Protein function annotation with Structurally Aligned Local Sites of Activity (SALSAs)

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Abstract

Background: The prediction of biochemical function from the 3D structure of a protein has proved to be much more difficult than was originally foreseen. A reliable method to test the likelihood of putative annotations and to predict function from structure would add tremendous value to structural genomics data. We report on a new method, Structurally Aligned Local Sites of Activity (SALSA), for the prediction of biochemical function based on a local structural match at the predicted catalytic or binding site.

Results: Implementation of the SALSA method is described. For the structural genomics protein PY01515 (PDB ID 2aqw) from Plasmodium yoelii, it is shown that the putative annotation, Orotidine 5'-monophosphate decarboxylase (OMPDC), is most likely correct. SALSA analysis of YP_001304206.1 (PDB ID 3h3l), a putative sugar hydrolase from Parabacteroides distasonis, shows that its active site does not bear close resemblance to any previously characterized member of its superfamily, the Concanavalin A-like lectins/glucanases. It is noted that three residues in the active site of the thermophilic beta-1,4-xylanase from Nonomuraea flexuosa (PDB ID 1m4w), Y78, E87, and E176, overlap with POOL-predicted residues of similar type, Y168, D153, and E232, in YP_001304206.1. The substrate recognition regions of the two proteins are rather different, suggesting that YP_001304206.1 is a new functional type within the superfamily. A structural genomics protein from Mycobacterium avium (PDB ID 3q1t) has been reported to be an enoyl-CoA hydratase (ECH), but SALSA analysis shows a poor match between the predicted residues for the SG protein and those of known ECHs. A better local structural match is obtained with Anabaena beta-diketone hydrolase (ABDH), a known beta-diketone hydrolase from Cyanobacterium anabaena (PDB ID 2j5s). This suggests that the reported ECH function of the SG protein is incorrect and that it is more likely a beta-diketone hydrolase.

Conclusions: A local site match provides a more compelling function prediction than that obtainable from a simple 3D structure match. The present method can confirm putative annotations, identify misannotation, and in some cases suggest a more probable annotation.

Background

There are currently over 11,000 structural genomics (SG) protein structures in the Protein Data Bank (PDB) [1] and most of them are of unknown or uncertain function, as the inference of function from structure has proved to be more difficult than anticipated. Furthermore, when new structures of unknown function are determined, it is common practice to make a tentative functional assignment from the closest sequence match or the best 3D structure match to an annotated protein. Such tentative functional assignments are often incorrect [2]. Furthermore, one annotation error can propagate or "percolate" [2-4] in databases as additional proteins are annotated by automated or semi-automated means.

Overviews of current methods for the functional annotation of proteins from their sequence and/or structure...
have been given in recent reviews [5-8]. The simplest, and most commonly employed [6] methods seek the closest sequence matches using a search program such as BLAST [9], or alternatively the closest 3D structure match obtained from e.g. Dali [10], Combinatorial Extension (CE) [11], or Topfit [12], and then just transfer the function from the closest match to the query protein. However, even relatively high sequence similarity does not necessarily imply similar function [13]. Other types of sequence-based methods employ motif searching, phylogenetic profiling, or genome context. The Critical Assessment of Function Annotation (CAFA) experiment (http://biofunctionprediction.org/) seeks to assess the state of the current art of function prediction, chiefly from sequence. The aim of this work is to exploit structural information, together with computed chemical properties, to enhance function prediction capabilities.

It was hoped that SG would provide functional annotations for the protein products of newly-sequenced coding genes, as indeed the 3D structure can sometimes be indicative of function. Simple protein fold comparison does work in some cases, as domains having a common fold sometimes do have the same function. However, many folds have multiple functions. For instance, the Rossman fold and the TIM barrel each represent more than 50 different functions. The use of local 3D structural motifs or templates, a feature of the present method, is now emerging as a more promising path for correct functional annotation from structure [14-19].

In spite of recent advances in protein function prediction, inference of biochemical function from the structure is difficult [20,21]. Hundreds of SG structures have no functional assignment at all and, for thousands of other SG proteins, functional hypotheses for SG proteins are putative and uncertain. Not all such hypotheses will prove in time to be correct, as examples below will illustrate. The ability to determine function from the 3D structure would add great value to this growing volume of SG data.

A different approach to functional annotation from 3D structure is presented here and is based on the combination of functional site prediction with local 3D structural alignment. Functional site predictions are obtained from Partial Order Optimum Likelihood (POOL) [22,23], a monotonicity-constrained maximum likelihood method, using computed chemical, electrostatic, and geometric properties, as well as phylogenetic information (if available), as input features. POOL places all of the residues in the input protein structure into an ordered list, ranked according to probability of participation in the active site. The top-ranked residues constitute the active site prediction. Structural alignments are obtained for sets of these local sites. Characteristic spatial patterns of predicted residues at the structurally aligned local sites of activity (SALSAs) are then used to identify specific types of biochemical function. The quality of the match of the predicted functional site in the query protein to functional sites in proteins of known function is measured using a scoring function. The present method can determine whether a putative functional assignment is likely to be correct or incorrect. In some cases where a protein is shown to be misannotated, a probable functional assignment is made.

**Methods**

Functional residue predictions were made using POOL [22,23]. Input features for each residue in a given structure include: electrostatics information, as contained in THEMATICS metrics [24,25]; phylogenetic information from INTREPID [26,27]; and geometric information from ConCavity (structure only version) [28]. The top-ranked residues in the POOL output constitute the functional site prediction. Cut-off limits are specified for each case.

Multiple structure alignments are made for each set of proteins. The structural alignment of multiple structures of diverse function can be difficult and therefore multiple alignment methods [11,12,29] may be needed for some cases. In the examples shown here, T-Coffee [29] is used. For present purposes, a full alignment is not necessary. A quality alignment is only required in the local spatial region of the predicted active site.

SALSA tables are constructed for the locally aligned residues in the predicted active site. In a SALSA table, the rows represent individual protein structures and the columns represent spatially aligned positions.

Consensus signatures for a given functional subclass are established using POOL predictions on a set of previously characterized proteins with the same biochemical function, usually with common fold. To maximize sequence diversity in this reference set, sets of structures are sought with the lowest possible sequence identity among them. POOL-predicted residues of the same amino acid type in the same spatial position for the majority of the previously characterized proteins of common biochemical function then constitute the consensus signature for that functional group. The consensus signature for a given biochemical function thus consists of a series of amino acid types in specified spatial positions.

SG proteins of unknown or uncertain function are analyzed by POOL and the predictions are aligned with those of proteins of known function, or with the consensus signature.

Scoring the match between the predicted active site for the query protein and that of the consensus signature is performed using the BLOSUM62 matrix [30]. Scores are reported as a percentage of the maximum value (i.e. the score for the perfect match, the consensus signature with itself).
Results and discussion

Confirmation of annotation for PY01515, a putative Orotidine 5’-monophosphate decarboxylase (OMPDC)

Orotidine 5’-monophosphate decarboxylase (OMPDC) catalyzes one step in the pyrimidine biosynthesis pathway. It catalyzes the metal ion dependent decarboxylation of orotidine monophosphate (OMP) to uridine monophosphate (UMP) and CO₂ [31,32]. OMPDC is a member of the ribulose phosphate binding barrel (RPBB) superfamily and has a TIM barrel [33] structure, with the active site located inside the beta barrel, spanning the eight beta strands. The structural genomics protein PY01515 (PDB ID 2aqw) is a putative OMPDC from *Plasmodium yoelii* [34].

The POOL-predicted functional site for PY01515 was aligned with eight different functional site types predicted by POOL for structures in the RPBB superfamily and a strong match was found with that of the OMPDCs and not with the other seven functional types. Five previously characterized OMPDC structures, those from *Bacillus subtilis* (PDB ID 1dbt), *Methanothermobacter thermautotrophicus* (PDB ID 1dvj), *Saccharomyces cerevisiae* (PDB ID 1dqw), *Escherichia coli* (PDB ID 1l2u), and *Plasmodium falciparum* (PDB ID 2za1), were used to establish the consensus signature of an OMPDC active site. These five previously characterized OMPDCs represent considerable sequence diversity, as shown in Table 1. With the exception of structures 1 and 4, which share sequence identity of 60%, all other pairs of structures have sequence identities in the 6% - 30% range.

For the five previously characterized OMPDCs, the important residues are predicted using the top 9% of the residues, as ranked by POOL, for each protein structure. When these five predicted active sites are structurally aligned, eight spatial positions are found to have common predicted residues across the five diverse, previously characterized OMPDCs. Table 2 shows this local structural alignment. The rows in Table 2 represent individual protein structures, with the five previously characterized OMPDCs listed first; the last row is the query protein from SG. The columns represent spatially coincident positions in the local structural alignment. The residues predicted by POOL are shown in uppercase; residues not predicted by POOL are shown in lowercase. The beta strand on which each position is located is given at the top of the column, above the position number. The good match between the SG protein and the known OMPDCs suggests common function.

| PDB ID | 1dbt | 1dvj | 1dqw | 1l2u | 2za1 | 2aqw |
|--------|------|------|------|------|------|------|
| 1      | 1dbt | 0.240| 0.600| 0.060| 0.240|      |
| 2      | 1dvj | 0.280| 0.120| 0.220|      |      |
| 3      | 1dqw | 0.280| 0.280| 0.080| 0.280|      |
| 4      | 1l2u | 0.300| 0.060| 0.200|      |      |
| 5      | 2za1 | 0.080| 0.060| 0.200|      |      |
| 6      | SG   | 0.280| 0.200| 0.020|      |      |

Table 1 Sequence identity matrix for five previously characterized OMPDCs (structures 1-5) and the SG protein PY01515 (PDB ID 2aqw).

The quality of a match with the consensus signature may be measured using a scoring matrix. Using the BLOSUM62 [30] matrix, the first four proteins listed in Table 2 have a score of 48 with the consensus signature; this score is 100% of the maximum value. The *Plasmodium falciparum* structure has a score of 39 (81% of the maximum value) against the consensus signature.

The structurally aligned residues for the SG protein PY01515 from *Plasmodium yoelii* are shown in the last row of Table 2. For seven out of the eight positions, POOL predicts residues that are identical to the consensus signature residues of the previously characterized OMPDCs. The only variation is in position 6, where there is an asparagine that is not predicted by POOL, just as in
the Plasmodium falciparum OMPDC. PY01515 has a score of 39 (81% of the maximum value) against the consensus signature, using the BLOSUM62 scoring matrix. The strong match between the predicted active site for PY01515 and those of the previously characterized OMPDCs indicates that the putative OMPDC functional assignment is correct.

**YP_001304206.1 - a probable new functional type in the Concanavalin A-like lectins/glucanases superfamily**

YP_001304206.1 (PDB ID 3h3l) is a putative sugar hydrolase from *Parabacteroides distasonis*, a commensal bacterium of the human intestinal tract. YP_001304206.1 is a member of the Concanavalin A-like lectins/glucanases superfamily. The POOL-predicted functionally important residues for YP_001304206.1 show poor spatial overlap with those of all of the enzymes of known function within the Concanavalin A-like lectins/glucanases superfamily. Figure 1 shows a structural alignment of the predicted residues for YP_001304206.1 with those of its closest Dali [10,37] structural match, endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase (PDB ID 2ayh), a representative member of the glycoside hydrolases family 16 (GH16). The residues for the query protein YP_001304206.1 are shown in gray and those for endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase are shown in pink. Table 3 shows an alignment at the 14 consensus signature positions of GH16 for the representative GH16, endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase, with the SG protein YP_001304206.1. Previously reported active site residues [38] are shown in boldface. POOL-predicted residues (top 8%) are shown in uppercase; residues not predicted are shown in lowercase. Note that the SG protein has a gap (no residue well aligned) at three of the consensus signature positions. For the alignment shown in Table 3, a negative BLOSUM62 score of -5 is obtained, corresponding to -5% of the maximum value of +97. The three catalytic residues for endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase, E105, D107, and E109 [38], form an EXDXE motif on a common beta sheet and are seen forming a vertical line through the center of Figure 1. An alignment of the consensus signature residues, represented by enoyl-CoA hydratase from *Rattus norvegicus* (PDB ID 1ey3), with the corresponding spatially overlapping residues of the query protein, is shown in Table 4. Again, the rows represent individual protein structures and the columns represent spatial positions in the alignment. The known catalytic residues, A98, G141, E144, and E164 [40,41], are shown in boldface. Residues predicted by POOL are shown in uppercase and residues not predicted are shown in lowercase. The BLOSUM62 score between the SG protein and the known ECH is only 11, or 22% of the maximum value of 51, for these nine positions. Note further that the SG protein is missing the catalytic residues that correspond to E144 and E164 in the *Rattus*
Table 3 Local structural alignment of the residues in the GH16 consensus signature positions for the known representative GH16, endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase, with the SG protein YP_001304206.1.

| Spatial Positions | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 |
|-------------------|---|---|---|---|---|---|---|---|---|----|----|----|----|----|
| 2ayh (GH16)       | F92| Y94| W103| d104| E105| D107| E109| L111| Q119| N121| Y123| Y147| W158| W184|
| 3h3l (SG)         | F125| -  | Y138| -  | s140| E142| q144| L146| A166| Y168| -   | t187| H198| H256|

Previously reported active site residues are shown in boldface. POOL-predicted residues (top 8%) are shown in uppercase; residues not predicted are shown in lowercase. The poor match suggests different functions.
norvegicus ECH structure. These results strongly suggest that the reported enoyl-CoA hydratase annotation is incorrect.

Comparison of the local site prediction for the SG protein with those of other members of the ClpP/crotonase superfamily reveals a much better match with ABDH (Anabaena beta diketone hydrolase), a known \(\beta\)-diketone hydrolase from Cyanobacterium anabaena (PDB ID 2j5s). The local alignment of the top POOL-predicted residues for the \(M.\ avium\) structure with residues from ABDH is shown in Table 5. The known catalytic residues for ABDH [42] are shown in boldface. Again, the columns represent overlapping spatial positions, but in Table 5 they are listed in order of the POOL rank for the \(M.\ avium\) structure (D155 is ranked first, H146 second, E244 third, \(\ldots\)). Thus all of the residues listed for the SG protein in Table 5 are predicted by POOL. Residues not predicted by POOL for ABDH are shown in lowercase. Notice that four of the top-ranked POOL residues for the SG protein are aligned with the known catalytic residues of ABDH: D153, H144, E243, and H43. The BLOSUM62 score between the SG protein and the known ABDH for these seven positions is 30, or 60% of the maximum value. These results suggest that the \(M.\ avium\) structure may be a \(\beta\)-diketone hydrolase.

Table 4 Local structural alignment of the predicted active site residues by SALSA for a known ECH from Rattus norvegicus (PDB ID 1ey3) with predicted residues for a Structural Genomics protein from Mycobacterium avium (PDB ID 3q1t), reported to be an ECH.

| Spatial Positions | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 |
|-------------------|---|---|---|---|---|---|---|---|---|
| Known ECH (1ey3)  | A98 | g141 | E144 | C149 | D150 | E164 | R178 | k241 | N245 |
| SG protein “ECH” (3q1t) | G76 | A123 | V126 | a131 | D132 | H146 | C160 | k223 | n227 |

Known catalytic residues are shown in boldface. Residues predicted by POOL are in uppercase; residues not predicted are in lowercase. Note the poor match between the residues of the SG protein with those of the representative ECH.

Table 5 Local structural alignment of the predicted residues for the SG protein from Mycobacterium avium (PDB ID 3q1t) with the corresponding residues of ABDH from Cyanobacterium anabaena.

| POOL Ranking | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
|--------------|---|---|---|---|---|---|---|
| SG protein “ECH” (3q1t) | D155 | H146 | E244 | D144 | H44 | C160 | H156 |
| Known ABDH (2j5s) | D153 | H144 | E243 | D141 | H43 | I158 | g154 |

The spatial positions 1 through 7 correspond to the ordinal values for the top seven residues in the POOL rank order for 3q1t. Known catalytic residues for ABDH are shown in boldface. Residues predicted by POOL are in uppercase; residues not predicted are in lowercase. Note that the match between the residues of the SG protein and of ABDH is better than that of Table 4.
hydrolase, but perhaps with a native substrate different from that of the *Cyanobacterium anabaena* protein.

Figure 3 illustrates the structural alignment of the top POOL-predicted residues for the SG *M. avium* structure (purple) with the corresponding residues from ABDH (green), showing that the known catalytic residues of ABDH have strong overlap with the top POOL-predicted residues for the SG protein.

**Conclusions**

Local structural matching, as implemented by the SALSA method, provides a more compelling prediction of biochemical function than a simple, global 3D structure match. SALSA can confirm putative annotations, identify misannotations, suggest correct annotations, and, in some cases of misannotation, predict a more probable functional annotation.

For any given protein structure of previously characterized function, the list of residues reported in the literature to be important for the biochemical function is a subset of the list of residues predicted by POOL. This longer list is a key advantage of the present method, as it enables better discrimination between the functional subclasses.

To date, one prediction made by local site matching using our electrostatics-based functional site prediction has been verified experimentally by direct biochemical assays [43]. Further experimental testing of SALSA function predictions is in progress.

The BLOSUM62 scoring matrix has been used to measure the quality of the match between two predicted active sites. Whether there exists a better scoring matrix for this purpose is currently under investigation. At the present time, there are too few SG proteins with experimentally verified biochemical function to be able to translate the

**Figure 3** Structural alignment of the top POOL-predicted residues for the SG protein (purple; PDB 3q1t), reported to be an enoyl-CoA hydratase, with those of ABDH (green). H43, H144, D153, and E243 are known catalytic residues in ABDH.
match score into a confidence metric, but as experimental testing progresses, this will become possible.

The SALSA method is amenable to automation and could be used to complement sequence-based function annotation methods, such as those evaluated in the CAFA experiments.

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Abbreviations
ABDH: Arabidopsis beta-diketone hydrolase; BLOSUM: Blocks of amino acid Substitution Matrix; CAFA: Critical Assessment of Function Annotation; CE: Combinatorial Extension; ECH: enoyl-CoA hydratase; GH16: glycoside hydrolase family 16; INTREPID: Information-theoretic TREe traversal for Protein functional site Identification; OMP: orotidine monophosphate; OMPDC: orotidine 5′-monophosphate decarboxylase; PDB: Protein Data Bank; POOL: Partial Order Optimum Likelihood; RPBB: Ribulose Phosphate Binding Barrel; SALSA: Structurally Aligned Local Sites of Activity; SG: Structural Genomics; THEMATICS: THEoretical Microscopic Anomalous Titration Curve Shapes; UMP: uridine monophosphate.

Authors’ contributions
All six authors performed the calculations, participated in the development of the methodology, and contributed to the writing of the manuscript. ZW had primary responsibility for the analysis of the Concanavalin A-like lectins/glucanases, PY for the Micrococcus luteus SG protein, and JSL for the OMPDCs. ZW, PY, and JSL contributed equally to this work.

Competing interests
The authors declare that they have no competing interests.

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Declarations
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References
1. Westbrook J, Feng Z, Chen L, Yang H, Berman HM: The Protein Data Bank and structural genomics. Nucleic Acids Res 2003, 31:489-491.
2. Schoen AM, Brown SD, Dodekels I, Babbitt PC: Annotation Error in Public Databases: Misannotation of Molecular Function in Enzyme Superfamilies. PLoS Comp Biol 2009, 5:e1000065.
3. Gilks WR, Audit B, de Angelis D, Tsoka S, Ouzounis CA: Percolation of annotation errors through hierarchically structured protein sequence databases. Math Biosci 2005, 193:233-234.
4. Llewellyn R, Eisenberg DS: Annotating proteins with generalized functional linkages. Proc Natl Acad Sci USA 2008, 105:17700-17705.
5. Lee D, Redfern O, Orengo C: Predicting protein function from sequence and structure. Nat Rev Mol Cell Biol 2007, 8:995-1005.
6. Looeweinstein Y, Raimondi D, Redfern O, Watson J, Frishman D, Linial M, Orengo C, Thornton J, Tramontano A: Protein function annotation by homology-based inference. Genome Biology 2009, 207.
7. Sleator RO, Walsh P: An overview of in silico protein function prediction. Arch Microbiol 2010, 192:151-155.
8. Chi X, Hou J, Erkin S, Lisewski AM, Lichartse O: An Iterative Approach of Protein Function Prediction: towards integration of similarity metrics. BMC Bioinformatics 2011, 12:437.
9. Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ: Gapped BLAST and PSIBLAST: A new generation of protein database search programs. Nucleic Acids Res 1997, 25:3389-3402.
10. Holm L, Kaarna-Roos S, Wilton C, Plewniak D: Using Dali for structural comparison of proteins. Curr Protoc Bioinformatics 2006, Chapter 5: Unit 5.5.
11. Shindyalov IN, Bourne PE: Protein structure alignment by incremental combinatorial extension (CE) of the optimal path. Protein Eng 1998, 11:737-747.
12. Ilyin VA, Abyzov A, Leskin CM: Structural alignment of proteins by a novel TOPOFIT method, as a superimposition of common volumes at a topomax point. Protein Sci 2004, 13:1866-1874.
13. Rost B: Enzyme function less conserved than anticipated. J Mol Biol 2002, 318:595-608.
14. Ivannenko VA, Pintus S, Grigorovich DA, Kolchanov NA: PDBSiteScan: a program for searching for active, binding and posttranslational modification sites in the 3D structures of proteins. Nucleic Acids Res 2004, 32:W549-W554.
15. Meng EC, Polacco BJ, Babbitt PC: Superfamily active site templates. Proteins 2004, 55:962-976.
16. Binkowski T, Joachimkam A, Liang J: Protein surface analysis for function annotation in high-throughput structural genomics pipeline. Protein Science 2005, 14:2972-2981.
17. Shulman-Peleg A, Nussinov R, Wolfson H: SiteEngines: recognition and comparison of binding sites and protein-protein interfaces. Nucleic Acids Res 2005, 33:W337-W341.
18. Laskowski RA, Watson JD, Thornton JM: ProFunc: a server for predicting protein function from 3D structure. Nucleic Acids Res 2005, 33:W89-W93.
19. Parasaram R, Lee JS, Yin P, Somarowthu S, Ondrechen MJ: Functional classification of 3D protein structures from predicted local interaction sites. J Bioinform Comput Biol 2010, 8(Suppl 1):11-15.
20. Goldsmith-Fischman S, Honig B: Structural genomics: computational methods for structure analysis. Protein Sci 2003, 12:1813-1821.
21. Laskowski RA, Watson JD, Thornton JM: From protein structure to biochemical function. J Struct Funct Genomics 2003, 4:167-177.
22. Tong W, Wei Y, Murga LF, Ondrechen MJ, Williams RJ: Partial Order Optimum Likelihood (POOL): Maximum Likelihood Prediction of Protein Active Site Residues Using 3D Structure and Sequence Properties. PLoS Comp Biol 2009, 5:e1000266.
23. Somarowthu S, Yang H, Hildebrandt DGC, Ondrechen MJ: High-performance prediction of functional residues in proteins with machine learning and computed input features. Biopolymers 2011, 95:390-400.
24. Ko J, Murga LF, André P, Yang H, Ondrechen MJ, Williams RJ, Agouwamba A, Buddé DE: Statistical criteria for the identification of protein active sites using theoretical microscopic titration curves. Proteins 2005, 59:183-195.
25. Wei Y, Ko J, Murga LF, Ondrechen MJ: Selective Prediction of Interaction Sites in Protein Structures with THEMATICS. BMC Bioinformatics 2007, 8:119.
26. Sankaranaman S, Spjølker K: INTREPID: Information-theoretic TREe traversal for Protein functional site IDentification. Bioinformatics 2008, 24:2445-2452.
27. Sankaranaman S, Kolarzowski B, Spjølker K: INTREPID: a web server for prediction of functionally important residues by evolutionary analysis. Nucleic Acids Res 2009, 37:W390-W395.
28. Capra JA, Laskowski RA, Thornton JM, Singh M, Funkhouser TA: Predicting Protein Ligand Binding Sites by Combining Evolutionary Sequence Conservation and 3D Structure. PLoS Comput Biol 2009, 5:e1000585.

29. Notredame C, Higgins DG, Heringa J: T-Coffee: A novel method for multiple sequence alignments. J Mol Biol 2000, 302:205-207.

30. Eddy SR: Where did the BLOSUM62 alignment score matrix come from? Nature Biotechnology 2004, 22:1035-1036.

31. Harris P, Poulsen JN, Jensen K, Larsen S: Substrate binding induces domain movements in orotidine 5'-monophosphate decarboxylase. J Mol Biol 2002, 318:1019-1029.

32. Wu N, Mo Y, Gao J, Pai E: Structure and mechanism of the enzyme orotidine monophosphate decarboxylase. Proc Natl Acad Sci (USA) 2000, 97:2017-2022.

33. Wierenga RK: The TIM-barrel fold: A versatile framework for efficient enzymes. FEBS Lett 2001, 492:193-198.

34. Vedadi M, Lew J, Arz J, Amemi M, Zhao Y, Dong A, Wasney G, Gao M, Hills T, Broks S et al: Genome-scale protein expression and structural biology of Plasmodium falciparum and related Apicomplexan organisms. Molecular and Biochemical Parasitology 2007, 151:100-110.

35. Appleby TC, Kinsland C, Begley TP, Ealick SE: The crystal structure and mechanism of orotidine 5'-monophosphate decarboxylase. Proc Natl Acad Sci USA 2000, 97:2005-2010.

36. Harris P, Navarro Poulsen JC, Jensen KF, Larsen S: Structural basis for the catalytic mechanism of a proficient enzyme: orotidine 5'-monophosphate decarboxylase. Biochemistry 2000, 39:4217-4224.

37. Holm L, Park J: DaliLite workbench for protein structure comparison. Bioinformatics 2000, 16:566-567.

38. Hahn M, Keitel T, Heinemann U: Crystal and molecular structure at 0.16-nm resolution of the hybrid Bacillus endo-1,3-1,4-beta-D-glucan 4-glucanohydrolase H(A16-M). Eur J Biochem 1995, 232:849-858.

39. Hakulinen N, Turunen O, Janis J, Lesola M, Rouvinen J: Three-dimensional structures of thermophilic beta-1,4-xylanases from Chaetomium thermophilum and Nonomuraea flexuosa. Eur J Biochem 2003, 270:1399-1412.

40. Muller-Newen G, Janssen U, Stoffel W: Enoyl-CoA hydratase and isomerase form a superfamily with a common active-site glutamate residue. Eur J Biochem 1995, 228:68-73.

41. Bell AF, Feng Y, Hofstein HA, Parikh S, Wu J, Rudolph MJ, Kisker C, Whitty A, Tonge PJ: Stereoselectivity of enoyl-CoA hydratase results from preferential activation of one of two bound substrate conformers. Chem Biol 2002, 9:1247-1255.

42. Bennett JP, Whittingham JL, Brazowski AM, Leonard PM, Grogan G: Structural characterization of a beta-diketone hydrolase from the cyanobacterium Anabaena sp. PCC 7120 in native and product-bound forms, a coenzyme A-independent member of the crotonase superfamily. Biochemistry 2007, 46:1317-144.

43. Han GW, Ko J, Farr CL, Deller MC, Xu Q, Chiu H-J, Miller MD, Sefcikova J, Somarowthu S, Beuning PJ et al: Crystal structure of a metal-dependent phosphoesterase (YP_910028.1) from Bifidobacterium adolescentis: Computational prediction and experimental validation of phosphoesterase activity. Proteins 2011, 79:2146-2160.

http://www.biomedcentral.com/1471-2105/14/S3/S13