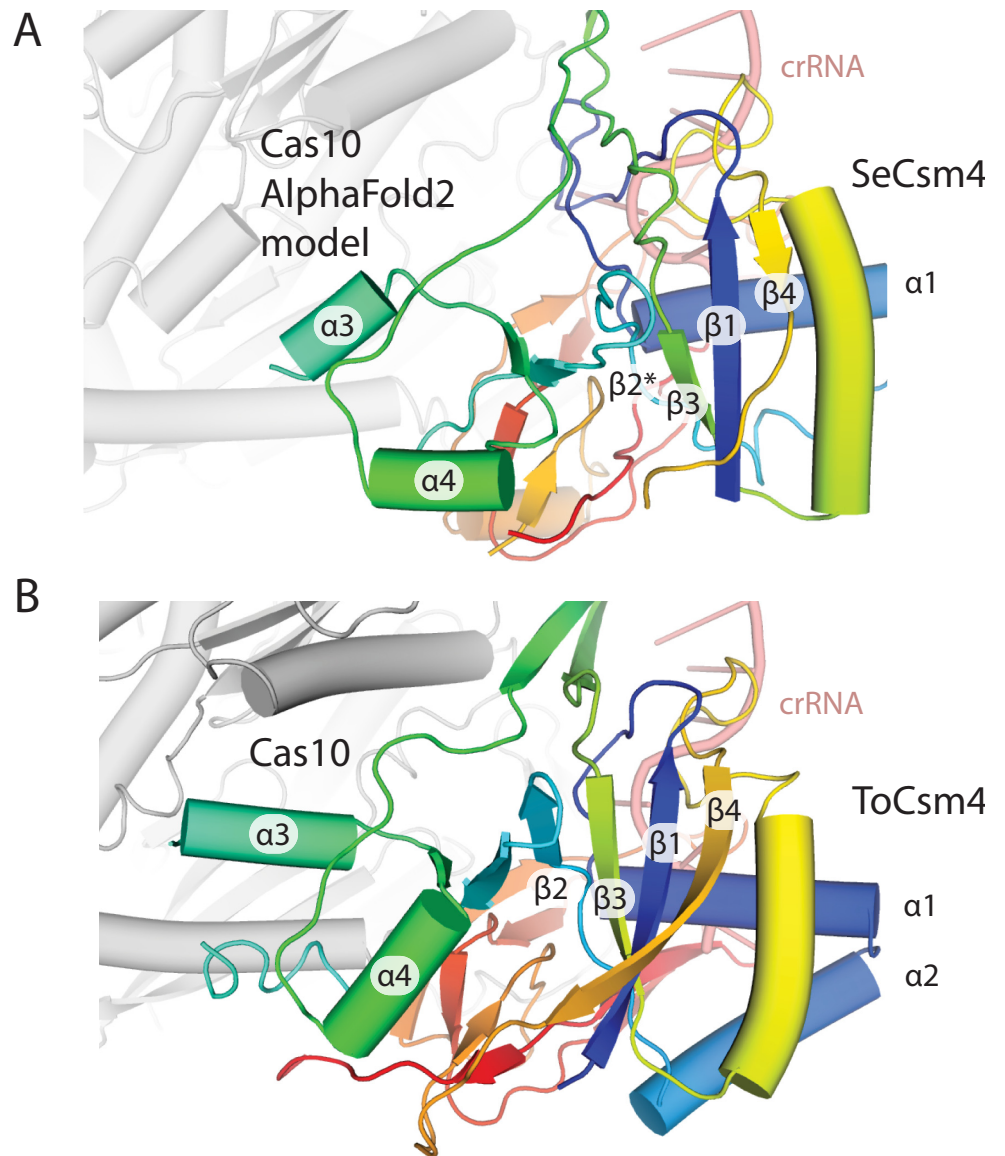


Fig 7. Csm4 possesses similar secondary structure elements in *T. onnurineus* and *S. epidermidis*. (A) The RNA recognition motif core of the Cas5 family member, SeCsm4, is elaborated to contact the 5' tag of crRNA and Cas10. An AlphaFold2 model of SeCas10 is docked into the density present in *S. epidermidis* cryo-EM maps. (B) The secondary structure elements of ToCsm4 are similar to SeCsm4 but $\alpha 3$ has more extensive interactions with Cas10.

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bacterial structures suggest likely mechanistic differences, the Csm4 structures suggest a conserved mode of interaction with crRNA.

Csm5 is a RAMP domain protein that binds the 3' end of the crRNA-target duplex. The position of Csm5 in the complex suggests it blocks extension of the Csm3 and Csm2 oligomers and promotes nucleolytic maturation of the 3' end of crRNA by cellular nucleases—two processes that are likely inter-related [50]. The RAMP domain of Csm5 blocks the extension of the Csm3 oligomer and the primarily α -helical capping domain blocks extension of the Csm2 oligomer (Figs 1D, 1E and 9A). Five regions of Csm5 interact with RNA including the $\beta 1$ -loop, the capping domain, the thumb, α -helix 1 and the $\beta 3'$ -loop (Fig 9A). The $\beta 1$ -loop possesses two conserved Gly residues allowing a sharp kink in the peptide backbone that hydrogen

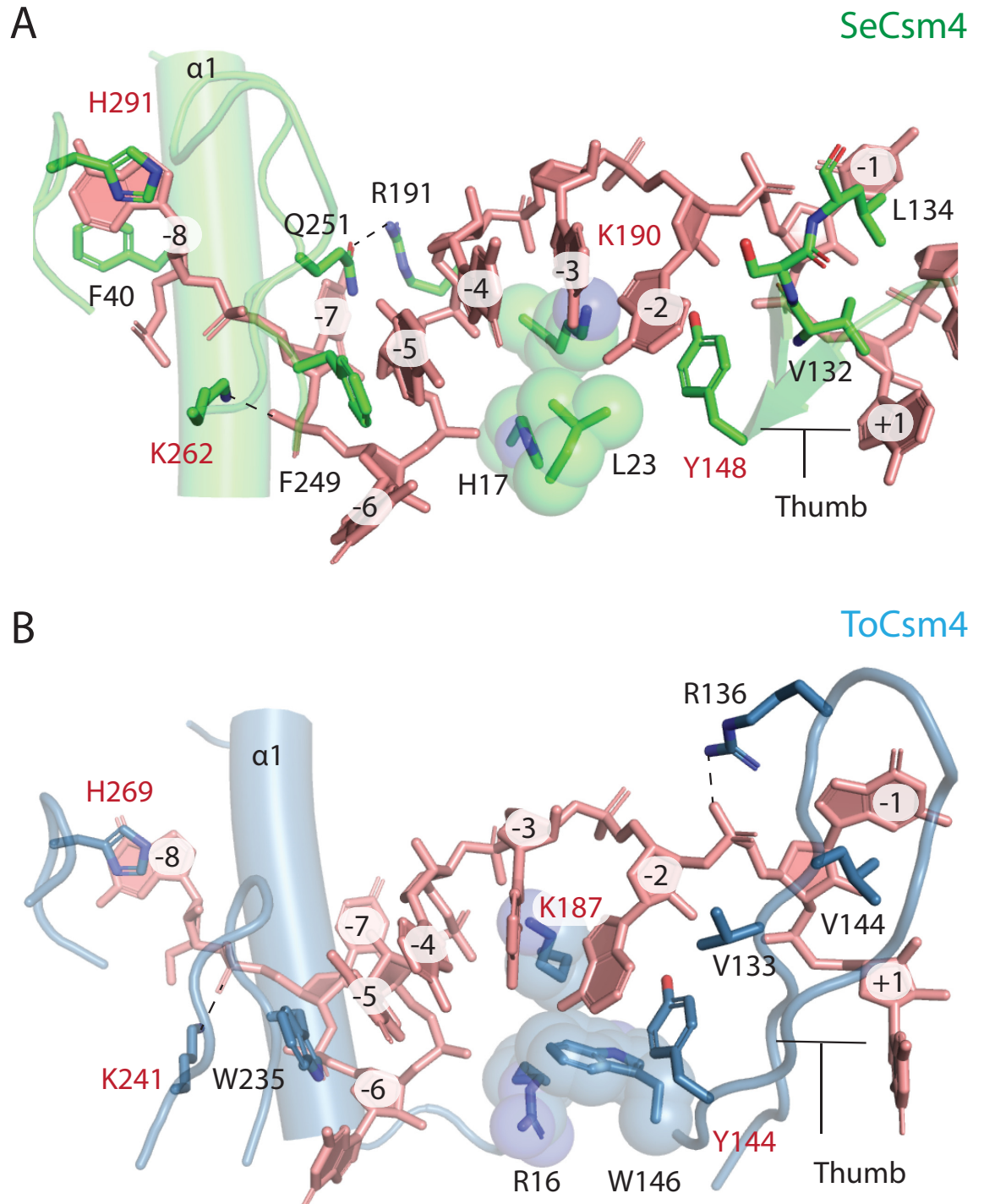


Fig 8. A detailed comparison of the interactions of SeCsm4 and ToCsm4 with crRNA. (A) SeCsm4 residues forming interactions with the crRNA 5'-tag that are likely critical for recognition shown as sticks. Three residues that form van der Waals interaction with crRNA and sterically position it are shown as spheres. Residues that are conserved between SeCsm4 and ToCsm4 are labeled in red. The thumb, β -hairpin-like element, that interrupts the co-axial nucleotide stack is labeled. The first α -helix of Csm4 is marked as $\alpha 1$. (B) ToCsm4 residues forming critical interactions with crRNA are shown as sticks or spheres. The β -hairpin-like element thumb is labeled as is α -helix 1, $\alpha 1$.

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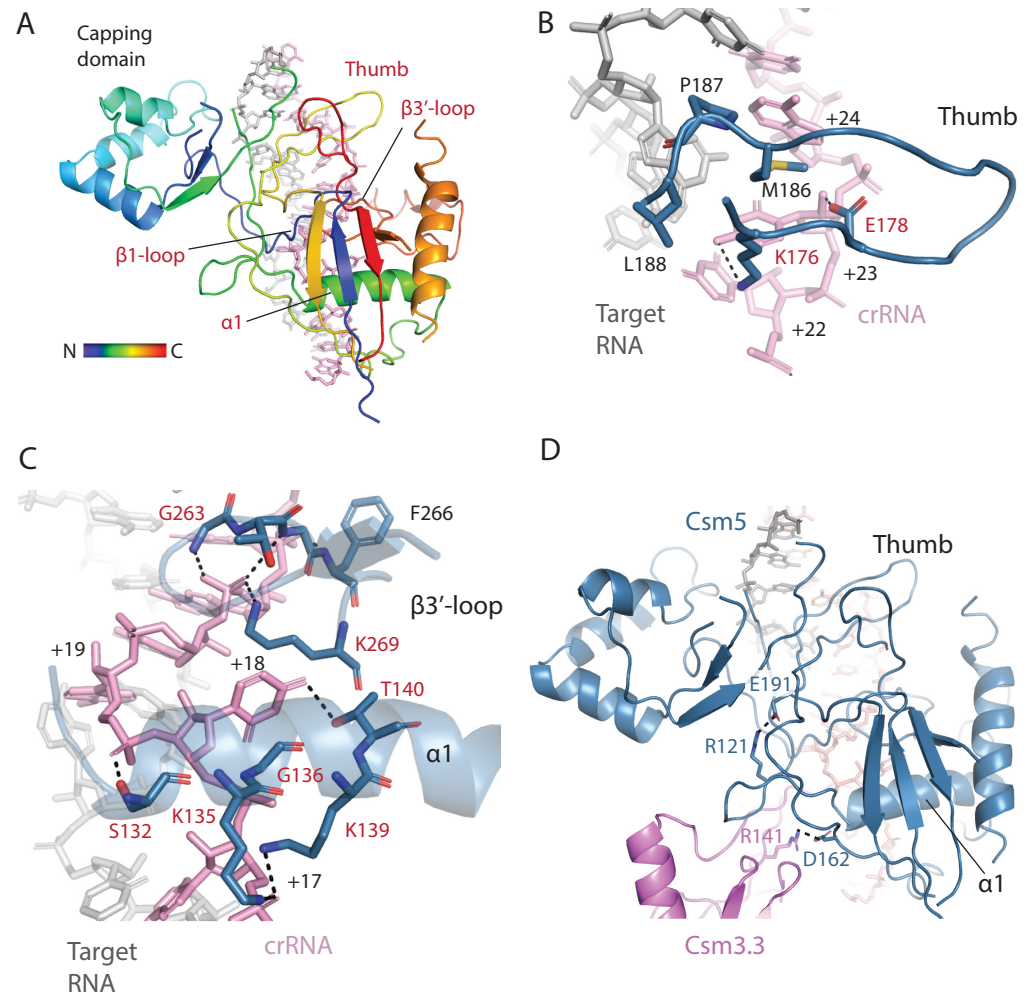


Fig 9. Most RNA-interacting regions of Csm5 are conserved and an electrostatic interaction may allow cross-talk between two regions. (A) SeCsm5 bound to the crRNA-target duplex is shown emphasizing regions that contact the duplex. Four of the five regions are conserved (red) across diverse Csm5 homologs. The secondary structure elements of Csm5 ($\beta 1$, $\alpha 1$ and $\beta 3'$) are named according to their position in the core, RRM domain—capping domain elements were bypassed. (B) The SeCsm5 Thumb region interacts with the major groove of the duplex by hydrogen-bonding and van der Waals interactions. Residues conserved across SeCsm5 homologs are colored red. (C) Helix- $\alpha 1$ and the $\beta 3'$ -loop interact with the duplex surrounding the flipped +18 nucleotide. (D) Electrostatic interactions between R121 and E191 may conformationally couple the helix- $\alpha 1$ and thumb components of SeCsm5. The interaction of SeCsm5 D162 with Csm3 R141 has been shown to be critical for crRNA maturation.

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bonds to crRNA (S10A Fig). The thumb interacts with the major groove of the RNA duplex but doesn't intercalate the duplex as the Csm3 thumb does (Fig 9B). The $\beta 3'$ -loop and $\alpha 1$ appear to work in unison gripping the crRNA in the vicinity of nucleotides +17 to +21 in the 276 kDa complex (Fig 9C). Interestingly, two SeCsm5 acidic residues, D162 and E191, that were previously shown to be required for crRNA maturation are observed participating in electrostatic interactions (Fig 9D) [50]. E191, which sits at the base of the thumb interacts with R121 that is positioned slightly C-terminal to α -helix 1. A potential explanation for the role of E191 in crRNA maturation is that the E191-R121 interaction allows cross-talk between the two crRNA binding regions, perhaps through an allosteric mechanism that enhances Csm5's affinity for crRNA. The interaction between Csm5 residue D162 and the Csm3 residue R141 might contribute to crRNA maturation by influencing how Csm5 is positioned relative to its

neighbors within the Cas10-Csm complex. Such positional and allosteric changes in Csm5 may impact the recruitment of PNPase and RNaseR, cellular nucleases which are now known to bind Csm5 and catalyze crRNA maturation [51,52].

Structural superpositions and multiple sequence alignments indicate that four of the five Csm5 regions that interact with RNA are conserved from bacteria to archaea (Figs 9 and S10). The capping domain, however, that exclusively interacts with target RNA, diverges in structure and sequence even among bacteria (Fig 10). SeCsm5 uses the basic residues K25 and K26 on the N-terminal end of the capping domain to form electrostatic interactions to target RNA (Fig 10A and 10B). These residues show modest conservation among bacterial Csm5 proteins but the additional SeCsm5 contacts to target RNA, such as R71, E72 or N114, display even less

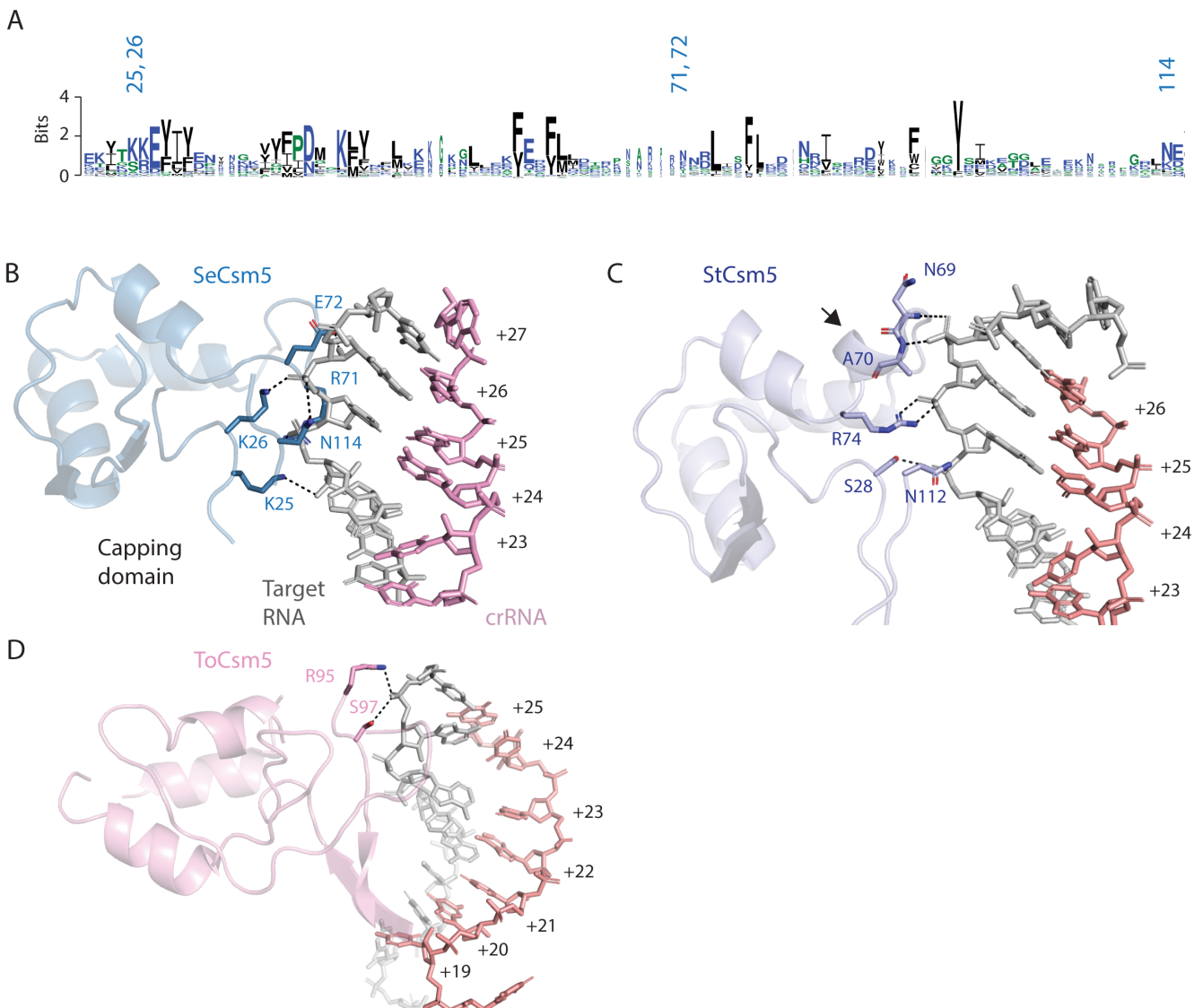


Fig 10. The capping domain of Csm5 diverges among Csm5 proteins. (A) A sequence logo of the capping domain with the positions of SeCsm5 residues that interact with target RNA numbered. The logo indicates there is limited sequence conservation within the capping domain. (B) Electrostatic and hydrogen bonding interactions between SeCsm5 and target RNA are indicated by dashed lines. (C) The interactions of the StCsm5 capping domain with target RNA are indicated. StCsm5 contains a helical segment, indicated by an arrow, absent in the other Csm5 structures. (D) Only two residues in the capping domain of the ToCsm5 structure reported in PDB code 6mus contact target.

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conservation (Fig 10A). Structural comparisons of SeCsm5 to StCsm5 confirm divergent contacts to target RNA are occurring and even show that StCsm5 possesses different secondary structure in the capping domain compared to SeCsm5 (See the arrow in Fig 10C). Inspections of the ToCsm5 capping domain, suggests minimal contacts to target RNA in the archaeal Csm5 (Fig 10D). In sum, while the RAMP domain of Csm5 possesses a pattern of interactions with crRNA consistent with that of other RAMP domain proteins (Csm3 and Csm4), the capping domain, which contacts target RNA, possesses idiosyncratic structures which suggest the possibility for functional differences among Csm5 proteins. Two potential areas for functional divergence could be in target RNA sensing or in the recruitment of cellular nucleases for crRNA 3' end maturation [51,52].

Discussion

We report cryo-EM data demonstrating that SeCas10-Csm forms two oligomerization states, a 276 kDa complex with the stoichiometry, Cas10₁ Csm2₂ Csm3₃ Csm4₁ Csm5₁, and a 318 kDa complex that is formed by addition of an extra copy of Csm2 and Csm3. These two stoichiometries were recently observed in the cryo-EM reconstruction of *L. lactis* Cas10-Csm [21]. Recent cryo-EM reconstructions of SeCas10-Csm by Smith and co-workers (published while this manuscript was in preparation) reported these two stoichiometries as well and observed a Cas10₁ Csm4₁ Csm3₅ stoichiometry; five copies of Csm3 were observed and Csm2 and Csm5 were not present [24]. Biochemical assays of SeCas10-Csm by Smith and co-workers demonstrated the presence of Csm2 and Csm5 in the as-isolated complex and the complexes were purified from *S. epidermidis* cells as ours also were. It is likely that in the absence of target RNA and under the conditions Smith and co-workers used for EM grid preparation, Csm2 and Csm5 weakly associate with the complex [24]. A weak association of Csm2 with apo *L. lactis* Cas10-Csm has been reported and this phenomenon is likely linked to conformational dynamics critical for regulating interference that will be discussed below [21]. The Csm3_n Csm2_{n-1} stoichiometry, that we report, has been observed for Cas10-Csm complexes from other species as well [20–22]. The existence of two stoichiometries of SeCas10-Csm differing by the presence of one Csm2-Csm3 unit was anticipated due to two previous observations: in the presence of a crRNA, in vitro, Csm3 assembles into oligomers with protomer number depending on crRNA length and as-isolated SeCas10-Csm contains primarily two lengths of crRNA, a 37-mer and 43-mer (Fig 1C) [27].

The identification of the two SeCas10-Csm states, 276 kDa and 318 kDa, is important because it has been recently argued that the co-existence of multiple oligomerization states in Type III CRISPR complexes imparts a unique interference behavior. Several researchers have shown that cOA synthesis is sensitive to mismatches in base-pairing between target and crRNA proximal to Cas10, the region denoted as positions +1 to +7 or the Cas10-activating region (Fig 1B) [12–14,29]. Steens and colleagues argue that the region of the target-crRNA duplex distal to Cas10, composed of positions +19 to +37, also performs a check on base-pairing complementarity by controlling initiation of duplex formation [13]. Mismatches in the vicinity of the 3' end crRNA disfavor target binding and thus interference. However, this effect is obscured when assayed in a population of mixed oligomerization states since the complexes contain different crRNA 3' ends [13]. The interaction of these phenomenon suggests that distinct oligomerization states of Type III CRISPR complexes are sensitive to mismatches at both the 5' and 3' end of crRNA but the effect of mismatches at the 3' end are obscured when pooled oligomerization states are assayed. The data indicating base-pairing at the crRNA 3' end critically controls interference were obtained in a Type III-B complex which contains the Cmr1 protein at this region, a protein absent from Type III-A complexes. Therefore, an important

future direction is to determine the precise mechanistic effects of mismatches throughout the crRNA-target duplex in Type III-A CRISPR and determine how Cas10-Csm oligomerization state affects sensitivity to mismatches. A complete understanding of how mismatches effect cOA synthesis in Type III CRISPR is required to understand the contributions of these complexes to bacterial physiology and to efficiently develop molecular diagnostics based on them.

The importance of Csm2 in Cas10-Csm mediated interference has been questioned because in vitro experiments showed that the critical biochemical activities remain in a *S. thermophilus* complex lacking Csm2 [53]. However, later experiments with *S. thermophilus* Cas10-Csm complex containing a Csm2 site-directed mutant showed a complete loss of target RNA cutting and experiments with *L. lactis* complexes showed severe defects in DNase activity, cleavage of target RNA and cOA synthesis in a complex lacking Csm2 [20,21]. Importantly, in vivo experiments in *S. epidermidis* and *L. lactis* revealed that interference falls to background levels in the absence of Csm2 [21,54]. A caveat of these studies is that deletion of Csm2 caused a depletion of Csm5 in the complex and the interference defects could be the combined effect of depletion of both proteins.

Csm2 contacts to cognate target RNA were well-resolved in our 3.1 Å SeCas10-Csm reconstruction. These contacts could not be resolved in previous reconstructions of SeCas10-Csm due to the lower resolution of those structures [24]. Interactions of Csm2 residues along α -helix 2 with target RNA are a common feature among structures from multiple organisms (Fig 4C) [20,21,23]. In the *S. thermophilus* complex Csm2 residue R41, which is structurally equivalent to SeCsm2 residue R49, has been shown to be required for target RNA cleavage [20]. A similar finding was made in *L. lactis* where Csm2 residue R48 is required for target RNA cleavage and the authors argue R48 acts as a general base on the 2' hydroxyl of the labile nucleotide in an RNase A like mechanism (Fig 6B) [21]. It is then unsurprising that this Arg is the most conserved residue in α -helix 2 (Fig 4B). While multiple studies observe interactions of Csm2 with target RNA similar to ours, the *T. onnurineus* Cas10-Csm structure diverges substantially (Fig 4D). ToCsm2 residue R96, which is equivalent to SeCsm2 R49 in multiple sequence alignments, does not contact the 2' hydroxyl of the labile, target nucleotide raising the question of why this residue is conserved in ToCsm2. One possibility is that the thermophilic ToCas10-Csm complex has not adopted its final, target bound conformation in the available cryo-EM reconstruction, a conformation that would involve a reorganization of ToCsm2 so that R96 contacts the labile target RNA residue [22].

Cas7 family members play a critical role in the function of Type I and Type III CRISPR systems by forming an oligomer that binds crRNA and presents it to solvent for base-pairing [48]. In Type III CRISPR systems, the Cas7 family member, Csm3, plays an additional role providing an Asp residue to the active site that cleaves target RNA [8,55]. The RNase loop containing the Asp residue binds in the major groove of the crRNA-target duplex but sits near the 2'OH of the labile nucleotide because this nucleotide is flipped out of the co-axial stack (Fig 6A and 6B). Structures of Cas10-Csm from *T. onnurineus* and *L. lactis* were determined using Asp to Ala mutants of the critical residue to prevent target RNA cleavage during structure determination [21,22]. We used the alternate approach of incubating the complex with EDTA to chelate Mg^{2+} allowing determination of a structure with wild type sequence. You and co-workers used a similar approach determining the structure of *S. thermophilus* Cas10-Csm with wild type sequence [20]. In both the structures with wild type sequence the Asp residue is pointed towards the 5'O of the reaction product at a distance of 4–5 Å (Fig 6B). The recent structure of SeCas10-Csm bound to cognate target (non-self) RNA by Smith and co-workers observed a distance of 10.2 Å between Csm3 D32 and the labile phosphate [24]. This large distance is explained by the fact that the investigators captured a product complex with cleaved target RNA. Sridhara and co-workers have proposed an RNase A-like mechanism for target RNA

cleavage and the position of the critical Asp is consistent with its role as the general acid in this mechanism but does not explain the importance of Mg^{2+} for target cleavage [21]. Future studies are needed to clarify the mechanism of target RNA cleavage but structural and biochemical data are consistent in implicating SeCsm2 R49 and SeCsm3 D32 as the critical residues for this activity [8,20,21,55].

Our analysis of Csm3 interactions with the minor groove of the crRNA-target duplex revealed a major difference between archaeal and bacterial Csm3 (Fig 6C). Archaeal Csm3 engages in interactions with the target RNA in the minor groove but the bacterial protein does not. Additionally, a Zn^{2+} ion binds adjacent to $\alpha 2$ and $\alpha 2'$ in the archaeal Csm3 but not in the bacterial protein. The functional significance of Csm3 interactions with the minor groove are not known but it seems reasonable to think they play a role in sensing bound target RNA, including a role in the sensitivity of Cas10-Csm to mismatches in base pairing. If this is the case, the differences in archaeal and bacterial Csm3 may mediate differences in mismatch sensitivity and there could be a role for the Zn^{2+} ion in target RNA detection in archaeal Cas10-Csm.

Functional studies have implicated a role for Csm5 in the maturation of the 3' end of crRNA [50], and it is now understood that Csm5 recruits two cellular nucleases, PNPase and RNaseR, which catalyze crRNA maturation [51,52]. Site-directed mutants of two acidic residues in SeCsm5 cause defects in crRNA maturation but the mechanism for this effect was unclear. Our structural results suggest an explanation. The R121-E191 interaction may allow cross-talk between the two crRNA binding regions of Csm5, the thumb region and the α -helix 1 region (Fig 9D). For example, binding of the α -helix 1 region to crRNA could influence nearby R121 to promote the formation of the electrostatic interaction with E191 enhancing the affinity of the thumb region for crRNA. While E191 has only been functionally investigated in *S. epidermidis*, this residue is conserved in Csm5 homologs and in the *S. thermophilus* structure reported by PDB code 6ifu the residue and its electrostatic interaction are conserved (S10D Fig). Therefore, the R121-E191 interaction may modulate Csm5 conformational changes and thus the recruitment of crRNA maturation nucleases in organisms harboring Type III systems. Our structure revealed that SeCsm5 residue D162 forms an electrostatic interaction with a Csm3 residue (Fig 9D). We hypothesize that the crRNA maturation defect arising from D162A is caused by a mis-positioning of Csm5 relative to Csm3 in this mutant. Further studies are needed to investigate this but it is notable that the D162-Csm3 R141 electrostatic interaction is also conserved in *S. thermophilus* (S10D Fig).

Csm5 has been found to influence the affinity of target RNA binding to *S. thermophilus* Cas10-Csm and in a Type III-B system it was found that mismatches between the 3' end of crRNA and target decreased affinity for target RNA leading to interference defects [13,53]. Both results suggest an important role for Csm5, or its Type III-B homolog Cmr6, in Type III CRISPR function. Importantly, we note in our structural analyses that the only region of Csm5 in Type III-A complexes that contacts target RNA is the capping domain, a region that diverges in structure and sequence among Csm5 homologs (Fig 10). Additionally, substantial conformational change in the Csm2 subunits has been observed upon target RNA binding to StCas10-Csm [20]. Since the capping domain of Csm5 contacts both target RNA and Csm2, we speculate that there may be an underlying connection between the phenomenon mentioned above: that interaction of the capping domain of Csm5 with target RNA promotes conformational changes in Csm2 that enhance its affinity for target RNA, potentially a critical early step in activating interference. Superpositions of target RNA bound versus unbound structures from *S. thermophilus* and *S. epidermidis* support this hypothesis (S11 Fig). The fact that the Csm5 capping domain diverges in structure and sequence appears to conflict with the argument that it plays a crucial role in interference, however, it may be that differences in the

capping domain are necessary for Csm5 to interface with non-Cas cellular nucleases specific to each organism.

Type III CRISPR systems are uniquely suited to serve as molecular diagnostics because these systems are adapted to perform specific detection of nucleic acids and amplify the detection event by a multiple-turnover enzymatic assay. This intrinsic potential of Type III CRISPR systems was realized in the last year with the publication of four examples of the deployment of a Type III complex to detect viral RNA [11–14]. We believe that the subtle variations in the structure of Type III-A complexes that we have described suggest there may be activity differences in these complexes, such as their intrinsic sensitivity to mismatches in base-pairing or affinity for target RNA, that would make one complex better suited to a role in molecular diagnostics than another. Additionally, we confirm that SeCas10-Csm forms two oligomeric states, a 276 kDa complex and a 318 kDa complex, and we believe this information will aid in investigating how mismatches at the 3' end of crRNA-target duplex affect interference, a question whose answer is important in the construction of molecular diagnostics and other biotechnologies based on Type III CRISPR-Cas systems.

Supporting information

S1 Fig. Processing workflow in cryoSPARC V2. At step 1, two initial maps with different stoichiometries were obtained using template matching. From particles making those maps a set of 5000 particles were used for Topaz training at step 2. Complexes I, II, and III denote the smallest to largest stoichiometries. At step 3, heterogeneous refinement was done on the full set of 1.4 million particles. Cas10 is grey, Csm2 is light green, Csm3 is magenta, Csm4 is dark green, Csm5 is blue and target RNA is black. At step 4, the map denoted as complex II and III was further classified with two input maps from variability analysis. Each of the complexes II and III were refined to high resolution with unbinned images at step 5. FSC curves and B-factor plots are included. Due to flexibility of the Cas10, this density is averaged out in the high-resolution refinement of complex II and to a lesser extent in complex III. Csm2 densities are also averaged out at high resolution in complex III because they were under-populated. Complex II was used for the molecular models presented in this manuscript.
(TIF)

S2 Fig. Cyclic oligoadenylate synthesis by SeCas10-Csm encoded by *pcrispr-spc1* or *pcrispr*. A 24% urea-PAGE gel was used to visualize production of ³²P-containing cyclic oligoadenylates produced by the incubation of target RNA (ssRNA-01) and ATP with *S. epidermidis* Cas10-Csm expressed from the *pcrispr spc1* (SeCas10-Csm *spc1*) plasmid which contains only one spacer gene or SeCas10-Csm expressed from the *pcrispr* plasmid which contains all three spacer genes found in the *S. epidermidis* genomic, CRISPR locus. The products from three technical replicates are shown.
(TIF)

S3 Fig. Cryo-EM map quality. Different subunits of the Cas10-Csm complex are shown in each panel. Each panel contains the density on the right and the model fit in the density on the left. Panel F shows the side chain densities for Csm2 along with the residue numbers.
(TIF)

S4 Fig. Local resolution plots of Complex II density. (A) A local resolution plot for the complex of Csm2-5 proteins, crRNA and target RNA. (B) Local resolution plots for the individual proteins and RNA in the complex. The label, RNA, denotes the crRNA-target duplex. (C) A histogram of voxels versus local resolution.
(TIF)

S5 Fig. Density for each domain of Cas10 exists in the map of the 276 kDa Cas10-Csm complex. An AlphaFold2 model of *S. epidermidis* Cas10 was docked into the density given by the map of the 276 kDa Cas10-Csm complex. The map is shown at $\sigma = 2.0$. Cas10 is color coded by domain: HD, HD nuclease domain, Palm1, Palm1 polymerase domain 1, Palm 2, Palm polymerase domain 2, D4, domain 4.

(TIF)

S6 Fig. Independently calculated rigid body models. Ten rigid body models calculated in SASREF are shown overlaid with each other; the two views are related by 90 degrees. The overall arrangement and shape of the models are highly consistent.

(TIF)

S7 Fig. Independently calculated *ab initio* models. *Ab initio* models were calculated using DAMMIN, with a D_{\max} of 160. The upper and lower panels show the same models rotated by 90 degrees.

(TIF)

S8 Fig. Deconvolution. EM models for the 276 kDa complex, the 318 kDa complex, and the 318 kDa complex without Csm2 were used to generate theoretical scattering curves, which were then used in OLIGOMER to deconvolute our experimental SAXS scattering curve. The resulting deconvolution strongly indicated that the 276 kDa complex alone was the best model for the experimental data. The composite fit is shown above as a line with the experimental data as open circles. The χ^2 for this fit was 1.75.

(TIF)

S9 Fig. Superposition of SeCas10-Csm models derived from cryo-EM and SAXS. (A) Superposition of the 276 kDa SeCas10-Csm complex (multi-colored, PDB code 8DO6) and the 318 kDa SeCas10-Csm complex (8DO6 chains docked to density, grey) reveals they differ in the number of Csm2 and Csm3 subunits and possess a modest shift of the Csm2 oligomer towards Cas10 (arrow). (B) A superposition of the 276 kDa molecular model (multi-colored, PDB code 8DO6) with a rigid-body model derived by SAXS is shown. Again, a modest shift of the Csm2 oligomer towards Cas10 is observed (arrow). Superpositions were performed using Csm4. D4, refers to Cas10 domain 4, the C-terminal domain of the protein.

(TIF)

S10 Fig. A comparison of the interactions of bacterial and archaeal Csm5 with target and crRNA. (A) A sharp kink in the peptide backbone of the loop region following β -strand 1 facilitates an interaction with crRNA. A sequence logo depicting the conservation of β 1-loop is shown above a superposition of *S. epidermidis* Csm5 (SeCsm5) and *S. thermophilus* Csm5 (SeCsm5) highlighting similarities in the interactions of the β 1-loop with crRNA. (B) The β 1-loop of *T. onnurineus* Csm5 (ToCsm5) interacts with crRNA in a conserved manner. (C) A sequence logo depicting the conservation of helix- α 1 which Csm5 interacts with the flipped +18 nucleotide in a similar manner in bacterial and archaeal Csm5 proteins. (D) Csm5 acidic residues required for crRNA maturation in *S. epidermidis* are conserved in StCsm5 and make similar electrostatic interactions.

(TIF)

S11 Fig. The structures of *S. thermophilus* Cas10-Csm display a conformational change upon target binding consistent with a role for the Csm5 capping domain in promoting a target-bound state. (A) The Cas10-Csm coordinates reported in the target RNA unbound structure, 6ifn, were superpositioned with the target bound structure, 6ifu, using Csm5

residues 111–355 as the basis for the superposition. (B) A detailed view of the boxed area in panel A, shows that the Csm5 capping domain (purple) moves toward the RNA cleft of the complex to interact with bound target RNA influencing the position of neighboring Csm2.2. (C) The Cas10-Csm coordinates reported in 7v02 were superpositioned with 7v01 using Csm5 residues 114–333. The comparison is limited by the fact that 7v01 has a partially cleaved target RNA which is also a non-cognate target, potentially affecting its interaction with the complex. (D) A detailed view of the boxed area in panel C shows a modest movement of the Csm5 capping domain toward bound target RNA.
(TIF)

S1 Table. Oligonucleotides used in the study.

(PDF)

S2 Table. Components of the atomic model of SeCas10-Csm (276 kDa complex) bound to target RNA.

(PDF)

S3 Table. SAXS rigid body components.

(PDF)

S4 Table. Cryo-EM Statistics for Data Collection and Model Quality.

(PDF)

S5 Table. Small Angle X-ray Scattering Model Calculation Statistics.

(PDF)

S6 Table. Agreement between SAXS *ab initio* models and molecular models.

(PDF)

S7 Table. CRYSOLO fitting statistics.

(PDF)

S8 Table. Sequence identities (% identity) among structurally characterized Cas10-Csm complexes.

(PDF)

S9 Table. SAXS rigid body components.

(PDF)

S1 Movie. Local resolution heat map of complex II with the Cas10 subunit excluded.

(MP4)

S2 Movie. Slice through the local resolution heat map of complex II with the Cas10 subunit excluded.

(MP4)

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