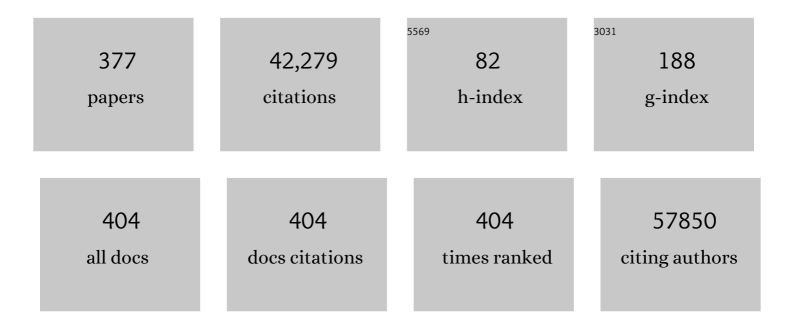
List of Publications by Year in descending order

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ADAM CODZIK

| #  | Article  | IF   | CITATIONS |
|----|--|------|-----------|
| 1  | Novel putative polyethylene terephthalate (PET) plastic degrading enzymes from the environmental metagenome. Proteins: Structure, Function and Bioinformatics, 2022, 90, 504-511.                              | 1.5  | 17        |
| 2  | Multiple expansions of globally uncommon SARS-CoV-2 lineages in Nigeria. Nature Communications, 2022, 13, 688.   | 5.8  | 23        |
| 3  | Defining the risk of SARS-CoV-2 variants on immune protection. Nature, 2022, 605, 640-652.   | 13.7 | 117       |
| 4  | Actin Cross-Linking Effector Domain of the <i>Vibrio vulnificus</i> F-Type MARTX Toxin Dominates Disease Progression During Intestinal Infection. Infection and Immunity, 2022, , e0062721.                    | 1.0  | 1         |
| 5  | A Genomic Island of Vibrio cholerae Encodes a Three-Component Cytotoxin with Monomer and<br>Protomer Forms Structurally Similar to Alpha-Pore-Forming Toxins. Journal of Bacteriology, 2022,<br>204, e0055521. | 1.0  | 3         |
| 6  | What the protein data bank tells us about the evolutionary conservation of protein conformational diversity. Protein Science, 2022, 31, .  | 3.1  | 2         |
| 7  | Identification and phylogenetic analysis of RNA binding domain abundant in apicomplexans or RAP proteins. Microbial Genomics, 2021, 7, .   | 1.0  | 5         |
| 8  | Detection of a SARS-CoV-2 variant of concern in South Africa. Nature, 2021, 592, 438-443.  | 13.7 | 1,381     |
| 9  | ModFlex: Towards Function Focused Protein Modeling. Journal of Molecular Biology, 2021, 433, 166828.   | 2.0  | 2         |
| 10 | HIV-1 and SARS-CoV-2: Patterns in the evolution of two pandemic pathogens. Cell Host and Microbe, 2021, 29, 1093-1110.   | 5.1  | 73        |
| 11 | The interplay of SARS-CoV-2 evolution and constraints imposed by the structure and functionality of its proteins. PLoS Computational Biology, 2021, 17, e1009147.  | 1.5  | 35        |
| 12 | NMR in structural genomics to increase structural coverage of the protein universe. , 2021, , 143-154.   |      | 0         |
| 13 | Dynamic changes in human single-cell transcriptional signatures during fatal sepsis. Journal of<br>Leukocyte Biology, 2021, 110, 1253-1268.  | 1.5  | 26        |
| 14 | Structure of galactarate dehydratase, a new fold in an enolase involved in bacterial fitness after antibiotic treatment. Protein Science, 2020, 29, 711-722.   | 3.1  | 4         |
| 15 | Protein structure, amino acid composition and sequence determine proteome vulnerability to oxidationâ€induced damage. EMBO Journal, 2020, 39, e104523.   | 3.5  | 34        |
| 16 | Microbial function and genital inflammation in young South African women at high risk of HIV infection. Microbiome, 2020, 8, 165.  | 4.9  | 23        |
| 17 | FATCAT 2.0: towards a better understanding of the structural diversity of proteins. Nucleic Acids Research, 2020, 48, W60-W64.   | 6.5  | 134       |
| 18 | Coronavirus3D: 3D structural visualization of COVID-19 genomic divergence. Bioinformatics, 2020, 36, 4360-4362.  | 1.8  | 39        |

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|----|--|------|-----------|
| 19 | Difference contact maps: From what to why in the analysis of the conformational flexibility of proteins. PLoS ONE, 2020, 15, e0226702.   | 1.1  | 11        |
| 20 | Understanding oncogenicity of cancer driver genes and mutations in the cancer genomics era. FEBS<br>Letters, 2020, 594, 4233-4246.   | 1.3  | 20        |
| 21 | Crystal structure of Nsp15 endoribonuclease <scp>NendoU</scp> from <scp>SARS oV</scp> â€2. Protein<br>Science, 2020, 29, 1596-1605.  | 3.1  | 294       |
| 22 | Comparison of metalâ€bound and unbound structures of aminopeptidase B proteins from<br><scp><i>Escherichia coli</i></scp> and <scp><i>Yersinia pestis</i></scp> . Protein Science, 2020, 29,<br>1618-1628.                 | 3.1  | 3         |
| 23 | Structures of singleâ€layer βâ€sheet proteins evolved from βâ€hairpin repeats. Protein Science, 2019, 28,<br>1676-1689.  | 3.1  | 4         |
| 24 | Stratification of amyotrophic lateral sclerosis patients: a crowdsourcing approach. Scientific<br>Reports, 2019, 9, 690.   | 1.6  | 46        |
| 25 | Community assessment to advance computational prediction of cancer drug combinations in a pharmacogenomic screen. Nature Communications, 2019, 10, 2674.   | 5.8  | 240       |
| 26 | Cancer3D 2.0: interactive analysis of 3D patterns of cancer mutations in cancer subsets. Nucleic Acids<br>Research, 2019, 47, D895-D899.   | 6.5  | 12        |
| 27 | Comprehensive Characterization of Cancer Driver Genes and Mutations. Cell, 2018, 173, 371-385.e18.   | 13.5 | 1,670     |
| 28 | Draft Genome Sequences of Two Vibrio parahaemolyticus Strains Associated with Gastroenteritis after Raw Seafood Ingestion in Colorado. Genome Announcements, 2018, 6, .  | 0.8  | 3         |
| 29 | The "Sticky Patch―Model of Crystallization and Modification of Proteins for Enhanced<br>Crystallizability. Methods in Molecular Biology, 2017, 1607, 77-115.   | 0.4  | 17        |
| 30 | The Functional Impact of Alternative Splicing in Cancer. Cell Reports, 2017, 20, 2215-2226.  | 2.9  | 517       |
| 31 | Comparison of algorithms for the detection of cancer drivers at subgene resolution. Nature Methods, 2017, 14, 782-788.   | 9.0  | 72        |
| 32 | Mapping genetic variations to three-dimensional protein structures to enhance variant interpretation: a proposed framework. Genome Medicine, 2017, 9, 113.   | 3.6  | 47        |
| 33 | Neutralizing antibody affords comparable protection against vaginal and rectal simian/human<br>immunodeficiency virus challenge in macaques. Aids, 2016, 30, 1543-1551.  | 1.0  | 47        |
| 34 | <scp>NMR</scp> in structural genomics to increase structural coverage of the protein universe. FEBS<br>Journal, 2016, 283, 3870-3881.  | 2.2  | 5         |
| 35 | Numerous proteins with unique characteristics are degraded by the 26S proteasome following<br>monoubiquitination. Proceedings of the National Academy of Sciences of the United States of<br>America, 2016, 113, E4639-47. | 3.3  | 127       |
| 36 | UHM–ULM interactions in the RBM39–U2AF65 splicing-factor complex. Acta Crystallographica Section<br>D: Structural Biology, 2016, 72, 497-511.  | 1.1  | 36        |

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|----|---|------|-----------|
| 37 | A Distinct Type of Pilus from the Human Microbiome. Cell, 2016, 165, 690-703.   | 13.5 | 78        |
| 38 | Mutation Drivers of Immunological Responses to Cancer. Cancer Immunology Research, 2016, 4, 789-798.  | 1.6  | 32        |
| 39 | Revealing aperiodic aspects of solenoid proteins from sequence information. Bioinformatics, 2016, 32, 2776-2782.  | 1.8  | 3         |
| 40 | PDBFlex: exploring flexibility in protein structures. Nucleic Acids Research, 2016, 44, D423-D428.  | 6.5  | 70        |
| 41 | Protael: protein data visualization library for the web. Bioinformatics, 2016, 32, 602-604.   | 1.8  | 9         |
| 42 | Crystal Structure and Activity Studies of the C11 Cysteine Peptidase from Parabacteroides merdae in the Human Gut Microbiome. Journal of Biological Chemistry, 2016, 291, 9482-9491.  | 1.6  | 15        |
| 43 | Broadly Neutralizing Antibody Responses in a Large Longitudinal Sub-Saharan HIV Primary Infection<br>Cohort. PLoS Pathogens, 2016, 12, e1005369.  | 2.1  | 241       |
| 44 | PROPER: Performance visualization for optimizing and comparing ranking classifiers in MATLAB.<br>Source Code for Biology and Medicine, 2015, 10, 15.  | 1.7  | 3         |
| 45 | Cofactorâ€induced reversible folding of <scp>F</scp> lavodoxinâ€4 from<br><scp><i>L</i>, 2015, 24, 1600-1608.</scp>   | 3.1  | 2         |
| 46 | A Pan-Cancer Catalogue of Cancer Driver Protein Interaction Interfaces. PLoS Computational Biology, 2015, 11, e1004518.   | 1.5  | 122       |
| 47 | Structure and sequence analyses of Bacteroides proteins BVU_4064 and BF1687 reveal presence of two novel predominantly-beta domains, predicted to be involved in lipid and cell surface interactions. BMC Bioinformatics, 2015, 16, 7.          | 1.2  | 8         |
| 48 | AIDA: <i>ab initio</i> domain assembly for automated multi-domain protein structure prediction and domain–domain interaction prediction. Bioinformatics, 2015, 31, 2098-2105.   | 1.8  | 59        |
| 49 | Analysis of Individual Protein Regions Provides Novel Insights on Cancer Pharmacogenomics. PLoS<br>Computational Biology, 2015, 11, e1004024.   | 1.5  | 10        |
| 50 | Structure-based discovery of NANOG variant with enhanced properties to promote self-renewal and reprogramming of pluripotent stem cells. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, 4666-4671. | 3.3  | 43        |
| 51 | Cancer3D: understanding cancer mutations through protein structures. Nucleic Acids Research, 2015,<br>43, D968-D973.  | 6.5  | 46        |
| 52 | Insights into Substrate Specificity of NlpC/P60 Cell Wall Hydrolases Containing Bacterial SH3<br>Domains. MBio, 2015, 6, e02327-14.   | 1.8  | 46        |
| 53 | POSA: a user-driven, interactive multiple protein structure alignment server. Nucleic Acids Research,<br>2014, 42, W240-W245.   | 6.5  | 50        |
| 54 | Improving the chances of successful protein structure determination with a random forest classifier. Acta Crystallographica Section D: Biological Crystallography, 2014, 70, 627-635.   | 2.5  | 46        |

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|----|---|-----|-----------|
| 55 | AIDA: ab initio domain assembly server. Nucleic Acids Research, 2014, 42, W308-W313.  | 6.5 | 47        |
| 56 | Crystal structure of a putative quorum sensingâ€regulated protein (PA3611) from the<br>Pseudomonasâ€specific DUF4146 family. Proteins: Structure, Function and Bioinformatics, 2014, 82,<br>1086-1092.  | 1.5 | 7         |
| 57 | Structure-Guided Functional Characterization of DUF1460 Reveals a Highly Specific NlpC/P60 Amidase<br>Family. Structure, 2014, 22, 1799-1809.   | 1.6 | 10        |
| 58 | Molecular characterization of novel pyridoxalâ€5′â€phosphateâ€dependent enzymes from the human<br>microbiome. Protein Science, 2014, 23, 1060-1076.   | 3.1 | 8         |
| 59 | Structural genomics analysis of uncharacterized protein families overrepresented in human gut<br>bacteria identifies a novel glycoside hydrolase. BMC Bioinformatics, 2014, 15, 112.  | 1.2 | 13        |
| 60 | Phylogenomic analysis of glycogen branching and debranching enzymatic duo. BMC Evolutionary<br>Biology, 2014, 14, 183.  | 3.2 | 27        |
| 61 | PubServer: literature searches by homology. Nucleic Acids Research, 2014, 42, W430-W435.  | 6.5 | 13        |
| 62 | Structural analysis of arabinose-5-phosphate isomerase fromBacteroides fragilisand functional implications. Acta Crystallographica Section D: Biological Crystallography, 2014, 70, 2640-2651.  | 2.5 | 0         |
| 63 | Structures of a Bifunctional Cell Wall Hydrolase CwlT Containing a Novel Bacterial Lysozyme and an<br>NlpC/P60 dl-Endopeptidase. Journal of Molecular Biology, 2014, 426, 169-184.  | 2.0 | 25        |
| 64 | bNAber: database of broadly neutralizing HIV antibodies. Nucleic Acids Research, 2014, 42, D1133-D1139.   | 6.5 | 69        |
| 65 | FFAS-3D: improving fold recognition by including optimized structural features and template re-ranking. Bioinformatics, 2014, 30, 660-667.  | 1.8 | 97        |
| 66 | Crystal structures of three representatives of a new <scp>P</scp> fam family PF14869 (DUF4488) suggest they function in sugar binding/uptake. Protein Science, 2014, 23, 1380-1391.   | 3.1 | 3         |
| 67 | Structure- and context-based analysis of the GxGYxYP family reveals a new putative class of Glycoside<br>Hydrolase. BMC Bioinformatics, 2014, 15, 196.  | 1.2 | 8         |
| 68 | Basis for substrate recognition and distinction by matrix metalloproteinases. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, E4148-55.   | 3.3 | 75        |
| 69 | S-Nitrosylation-Mediated Redox Transcriptional Switch Modulates Neurogenesis and Neuronal Cell<br>Death. Cell Reports, 2014, 8, 217-228.  | 2.9 | 58        |
| 70 | ConSole: using modularity of Contact maps to locate Solenoid domains in protein structures. BMC<br>Bioinformatics, 2014, 15, 119.   | 1.2 | 23        |
| 71 | Structure and computational analysis of a novel protein with metallopeptidase-like and circularly permuted winged-helix-turn-helix domains reveals a possible role in modified polysaccharide biosynthesis. BMC Bioinformatics, 2014, 15, 75. | 1.2 | 1         |
| 72 | Structural systems biology: from bacterial to cancer networks. BMC Genomics, 2014, 15, O14.   | 1.2 | 0         |

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| 73 | e-Driver: a novel method to identify protein regions driving cancer. Bioinformatics, 2014, 30, 3109-3114.  | 1.8 | 116       |
| 74 | Crystal structure of a member of a novel family of dioxygenases (PF10014) reveals a conserved cupin fold and active site. Proteins: Structure, Function and Bioinformatics, 2014, 82, 164-170. | 1.5 | 15        |
| 75 | OUR EXPANDING PROTEIN UNIVERSE. , 2014, , .  |     | 0         |
| 76 | THE MICROBIOME(S): MICROBIOTA, FAMILIES, FUNCTIONS. , 2014, , .  |     | 0         |
| 77 | JCSC – Adapting Structural Genomics to Eukaryotic Complexes. Acta Crystallographica Section A:<br>Foundations and Advances, 2014, 70, C1148-C1148.   | 0.0 | 0         |
| 78 | MORPH-PRO: a novel algorithm and web server for protein morphing. Algorithms for Molecular<br>Biology, 2013, 8, 19.  | 0.3 | 9         |
| 79 | Evolution of the Animal Apoptosis Network. Cold Spring Harbor Perspectives in Biology, 2013, 5, a008649-a008649.   | 2.3 | 63        |
| 80 | Two Pfam protein families characterized by a crystal structure of protein lpg2210 from Legionella pneumophila. BMC Bioinformatics, 2013, 14, 265.  | 1.2 | 3         |
| 81 | Polysaccharides utilization in human gut bacterium Bacteroides thetaiotaomicron: comparative genomics reconstruction of metabolic and regulatory networks. BMC Genomics, 2013, 14, 873.        | 1.2 | 122       |
| 82 | Divergent evolution of protein conformational dynamics in dihydrofolate reductase. Nature<br>Structural and Molecular Biology, 2013, 20, 1243-1249.  | 3.6 | 153       |
| 83 | Structural Systems Biology Evaluation of Metabolic Thermotolerance in <i>Escherichia coli</i> .<br>Science, 2013, 340, 1220-1223.  | 6.0 | 111       |
| 84 | LUD, a new protein domain associated with lactate utilization. BMC Bioinformatics, 2013, 14, 341.  | 1.2 | 8         |
| 85 | Structure and Function of a Novel <scp>ld</scp> -Carboxypeptidase A Involved in Peptidoglycan<br>Recycling. Journal of Bacteriology, 2013, 195, 5555-5566.                                     | 1.0 | 16        |
| 86 | Structure and Function of the DUF2233 Domain in Bacteria and in the Human Mannose 6-Phosphate<br>Uncovering Enzyme. Journal of Biological Chemistry, 2013, 288, 16789-16799.                   | 1.6 | 7         |
| 87 | This Déjà Vu Feeling—Analysis of Multidomain Protein Evolution in Eukaryotic Genomes. PLoS<br>Computational Biology, 2012, 8, e1002701.  | 1.5 | 37        |
| 88 | The Structure of Mlc Titration Factor A (MtfA/Yeel) Reveals a Prototypical Zinc Metallopeptidase<br>Related to Anthrax Lethal Factor. Journal of Bacteriology, 2012, 194, 2987-2999.           | 1.0 | 14        |
| 89 | A Blueprint for HIV Vaccine Discovery. Cell Host and Microbe, 2012, 12, 396-407.   | 5.1 | 348       |
| 90 | MORPH-PRO: A Novel Algorithm and Web Server for Protein Morphing. Lecture Notes in Computer Science, 2012, , 262-273.  | 1.0 | 0         |

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|-----|--|------|-----------|
| 91  | Structure of a Novel Winged-Helix Like Domain from Human NFRKB Protein. PLoS ONE, 2012, 7, e43761.   | 1.1  | 5         |
| 92  | Structure of the pilus assembly protein TadZ from <i>Eubacterium rectale</i> : implications for polar localization. Molecular Microbiology, 2012, 83, 712-727.   | 1.2  | 22        |
| 93  | Structure of an MmyB-Like Regulator from C. aurantiacus, Member of a New Transcription Factor<br>Family Linked to Antibiotic Metabolism in Actinomycetes. PLoS ONE, 2012, 7, e41359.   | 1.1  | 14        |
| 94  | Structural Determinants of Limited Proteolysis. Journal of Proteome Research, 2011, 10, 3642-3651.   | 1.8  | 68        |
| 95  | Strong functional patterns in the evolution of eukaryotic genomes revealed by the reconstruction of ancestral protein domain repertoires. Genome Biology, 2011, 12, R4.  | 13.9 | 84        |
| 96  | TIR domain-containing adaptor SARM is a late addition to the ongoing microbe–host dialog.<br>Developmental and Comparative Immunology, 2011, 35, 461-468.  | 1.0  | 66        |
| 97  | Dynamics of coregulator-induced conformational perturbations in androgen receptor ligand binding domain. Molecular and Cellular Endocrinology, 2011, 341, 1-8.   | 1.6  | 8         |
| 98  | Structural and Sequence Analysis of Imelysin-Like Proteins Implicated in Bacterial Iron Uptake. PLoS<br>ONE, 2011, 6, e21875.  | 1.1  | 17        |
| 99  | Metagenomics and the protein universe. Current Opinion in Structural Biology, 2011, 21, 398-403.   | 2.6  | 62        |
| 100 | Crystal structure of a metalâ€dependent phosphoesterase (YP_910028.1) from <i>Bifidobacterium<br/>adolescentis</i> : Computational prediction and experimental validation of phosphoesterase activity.<br>Proteins: Structure, Function and Bioinformatics, 2011, 79, 2146-2160. | 1.5  | 11        |
| 101 | Internal organization of large protein families: Relationship between the sequence, structure, and functionâ€based clustering. Proteins: Structure, Function and Bioinformatics, 2011, 79, 2389-2402.  | 1.5  | 4         |
| 102 | TOPSAN: a dynamic web database for structural genomics. Nucleic Acids Research, 2011, 39, D494-D496.   | 6.5  | 17        |
| 103 | Structural Insights into Inhibition of Bacillus anthracis Sporulation by a Novel Class of Non-heme<br>Globin Sensor Domains. Journal of Biological Chemistry, 2011, 286, 8448-8458.  | 1.6  | 22        |
| 104 | The FGGY Carbohydrate Kinase Family: Insights into the Evolution of Functional Specificities. PLoS<br>Computational Biology, 2011, 7, e1002318.  | 1.5  | 48        |
| 105 | FFAS server: novel features and applications. Nucleic Acids Research, 2011, 39, W38-W44.   | 6.5  | 130       |
| 106 | Structure and Function of the First Full-Length Murein Peptide Ligase (Mpl) Cell Wall Recycling<br>Protein. PLoS ONE, 2011, 6, e17624.   | 1.1  | 30        |
| 107 | Structural Analysis of Papain-Like NlpC/P60 Superfamily Enzymes with a Circularly Permuted Topology<br>Reveals Potential Lipid Binding Sites. PLoS ONE, 2011, 6, e22013.   | 1.1  | 22        |
| 108 | CARD8 and NLRP1 Undergo Autoproteolytic Processing through a ZU5-Like Domain. PLoS ONE, 2011, 6, e27396.   | 1.1  | 168       |

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|-----|--|-----|-----------|
| 109 | Domain architecture evolution of pattern-recognition receptors. Immunogenetics, 2010, 62, 263-272.   | 1.2 | 68        |
| 110 | Structure of a Virulence Regulatory Factor CvfB Reveals a Novel Winged Helix RNA Binding Module.<br>Structure, 2010, 18, 537-547.  | 1.6 | 23        |
| 111 | TOPSAN: a collaborative annotation environment for structural genomics. BMC Bioinformatics, 2010, 11, 426.   | 1.2 | 19        |
| 112 | A widespread peroxiredoxin-like domain present in tumor suppression- and progression-implicated proteins. BMC Genomics, 2010, 11, 590.   | 1.2 | 26        |
| 113 | The structure of KPN03535 (gi 152972051), a novel putative lipoprotein fromKlebsiella pneumoniae,<br>reveals an OB-fold. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66,<br>1254-1260.  | 0.7 | 3         |
| 114 | Conformational changes associated with the binding of zinc acetate at the putative active site<br>ofXcTcmJ, a cupin fromXanthomonas campestrispv.campestris. Acta Crystallographica Section F:<br>Structural Biology Communications, 2010, 66, 1347-1353.                                  | 0.7 | 5         |
| 115 | Structure of an essential bacterial protein YeaZ (TM0874) from <i>Thermotoga maritima</i> at 2.5â€Ã<br>resolution. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1230-1236.   | 0.7 | 17        |
| 116 | The structure of the first representative of Pfam family PF09836 reveals a two-domain organization<br>and suggests involvement in transcriptional regulation. Acta Crystallographica Section F: Structural<br>Biology Communications, 2010, 66, 1174-1181.                                 | 0.7 | 8         |
| 117 | The structure of the first representative of Pfam family PF06475 reveals a new fold with possible involvement in glycolipid metabolism. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1211-1217.  | 0.7 | 10        |
| 118 | Structure of LP2179, the first representative of Pfam family PF08866, suggests a new fold with a role<br>in amino-acid metabolism. Acta Crystallographica Section F: Structural Biology Communications, 2010,<br>66, 1205-1210.  | 0.7 | 3         |
| 119 | The structure of Jann_2411 (DUF1470) from <i>Jannaschia</i> sp. at 1.45â€Ã resolution reveals a new fold<br>(the ABATE domain) and suggests its possible role as a transcription regulator. Acta<br>Crystallographica Section F: Structural Biology Communications, 2010, 66, 1198-1204.   | 0.7 | 9         |
| 120 | Structure of the first representative of Pfam family PF09410 (DUF2006) reveals a structural signature of the calycin superfamily that suggests a role in lipid metabolism. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1153-1159.                       | 0.7 | 18        |
| 121 | Open and closed conformations of two SpoIIAA-like proteins (YP_749275.1 and YP_001095227.1) provide insights into membrane association and ligand binding. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1245-1253.                                       | 0.7 | 8         |
| 122 | The structure of BVU2987 fromBacteroides vulgatusreveals a superfamily of bacterial periplasmic proteins with possible inhibitory function. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1265-1273.  | 0.7 | 8         |
| 123 | Structures of the first representatives of Pfam family PF06938 (DUF1285) reveal a new fold with repeated structural motifs and possible involvement in signal transduction. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1218-1225.                      | 0.7 | 8         |
| 124 | Structures of the first representatives of Pfam family PF06684 (DUF1185) reveal a novel variant of theBacilluschorismate mutase fold and suggest a role in amino-acid metabolism. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1182-1189.                | 0.7 | 3         |
| 125 | The structure of SSO2064, the first representative of Pfam family PF01796, reveals a novel two-domain zinc-ribbon OB-fold architecture with a potential acyl-CoA-binding role. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1160-1166.                   | 0.7 | 20        |
| 126 | A conserved fold for fimbrial components revealed by the crystal structure of a putative fimbrial<br>assembly protein (BT1062) from <i>Bacteroides thetaiotaomicron</i> at 2.2â€Ã resolution. Acta<br>Crystallographica Section F: Structural Biology Communications, 2010, 66, 1281-1286. | 0.7 | 9         |

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|-----|---|-----|-----------|
| 127 | Structure of the first representative of Pfam family PF04016 (DUF364) reveals enolase and<br>Rossmann-like folds that combine to form a unique active site with a possible role in heavy-metal<br>chelation. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1167-1173.  | 0.7 | 3         |
| 128 | Structures of three members of Pfam PF02663 (FmdE) implicated in microbial methanogenesis reveal a<br>conserved α+β core domain and an auxiliary C-terminal treble-clef zinc finger. Acta Crystallographica<br>Section F: Structural Biology Communications, 2010, 66, 1335-1346.   | 0.7 | 8         |
| 129 | Structure of the Î <sup>3</sup> - <scp>D</scp> -glutamyl- <scp>L</scp> -diamino acid endopeptidase YkfC<br>from <i>Bacillus cereus</i> in complex with <scp>L</scp> -Ala-Î <sup>3</sup> - <scp>D</scp> -Clu: insights into substrate<br>recognition by NlpC/P60 cysteine peptidases. Acta Crystallographica Section F: Structural Biology<br>Communications. 2010. 66. 1354-1364. | 0.7 | 64        |
| 130 | The structure ofHaemophilus influenzaeprephenate dehydrogenase suggests unique features of<br>bifunctional TyrA enzymes. Acta Crystallographica Section F: Structural Biology Communications,<br>2010, 66, 1317-1325.   | 0.7 | 11        |
| 131 | Structure of a membrane-attack complex/perforin (MACPF) family protein from the human gut<br>symbiontBacteroides thetaiotaomicron. Acta Crystallographica Section F: Structural Biology<br>Communications, 2010, 66, 1297-1305.   | 0.7 | 30        |
| 132 | Structure of a putative NTP pyrophosphohydrolase: YP_001813558.1 fromExiguobacterium sibiricum255-15. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1237-1244.   | 0.7 | 2         |
| 133 | Structure of <i>Bacteroides thetaiotaomicron</i> BT2081 at 2.05â€Ã resolution: the first structural representative of a new protein family that may play a role in carbohydrate metabolism. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1287-1296.   | 0.7 | 2         |
| 134 | Structure of BT_3984, a member of the SusD/RagB family of nutrient-binding molecules. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1274-1280.   | 0.7 | 11        |
| 135 | TOPSAN: use of a collaborative environment for annotating, analyzing and disseminating data on JCSG<br>and PSI structures. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66,<br>1143-1147.   | 0.7 | 33        |
| 136 | Structure of a tryptophanyl-tRNA synthetase containing an iron–sulfur cluster. Acta<br>Crystallographica Section F: Structural Biology Communications, 2010, 66, 1326-1334.   | 0.7 | 19        |
| 137 | The JCSC high-throughput structural biology pipeline. Acta Crystallographica Section F: Structural Biology Communications, 2010, 66, 1137-1142.   | 0.7 | 99        |
| 138 | The crystal structure of a bacterial Sufuâ€like protein defines a novel group of bacterial proteins that<br>are similar to the Nâ€terminal domain of human Sufu. Protein Science, 2010, 19, 2131-2140.  | 3.1 | 12        |
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