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Crystal structures of three representatives of a new Pfam family PF14869 (DUF4488) suggest they function in sugar binding/uptake

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Abstract: Crystal structures of three members (BACOVA_00364 from *Bacteroides ovatus*, BACUNI_03039 from *Bacteroides uniformis* and BACEGG_00036 from *Bacteroides eggerthii*) of the Pfam domain of unknown function (DUF4488) were determined to 1.95, 1.66, and 1.81 Å resolutions, respectively. The protein structures adopt an eight-stranded, calycin-like, β -barrel fold and bind an endogenous unknown ligand at one end of the β -barrel. The amino acids interacting with the ligand are not conserved in any other protein of known structure with this particular fold. The size and chemical environment of the bound ligand suggest binding or transport of a small polar molecule(s)

Abbreviations: ALS, Advanced Light Source; JCSG, Joint Center for Structural Genomics; LB, Luria-Bertani broth; MAD, multi-wavelength anomalous diffraction; MES, 2-(N-morpholino)ethanesulfonic acid; MgSO₄, magnesium sulfate; NCS, non-crystallographic symmetry; NaCl, sodium chloride; PCR, polymerase chain reaction; PIPE, polymerase incomplete primer extension; SAD, single-wavelength anomalous diffraction; SSRL, Stanford Synchrotron Radiation Lightsource; TCEP, tris(2-carboxyethyl)-phosphine; TEV, tobacco etch virus; TLS, translation, libration, screw; Tris, tris(hydroxymethyl)aminomethane; UNL, unknown ligand.

Additional Supporting Information may be found in the online version of this article.

Bacteroides are Gram-negative anaerobic bacteria that inhabit the mammalian gastrointestinal tract and comprise a significant portion of the human gut microbiome. Here we report the crystal structures of three homologous proteins from *Bacteroides*. An unknown ligand is bound in the same location in each of the structures. The general nature of the ligand and its interactions with the protein indicate that it is a small polar, ring-like molecule, which suggests a binding/acquisition/transport like function for this protein.

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as a potential function for these proteins. These are the first structural representatives of a newly defined PF14869 (DUF4488) Pfam family.

Keywords: *Bacteroides*; DUF4488; sugar binding; calycins; unknown ligand; crystal structure

Introduction

Bacteroides are anaerobic, bile-resistant, non-spore forming, Gram-negative bacteria that inhabit the mammalian gastrointestinal tract. *Bacteroides* comprise nearly 25% of the 10^{11} organisms per gram of content typically found in the human gut.¹ These bacteria maintain a commensal or mutualistic² relationship with the host, playing a fundamental role in the processing of complex nutrients into simpler ones that can be readily processed by the host.^{3–5} However, some species can cause disease, such as sepsis, abscess formation in multiple organs, and bacteremia, when they escape the host intestine.⁴ Genomic and subsequent proteomic analyses of two *Bacteroides* species, *B. thetaiotaomicron*, and *B. fragilis*, reveal that a significant proportion of their genome is dedicated to nutrient-sensing and nutrient-metabolizing machinery, mainly carbohydrate degradation/acquisition/utilization systems.^{6–8} For example, the *B. thetaiotaomicron* genome encodes 172 glycosyl hydrolases and 163 starch-binding proteins (SusC and SusD homologs), which are involved in the breakdown of complex polysaccharides.^{3,6} As part of our efforts to explore and complement genomic studies of over-represented protein families in the human gut microbiome and expand the structural coverage of these proteins, the Joint Center for Structural Genomics (JCSG) has to date determined structures of 239 of a total of 544 *Bacteroides* protein structures in the PDB as of May 2014. Here, we report crystal structures of three homologous proteins of unknown function, BACOVA_00364 (ZP_02063416.1) from *Bacteroides ovatus* (*B. ovatus*), BACUNI_03039 (ZP_02071597.1) from *Bacteroides uniformis* (*B. uniformis*) and BACEGG_00036 (ZP_03457270.1) from *Bacteroides eggerthii* (*B. eggerthii*). These proteins share a sequence identity of 80% and are conserved in at least 22 of the 33 known *Bacteroides* species (<http://www.ncbi.nlm.nih.gov/genome>). The structures reveal an eight-stranded, β -barrel fold with a putative ligand binding site located at one end of the barrel, reminiscent of a group of proteins known as calycins. Interestingly, an unknown ligand (UNL) is bound at this site in each of the three structures. The nature of the ligand and its interactions with the proteins suggest that these proteins bind small polar molecules, such as carbohydrates, thereby indicating a possible function in nutrient binding/acquisition/transport. Further analysis of these structures indicated that they belong to a separate protein family and resulted in the creation of a new Pfam family, PF14869 (DUF4488).

Results

Overall structure

The crystal structure of BACOVA_00364 contains eight protein molecules (residues 25–163 in chain A, 31–163 in chain B, 28–163 in chain C, 31–163 in chain D, 23–163 in chain E, 28–163 in chain F, 28–163 in chain G, and 28–133, 140–163 in chain H), one sodium ion, one acetate ion, four glycerol molecules, five UNLs and 555 water molecules in the crystallographic asymmetric unit (asu). The BACUNI_03039 structure contains four protein molecules (residues 23–163 in chain A, 28–163 in chain B, 28–163 in chain C and 31–163 in chain D), one glycerol molecule, five polyethylene glycol fragments, four UNLs and 628 water molecules in the asu. The BACEGG_00036 structure contains two protein molecules (residues 27–163 in chain A and 27–163 in chain B), two UNLs and 293 water molecules in the asu. (n.b. possible oligomeric assemblies relevant for biological function *in vivo* are discussed below). A few residues at the N-terminus of most chains in BACOVA_00364 and BACUNI_03039 were not modeled in the structures due to lack of interpretable electron density. The Matthews' coefficient (V_M)⁹ and the estimated solvent content are 2.36 Å³/Da and 47.8% for BACOVA_00364, 2.73 Å³/Da and 54.9% for BACUNI_03039, and 2.69 Å³/Da and 54.3% for BACEGG_00036, respectively. The Ramachandran plots produced by MolProbity¹⁰ show more than 98.0% of the residues are in favored regions with no outliers for all three structures.

All structures adopt a β -barrel fold, comprised of eight anti-parallel β -strands and one 3_{10} -helix (residue 105–109) (Fig. 1). BACOVA_00364 contains an additional 3_{10} -helix at the N-terminus in one of its chains. A UNL is bound at the more open end of the β -barrel in all three structures.

Similarity among the three structures

As expected, the structures of the three proteins are essentially identical [Fig. 2(A)]. A multiple structure alignment of the proteins by EBI-SSM server¹⁴ returns an overall C α atom RMSD of 0.84 Å, overall Q-score of 0.91, and sequence identity of 80% using 135 equivalent residues in the alignment [Fig. 1(B)]. The presence of an additional 3_{10} -helix at the N-terminus and a different orientation of this region in BACOVA_00364 in one chain (likely due to crystal packing interactions that influence the local structure of this region in this chain but not the others) and some variations in loop orientations constitute

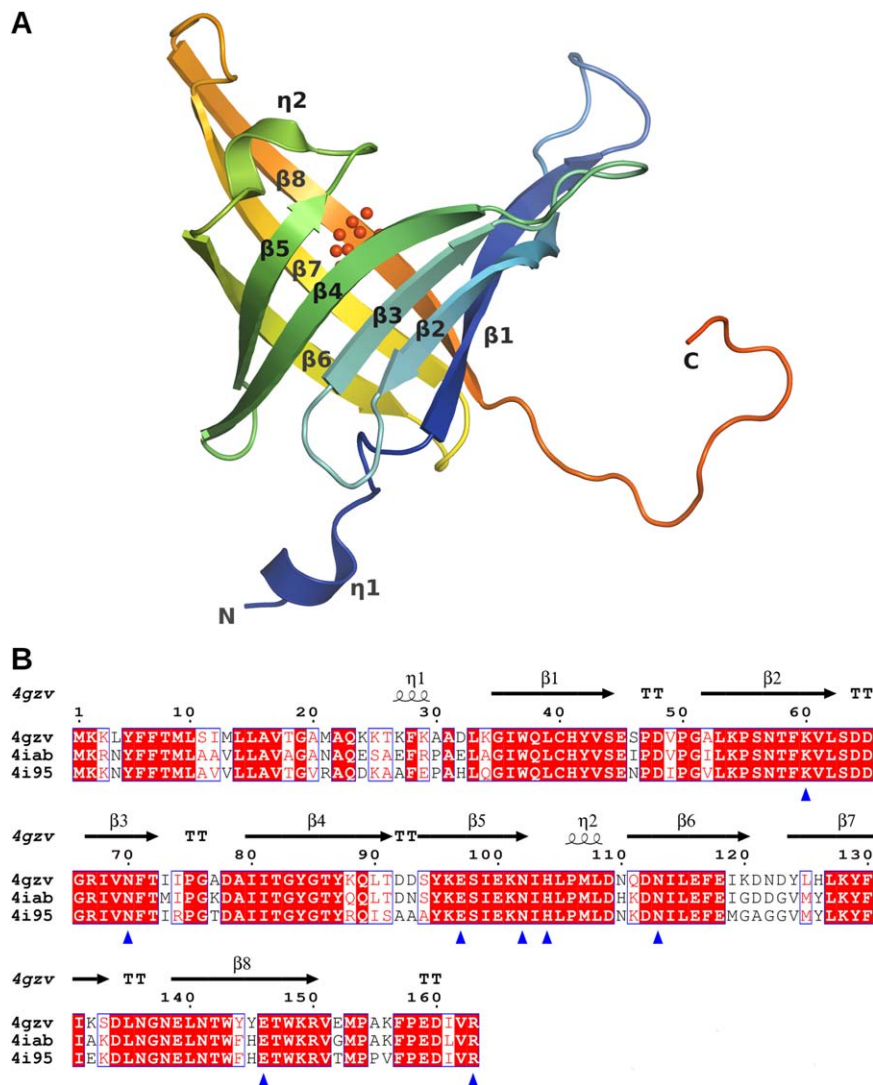


Figure 1. Overall structure and sequence alignment of the PF14869 monomer. (A) The eight-stranded β -barrel fold of the BACOVA_00364 protomer (pdb code 4gzv, chain A) is shown in a rainbow cartoon representation from blue to red with the secondary structure elements labeled. The N- and C-termini are also labeled. BACOVA_00364 has an additional 3_{10} -helix near the N-terminus that is absent in BACUNI_03039 (pdb code 4iab) and BACEGG_00036 (pdb code 4i95). The UNL is shown as red spheres. This and other figures were prepared with PyMOL.¹¹ (B) The sequences were aligned using ClustalW¹² and rendered using EsPrint.¹³ The secondary structural elements corresponding to the 4gzv structure are identified at the top of the sequence. Identical residues are in bold white font on a red background while similar residues are in red font against a white background. The residues interacting with the ligand are indicated by blue triangles.

the only structural variability among these three proteins [Fig. 2(B)].

Ligand binding site

A putative ligand binding site consisting of a small cavity lined by conserved polar residues (Lys60, Asn70, Glu97, Asn102, His104, Asn113, and Glu146) was initially identified based on the presence of unaccounted for electron density, as illustrated in Figures 1(B) and 3(A). The volume of this cavity calculated with the CASTp server¹⁵ ranges from 235 to 250 Å³, depending on the specific protein and chain. A virtual library screen containing 144,110 small molecules of less than 20 non-hydrogen atoms identified many potential ligands, with the highest scor-

ing candidate (binding affinity of -6.9 kcal/mol) 4,8-dihydro-6H-1,2,5-oxadiazolo[3,4-e]1,2,3-triazolo[4,5-b]pyrazine. However, closer inspection of the top 100 hits did not identify any ligand with a good fit to the electron density. Thus, a UNL was modeled at corresponding sites when warranted by the density (five out of eight chains in BACOVA_00364 and all chains in BACUNI_03039 and BACEGG_00036); a glycerol molecule was modeled at this site in the other three chains of BACOVA_00364 as this gave the best fit to the electron density.

Interestingly, the C-terminus (Arg150–Arg163) reaches over and inserts its tail into the putative ligand binding site pocket of the adjacent molecule in the tetramer and the terminal Arg163 residue

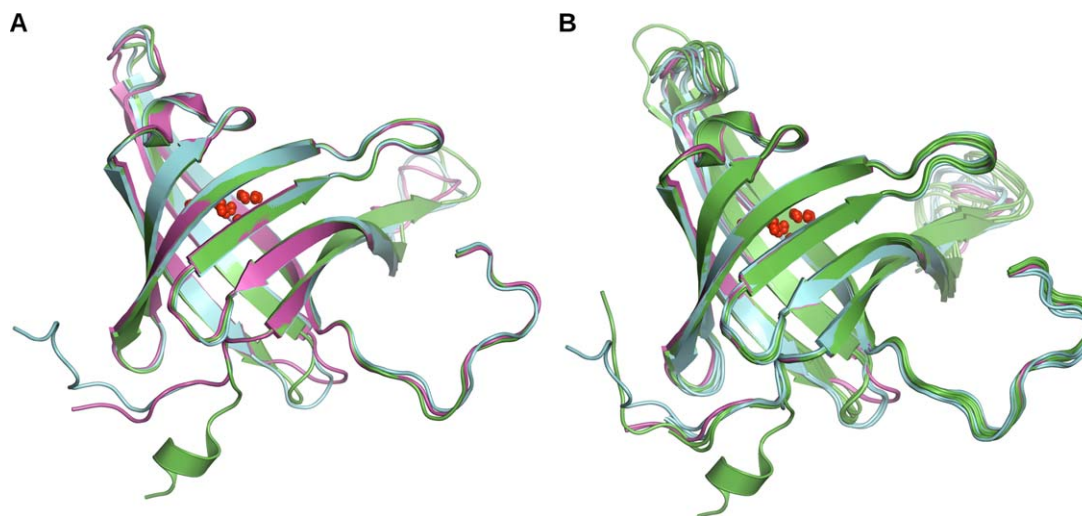


Figure 2. Structural comparison of BACOVA_00364, BACUNI_03039, and BACEGG_00036. (A) Cartoon representation of the superposition of the three structures (chain A) showing the high structural similarity with only minor differences at the N-terminus and a couple of loops. BACOVA_00364 is in green, BACUNI_03039 in cyan and BACEGG_00036 in magenta. (B) The structural similarities and differences are highlighted by a superposition of all chains in the asu in all three structures.

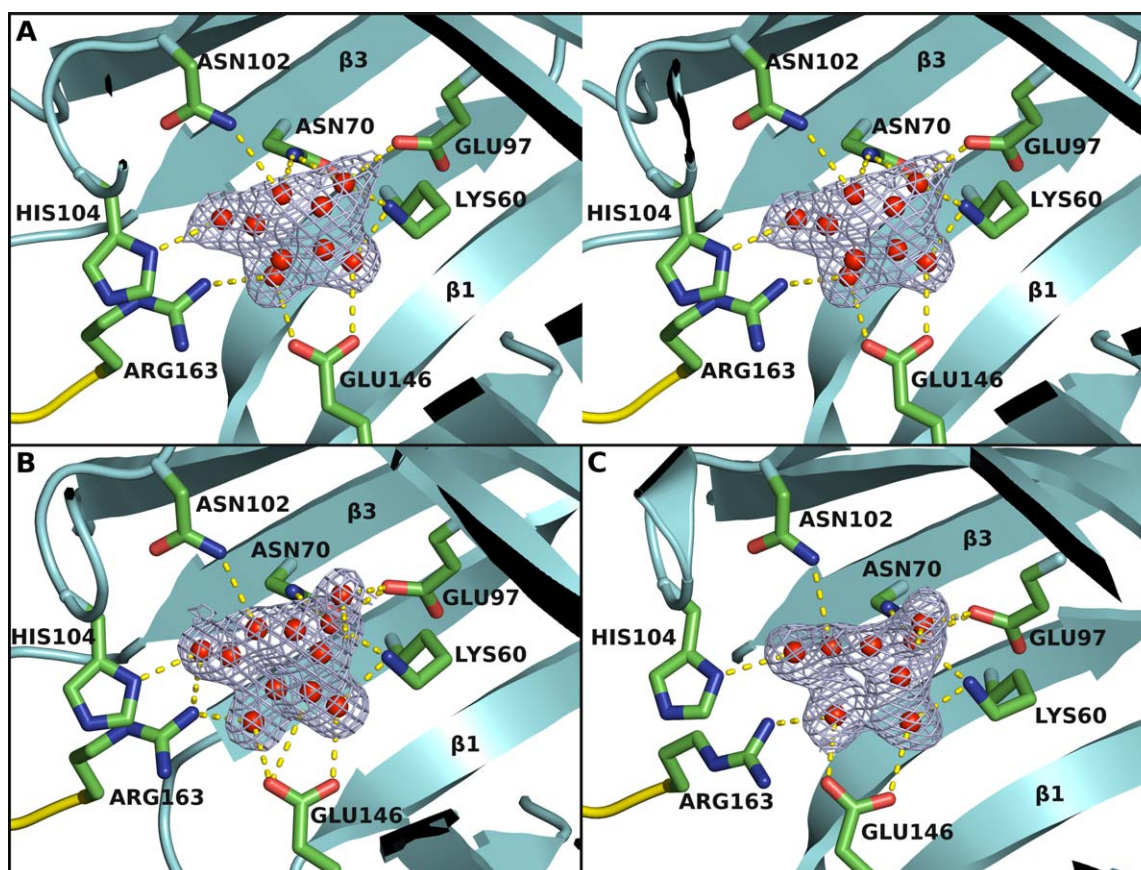


Figure 3. The ligand binding site. The binding site is located at one end of the β -barrel as identified from binding of a UNL (red spheres). (A) Stereo view of the UNL and its interaction with the protein in BACOVA_00364 (pdb code 4gzv). Omit map is contoured at 1.25σ level above the mean density. The residues interacting with the UNL are shown in stick representation. Arg163 that interacts with the UNL comes from an adjacent protomer in the biological tetramer. (B) and (C) are the corresponding UNL binding sites for BACUNI_03039 (pdb code 4iab) and BACEGG_00036 (pdb code 4i95), respectively, where the same or similar UNL is bound.

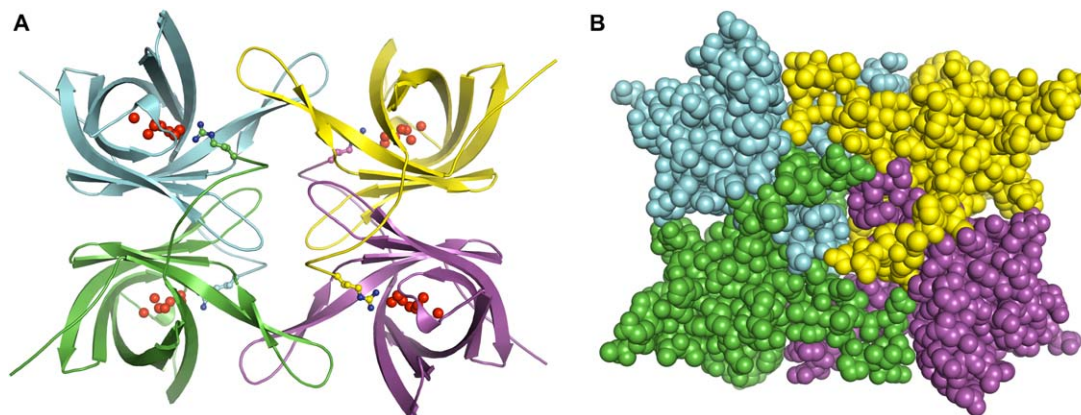


Figure 4. Putative biologically relevant oligomeric state. The proteins associate as a tetramer (dimer of dimers) shown in cartoon (A) and space-filling (B) representations and colored by chain in all three structures even when crystallized in different space groups. The UNL is shown as red spheres. Arginine 163 from a neighboring protomer that interact with the UNL is shown in ball and stick representation.

interacts with the UNL (Fig. 3). The shape of the UNL and the residues interacting with it suggest the same small molecule is bound in all three proteins. As nothing similar was present in the purification or crystallization reagents, this compound likely originated from the bacterial growth medium used during protein expression. Despite the possibility that this may not be the natural substrate, the ligand shape, interatomic distances, and chemical environment suggest a ribose/erythofuranose derivative, thus hinting at a sugar binding/transport/uptake-like function for these proteins. However, we cannot exclude the possibility that the Arg163 side chain could be acting as a partial surrogate for a larger substrate and could be displaced in the presence of the natural ligand.

Oligomeric state

Analytical size exclusion chromatography indicates BACEGG_00036 is a tetramer in solution while BACOVA_00364 and BACUNI_03039 are mixtures of monomers and dimers. Crystal packing analysis of all three proteins using the PISA server¹⁶ suggests that the proteins most likely exist as a tetramer (dimer of dimers; Fig. 4), with dimeric arrangements predicted as well. For the tetrameric association, the total buried surface area and change in solvent free energy are 11,810 Å² and -70.0 kcal/mol for BACOVA_00364, 14,680 Å² and -66.0 kcal/mol for BACUNI_03039, and 13,110 Å² and -60.0 kcal/mol for BACEGG_00036. The proteins' biologically relevant oligomeric state must be at least dimeric since the C-terminus (Arg163) of one protomer forms an integral part of the active site of the neighboring protomer (Fig. 3). This dimeric interface buries a total surface area of 1967 Å² in BACEGG_00036 (the other dimeric interface between two subunits buries a surface area of 322 Å²). However, the tetrameric association seems prob-

able because of the conservation of residues at the oligomeric interface and the fact that this arrangement is conserved in the different space groups (P1, C222₁, P6₃22) in which these proteins were crystallized.

Sequence and structural similarities to proteins in UniProtKB and in the Protein Data Bank

BACOVA_00364, BACUNI_03039, and BACEGG_00036 are members of the Pfam family DUF4488 (PF14869).¹⁷ We ran the most recent version of the PF14869 family HMM (Supporting Information "PF14869-HMM.docx") against the UniProtKB¹⁸ (March 2014 version) using the hmmer website (<http://hmmer.janelia.org/>) (hmmsearch with Pfam manually curated significance bit score thresholds of 27.0). This analysis identified 146 distinct, significant matches and uncovered an overlap with a region assigned by Pfam to the lipocalin_5-PF13924 family in the calycin clan (CL0116), indicating PF14869 may also belong to this clan (note: PF14869 [Pfam release 27.0 March 2013] contains only 63 sequences and no clan assignment, but this information will be updated in the next release). Except for four regions found in proteins from unclassified organisms (from metagenomics data), all other matches were with proteins from the phylum Bacteroidetes [Fig. 5(A)]. Most of these appear to be single-domain PF14869 proteins [Fig. 5(B)]. However, in 19 sequences, the PF14869 domain is found downstream of a domain that is a member of the TonB_C-PF03544 family. TonB_C is generally located at the C-terminus of TonB, a protein mostly involved in iron transport in Gram-negative bacteria. In one instance, F9YU00 (UniProt Id: F9YU00_CAPCC), the PF14869 domain is found upstream of a peptidase M60-like-PF13402 domain and, in another, R5PRP8 (UniProt Id: R5PRP8_9BACT), downstream of a FGE-sulfatase-PF03781 domain.

Indeed, structural comparisons revealed further similarities between the three structures presented here and members of PF14869 family, and

A

UniProtKB id	Position	Identity (%)	Sequence
A7LRD6_BACOV	32 163	100.0	-DLKGIWQLGHYVSESP.DVPGALKPSNTFKVLSDDGRIVNFTI-...IPGDAIIT.GYGYKQLTDDSYKESIEKN
U2DU09_9BACE	32 163	79.5	-ELKGIWQLGHYVSEAA.DLPGKLPKSNFTKVLSEDDGRIVNFTI-...IPGSDAIIT.GYGAVQLSDITTKESIEKN
E6SSK7_BACT6	32 163	78.8	-NLAGIWQLGHYVSETP.DVPGTLKPSNTFKVLSDDGRIVNFTI-...RPGSDAIIT.GYGYQOISDYAYKESIEKN
R7NSA5_9BACE	33 164	61.4	-DLKGIWQMGFYVSGVP.NTPGELKPSNSFKVLSDDGKFTNMTV-...IPNHGAIIT.GAGTYKQTAENAYTEHVEKN
S8GW95_9BACT	1 126	58.4	-----MCFYVSASP.DMIGELKPSNSFKVLSDDGKFTNMTV-...IPNRGAIIT.GSGTYEISSDSIYVEHVEKS
R6CHJ7_9BACE	32 163	55.0	-TLSGIWQMGFYRSSQP.GIPGELKTGNTLKLSDDGGRFNSVVM-...MP-TGAVII.GYGYTIDSDSTYVENVEKN
R6S3Z0_9BACE	32 162	55.0	-DLRGWQMGFYRSDAP.GVAGQLKTSNSLKLSDDGKFTNLMV-...MP-QGAIIT.GEGTYEQTSGDSYTEMVEKN
R6YI88_9BACE	33 164	54.2	-TLAGWQMGFYRSASP.DVPGELKTSNSLKLSDDGHTNIVM-...MQ-TGAIIT.GYGDYILKSPGVYTEVEIKN
B3JJX4_9BACE	32 163	50.4	-TLNGWQMGFYRSSSP.NVVGELKTGNTLKLSDDGGRFINLVM-...MQ-TGAIIT.GYGYKQESNSYTEYVEKN
R7A830_9BACE	32 161	48.1	-TLHGWQMGFYRSNSP.DIPGELKTSNSLKLSDDGHTNLLM-...MQ-TGAVIL.GYGYEINNEGLTYEYVEKN
F0R2C8_BACSH	32 160	44.1	ESLKGWQMGFYSSSLP.ALPGELKPSNSLKLGSSEGEFTNVVM-...MP-AGAIIT.GSGTYTLNSDSTYTEHVKN
I9ESJ6_9BACE	67 188	31.4	-SLEGWQIGTHVEPLG.YGKFDIQTGPYMKLSDGKFFNIHLA...ITDERAVIT.AMGTYKVTSDCTYVEKIFKS
R7PAB7_9BACT	46 162	31.0	-KLCGIWQMGFYISIEK.NK-PVVHFTPYMKALNSDGKFFVNMGLE...TGRSCCFIA.TMGESIVSDSVYVEHIDRS
I3ZZK4_ORNRL	30 142	30.4	KNLVGIVQHV-YPIISG.NY----IKTGNYSITPDGTFSLVFI-...-GKNKTTLT.GYGYKITSDSSTFTKEMISH
J9FYR7_9ZZZZ	2 88	30.2	-----QDGSFCTFLIA...NQSCKSIIT.NEGTYKVTSDSTYVEHVTGS
R5P981_9PORP	81 199	29.7	-TLQGIWQIGRLNNTSD.NEKYSITAGPYMKLSDGKFFNIHLA...TSGHVSIVT.SNGTYTQSDSTYVESVSES
I3ZZK3_ORNRL	31 148	29.2	-DLVGFWQQR-YIIQTA.NGEKLLKSSGNYKVINPDGTYTFMIVaprnSDIQVPIIL.QYGYELIDDTFHNEHIIHD
F9YU00_CAPCC	10 103	29.0	-----MFKVFFONGTFRNYMFG...LP--KSIIT.TKGYKVTSDSTYVEHVTGS
R9I4T1_9BACE	55 168	28.3	HELAGIWQIG--LLEQK.ADGVHLKLLPVMKFLSPDQTFVNVLA...-ANNRIVVT.NQGYTYKSEFLYVEEILKT
R5PRP8_9BACT	292 402	27.8	-DLAGWQYI----SF.DANGKRKYHVALKFLNADGTFQNLQFS...QSGNGQIMYKAGTWWLKL-DGCTIVQYKYG
R7LJK2_9BACT	27 139	26.8	-KLCGIWQVQV---QQA.KDGSRVVRLPVMKFLSDGKFFNIHLA...NEQAKSIIT.NRGEYKVTSDSTYVEHVTGS
G5GA52_9BACT	26 139	24.8	-PLLGWQOQEM-ITQEG.NR---ILIPVWKLISDGTFVSVILMT...NKDGRVTQT.IEGSYVINSQNTYTEOVKTN
F9YVZ3_CAPCC	33 150	24.3	-SVVGVWQVGLERMG.EV--KFIPLPMKLYNEDGTFVLRMF...EKSSFFAIT.LYGNVEISEDGMKEYIKGT
A5ZK07_9BACE	157 278	24.2	KRMQGWQVTC--MVESV.EKGYRLSLGPKLKFISVDNFMNIID...TNKMGSVIL.VQGEYKVTSDSTYVEHVTGS
S7VP50_9FLA0	112 221	24.1	-KLOGAWLMSGRVRDgkTETRDTSRPRKTKLISG-TRFOWIAYN...TETKQFMGT.GGGYTYT-INGVYTESIEFF
I3ZZM5_ORNRL	36 149	22.8	--LAGFWQIMPYDYQG.DA--RIMRSGQKIMNPDGTFYCFLT...DNKGVQGIS.FYGNVVTSDSTYVEHIVNS
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G6IPT9_9BACE	159 281	22.5	-SLQGWQSC--VVQPG.EHGKILLPVLKLVSPDQTFMNIIMTAg.mNGRSNAIIF.CQGEYKVTSDSTYVEHVTGS
E6SUV5_BACT6	157 278	22.5	KTMQGWQVTC--RVDSV.ELGYISLLPVLKLVSDGNTFMNIIVQ...VEMGGSMIL.AQGEYKVTSDSTYVEHVTGS
C3QY78_9BACE	62 180	22.2	KSLEGWQVTC-EVAKDS.ADNYIVTSGNALKVISADKTFKNLLLS...AGSANSTIF.MEGNYKVTSDSTYVEHVTGS
R5GJA6_9BACT	47 138	17.6	-----YKLTIEPDGGRFLNLPAD...EATGIYYS.RQGYTLEGNHYVETITHE

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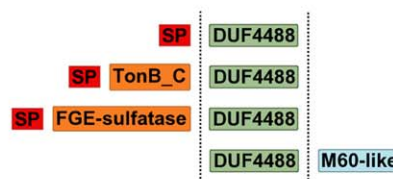
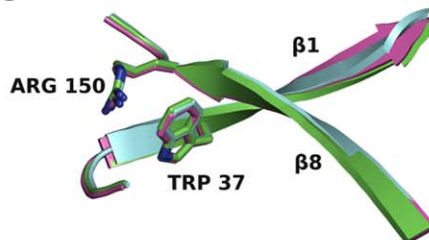
B**C**

Figure 5. Sequence alignment, family architecture and calycin signature residues in the DUF4488/PF14869 Pfam family. (A) Multiple sequence alignment of DUF4488/PF14869 Pfam family members using a redundancy cutoff of 80% sequence identity. For each sequence, we report the UniProtKB id, the position of first and last residue in the alignment, the percent sequence identity with respect to A7LRD6_BACOV (i.e., BACOVA_00364) and the amino acid sequence. Shades of gray in alignment columns reflect average similarity at a given position as calculated from the BLOSUM62 amino acid substitution matrix (black most conserved, white least conserved). Dashes (-) represent deletions. Lower case letters represent inserted residues and dots (.) in the same columns are fillers for sequences lacking the inserted residues. The red boxes mark the position of residues involved in the calycin motif (GxW/R). Alignment visualized with Belvu (<http://sonnhammer.sbc.su.se/Belvu.html>). (B) General architecture of DUF4488/PF14869 family members. Out of 146 members, 125 contain a single DUF4488 domain; 19 also contain a TonB_C domain, one contains a M60-like domain, and one a FGE-sulfatase domain. 'SP' indicates a predicted signal peptide (according to PHOBIUS¹⁹). (C) Superposition of strand 1 and strand 8 of the β -barrel of 4gzv (green), 4iab (cyan), and 4i95 (magenta) in cartoon representation, highlighting the calycin motif residues tryptophan and arginine packing against each other.

a group of proteins known as calycons,²⁰ which include lipocalins, streptavidins/avidins, triabins, fatty acid-binding proteins and metallopeptidase inhibitors (part of the IK MEROPS clan²¹; referred to simply as IK inhibitors from now on). These are all characterized by a calyx-like β -barrel constituted from eight strands (10 strands in fatty acid-binding proteins) with an up-and-down +1 topology (except for triabin, which features a strand swap). The shear number, a measure of the barrel stagger,²² is 12 for lipocalins, triabin and fatty acid-binding proteins, whereas streptavidins, avidins and IK inhibitors have a shear number of 10, reflecting a generally straighter β -barrel. Accordingly, the structural classification database SCOP²³ puts streptavidins, avidins, and IK inhibitors in separate folds (*streptavidin-like* fold versus *lipocalins* fold). Pfam groups the lipocalins, fatty acid-binding proteins and triabin into the calycin-CL0116 clan, IK inhibitors into the bBprotInhib-CL0354 clan, whereas avidins and streptavidins are in a family that is not assigned to any clan (Avidin-PF01382).

Although calycons are generally very diverse at the sequence level, most feature a conserved signature motif, typically Gx[WY]/[RK]. The first part of this motif (Gx[WY]) is located on the first strand of the β -barrel, whereas the second ([RK]) is located on the last strand. Typically, the positively charged residue located on the last strand of the β -barrel interacts with an aromatic residue on the first strand engaging in a cation- π interaction.^{24,25} The importance of this motif in β -barrel stability has been demonstrated.^{26,27} Conservation of this motif and other structural elements are often key to identifying calycons. All members of Pfam family DUF4488-PF14869 feature the calycin signature motif (GxW/R), except for a few proteins that appear to lack the N-terminal portion of the domain altogether (F9YU00 and A5ZKH0) and, thus, only have the arginine residue [Fig. 5(A)]; one protein (S7VPS0) lacks the final arginine. Additionally, four members in the family have the arginine residue substituted with lysine. Other conserved residues in the Pfam alignment [Fig. 5(A)] are identified with important structural/functional roles by mapping the sequence to the structure. For example, Lys60, Asn70, Glu97, Glu146 interact with the UNL, Gln38, Ile81, Val151 are involved in oligomerization, while Tyr95, Tyr129, Trp148 line the binding cavity (without making direct interaction with UNL). Interaction between the tryptophan and the arginine is observed in all three of our structures [Fig. 5(C)].

A structural similarity search using DALI²⁸ provides several significant hits. All annotated proteins at the top of the list are calycons. These include, among others, bilin-binding protein 1bbp (Z-score 8.3), apolipoprotein D 2hzq (Z-score 7.6) and retinol-

binding protein 1fem (Z-score 7.1) (all lipocalins), avidin 1avd (Z-score 7.1) and fatty acid-binding protein 1o8v (Z-score 7.0). Although the location of the UNL in our structures of PF14869 members coincides with that of the ligand in the bilin-binding protein structure (1bbp), the residues interacting with the UNL in our structures are not conserved in any of these other proteins (Fig. 6). The eight-stranded β -barrel in the PF14869 structures has a shear value of 12 (based on the 4gzv structure), suggesting a shape more reminiscent of structures in the *lipocalins* SCOP fold than those in the *streptavidin-like* SCOP fold. Lipocalins feature an additional helix-strand structural motif at the C-terminus [Fig. 6(B)] that is not generally found in other calycons or in the PF14869 proteins.

Discussion

The BACOVA_00364, BACUNI_03039, and BACEGG_00036 structures adopt an eight-stranded, β -barrel fold similar to calycons. Neither sequence nor structure searches reveal any clear function for these proteins despite overall structural similarity to proteins of known function. Likewise, genome context and potential protein-protein interactions analysis (using STRING²⁹ and SEED³⁰) did not provide any clear insight into the potential functions of these *Bacteroides* proteins. Calycin function is generally heavily influenced by loop conformation at the end of the β -barrel and conservation of specific binding residues.³¹ As a consequence, the overall β -barrel structural similarity alone is not sufficient to infer function, as confirmed by the diversity of functions represented within a very small range of Z-scores for the proteins returned by a DALI search. However, the presence of a UNL in the crystal structures helps identify the putative ligand binding site. The size and composition of the binding pocket suggests that the ligand could be a small polar, cyclical molecule, such as a sugar. The proteins likely assemble as a tetramer as indicated by crystal packing analysis in three independent space groups. These proteins provide the first representative structures of a newly defined Pfam family PF14869, (Pfam 27.0, March 2013).

Materials and Methods

Cloning, expression, purification and crystallization

The *B. ovatus* genomic DNA was extracted from cells (ATCC Number 8483) obtained from the American Type Culture Collection (ATCC), that for *B. uniformis* extracted from cells (ATCC 8492) provided by The Human Microbiome Project, and genomic *B. eggerthii* DNA was extracted from cells (DSM 20697) obtained from the DSMZ (The German Collection of Microorganisms and Cell Cultures). Clones were generated using the Polymerase Incomplete Primer

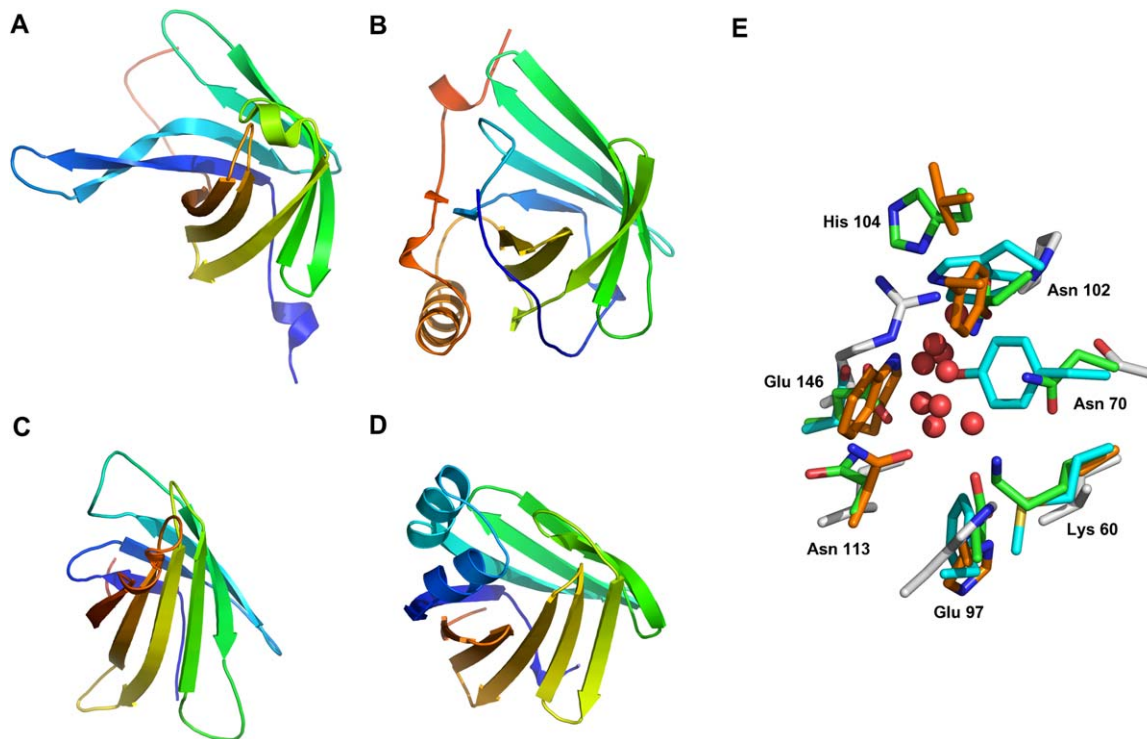


Figure 6. Comparison of the overall architecture and binding site of representative members of the calycin family. (A) BACOVA_00364 (pdb code 4gzv, chain A) in rainbow representation (N-terminus blue and C-terminus red). The structures of (B) Lipocalin (pdb code 1bpp), (C) Avidin (pdb code 1avd), and (D) fatty acid-binding protein (pdb code 1o8v) are shown in same orientation to illustrate the similarity of their overall structures. Lipocalins feature an additional helix+strand structure near the C-terminus. (E) The residues in the binding site are not conserved among members of the calycin family. The residues (green sticks) interacting with UNL (red spheres) in BACOVA_00364 are labeled and the corresponding residues in Lipocalin (orange), 1avd (cyan) and 1o8v (gray) are shown.

Extension (PIPE) cloning method.³² The genes encoding the three proteins were amplified by polymerase chain reaction (PCR) from genomic DNA using *PfuTurbo* DNA polymerase (Stratagene) and I-PIPE (Insert) primers that included sequences for the predicted 5' and 3' ends. The primers used were forward primer, 5'-ctgtacttcagggcCAGAAGAAACC AAATTCAAAGCGGCCG-3'; reverse primer, 5'-aatt aagtcgcttaTCTCACAATATCTTCCGGAAATTTAGCC-3' for BACOVA_00364 from *B. ovatus*, forward primer, 5'-ctgtacttcagggcCAGGAGAGCGCAGAGTTT AGGCCTGCGG-3'; reverse primer, 5'-aattaagtcgct taACGGACAAGGTCCTCGGGGAATTTGCGG-3' for BACUNI_03039 from *B. uniformis*, and forward primer, 5'-ctgtacttcagggcCAGGATAAGGCCGCTTTT GAGCCTGCGG-3'; reverse primer, 5'-aattaagtcgct ttaCCGGACAATGTCTTCCGGAAACACCGGC-3' for BACEGG_00036 from *B. eggerthii*; the target sequences are in upper case. The expression vector, pSpeeDET, which encodes an amino-terminal tobacco etch virus (TEV) protease-cleavable expression and purification tag (MGSDKIHHHHHHENLYFQ/G), was PCR amplified with V-PIPE (Vector) primers (forward primer: 5'-taacgcgacttaattaactgctttaaaggtctccagc-3', reverse primer: 5'-gccctggaagtacaggttttcgtgatgatgatgatgatg-3'). V-PIPE and I-PIPE PCR products were

mixed to anneal the amplified DNA fragments together. *Escherichia coli* GeneHogs (Invitrogen) competent cells were transformed with the I-PIPE/V-PIPE mixture and dispensed on selective LB-agar plates. The cloning junctions were confirmed by DNA sequencing. Using the PIPE method, the gene segment encoding residues M1-A22 were deleted from expression construct to produce soluble protein since it is predicted to contain either a signal peptide using SignalP³³ or transmembrane helices using TMHMM-2.0.³⁴

Expression was performed in a selenomethionine-containing medium at 37°C and selenomethionine was incorporated via inhibition of methionine biosynthesis.³⁵ At the end of fermentation, lysozyme was added to the culture to a final concentration of 250 µg/mL, and the cells were harvested and frozen. After one freeze/thaw cycle, the cells were homogenized and sonicated in lysis buffer [40 mM Tris-HCl, 300 mM NaCl, 10 mM imidazole, 1 mM Tris (2-carboxyethyl) phosphine-HCl (TCEP)-HCl, pH 8.0]. Any remaining nucleic acids were digested with the addition of 0.4 mM MgSO₄ and 1 µL of 250 U/µL benzonase (Sigma) to the lysate. The lysate was clarified by centrifugation at 32,500g for 25 min. The soluble fraction was passed over nickel-chelating resin (GE

Healthcare) pre-equilibrated with lysis buffer, the resin was washed with wash buffer [40 mM Tris-HCl, 300 mM NaCl, 40 mM imidazole, 10% (v/v) glycerol and 1 mM TCEP-HCl, pH 8.0], and the protein was eluted with elution buffer [20 mM Tris, 300 mM imidazole, 10% (v/v) glycerol, 150 mM NaCl and 1 mM TCEP-HCl, pH 8.0]. The eluate buffer was exchanged with TEV buffer [20 mM Tris-HCl, 150 mM NaCl, 30 mM imidazole, 1 mM TCEP-HCl, pH 8.0] using a PD-10 column (GE Healthcare), and incubated with 1 mg of TEV protease per 15 mg of eluted protein for 2 h at ambient temperature followed by overnight at 4°C. The protease-treated eluate was passed over nickel-chelating resin (GE Healthcare) pre-equilibrated with crystallization buffer [20 mM Tris, 150 mM NaCl, 30 mM imidazole, and 1 mM TCEP-HCl, pH 8.0] and the resin was washed with the same buffer. The flow-through and wash fractions were combined and concentrated to ~20 mg/mL by centrifugal ultrafiltration (Millipore) for crystallization trials.

The proteins were crystallized using the nanodroplet vapor diffusion method³⁶ with standard JCSG crystallization protocols.³⁷ Sitting drops composed of 100 nL protein solution mixed with 100 nL crystallization solution (0.1M Tris-HCl pH 8.5, 15% glycerol, 25.5% polyethylene glycol 4000, 0.17M sodium acetate for BACOVA_00364, and 1.0M lithium chloride, 20% polyethylene glycol 6000, 0.1M MES, pH 6.0 for BACUNI_03039 and BACEGG_00036) were equilibrated against a 35–50 μ L reservoir at 277 K for 27–43 days prior to harvest. 20% (v/v) glycerol was added to the crystals of BACUNI_03039 and BACEGG_00036 as a cryoprotectant while no additional cryoprotectant was added to the crystal of BACOVA_00364. Initial screening for diffraction was carried out using the Stanford Automated Mounting system³⁸ at the Stanford Synchrotron Radiation Lightsource (SSRL, Menlo Park, CA).

Analytical size exclusion chromatography

The oligomeric state of the proteins in solution was determined using a 0.8 cm \times 30 cm Shodex Protein KW-803 size exclusion column (Thomson Instruments)³² pre-calibrated with gel filtration standards (Bio-Rad). The mobile phase consisted of 20 mM Tris-HCl pH 8.0, 150 mM NaCl, and 0.02% (w/v) sodium azide.

Data collection, structure solution, refinement

Multi-wavelength anomalous diffraction (MAD) data were collected to 1.95 Å resolution at wavelengths corresponding to inflection, peak, and high energy remote of the selenium edge on beamline BL9-2 at SSRL for BACOVA_00364. A similar MAD dataset for BACUNI_03039 was collected to 1.70 Å resolution at beamline BL8.2.2 at Advanced Light Source (ALS, Berkeley, CA) and a SAD dataset for

BACEGG_00036 was collected to 1.81 Å resolution at beamline BL12-2 at SSRL. The data sets were collected at 100 K using a MAR325 CCD detector (BL9-2) or a Dectris Pilatus 6M pixel detector (BL12-2) with the BLU-ICE data collection environment³⁹ at SSRL and an ADSC Q315 CCD detector at ALS. The data were processed with MOSFLM⁴⁰ and scaled with SCALA⁴¹ for BACOVA_00364 while they were processed with XDS⁴² and scaled with XSCALE⁴³ for BACUNI_03039 and BACEGG_00036. Phasing was performed with SHELXD⁴⁴ and autoSHARP⁴⁵ with a mean figure of merit of 0.27 for BACOVA_00364 (with two selenium sites per protein chain), 0.52 for BACUNI_03039 (with six sites per chain), and 0.33 for BACEGG_00036 (with four sites per chain). Automatic model building was performed with BUC-CANEER.⁴⁶ Model completion and refinement were performed with COOT⁴⁷ and REFMAC.⁴⁸ Experimental phase restraints in the form of Hendrickson-Lattman coefficients from SHARP, NCS restraints (except for BACUNI_03039), and TLS parameters were used during refinement. Data collection and refinement statistics are summarized in Table I.

Virtual library screen

A virtual ligand screen was performed at the putative ligand binding site in chain A of BACOVA_00364 (pdb 4gzv) with AutoDock Vina⁵⁵ against the “ChemBridge_FullLibrary2011” from ZINC.¹² Hydrogens and partial charges were added using MGLTools⁵⁵ and the search limited to a $10 \times 10 \times 10$ Å³ box around the ligand site. Based on the size of the binding site, the docking was limited to library entries ranging from 5 to 20 non-hydrogen atoms. The docking results (docked poses) were visually analyzed in COOT to identify candidates with acceptable fit to the electron density.

Validation and deposition

The quality of the crystal structures was analyzed using the JCSG Quality Control server (<http://smb.slac.stanford.edu/jcsg/QC/>). This server reports the stereochemical quality of the model using AutoDepInputTool,⁵⁶ MolProbity,¹⁰ and Phenix,⁵⁷ the agreement between the atomic model and the data using RESOLVE,²¹ the protein sequence using CLUSTALW,⁵⁸ the ADP distribution using Phenix, and differences in $R_{\text{cryst}}/R_{\text{free}}$, expected $R_{\text{free}}/R_{\text{cryst}}$ and various other items including nomenclature, atom occupancies, consistency of NCS pairs, ligand interactions, special positions, and so forth, using in-house scripts to analyze refinement log file and PDB header. Protein quaternary structure analysis was carried out using the PISA server.¹⁶ Atomic coordinates and experimental structure factors have been deposited in the PDB and are accessible under the codes 4gzv (BACOVA_00364), 4iab (BACUNI_03039), and 4i95 (BACEGG_00036).

Table I. Crystallographic data and refinement statistics for BACOVA_00364 (4gzv), BACUNI_03039 (4iab), and BACEGG_00036 (4i95)

Protein (PDB ID)	BACOVA_00364 (4gzv)			BACUNI_03039 (4iab)			BACEGG_00036 (4i95)
	λ_1 (remote)	λ_2 (inflection)	λ_3 (peak)	λ_1 (inflection)	λ_2 (remote)	λ_3 (peak)	λ_1 (peak)
Data collection							
Space group	P1			C222 ₁			P6 ₃ 22
Unit cell parameters (Å)	$a = 44.78, b = 66.32, c = 109.74,$ $\langle \alpha = 88.2^\circ, \beta = 82.3^\circ, \gamma = 74.8^\circ$			$a = 43.73, b = 213.88,$ $c = 153.64$			$a = 103.07,$ $b = 103.07,$ $c = 114.98$
Wavelength (Å)	0.9116	0.9792	0.9791	0.9795	0.9184	0.9793	0.9795
Resolution range (Å)	29.85–1.95 (2.00–1.95)	29.85–1.95 (2.00–1.95)	29.83–1.95 (2.00–1.95)	46.19–1.66 (1.72–1.66)	46.19–1.70 (1.76–1.70)	46.22–1.73 (1.79–1.73)	48.33–1.81 (1.87–1.81)
No. of observations	250,141	242,093	243,550	306,941	286,554	271,057	1,307,232
No. of unique reflections	86,141	85,819	85,941	84,998	79,136	75,282	33,477
Completeness (%)	97.8 (96.7)	97.4 (95.7)	97.6 (96.4)	99.1 (98.9)	99.0 (98.7)	99.0 (100.0)	100.0 (100.0)
Mean I/σ (I)	10.5 (1.8)	8.1 (1.7)	9.4 (1.6)	15.5 (2.2)	15.0 (2.3)	12.3 (2.1)	15.7 (2.9)
R_{merge} on I^a (%)	7.1 (44.5)	7.4 (45.7)	6.6 (50.7)	4.6 (56.0)	5.0 (53.6)	5.9 (53.8)	19.3 (156.2)
R_{meas} on I^b (%)	8.7 (62.8)	9.2 (64.7)	8.1 (71.7)	5.3 (66.0)	5.9 (63.0)	7.0 (63.3)	19.6 (158.2)
R_{pim} on I^c (%)	4.9 (44.2)	5.3 (45.7)	4.7 (50.7)	2.7 (35.8)	3.0 (34.3)	3.6 (34.4)	3.1 (25.8)
$CC_{1/2}^d$	0.997 (0.572)	0.997 (–) ⁱ	0.998 (–) ⁱ	0.999 (0.732)	0.999 (0.780)	0.998 (0.761)	0.998 (0.907)
Wilson B (Å ²)	27.5	27.6	29.0	21.7	20.6	20.8	21.9
Model and refinement statistics							
Resolution range (Å)		29.85–1.95			46.19–1.66		48.33–1.81
No. of reflections (total) ^e		86,030			84,954		33,450
No. of reflections (test)		4305			4249		1693
Completeness (%)		97.7			99.1		100.0
Data set used in refinement		λ_1			λ_1		λ_1
Cutoff criteria		$ F > 0$			$ F > 0$		$ F > 0$
R_{cryst}^f		0.189			0.163		0.174
R_{free}^f		0.220			0.184		0.204
Ramachandran stats							
Favored (%)		98.0			98.5		98.9
Outliers		0			0		0
Restraints (r.m.s.d. observed)							
Bond angles (°)		1.80			1.57		1.76
Bond lengths (Å)		0.012			0.012		0.016
Mean isotropic B value ^g (Å ²)							
All		48.3			27.6		27.0
Protein		48.4			25.8		26.0
ESU ^h based on R_{free} (Å)		0.15			0.08		0.11
Protein residues/atoms		1084/8763			547/4627		274/2273
Waters/solvent/UNLs		555/6/5			628/6/4		293/0/2

Values in parentheses are for the highest resolution shell.

^a $R_{\text{merge}} = \sum_{hkl} \sum_i |I_i(hkl) - \langle I(hkl) \rangle| / \sum_{hkl} \sum_i I_i(hkl)$.

^b $R_{\text{meas}} = \sum_{hkl} [N / (N - 1)]^{1/2} \sum_i |I_i(hkl) - \langle I(hkl) \rangle| / \sum_{hkl} \sum_i I_i(hkl)$.⁴⁹

^c $R_{\text{pim}} = \sum_{hkl} [1 / (N - 1)]^{1/2} \sum_i |I_i(hkl) - \langle I(hkl) \rangle| / \sum_{hkl} \sum_i I_i(hkl)$.^{50,51}

^d $CC_{1/2}$ is the correlation of an observed dataset with the underlying true signal.⁵²

^e Typically, the number of unique reflections used in refinement is slightly less than the total number that were integrated and scaled. Reflections are excluded owing to negative intensities and rounding errors in the resolution limits and unit-cell parameters.

^f $R_{\text{cryst}} = \sum_{hkl} (|F_{\text{obs}}| - |F_{\text{calc}}|) / \sum_{hkl} |F_{\text{obs}}|$, where F_{calc} and F_{obs} are the calculated and observed structure-factor amplitudes, respectively. R_{free} is the same as R_{cryst} but for 5% of the total reflections chosen at random and omitted from refinement.

^g This value represents the total B that includes TLS and residual B components.

^h Estimated overall coordinate error.^{53,54}

ⁱ The number of replicates for λ_2 and λ_3 in the high resolution shell was too small for scala to compute the $CC_{1/2}$ value, but that the $CC_{1/2}$ value in the 2.06–2.12 shell was 0.864, 0.860, 0.847 for λ_1, λ_2 and λ_3 respectively.

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