NMR Structure of Lipoprotein YxeF from \textit{Bacillus subtilis} Reveals a Calycin Fold and Distant Homology with the Lipocalin Blc from \textit{Escherichia coli}

Yibing Wu\textsuperscript{1,6,a}, Marco Punta\textsuperscript{2,6,b}, Rong Xiao\textsuperscript{3,6}, Thomas B. Acton\textsuperscript{3,6}, Bharathwaj Sathyamoorthy\textsuperscript{1}, Fabian Dey\textsuperscript{4,6,a}, Markus Fischer\textsuperscript{4,6,d}, Arne Skerra\textsuperscript{5}, Burkhard Rost\textsuperscript{2,6}, Gaetano T. Montelione\textsuperscript{3,6}, Thomas Szyperski\textsuperscript{1,6,a}

1 Department of Chemistry, State University of New York at Buffalo, Buffalo, New York, United States of America, 2 Department of Computer Science and Institute for Advanced Study, Technical University of Munich, Munich, Germany, 3 Center of Advanced Biotechnology and Medicine, Department of Molecular Biology and Biochemistry, Robert Wood Johnson Medical School, The State University of New Jersey, Piscataway, New Jersey, United States of America, 4 Howard Hughes Medical Institute, Department of Biochemistry and Molecular Biophysics, Center for Computational Biology and Bioinformatics, Columbia University, New York, New York, United States of America, 5 Munich Center for Integrated Protein Science, CIPS-M, and Lehrstuhl für Biologische Chemie, Technische Universität München, Freising-Weihenstephan, Germany, 6 Northeast Structural Genomics Consortium

Abstract

The soluble monomeric domain of lipoprotein YxeF from the Gram positive bacterium \textit{B. subtilis} was selected by the Northeast Structural Genomics Consortium (NESG) as a target of a biomedical theme project focusing on the structure determination of the soluble domains of bacterial lipoproteins. The solution NMR structure of YxeF reveals a calycin fold and distant homology with the lipocalin Blc from the Gram-negative bacterium \textit{E.coli}. In particular, the characteristic \(\beta\)-barrel, which is open to the solvent at one end, is extremely well conserved in YxeF with respect to Blc. The identification of YxeF as the first lipocalin homologue occurring in a Gram-positive bacterium suggests that lipocalins emerged before the evolutionary divergence of Gram positive and Gram negative bacteria. Since YxeF is devoid of the \(\alpha\)-helix that packs in all lipocalins with known structure against the \(\beta\)-barrel to form a second hydrophobic core, we propose to introduce a new lipocalin sub-family named ‘slim lipocalins’, with YxeF and the other members of Pfam family PF11631 to which YxeF belongs constituting the first representatives. The results presented here exemplify the impact of structural genomics to enhance our understanding of biology and to generate new biological hypotheses.

Introduction

The lipoprotein YxeF from \textit{Bacillus subtilis} was selected by the Northeast Structural Genomics Consortium (NESG; http://www.nesg.org) as a target \(\{\text{gi\|55674274, SwissProt/TriEMBL ID YXEF_BACSU, access number P54945, NESG target ID SR500A}\}\) of a biomedical theme project focusing on the structure determination of the soluble domains of bacterial lipoproteins [1,2]. YxeF exhibits no significant sequence similarity with any protein with known three-dimensional structure and is one of only eight members forming Pfam [3] family PF11631 for which no functional annotation is available (Pfam 26.0 release). All members of the family are from the genus \textit{Bacillus} and present high sequence similarity to YxeF, with A7ZAF5 and E1UTS8 from \textit{B. amyloliquefaciens} being the most distant homologues (61\% sequence identity to YxeF over 129 residues).

Bacterial lipoproteins represent a class of secreted, membrane-anchored proteins that are conserved throughout bacteria and play critical roles in a wide range of biological processes, including bacterial pathogenesis and host immune response [1]. They contain a conserved N-terminal type II signal peptide also known as the ‘lipobox’, which is immediately followed by an invariant Cys residue. After cleavage of the signal peptide, the lipoprotein is anchored into the bacterial membrane via a diacylglycerol moiety forming a thioredoxin linkage to the Cys side chain as well as a fatty acid moiety coupled to the N-terminus.

In general, lipobox sequences exhibit the consensus sequence \([\text{Leu/Val/Ile}}\text{-}[\text{Ala/Ser/Thr/Va}]\text{l/Ile}\text{]-[Gly/Ala/Ser]}\text{-}[\text{Cys]}\text{[Val-}
Ser-Gly-Cys in YxeF). The residue following the Cys is important for localization of the lipoprotein [1]. In Gram-negative bacteria the lipoprotein is usually anchored in the inner membrane if Cys is followed by Asp, whereas otherwise it is anchored in the inner leaflet of the outer membrane [2]. Although Cys is followed by Glu in YxeF, it is supposedly anchored in the only lipid membrane of the Gram-positive Bacillus subtilis cell. Here we report the high-quality NMR solution structure of the soluble domain of protein YxeF comprising residues 19–144, along with a structural bioinformatics analysis to classify its structure and to gain new insights into its evolutionary origin.

**Results and Discussion**

**NMR Structure of the Soluble Domain of Lipoprotein YxeF**

A high-quality NMR structure of the soluble domain (comprising residues 19–144) of lipoprotein YxeF was obtained by multidimensional NMR spectroscopy [Figures 1A, 2; Table 1; PDB (protein data bank) accession number 2JOZ]. For simplicity, we henceforth refer to the structure of this domain as to the ‘YxeF structure’. YxeF contains nine β-strands (A to D and D’ to H) comprising residues 36–40, 52–57, 61–68, 72–75, 79–86, 91–97, 104–109, 114–119, and 122–129, respectively [Figure 2; strand assignment according to STRIDE [4]]. Since strands D and D’ point into the same direction and are connected by a short coil region in a mostly extended conformation (comprising Pro 78), we will refer to this entire polypeptide segment as strand D (Figure 2). All β-strands are then arranged in anti-parallel fashion forming a +1 up-and-down β-barrel and are connected by seven loops L1 to L7. The β-barrel is closed on one side (loops L2, L4, L6), primarily by dense side chain packing of a number of hydrophobic and aromatic residues, including Phe 33, Tyr 34, Tyr 35, Trp 38, which includes lipocalins, fatty acid binding proteins, triabin, avidins/streptavidins and a class of metalloprotease inhibitors. All similar proteins are calycins from all groups alluded to above. Top hits span a quasi-continuum of Z-scores between 7.7 (top hit) and 2.1 (taken as a lower limit of significance). Among the structurally similar proteins are calycins as well as lipocalins, fatty acid-binding proteins and triabin. Avidin/streptavidin and some metalloprotease inhibitors are not considered to be part of the ‘calycin superfamily’. These classifications are (i) based on both sequence and structure comparisons, (ii) rely, at least to some degree, on manual curation, and (iii) favor the hypothesis that an evolutionary link exists between lipocalins, fatty acid binding proteins and triabin. They leave, however, the tetrameric avidins/streptavidins and some metalloprotease inhibitors in limbo with respect to their relationship to the other proteins alluded to above.

**YxeF Structure Belongs to the Calycin Superfamily**

Calycins feature a conserved Gly-X-Trp/Arg signature motif (Figure 5), in which the Arg side chain is located on strand H, interacts with the Trp side chain located on strand A and also forms hydrogen bonds with the backbone carbonyl groups of some other N-terminal residues [5,11,13]. Consistently, this motif has been shown to be important for protein stability in the retinol-binding protein, a prototypic member of the lipocalin family [14,15]. In YxeF, the motif is entirely conserved (Gly 36, Trp 38 and Arg 128), although the conformation of the side chain of Arg 128 is rather poorly defined in the NMR structure (Figure 5B). Conservation of the calycin signature motif and of the β-barrel topology (Figure 4) renders straightforward the classification of YxeF as a ‘calycin’.

**DALI-based Search for Similar Structures**

A search of the PDB using the program DALI [16] for proteins which are structurally similar to YxeF yielded more than 700 significant hits (152 when considering PDB90 or a set of PDB proteins redundancy reduced at 90% sequence identity). These hits span a quasi-continuum of Z-scores between 7.7 (top hit) and 2.1 (taken as a lower limit of significance). Among the structurally similar proteins are calycins from all groups alluded to above. Top hits include the cysteine protease inhibitor staphostatin B (Z-score = 7.7), several 10-stranded fatty acid binding proteins (top Z-score = 7.5), the C-terminal domain of a self-compartmentalizing protease PabB from Pyrococcus abyssi (Z-score = 7.2) which has been claimed [17] to be the first domain with lipocalin-like architecture from Archaea, and several lipocalins (top Z-score 6.9).
Figure 1. 2D \([^{15}N,^{1}H]\) HSQC spectra of lipoprotein YxeF. (A) Spectrum recorded for the sample used for NMR structure determination at 750 MHz \(^1H\) resonance frequency. Resonance assignments are indicated using the one-letter amino acid code. Signals arising from side chains (Asn \(H_d^{\alpha}/N_d^{\alpha}\), Gln \(H_e/N_e\), Arg \(H_e/N_e\) and Trp \(H_e^{\alpha}/N_e^{\alpha}\)) are labeled with (*) and folded signals are designated with (f) next to the residue number. Signals arising from the His purification tag were not sequence specifically assigned. The spectral region indicated by dotted lines comprises most of the signals arising from the \(\beta\)-barrel (Figure 2) and is displayed for the spectra shown in (B). Those were recorded at different temperatures at 500 MHz \(^1H\) resonance frequency (see text).

doi:10.1371/journal.pone.0037404.g001

Figure 2. NMR structure of the soluble domain of lipoprotein YxeF. Only residues 34–132 are shown because the terminal polypeptide segments are flexibly disordered in solution. The N-terminal residue Phe 34 and the C-terminal residue Ser 132 are labeled. Lines point at the \(\Omega\)-type loop, which connects \(\beta\)-strands A and B and is poorly defined in the NMR structure. (A) Ribbon drawing of the first conformer of PDB entry 2JOZ with \(\beta\)-strands being depicted in cyan. (B) Superposition of the 20 conformers representing the NMR solution structure obtained after superposition of the backbone heavy atoms (N, C\(^{\alpha}\) and C\(^\circ\)) of the \(\beta\)-strands. (C) "Sausage" representation of backbone and best-defined side chains; a spline function was drawn through the mean positions of C\(^\circ\) atoms with the thickness being proportional to their mean global displacement in the 20 conformers after superposition as in B). The figure was generated using the program MOLMOL [59].

doi:10.1371/journal.pone.0037404.g002
Staphostatin B is classified in SCOP as having a ‘Streptavidin-like’ fold but high structural similarity to lipocalins was recognized previously [18]. It should be noted, however, that in spite of the high Z-score returned by overall structure superposition, topology of strand A to C in staphostatin’s \( b \)-barrel is very different with respect to the one observed in both streptavidins and lipocalins. Other calycin structures result in lower DALI Z-scores (avidin’s Z-score is 5.5).

Overall structural similarity identified with DALI thus seems to clearly indicate evolutionary relatedness between YxeF, fatty-acid binding proteins and lipocalins. Two structural features shared by YxeF and lipocalins, however, strongly suggest that YxeF is closer to the latter than to fatty-acid binding proteins (Figure 4): (i) the \( b \) strands forming the \( b \)-barrel (versus 10 in fatty-acid-binding proteins), and (ii) the presence of an \( \Omega \)-type loop’ connecting \( b \)-strands A and B (in fatty-acid binding proteins two short \( \alpha \)-helices are inserted between strands A and B). Notably, the \( \Omega \)-type loop that often acts as a flexible lid for the open end of the \( b \)-barrel in lipocalins appears to be disordered in YxeF (Figure 2). On the other hand, as mentioned above, YxeF lacks two structural features conserved in the lipocalins (Figure 4): an N-terminal 310-helix, which is sometimes replaced by a longer \( \alpha \)-helix [19,20], and a C-terminally located \( \alpha \)-helix followed by an additional \( b \)-strand [21,22].

### Table 1. Statistics of YxeF(19–144) NMR Structure.

|                     | Value                  |
|---------------------|------------------------|
| Completeness of stereo-specific assignments* [%] |                       |
| \( \text{CH}_2 \) of Gly | 8 (1/13)               |
| \( \text{CH}_3 \)             | 20 (16/80)             |
| Val and Leu methyl groups | 79 (11/14)            |
| Conformationally-restricting distance constraints |                       |
| Intraresidue \(| i = j | 462                  |
| Sequential \(| i = j | 556                  |
| Medium Range \(| 1 < | i | j | < 5 | 148                |
| Long Range \(| i = j | > 5 | 662                |
| Total                | 1828                  |
| Dihedral angle constraints \((\phi/\psi| 49/49               |
| Number of constraints per residue | 15.2               |
| Number of long-range distance constraints per residue | 5.2               |
| CYANA target function \([	ext{Å}^2]| 1.14±0.15            |
| Average number of distance constraints violations per CYANA conformer |                       |
| 0.2–0.5 Å | 0                      |
| >0.5 Å   | 0                      |
| Average number of dihedral-angle constraint violations per CYANA conformer >5° | 1.0                  |
| Average r.m.s.d. to the mean CNS coordinates \([	ext{Å}| 0.51±0.13            |
| Regular secondary structure elementsa, backbone heavy atoms |                       |
| Ordered residuesb, backbone heavy atoms | 0.81±0.14            |
| Ordered residesc, all heavy atoms | 1.26±0.15            |
| Heavy atoms of molecular core including best-defined side chainsd | 0.55±0.09            |
| PROCHECK\({^e}\) G-factors raw score \((\phi \text{ and } \psi/\text{all dihedral angles}) | −0.51/−0.38          |
| PROCHECK\({^e}\) G-factors Z-score \((\phi \text{ and } \psi/\text{all dihedral angles}) | −1.69/−2.25          |
| MOLPROBITY\({^f}\) clash score \((\text{raw/Z-score}) | 17.34/1.45            |
| PRF R/P/DP scores\({^g} [%| 0.99/0.87/0.85          |
| Ramachandran plot summary c [%] |                       |
| most favored regions | 93.0                  |
| Additionally allowed regions | 6.5                |
| generously allowed regions | 0.2                |
| disallowed regions | 0.2                  |

*Relative to pairs with non-degenerate chemical shifts.

aResidues 37–40, 53–58, 62–69, 73–76, 80–86, 92–98, 105–110, 116–120, and 123–128.

bResidues 33–40, 45–48, 51–87, 93–96, 99–101, 104–111, 115–131.

cBackbone and side-chain heavy atoms of residues 37–39, 54–59, 63–64, 66, 68, 75, 77, 81–82, 84–86, 93–96, 98, 105–108, 115–118. Best-defined side chains are those exhibiting a displacement of less than 1 Å for their side chain heavy atoms after superposition of the \( b \)-strands for minimal r.m.s.d.

dBackbone and side-chain heavy atoms of residues 37–39, 54–59, 63–64, 66, 68, 75, 77, 81–82, 84–86, 93–96, 98, 105–108, 115–118. Best-defined side chains are those exhibiting a displacement of less than 1 Å for their side chain heavy atoms after superposition of the \( b \)-strands for minimal r.m.s.d.

dBackbone and side-chain heavy atoms of residues 37–39, 54–59, 63–64, 66, 68, 75, 77, 81–82, 84–86, 93–96, 98, 105–108, 115–118. Best-defined side chains are those exhibiting a displacement of less than 1 Å for their side chain heavy atoms after superposition of the \( b \)-strands for minimal r.m.s.d.

eResidues 30–40, 45–48, 51–87, 93–96, 99–101, 104–111, 115–131.

fResidues 37–39, 54–59, 63–64, 66, 68, 75, 77, 81–82, 84–86, 93–96, 98, 105–108, 115–118. Best-defined side chains are those exhibiting a displacement of less than 1 Å for their side chain heavy atoms after superposition of the \( b \)-strands for minimal r.m.s.d.

gResidues 37–39, 54–59, 63–64, 66, 68, 75, 77, 81–82, 84–86, 93–96, 98, 105–108, 115–118. Best-defined side chains are those exhibiting a displacement of less than 1 Å for their side chain heavy atoms after superposition of the \( b \)-strands for minimal r.m.s.d.

**doi:**10.1371/journal.pone.0037404.t001

Related to this is the observation that the \( b \)-barrel in lipocalins is flanked by a globular domain containing the binding cavity, whereas in YxeF (Figure 2) these domains are disordered.
absence of the C-terminal α-helix and β-strand remain as a stark structural difference when comparing the structure of YxeF with known lipocalin structures.

YxeF and Lipocalin Blc from E. coli are Distant Homologues

To further refine our structural analysis, we compared in detail the structure of YxeF with that of lipoprotein Blc from E. coli (Figure 5), the only bacterial lipocalin for which an atomic-resolution structure is currently available (PDB ID: 3MBT) [23]. It has been suggested that lipocalins can be grouped into ‘kernel’ or ‘outlier’ lipocalins [11] depending on the presence or absence of three so-called ‘structurally conserved regions’ (SCRs) which correlate to some degree with sequence conservation (Figure 5A). The most conserved SCR1 comprises the N-terminal Gly-X-Trp segment of the calycin Gly-X-Trp/Arg signature motif located on β-strand A, while SCR3 (formed primarily by residues found on β-strand H) contains the Arg residue as part of the same motif. SCR2, instead, spans the region between the termini of β-strands F and G including the loop L6 connecting the two β-strands. Because the SCR2 and SCR3 sequence motifs are not conserved in Blc, it was initially classified as an ‘outlier lipocalin’ [24]. However, this sequence-based classification does not appear to be well justified when considering that the X-ray structure of Blc revealed a remarkably close structural similarity with ‘kernel’ lipocalins [23].

Like YxeF, Blc is monomeric and does not contain disulfide bridges, which are frequently found in other lipocalins. While the overall structural alignment of YxeF and Blc results in a highly significant, but not the highest DALI Z-score ([Z-score = 6.9; 2.7 Å r.m.s.d. (root mean square deviation) for superposition of the Cα atoms of 88 aligned residues exhibiting 17% sequence identity], the structural similarity of the two β-barrels (48 residues) is truly striking: they superimpose with a backbone r.m.s.d. of only 1.8 Å (Figure 5). Indeed, the structural alignment of only the β-barrels yields the highest Z-score for Blc among a selection of full structure alignments.
alignment top hits (Table 2; note that in these comparisons we consider conformer 1 of the ensemble representing the YxeF solution structure 2JOZ). The corresponding values calculated for avidin, which contains a β-barrel of evidently different shape (Figure 6), are also provided in Table 2 for comparison. The only minor structural difference between the YxeF and Blc β-barrels, possibly reflecting different ligand specificities, relates to β-strand A at the base of the Ω-type loop. This strand is shorter in YxeF where it creates a small V-shaped aperture on the side of the β-barrel (Figure 6).
As indicated above, the C-terminally located α-helix characteristic of lipocalins is not present in YxeF (Figure 4). In Blc, this α-helix packs against the outside of the β-barrel, primarily against β-strands G and H (Figure 5C). As a result, a second hydrophobic core is formed, adding to the one found in the lower, closed part of the β-barrel itself (Figure 5D). In YxeF, the corresponding C-terminally located polypeptide segment is highly polar and flexibly disordered in solution. The absence of hydrophobic residues in this segment and also on the exterior of β-strands G and H apparently prevents the formation of a second hydrophobic core. Since this core has been shown to be important for the stability of lipocalins, and Blc exhibits a comparably low melting temperature of ~45°C even with this second core, the stability of the well-defined fold of YxeF evidenced by our high-quality structure (Figure 2) is somewhat unexpected. Intriguingly, although the onset of protein precipitation at higher temperature prevented us from accurately determining a heat denaturation ‘melting’ temperature, the inspection of 2D [15N,1H] HSQC (heteronuclear single quantum coherence) spectra recorded up to ~50°C revealed that YxeF’s β-barrel is intact even at such elevated temperatures (Figure 1B).

Taken together, in spite of the lack of the C-terminally located α-helix, the strong structural similarity of the β-barrels of YxeF and Blc (Table 2), together with a remarkably similar relative spatial orientation of the Trp and Arg residues of the Gly-X-Trp/Arg signature motif (Figure 5B), reveals that YxeF and lipocalin Blc are distant homologues. This is consistent with the fact that both YxeF and Blc are secretory lipoproteins, a characteristic common to most predicted lipocalins in Gram-negative Bacteria [24].

### Homology Model of Protein A7ZAF5: Insights into Putative Ligand Binding

The only proteins known to show significant sequence identity with YxeF are found in the Bacillus genus, e.g., D5MWC1 from B. subtilis strain ATCC 6633, E3E109 form B. atrophaeus strain 1942, and the somewhat more distant homolog A7ZAF5 from B. amyloliquefaciens (61% identity; member of PF11631; see Figure 3E for a multiple sequence alignment of all members). In D5MWC1 and E3E109, the three glutamate residues that confer a negative charge distribution to YxeF’s putative ligand binding site (Figure 3A,C) are conserved. Despite the high overall sequence similarity, these glutamates are absent in A7ZAF5. Interestingly, however, in a homology model created for the soluble domain of A7ZAF5 based on conformer 1 of the YxeF NMR structure (2JOZ), it appears that other negatively charged residues, i.e., Glu 52, Asp 78, Glu 121 (residue numbers as in the UniProt [25] sequence of A7ZAF5) create a similar surface charge distribution within the corresponding cavity of A7ZAF5 (Figure 3B,D,E; note that Glu 121 is replaced by Gly in E1UTS8, while additionally conserved in E3E109). Considering the robustness of homology modeling at ~60% pairwise sequence identity and the absence of any gaps in the alignment, this finding suggests that these proteins may function by binding yet to be identified ligands that share similar electrostatic properties.

### Evolutionary Origin of Lipocalins and Protein YxeF

Lipocalins are a functionally diverse group of proteins that usually bind small hydrophobic molecules. Bacterial lipocalins, in particular, have been proposed to be implicated in biogenesis of the outer membrane of Gram-negative bacteria based on three findings: (i) Blc from E. coli has been shown to bind fatty acids and (lyso-)phospholipids [26], suggesting its participation in lipid metabolism [26], (ii) no lipocalin had so far been identified in microorganisms that lack an outer lipid membrane, that is, Gram-positive bacteria or Archaea, and (iii) some integral outer membrane proteins exhibit significant structural similarity with lipocalins [21,27].

Our finding that YxeF and Blc are distant homologues suggests to classify YxeF (and thus all currently known members of PF11631 exhibiting high sequence similarity; Figure 3E) as a hitherto unknown type of lipocalin constituting a new sub-family characterized by its distinct ‘β-barrel only’ architecture. We suggest to name this new sub-family ‘slim lipocalins’. It is then of key importance for our understanding of the evolution of lipocalins.

### Table 2. Comparison of β-barrels occurring in selected structures* yielding top DALI Z-scores after full structure alignment with YxeF.

| Structure compared with YxeF structure | β-barrel only | DALI Z-score | r.m.s.d.(Å) | Number of residues | Sequence identity | DALI Z-score for entire structure |
|---------------------------------------|--------------|--------------|-------------|-------------------|------------------|---------------------------------|
| 1Y4H(C)                               | 4.4          | 1.7          | 40          | 13%               | 7.7              |                                 |
| 3D95(B)                               | 4.2          | 2.7          | 49          | 18%               | 7.5              |                                 |
| 1JJJA(A)                              | 4.2          | 2.9          | 50          | 12%               | 7.4              |                                 |
| 2OML_C(B)                             | 3.9          | 1.9          | 40          | 8%                | 7.2              |                                 |
| 3AKM(A)                               | 4.2          | 3.0          | 50          | 12%               | 7.1              |                                 |
| 1P6P                                  | 4.5          | 2.9          | 49          | 8%                | 7.1              |                                 |
| Blc                                   | 5.7          | 1.8          | 48          | 15%               | 6.9              |                                 |
| 2CAM(A)                               | 3.8          | 2.3          | 46          | 13%               | 5.3              |                                 |

*Structures were selected among those with the top DALI Z-score (see text) when aligned with the entire folded YxeF domain (residues 32–132), and they are ranked according to the Z-score from that comparison (right-most column). The location of the β-strands forming the β-barrels were identified using the program STRIDE and pairwise structural comparisons of the β-barrels was again performed using the program DALI. R.m.s.d. values were calculated after superposition of the Cα atoms. Structures of proteins other than E. coli Blc (seventh row, highlighted in bold) are designated by their PDB IDs: 1Y4H (chain C is staphylin B; 3D95 (chain B) is the cellular retinoic acid-binding protein β; 1JJJA (chain A) is the human epidermal-type fatty acid-binding protein; 2OML_C (chain B) is the C-terminal domain of protease Pab87; 3AKM (chain A) is the human intestinal fatty acid binding protein; 1P6P is the toad liver basic fatty acid-binding protein. For comparison, the values for avidin (PDB-ID 2CAM, chain A) are also provided (bottom row; see also Figure 6).
that YxeF is from the Gram-positive bacterium *B. subtilis*. It has been suggested that lipocalins first emerged along with the evolution of the outer membrane in Gram-negative bacteria [21]. The identification of distant homology between Blc and YxeF now suggests instead that lipocalins may have emerged before the divergence of Gram-positive and Gram-negative bacteria [28]. Although the presence of YxeF in *B. subtilis* might also have resulted from horizontal gene transfer, the genomic island predictor Alien_hunter [29] applied to genomic regions in and around the yxeF gene in both *B. subtilis* and *B. amyloliquefaciens* did not provide any support for this hypothesis. Hence, based on the evidence collected so far, it is thus indeed tempting to speculate that an ancient lipocalin, like protein YxeF devoid of a second hydrophobic core, evolved into both YxeF-like proteins of Gram-positive bacteria and the lipocalins present nowadays in Gram-negative bacteria and Eukaryotes. Additional structural variability of the β-barrel, required to convey specificity for other physiological ligands, might have co-evolved with the formation of the second hydrophobic core and, possibly, the disulfide bridges present in many eukaryotic lipocalins in order to maintain, or even increase protein stability [30].

As indicated above, the C-terminal domain of a self-compartmentalizing protease Pab87 from *Pyrococcus abyssi* has been suggested to be the first domain with lipocalin-like architecture from Archaea [17]. Notably, the Pab87 structure (i) ranks high among the top DALI structural matches for YxeF (Z-score 7.2; r.m.s.d. 2.8 Å, 76 aligned residues, 5% sequence identity), (ii) likewise represents a slim ‘β-barrel only’ structure, and (iii) contains a calycin signature motif (Gly-X-Tyr/Lys). However, the DALI Z-score for comparison of the β-barrels (Table 2) is comparably low, and the domain does not exhibit the ‘Ω-type’ loop characteristic of lipocalins. Consistently, its function was linked to protein oligomerization (and thus compartmentalization of the active site of the full-length protein) and not to the binding of a ligand in a fashion typically observed for lipocalins. Intriguingly, however, comparison of the structures of YxeF and the C-terminal domain of Pab87 indicates that possibly a very ancient lipocalin may have existed, even before the divergence of Bacteria and Archaea occurred.
Prospects for Protein Design

The discovery of the ‘slim’ lipocalin YxeF devoid of the C-terminal α-helix (and of the entire second hydrophobic core) as well as disulfide bridges also provides a promising new scaffold for the design of lipocalins with novel ligand binding functions, so-called ‘anticalins’ [31]. The four structurally variable loops that form the entrance to the ligand pocket at the open end of a lipocalin’s β-barrel share functional similarity with the six hypervariable loops (CDRs) of antibodies. When compared with immunoglobulins, however, lipocalins are much smaller (~160–180 residues), comprise only a single polypeptide chain, and can be produced at high yields in microbial host cells. Using targeted randomization of the structurally variable loop region in combination with phage display selection, anticalins with novel specificities have been engineered for the high affinity complexation of both low molecular weight compounds and protein antigens [7,31,32].

In many cases the lipocalin β-barrel is thermally rather stable and tolerates a wide range of amino acid substitutions at the ligand-binding site. Melting temperatures of natural lipocalins are often above 70°C [30] and can range beyond 95°C, for example, for human tear lipocalin. As mentioned above, Bcl is actually a notable exception having a melting temperature of just ~45°C. Attempts to engineer anticalins that lack the C-terminal α-helix were thus far not successful. Hence, the soluble domain of YxeF may turn out to be a promising target for the design of a novel line of minimal ‘β-barrel-only’, or ‘slim’ anticalins that can be used as reagents for bioanalytical purposes or separation tasks. It is evident that NMR and X-ray crystallographic studies [23,33] will continue to be of key importance for this endeavor. Finally, since endogenous lipocalins are believed to play a role in antibiotic resistance and activation of immunity in Gram-negative bacteria, lipocalins of Gram-positive bacteria might turn out to be relevant biomedical targets themselves, e.g., for the development of new antibiotics [24].

Conclusions

The structure of the soluble domain of lipoprotein YxeF from the Gram-positive B. subtilis revealed an unexpected distant homology with lipocalin Bcl from Gram-negative E. coli. Because YxeF is devoid of a second hydrophobic core typical for all lipocalins, we propose to introduce a new lipocalin sub-family named the ‘slim lipocalins’, with the members of Psm family PF11631 being the first known representatives. The identification of YxeF as the first lipocalin homologue from a Gram-positive bacterium has far reaching consequences for our understanding of the evolution of this important class of proteins: lipocalins may have emerged well before the evolutionary divergence of Gram-positive and Gram-negative bacteria. Furthermore, we expect that the discovery of the ‘slim lipocalin’ YxeF will impact design of new anticalins with prescribed binding specificities. The results presented in this publication thus exemplify the role of structural genomics to generate new biological hypotheses and to support protein design efforts.

Materials and Methods

NMR Sample Preparation

The soluble domain of protein YxeF (excluding the ‘lipobox’ signal sequence) was cloned, expressed and purified following standard protocols developed by the NESG for production of uniformly 13C,15N and 5%-13C,15N-labeled protein samples [34,35]. Briefly, the truncated yxeF gene from Bacillus subtilis containing residues 19–144 was cloned into a pET21 (Novagen) derivative, yielding plasmid SR500A-21.2. The resulting construct contains eight nonnative residues at the C-terminus (LEHHHHHHH) that facilitate protein purification. This expression vector, NESG SR500A-21.2, has been deposited in the PSI Materials Repository (http://psimr.asu.edu/).

Using BL21 (DE3) pMGK cells, a codon enhanced strain, were transformed with SR500A-21.2 and cultured in M9 minimal medium containing 15NH4Cl and 13C-glucose or 5%-13C-glucose/95% unlabeled glucose [36]. U-13C,15N and 5%-13C,15N-labeled yxeF protein was purified using an AKTAxpress (GE Healthcare) based two-step protocol consisting of IMAC (HiTrap HP), and gel filtration (HiLoad 26/60 Superdex 75) chromatography. The final yield of purified U-13C,15N and 5%-13C,15N protein YxeF (>98% homogenous by SDS-PAGE; 16.1 kDa by MALDI-TOF mass spectrometry) was ~63 mg/L and ~23 mg/L, respectively. In addition, U-15N, 5% biosynthetically directed fractionally 13C-labeled samples were generated to stereo-specifically assign Val and Leu methyl groups [37]. The final concentrations of U-13C,15N and 5%-13C,15N labeled protein samples were about 1.1 mM and 1.2 mM, respectively, in a 90% H2O/10% D2O solution containing 100 mM NaCl, 5 mM CaCl2, 10 mM DTT, 0.02% NaN3, 50 mM 4,4-dimethyl-4-silapentane-1-sulfonic acid (DSS) and 20 mM 2-(N-morpholino)ethanesulfonic acid (MES) at pH = 6.5. An isotropic overall rotational correlation time of 8.6 ns was inferred from 15N spin relaxation times indicating that the protein yxeF is monomeric in solution. This conclusion was confirmed by analytic gel-filtration (Agilent Technologies) followed by a combination of static light scattering and refractive index (Wyatt Technology).

NMR Spectroscopy

NMR spectra were recorded at 25°C on Varian INOVA 600 or 750 spectrometers equipped with cryogenic probes. Five through-bond correlated G-matrix Fourier transform (GFT) NMR experiments complemented by three-dimensional (3D) HNCO as described [38–41], were collected for backbone and side chain resonance assignment (total measurement time: 81 hours). Simultaneous 3D 15N/13C-labeled 13C-caromate-resolved [1H,1H] NOESY (nuclear Overhauser enhancement spectroscopy; mixing time: 60 ms; measurement time: 26 hours) was acquired to derive 1H–1H distance constraints [40]. Two-dimensional (2D) constant-time 13C[1H]-HSQC spectra were recorded as was described for the 5% fractionally 13C-labeled samples to obtain stereo-specific assignments for isopropyl groups of Val and Leu [41]. In order to assess thermal stability of protein YxeF, a series of 2D [13C,1H]-HSQC spectra were recorded for a ~150 μM protein solution between 25°C and about 50°C on a Varian INOVA 500 spectrometer equipped with a conventional probe (total measurement time: 30 hours; Figure 1B). All spectra were processed and analyzed with the programs NMRPipe and XEASY, respectively [42,43].

Sequence specific backbone (1H N, 13N, 1H α, 13C α) and 1Hβ/13Cβ resonance assignments were obtained by using (4,3)D HNNCCα/β/γ/δ/ε/ζ/η/ι/κ/λ/μ/ν/ξ/ω CO/NHN and (4,3)D HNCCα/β/γ/δ/ε/ζ/η/ι/κ/λ/μ/ν/ξ/ω CO/NHN along with the program AUTOASSIGN [44], and polypeptide backbone 13C resonances were assigned using 3D HNCO. More peripheral side chain chemical shifts were assigned with aliphatic (4,3)D HHCH and 3D 13Cα-labeled 13C-caromate-resolved [1H,1H]-NOESY (for details of NESG NMR protocols, see http://www.nmr2.buffalo.edu/nesg.wiki). Overall, assignments were obtained for 99% of the backbone (excluding the N-terminal NHδ), the Pro \(\delta\)N and the \(\gamma\)C preceding prolyl residues; Figure 1A) and 13Cδ, and for 98% of the side chain chemical shifts (excluding Lys NHε, Arg NHε, OH, side chain

PLOS ONE | www.plosone.org 10 June 2012 | Volume 7 | Issue 6 | e37404
including the programs DALI and Skan for identification of MarkUs server integrating a variety of computational tools [55], throughout the paper.

P54945 to residues 19–144, which is the numbering used being the Met start residue of the recombinant protein). The soluble domain only, numbered as residues 2–127 (with residue 1 Amino acid numbers in the PDB coordinate file are those of the RCSB Protein Data Bank (PDB) with accession code 2JOZ [54].

structure quality was assessed with the Protein Structure Validation Software Suite (PSVS) and evaluated by structural genomics

water bath refinement using the program CNS [51]. NMR calculations were performed using CYANA followed by explicit parallel to assign long-range NOEs [47–50]. The final structure derived from chemical shifts using the program TALOS for 16. Holm L, Kaariainen S, Rosenstrom P, Schenkel A (2008) Searching protein 15. Katakura Y, Totsuka M, Ametani A, Kaminogawa S (1994) Tryptophan-19 of 14. Greene LH, Chrysina ED, Irons LI, Papageorgiou AC, Acharya KR, et al. 13C' and aromatic 13C(b) which are assignable with the set of NMR experiments provided above. Furthermore, 79% of Val and Leu isopropyl moieties and 20% of b2-methylene groups with non-degenerate proton chemical shifts were stereo-specifically assigned (Table 1). Chemical shifts were deposited in the BioMagResBank (accession code: 15211) [45]. 1H–1H upper distance limit constraints for structure calculations were extracted from NOESY (Table 1). In addition, backbone dihedral angle constraints were derived from chemical shifts using the program TALOS for residues located in well-defined secondary structure elements [46]. The programs CYANA and AUTOSTRUCTURE were used in parallel to assign long-range NOEs [47–50]. The final structure calculations were performed using CYANA followed by explicit water bath refinement using the program CNS [51], NMR structure quality was assessed with the Protein Structure Validation

Author Contributions

Conceived and designed the experiments: YW RX TBA BS. Analyzed the data: YW MP RX TBA MF AS BR GTM TS. Wrote the paper: YW MP AS BR GTM TS.

Structural Bioinformatics

In silico studies of YxeF were primarily performed using the MarkUs server integrating a variety of computational tools [55], including the programs DALI and Skan for identification of structural similarities and calculation of structural alignments [16,56,57]. Moreover, the DALI pairwise alignment server was used to refine structural comparisons [58]. The programs MOLMOL [59] and STRIDE [4] were used to identify the location of regular secondary structure elements. A homology model was obtained for protein A7ZAF5, that is, one of the most distant known sequence homolog of YxeF, by submitting the YxeF-A7ZAF5 BLAST pairwise sequence alignment to the SWISS-MODEL server in alignment mode [60,61]. Given the high sequence identity between template and target (61%) and the absence of any gaps, the comparative spatial localization of acidic residues inside the b-barrel appears to be robust. Alien_hunter genomic island predictions were obtained via the EnsemblBacteria website [29,62].

Acknowledgments

We thank R. Shastry, C. Ciccossanti, H. Janjua, and G.V.T. Swapna for contributions in sample preparation, and Dr. Masayori Inouye, Rutgers University, for advice regarding the biology and biochemistry of bacterial lipoproteins. Support was also obtained from the University at Buffalo's Center for Computational Research.

References

1. Bahu MM, Priya ML, Selvan AT, Madera M, Gough J, et al. (2006) A database of bacterial lipoprotein (DOLOP), with functional assignments to predicted lipoproteins. J Bacteriol 188: 2761–2773.
2. Tokuda H, Masuyama S (2004) Sorting of lipoproteins to the outer membrane in E. coli. Biochim Biophys Acta 1693: 3–13.
3. Punta M, Coggill PC, Eberhardt RY, Mistry J, Tate J, et al. (2012) The Pfam protein families database. Nucleic Acids Res 40: D290–301.
4. Heinig M, Frishman D (2004) STRIDE: a web server for secondary structure prediction. Nucleic Acids Res 32: W549–560.
5. Flower DR, North AG, Sansom CE (2000) The lipocalin protein family, structural and sequence overview. Biochim Biophys Acta 1482: 9–24.
6. Grzhy Z, Latoszki D, Strzałka K (2006) Lipocalins - a family portrait. J Plant Physiol 163: 895–915.
7. Best G, Schmidt FS, Suhora T, Skerra A (1999) Small antibody-like proteins with prescribed ligand specificities derived from the lipocalin fold. Proc Natl Acad Sci U S A 96: 1908–1913.
8. Greene LH, Lewis TE, Aldous S, Cuff A, Dallman T, et al. (2007) The CATH domain structure database: new protocols and classification levels give a more comprehensive resource for exploring evolution. Nucleic Acids Res 35: D291–297.
9. Andreeva A, Howorth D, Chandona JM, Brenner SE, Hubbard TJ, et al. (2008) Data growth and its impact on the SCOP database: new developments. Nucleic Acids Res 36: D419–425.
10. Finn RD, Mistry J, Tate J, Coggill P, Heger A, et al. (2010) The Pfam protein families database. Nucleic Acids Res 38: D211–222.
11. Flower DR, North AG, Atwood TK (1994) Structure and sequence relationships in the lipocalins and related proteins. Protein Sci 2: 733–761.
12. Sadreyev RI, Kim BH, Grishin NV (2009) Discrete-continuous duality of protein structure space. Curr Opin Struct Biol 19: 321–328.
13. Andreeva A, Howorth D, Chandona JM, Brenner SE, Hubbard TJ, et al. (2008) Data growth and its impact on the SCOP database: new developments. Nucleic Acids Res 36: D419–425.
14. Finn RD, Mistry J, Tate J, Coggill P, Heger A, et al. (2010) The Pfam protein families database. Nucleic Acids Res 38: D211–222.
15. Flower DR, North AG, Atwood TK (1994) Structure and sequence relationships in the lipocalins and related proteins. Protein Sci 2: 733–761.
16. Greene LH, Chrysina ED, Irons LI, Papageorgiou AC, Acharya KR, et al. (2001) Role of conserved residues in structure and stability: tryptophans of human serum retinol-binding protein, a model for the lipocalin superfamily. Protein Sci 10: 2301–2316.
17. Délöuse V, Giraud E, Birk C, Delmarche M, Delarue M, et al. (2009) Structure of the archaeal pabL77 peptide reveals a novel self-compartmentalizing protose family. PLoS One 4: e4712.
18. Raychon M, Filipek R, Sahat A, Kosowska K, Dubin A, et al. (2003) Saghostainis resemble lipocalins, not cytokinins in fold. Protein Sci 12: 2222–2226.
19. Mans BJ, Ribeiro JM, Andersen JF (2008) Structure, function, and evolution of biogenic amine-binding proteins in soft ticks. J Biol Chem 283: 18721–18733.
20. Paesen GC, Adams PL, Harlos K, Nutta11 PA, Stuart DJ (1999) Tick histamine-binding proteins: isolation, cloning, and three-dimensional structure. Mol Cell 3: 661–671.
21. Bishop R, Cambillau C, Privé G, Hui D, Tilko D, et al. (2000) Bacterial lipocalins: Origin, Structure, and Function. Austin (TX): Landes Bioscience.
22. Skerra A (2000) Lipocalins as a scaffold. Biochim Biophys Acta 1482: 337–350.
23. Schirner A, Chatter L, Duretelt DA, Skerra A (2010) Structural and biochemical analyses reveal a monomeric state of the bacterial lipocalin Blc. Acta Crystallography D Biol Crystallogr 66: 1308–1315.
24. Bishop RE (2000) The bacterial lipocalins. Biochim Biophys Acta 1482: 73–83.
25. UniProt Consortium (2010) The Universal Protein Resource (UniProt) in 2010. Nucleic Acids Res 38: D142–148.
26. Campanacci V, Bishop RE, Blangy S, Tegoni M, Camblanill C (2006) The membrane bound bacterial lipocalin Blc is a functional dimer with binding preference for lypoxygenophilic. FEMS Lett 380: 4077–4083.
27. Gaufoiuina MD, Gutierrez G, Bastioni M, Sanchez D (2000) A phylogenetic analysis of the lipocalin protein family. Mol Biol Evol 17: 114–126.
28. Woese CR (1987) Bacterial evolution. Microbiol Rev 51: 221–271.
29. Vernikos GS, Parkhill J (2006) Interpolated variable order motifs for identification of horizontally acquired DNA: revisiting the Salmonella pathogenicity islands. Bioinformatics 22: 2196–2203.
30. Schelhuber S, Skerra A (2002) Tuning ligand affinity, specificity, and folding stability of an engineered lipocalin variant – a so-called ‘anticalin’ – using a molecular random approach. Biophys Chem 96: 213–220.
31. Schonfeld D, Matschiner G, Chatter L, Trensmann S, Gille H, et al. (2009) An engineered lipocalin specific for CTLA-4 reveals a combining site with structural and conformational features similar to antibodies. Proc Natl Acad Sci U S A 106: 6198–6203.
32. Kim HJ, Eichinger A, Skerra A (2009) High-affinity recognition of lantahani de(III) chelate complexes by a reprogrammed human lipocalin 2. J Am Chem Soc 131: 3565–3576. NMR Structure of Protein YxeF
