Structure of the archaeal Cascade subunit Csa5
Relating the small subunits of CRISPR effector complexes

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Abbreviations: αCascade, Type I-A Cascade; ASU, asymmetric unit; Cas, CRISPR-associated; Cascade, CRISPR-associated complex for antiviral defence; CRISPR, clusters of regularly interspaced short palindromic repeats; crRNA, CRISPR RNA; Cmr, CRISPR module RAMP; Csa, CRISPR subtype Apern; Cse, CRISPR subtype Ecoli; CSM, CRISPR subtype Mtube; ds, double-stranded; eCascade, Type I-E Cascade; HGT, horizontal gene transfer; ITC, isothermal calorimetry; ss, single-stranded; Sso, Sulfolobus solfataricus

The Cascade complex for CRISPR-mediated antiviral immunity uses CRISPR RNA (crRNA) to target invading DNA species from mobile elements such as viruses, leading to their destruction. The core of the Cascade effector complex consists of the Cas5 and Cas7 subunits, which are widely conserved in prokaryotes. Cas7 binds crRNA and forms the helical backbone of Cascade. Many archaea encode a version of the Cascade complex (denoted Type I-A) that includes a Csa5 (or small) subunit, which interacts weakly with the core proteins. Here, we report the crystal structure of the Csa5 protein from Sulfolobus solfataricus. Csa5 comprises a conserved α-helical domain with a small insertion consisting of a weakly conserved β-strand domain. In the crystal, the Csa5 monomers have multimerized into infinite helical threads. At each interface is a strictly conserved intersubunit salt bridge, deletion of which disrupts multimerization. Structural analysis indicates a shared evolutionary history among the small subunits of the CRISPR effector complexes. The same α-helical domain is found in the C-terminal domain of Cse2 (from Type I-E Cascade), while the N-terminal domain of Cse2 is found in Cmr5 of the Cmr (Type III-B) effector complex. As Cmr5 shares no match with Csa5, two possibilities present themselves: selective domain loss from an ancestral Cse2 to create two new subfamilies or domain fusion of two separate families to create a new Cse2 family. A definitive answer awaits structural studies of further small subunits from other CRISPR effector complexes.

Introduction

Clustered regularly interspaced short palindromic repeats (CRISPRs) are a prokaryotic mechanism for preventing horizontal gene transfer (HGT) and infection by mobile genetic elements. Cas genes encode the proteins that mediate the CRISPR response to the insertion of foreign double-stranded DNA (dsDNA) or single-stranded RNA (ssRNA) into the cell. The response can be divided into three stages: adaptation, the transcription of the array and processing of the transcript to short, mature CRISPR RNAs (crRNA); interference, the recognition and degradation of foreign dsDNA or ssRNA.

cas genes are arranged in loci, which can be classified into one of three main systems (types I, II and III) based on the identity and order of genes in the locus. The CRISPR/Cas systems can be further subdivided into a total of 10 subtypes (Fig. S1), all of which, except type III-B, target dsDNA. Uniquely type III-B targets ssRNA. The type I interference complex is known as the CRISPR-associated complex for antiviral defense (Cascade). The exemplar is the type I-E complex (αCascade) from E. coli, which consists of Cas5, Cas6e, Cas7, Cse1 and Cse2 in a 1:1:6:1:2 stoichiometry with the six Cas7 subunits forming a central helical spine to which crRNA and all the other subunits bind. This topology appears to be conserved in the type I-C and I-F systems.

Significantly less is known about the effector complex of type I-A systems (αCascade, also known as the archaeal Cascade), which consists of Cas5, Cas7, Csa5 and in some systems Cas8a1 or Cas8a2. It has not yet been possible to purify a full αCascade complex; the Sulfolobus solfataricus (Sso) αCascade partly dissociates during purification, while the Thermoproteus tenax complex is insoluble after overexpression, with minimal soluble material after refolding. Only a stable core complex of Cas5 and Cas7 remains after the partial dissociation of Sso αCascade. The Cas5/Cas7 complex forms helical oligomers reminiscent of the Cas7 backbone of αCascade. Interestingly this core complex

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Csa5 interacts with the Cas5/Cas7 complex. The two copies of \textit{S. solfataricus} \textit{aCascade} studied in this paper are encoded by the genes \textit{sso}1398-1400 and \textit{sso}1441-1443 (Fig. 1A). The two Csa5 paralogs share 11% sequence identity. Experiments described here utilize the Sso1441-1443 proteins unless otherwise indicated. Immunoprecipitation of the Cas5/Cas7 core complex from \textit{S. solfataricus} cell lysates using affinity-purified antibodies raised against the recombinant complex co-precipitated small amounts of Csa5 (Fig. 1B). The reciprocal experiment using antibodies against Csa5 pulled down both Cas5 and Cas7. The small amounts that are co-immunoprecipitated are indicative of a weak interaction between Csa5 and Cas5/Cas7, consistent with the dissociation of Csa5 from the complex during purification. \textsuperscript{19}

Lintner et al.\textsuperscript{19} showed that cellular Cas5/Cas7 (Sso1441-1442) bound to crRNA, raising the possibility that Csa5 only binds to Cas5/Cas7 in the presence of crRNA. Fluorescently labeled oligonucleotides were used for fluorescence anisotropy experiments to determine interactions between Cas5/Cas7 and Csa5. Consistent with previous studies,\textsuperscript{19} the Cas5/Cas7 complex bound to fluorescently labeled crRNA with a dissociation constant (K\textsubscript{D}) of 1.38 ± 0.14 \textmuM (Fig. 2A). However, Csa5 itself did not form a stable interaction with crRNA, ssDNA or a heteroduplex formed from crRNA and ssDNA containing a protospacer in vitro (Fig. 2B), nor did we observe convincing evidence for binding to a pre-formed complex of Cas5/Cas7/crRNA (Fig. 2C).

The crystal structure of Csa5. The limited solubility of Csa5 encoded by Sso1443 hindered its crystallization and so its paralog, Sso1398, was crystallized. The structure was solved using a selenomethionyl variant with diffraction recorded to a resolution of 2.7 Å (Fig. 3A and Table 1). The asymmetric unit (ASU) contains nine monomers and each protein monomer is constructed of two domains, denoted \alpha and \beta (Fig. 3B). The \alpha-domain is an all \alpha-helical arrangement consisting of \alpha\textsubscript{1}, \alpha\textsubscript{2}, formed by residues 5–64 and \alpha\textsubscript{3} and \alpha\textsubscript{4}, formed by residues 130–162. The \beta-domain is formed by residues 65–129 and is an insertion between \alpha\textsubscript{4} and \alpha\textsubscript{5}. The \beta-domain contains a single helix (\alpha\textsubscript{6}) and six \beta-strands (\beta\textsubscript{1}–\beta\textsubscript{6}) arranged as three anti-parallel two-stranded \beta-sheets (Fig. 3A). Residues 1–4 were disordered in all monomers; the loops connecting \beta\textsubscript{1}–\beta\textsubscript{2} and \beta\textsubscript{5}–\beta\textsubscript{6} were disordered in some monomers. The residues that are conserved across the Csa5 family are located in the \alpha-domain, although the short \beta\textsubscript{6} strand may show some conservation (Fig. S2). The \beta-domain insertions of Csa5 homologs vary in length from 20–67 residues and show minimal sequence conservation, even between closely related strains (Fig. S2).

Strikingly the nine monomers of the ASU are assembled in a continuous helical oligomer that, due to crystal symmetry, creates
Dynamic light scattering experiments showed that the mutant had a smaller radius of gyration than the wild-type, supporting the gel filtration data (Fig. 4B). As both residues are conserved, we reasoned the salt bridge may be a conserved feature, so we examined the effect of the corresponding D32A mutation in Sso1443 (Csa5 itself). Recombinant wild-type Csa5 had a solubility limit of 1 mg/ml, whereas Csa5D32A exhibited a >10-fold increase in solubility. Csa5D32A eluted from a gel filtration column consistent with a monomeric species, an observation independently supported by isothermal calorimetry (Fig. S4).

Structural homology to other Cas proteins. The structure of Sso1398 was compared with known structures and while the PDBeFold server did not find any significant matches, the DALI server detected homology for the \( \alpha \)-domain while the \( \beta \)-domain does appear to be unique. The \( \alpha \)-domain (excluding \( \alpha_1 \)) matched the C-terminal domain of Cse2 from *Thermus thermophilus* (PDB 2ZCA, Z score 4.5, 2.7 Å over 73 residues) and to a lesser extent Cse2 from *Thermobifida fusca* (PDB 4H79, Z-score 2.9, 2.9 Å over 73 residues). It is also similar to the N-terminal domain of StySJI M, a Type I HsdM subunit from a restriction enzyme of *Bacteroides thetaiotaomicron* (PDB 2OKC, Z score 3.3, 2.6 Å over 68 residues); the domain itself has no assigned function.

Oligomerization of Csa5 proteins. The interfacing residues are predominantly located in the \( \alpha \)-domain with a few contacts from \( \beta_1 \), \( \beta_2 \), and \( \beta_6 \). Analysis of the interfaces within the helices using the PDBe PISA server showed that 1100 Å\(^2\) of surface area from each monomer was buried within a “dimer.” The interfaces are hydrophilic and are scored in PISA as unstable in solution. In our analysis, we noted an intersubunit salt bridge between two conserved residues, Asp35 and Arg55 (Figs. 3D; Fig. S2). Disrupting this salt bridge (Sso1398D35A) led to the protein eluting from a calibrated gel filtration column with a larger retention volume than the wild-type protein (16.4 ml and 13.9 ml, respectively) (Fig. 4A), indicating that the mutant is smaller. The estimated weights of both native and mutant are physically unrealistic (26 and 6 kDa, respectively, where a monomer is 19 kDa) and we attribute this to the unusual shape of the protein. Dynamic light scattering experiments showed that the mutant had a smaller radius of gyration than the wild-type, supporting the gel filtration data (Fig. 4B).

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**Figure 2.** Analysis of the interactions of a Cascade with nucleic acids using fluorescence anisotropy. (A) The average change in anisotropy upon binding of the Cas5/Cas7 complex to fluorescent crRNA. Experiments were performed in triplicate and error bars represent the standard error of the data. The data were fitted to a binding curve with a \( K_D \) of 1.38 ± 0.14 μM (error estimated from curve fitting). (B) Csa5 does not interact with crRNA, ssDNA or a heteroduplex of crRNA and protospacer. (C) Csa5 does not interact with a pre-formed complex of Cas5/Cas7/crRNA.
Cse2 forms a dimer in εCascade, which creates a positively charged surface for interacting with nucleic acids. This face is formed by both domains of the protein and, therefore, is only partially conserved in Csa5. This surface is not significantly positively charged in Sso1398 or in a homology model of Csa5. Although Csa5 possesses a weakly positively charged face at the base of the α-domain (Fig. S5), the residues that form it are not well-conserved across the family. Cse2 itself is capable of binding both dsDNA and ssRNA non-specifically and an εCascade-crRNA complex lacking Cse2 shows reduced binding affinity for target plasmid DNA. It has been hypothesized that Cse2 stabilizes the R-loop nucleotide structure by either binding the RNA/DNA heteroduplex directly or indirectly by binding to non-complementary DNA strand. In contrast, Csa5 did not bind to any nucleic acid tested, including a DNA/RNA heteroduplex and ssDNA, both of which are present in an R-loop. Therefore, despite the structural homology between Cse2 and Csa5, these two proteins may play different roles within their respective Cascade complexes.

Discussion

Our data show that the interaction of the Csa5 subunit with the core Cascade structure is below the detection limit of the biophysical techniques we employed. Pull-down experiments also point to the interaction being weak. Of course, we cannot exclude the possibility that additional factors such as Cas6 and/or the presence of target DNA are required for assembly. However, in general our data fit with an emerging picture that Csa5 easily dissociates from the εCascade complex.

The structure of the paralog Sso1398 shows a remarkable arrangement in the crystal. The continuous helical oligomers are reminiscent of the Ca7 helical backbone that appears to be a conserved feature of Cascade complexes. The S. solfataricus Cas7 oligomer exhibits a helical pitch of 14 nm, which is broadly compatible with the Csa5 pitch of 18 nm. The helical threads are stabilized by a conserved pair of residues that form an intersubunit salt bridge. The absolute conservation of these residues is striking, as only one other residue (Ala158, located in the hydrophobic core) is completely conserved across the Csa5 family (Fig. S2). Mutation of the salt bridge both in the paralog and in Csa5 itself results in a profound change in the biophysical properties of the proteins. In both proteins, the mutant is smaller and less prone to aggregation. Since the change is seen in both proteins and the salt bridge is conserved, we hypothesize that the interface (but not the infinite threads) has a functional significance that will be revealed when the structure of the εCascade complex is determined.

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In a recent bioinformatics study, Makarova et al. suggested that several small Cas proteins, which were predicted to be largely helical, could be combined into a new family (Cas11). The putative Cas11 family has been proposed to comprise Csa5 (type I-A), Cse2 (type I-E), Csm2 (type III-A) and Cmr5 (type III-B) as well as the C-terminal helical domain of Cas8 (various subtypes). Crystal structures are now available for three of these protein families: Csa5 (Sso1398), Cse2 (TphCse2 and TfuCse2) and Cmr5 “T. thermophilus” (PDB 2ZOP) and Archaeoglobus fulgidus (PDB 2OEB),
allowing an assessment of these predictions. The α-domain of Sso1398 is indeed homologous to the C-terminal domain of Cse2 (Fig. 5) but neither it nor the β-domain bears any resemblance to either Cmr5 structure. However, the N-terminal domain of Cse2 is homologous to four helices of Cmr5 (Cα rmsds of 3.2 Å over 50 residues and 3.0 Å over 60 residues to AfuCmr5 and TthCmr5, respectively) (Fig. 5). Sequence conservation between the structures is limited to residues of the hydrophobic core; the lack of conservation in exposed residues would argue against a common function. This is supported by the different properties of Csa5 and Cse2 combined with the observation that Cmr5 is not essential for the function of the CMR complex, whereas loss of Cse2 significantly impacts the activity of eCascade.27

Analysis of the structures reveals a nuanced set of relationships among the small subunits. Of the two domains of Cse2, one is found in the Cmr5 family and the other in the Csa5 family; there is no structural relationship between the Cmr5 and Csa5 families. This is consistent with either selective loss of one or other domain from the ancestral Cse2 (Cas11) family during the diversification of the CRISPR/Cas systems to form the Cmr5 and Csa5 lineages, or alternatively a fusion of the Cmr5 and Csa5 families to create the Cse2 lineage as the Cmr5 and Csa5 proteins have nothing in common beyond a potential shared ancestor in Cse2. To further elucidate the evolutionary relationships between the so-called small subunits of the interference complexes, the structure of Csm2 would be particularly helpful, as it is a key remaining missing piece. The full-length structure of Cas8 is also likely to be useful in firmly establishing such relationships.

Materials and Methods

Cloning. sso1398 was cloned into the pDEST14 plasmid and sso1443 into pET151/D-TOPO (Invitrogen) with cleavable N-terminal His6-tags. Mutations were introduced using published or standard QuikChange (Stratagene) protocols. The oligonucleotide sequences used for cloning and mutagenesis are available on request.

Expression and purification. Sso1398, Sso1443 and their variants were expressed in C43 (DE3) E. coli in LB medium using 1.0 or 0.4 mM isopropyl β-D-1-thiogalactopyranoside (IPTG), respectively, to induce protein expression. The cultures were incubated at 37°C throughout (Sso1398) or 25°C (Sso1443) after induction and the cells harvested after overnight incubation. The selenomethionyl derivative of Sso1398 was expressed in B834 (DE3) E. coli grown in M9 minimal medium supplemented with Selenomethionine Nutrient Mix (Molecular Dimensions) and 50 mgL⁻¹ (L)-selenomethionine. The expression conditions were as with the native protein except the cells were harvested after 30 h. Sso1441 and Sso1442 were co-expressed in E. coli as described previously.19

All proteins were purified using previously published protocols.19,29 Briefly, cell pellets were resuspended and then lysed by sonication (MSE Soniprep) or with a cell disruptor (Constant Systems Ltd.). The lysate was cleared by centrifugation and passed over a Ni-NTA column (Qiagen). The target protein (Cas5/Cas7 or Csa5) and resin were incubated allowing an assessment of these predictions. The α-domain of Sso1398 is indeed homologous to the C-terminal domain of Cse2 (Fig. 5) but neither it nor the β-domain bears any resemblance to either Cmr5 structure. However, the N-terminal domain of Cse2 is homologous to four helices of Cmr5 (Cα rmsds of 3.2 Å over 50 residues and 3.0 Å over 60 residues to AfuCmr5 and TthCmr5, respectively) (Fig. 5). Sequence conservation between the structures is limited to residues of the hydrophobic core; the lack of conservation in exposed residues would argue against a common function. This is supported by the different properties of Csa5 and Cse2 combined with the observation that Cmr5 is not essential for the function of the CMR complex, whereas loss of Cse2 significantly impacts the activity of eCascade.27

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Pull-down interactions. For the production of antibodies, 1 mg of purified recombinant Csa5 or Cas5/Cas7 protein was used to raise polyclonal antibodies in sheep (Scottish National Blood Transfusion Service). The antibodies were affinity purified using cyanogen bromide-activated-Sepharose 4B resin (Sigma). The target protein (Cas5/Cas7 or Csa5) and resin were incubated

Figure 4. The oligomerization of Csa5. (A) Gel filtration elution profiles for Sso1398WT and Sso1398D35A using 2 mgml⁻¹ protein loaded onto a Superose 12 10/300 column. The proteins ran aberrantly on gel filtration and so the masses of the oligomers could not be accurately estimated. (B) Representative radius distributions of Sso1398WT and Sso1398D35A determined by dynamic light scattering.
of which was 100 mM glycine pH 2.5 directly into 1 M Tris pH 8.0. The resin was further washed with 10 mM Tris pH 7.5 and then 10 mM Tris pH 7.5, 0.5 M NaCl. The antibodies were eluted in two stages, the first of which was 100 mM triethylamine pH 11.5 into 1 M Tris pH 8.0. The resin was further washed with 10 mM Tris pH 8.8 and the antibodies then eluted with 100 mM triethylamine pH 11.5 into 1 M Tris pH 8.0. The purified antibodies were dialyzed into PBS and then cross-linked to Protein G Sepharose (Sigma) by incubating together in TBS buffer followed by cross-linking with 20 mM dimethyl pimelimidate in 0.2 M triethanolamine. The beads were washed with trithenaolamine and ethanolamine before overnight in 0.1 M NaHCO$_3$ pH 8.3, 0.5 M NaCl at 4°C. The resin was washed with buffer, resuspended in 1 M ethanolamine pH 8.0 and incubated at 4°C overnight. 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The resin was washed with 0.1 M NaHCO$_3$ pH 8.3, 0.5 M NaCl and then 10 mM Tris pH 4.0 for four cycles followed by incubation with the immune serum in 10 mM Tris pH 7.5 at 4°C overnight. The resin was washed with 10 mM Tris pH 7.5 and then 10 mM Tris pH 7.5, 0.5 M NaCl. The antibodies were eluted in two stages, the first
incubation with *S. solfataricus* P1 cell lysate prepared as described previously.\(^9\) After overnight incubation at 4°C, the beads were washed with TBS and the bound protein eluted with sodium citrate pH 2.5. Samples were run on NuPAGE SDS-PAGE gels (Invitrogen) and blotted onto iBlot stacks (Invitrogen). The samples were blocked with 5% milk powder in PBS, 0.05% TWEEN 20 for 1 h and then incubated with the primary antibody of choice. The blots were incubated with 1:10,000 Immunopure rabbit anti-goat HRP (Thermo Scientific) in PBS, 0.1% TWEEN 20 for 1 h and then visualized with SuperSignal West Pico kit (Pierce) according to the manufacturer’s instructions.

**Fluorescence anisotropy.** Synthetic oligonucleotides were purchased from Integrated DNA Technologies of fluorescently labeled crRNA from spacer 26 of CRISPR locus A of *S. solfataricus* P1, non-labeled complementary target ssDNA with PAM sequence and fluorescently labeled non-target ssDNA (see Table S1 for sequences). crRNA and protospacer were mixed 1:1, heated at 80°C for 5 min and cooled slowly to form a heteroduplex. Fluorescence anisotropy measurements (\(\lambda\_\text{ex} 490 \text{ nm}, \lambda\_\text{em} 535 \text{ nm}\)) were recorded using a Varian Cary Eclipse fluorescence spectrophotometer at room temperature. Protein solutions were titrated into 20 nM crRNA, 20 nM non-target ssDNA or 20 nM heteroduplex in 20 mM Tris pH 7.5, 150 mM NaCl, 5% glycerol, 1 mM EDTA. Measurements were recorded over a range of protein concentrations (0.04 –11 \(\mu\text{M}\)). The data were analyzed using KaleidaGraph (Synergy Software) and, where appropriate, fitted to binding curves.

**Crystallization of Sso1398.** The selenomethionyl derivative of Sso1398 was crystallized at 2 mg ml\(^{-1}\) in vapor diffusion sitting drop experiments at 20°C. The crystals grew in a condition of 1:1 protein-to-precipitant with a reservoir of 0.1 M sodium acetate pH 4.0, 0.23 M potassium chloride and 39% PEG500. The crystals were cryo-cooled in liquid nitrogen directly from the drop. A single-wavelength anomalous diffraction data set was collected at the Se-K absorption edge at 100 K on the microfocus beamline I24, Diamond Light Source. The data were processed with xia\(^2\) and the phases solved using Phenix AutoSolve.\(^{32}\) The electron density map was further modified using Parrot\(^33\) before chain tracing with Buccaneer,\(^34\) both as part of the CCP4 suite.\(^35\) The chains were manually assigned followed by automated model building with Phenix AutoBuild.\(^32\) The model was refined by cycles of manual correction in COOT\(^9\) and refinement with REFMACv5.6\(^37\) using TLS constraints generated using the TLSMD web server.\(^38,39\) The structure was validated with MolProbity.\(^40\)

**Table 1. Data collection and refinement statistics**

| Data collection | Sso1398\(^{\text{Selen}}\) | Refinement | Sso1398\(^{\text{Selen}}\) |
|-----------------|--------------------------|------------|--------------------------|
| Wavelength (Å)  | 0.98                     | \(R\_\text{work}/R\_\text{free}\) | 0.20/0.22 |
| Space group     | C 1 2 1                  | Mean B-value (Å\(^2\)) | 43 |
| \(a, b, c\) (Å) | 285.0, 337.7, 216.7      | All atoms | 43 |
| \(\alpha, \beta, \gamma\) (°) | 90.0, 124.7, 90.0 | Protein | 43 |
| Resolution (Å)  | 2.72                     | Water      | 26 |
| I/\(\sigma\)     | 12.3 (2.3)               | PEG        | 55 |
| \(R\_\text{merge}\) | 0.109 (0.814)          | Rmsd       | |
| Completeness    | 100.0 (100.0)            | Bond lengths (Å) | 0.01 |
| Multiplicity    | 6.7 (7.0)                | Angles (°) | 1.13 |
| Anomalous       | 99.5 (99.8)              |            | |
| Anomalous       | 3.5 (3.6)                |            | |

Data are presented as averages with statistics for the highest resolution shell in parentheses.

**Dynamic light scattering.** DLS experiments were performed using a Zetasizer \(\mu\text{V}\) system (Malvern Instruments) with a quartz SUPRASIL\(^6\) cuvette. The experiments were performed at room temperature and in triplicate. The data were analyzed using the Zetasizer software.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

The data and coordinates of the structure of Sso1398 were deposited in the Protein Data Bank under the accession code 3ZC4.

**Supplemental Material**

Supplemental material may be found here: www.landesbioscience.com/journals/rnabiology/article/23854

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