The Thalidomide-Binding Domain of Cereblon Defines the CULT Domain Family and Is a New Member of the β-Tent Fold

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Abstract

Despite having caused one of the greatest medical catastrophes of the last century through its teratogenic side-effects, thalidomide continues to be an important agent in the treatment of leprosy and cancer. The protein cereblon, which forms an E3 ubiquitin ligase complex together with damaged DNA-binding protein 1 (DDB1) and cullin 4A, has been recently indentified as a primary target of thalidomide and its C-terminal part as responsible for binding thalidomide within a domain carrying several invariant cysteine and tryptophan residues. This domain, which we name CULT (cereblon domain of unknown activity, binding cellular ligands and thalidomide), is also found in a family of secreted proteins from animals and in a family of bacterial proteins occurring primarily in β-proteobacteria. Its nearest relatives are yippee, a highly conserved eukaryotic protein of unknown function, and Mis18, a protein involved in the priming of centromeres for recruitment of CENP-A. Searches for distant homologs point to an evolutionary relationship of CULT, yippee, and Mis18 to proteins sharing a common fold, which consists of two four-stranded β-meanders packing at a roughly right angle and coordinating a zinc ion at their apex. A β-hairpin inserted into the first β-meander extends across the bottom of the structure towards the C-terminal edge of the second β-meander, with which it forms a cradle-shaped binding site that is topologically conserved in all members of this fold. We name this the β-tent fold for the striking arrangement of its constituent β-sheets. The fold has internal pseudosymmetry, raising the possibility that it arose by duplication of a subdomain-sized fragment.

Introduction

Thalidomide was provided to pregnant women as an anti-nausea and sedative drug from 1957 to 1962, and was available over the counter in many countries. It was withdrawn after it became apparent that it had caused a range of birth defects in many newborns, with over 10,000 cases reported from more than 46 countries [1,2]. Soon after its ban, however, it was reintroduced as an agent against a complication of leprosy [2–4], due to its anti-inflammatory and immunomodulatory activity, and has since then also been evaluated for treatment of, among others, AIDS and Crohn’s Disease [3]. In 1994, evidence of its antiangiogenic activity led to its consideration for cancer therapy [6] and it is one of the main drugs available today against multiple myeloma [7,8]. Despite strict controls on its use, its value in the treatment of leprosy leads to the ongoing birth of babies with thalidomide-induced malformations in developing countries [2,9].

Because of its antiangiogenic and immunomodulatory activities, pharmacological interest in thalidomide continues to be very high [5], but until recently, its molecular mechanism of action – both with respect to its positive and its negative effects – remained unclear due to a lack of known targets. In 2010, Handa and co-workers showed in a landmark study that cereblon, a protein originally identified in a screen for mutations causing mild mental retardation [10], is a major target of thalidomide and is responsible for the teratogenic effects of the drug [11]. Cereblon owes its name to its involvement in brain development and to its central LON domain, which led to its initial annotation as an ATP-dependent Lon protease [10]. In the 2010 study, Handa and co-workers showed that cereblon is a cofactor of damaged DNA-binding protein 1 (DDB1), which acts as the central component of an E3 ubiquitin ligase complex and regulates the selective degradation of key proteins in DNA repair, replication and transcription [12]. Binding of thalidomide to a C-terminal region in cereblon inhibits the E3 ubiquitin ligase activity of the complex and leads to developmental limb defects in chicks and zebrafish [11]. Point mutations in this region, which abolish thalidomide binding, but allow the continued formation of the E3 complex, restore ubiquitination and prevent the teratogenic activities of thalidomide.

Despite these advances, there has been little progress in understanding the mechanism of thalidomide binding and teratogenicity, due to the difficulties in preparing cereblon protein for biochemical and biophysical studies. Such progress would
Results

The domain structure of cereblon

Cereblon proteins occur throughout eukaryotes, however not in fungi. They are typically 400–600 residues long (442 in the case of human cereblon) and their genes occur in single copy per genome. Their most salient feature is the presence of a central LON domain (residues 80–317 in human cereblon, Fig. 1). As defined by secondary structure prediction and length of the domain (LON−C), all these domains are seen to establish experimentally, all these domains are present in several protein families, and that this domain is related to several domains with known functions. Where

![Diagram of the domain structure of cereblon](image)

Fig. 1. Domain structure of cereblon. (a) Schematic view of the domains and motifs in cereblon. The Tyr and Trp residues of the CULT domain whose mutation abolishes thalidomide binding (Ito et al., 2010) are highlighted in grey. Sequences with high helical propensity in the putative DDB1-binding region are underlined. (c) Model of cereblon. The LON domain was modeled on PDB:1ZBO. The N-terminal extension and the large connector between the LON subdomains are marked by dotted lines. The CULT domain was modeled on the structure of a bacterial CULT domain which we have determined [18]. Note that the CULT domain of human cereblon can also be modeled with good accuracy on the homologous structures of MsrB and RIG-I (Fig. S1). After submission of this manuscript, two cereblon-DDB1 complex structures [15,16] have confirmed the domain arrangement proposed in this model.

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interaction. After submission of this manuscript, two studies presented the structure of cereblon in complex with DDB1 [15,16], which showed that the interaction between the two proteins is topologically novel and not mediated by an H-box. All three regions with elevated helical propensity in the connector (Fig. 1) are indeed helical, albeit the first one as a 3_{10} helix. The main interactions are formed between the first region and DDB1 \beta-propeller A, and between the third region (containing the conserved motif) and DDB1 \beta-propeller C.

N-terminally to the LON domain, cereblon proteins have an extension of typically 50–100 residues (79 in the case of human cereblon), of which the front part is not conserved across phyla and generally predicted as intrinsically unstructured (Fig. 1). Starting about 35 residues prior to the beginning of the LON domain, the extension becomes well-conserved and particularly a motif FDxxLPxxHxYLG is recognizable in most cereblon homologs, from humans and plants to basal eukaryotes. Since the extension is adjacent to the connector between LON-N and LON-C in the LON domain model (Fig. 1) and experimental evidence suggested the binding interactions with DDB1 to be bipartite [14], we considered that this motif could contribute to DDB1 binding. The cereblon-DDB1 complex structures however show that the extension interacts with the C-terminal region of cereblon via the conserved motif [15,16]. Indeed, since it runs alongside the thalidomide-binding site, it may relay information on the occupancy of the site to the LON-N domain. We therefore conjecture that deletion of the extension could uncouple the C-terminal region from the rest of the E3 ligase complex, alleviating or even entirely abolishing the effects of thalidomide binding.

The C-terminal region of cereblon, comprising about 100–130 residues (125 in the case of human cereblon), represents the best-conserved part. It has multiple invariant residues and encompasses completely the part of the protein identified through deletion analysis as responsible for thalidomide binding [11]. Sequence similarity searches show that this region also occurs in much shorter proteins than cereblon, where it essentially covers the entire length of the protein, identifying it as a domain. We name this domain CULT, for cereblon domain of unknown activity, binding cellular ligands and thalidomide.

The CULT domain

With the CULT domain of human cereblon as a starting point, PSI-Blast searches of the non-redundant protein database at NCBI converge in three iterations. Analysis of the results, for example using clustering by pairwise sequence similarity in CLANS (Fig. 2), shows that most of the search space consists of cereblon sequences, recognizable by their LON domain, but that several other groups of CULT domain-containing proteins are identifiable. The two main groups are: (I) prokaryotic proteins, mainly from \sigma- and \gamma-proteobacteria, but with a few representatives from \alpha- and \beta-proteobacteria and one sequence from a spirochete; these proteins consist entirely of the CULT domain; and (II) animal proteins from placozoans to vertebrates, but not occurring beyond fishes; these proteins also consist of the CULT domain, but carry an N-terminal secretion signal sequence. Indeed, the homolog from the sand fly Phlebotomus arabicus has been identified experimentally as a salivary protein [17]. Two further, more divergent groups are also apparent: one from oomycetes, with an N-terminal signal sequence followed by a CULT domain and ending with a carbohydrate-binding domain (SCOP: b.64); and the second from kinetoplastids, with an N-terminal CULT domain followed by a C-terminal region that cannot be assigned to a known domain family at present. The remaining few sequences from the PSI-Blast search do not recognizably belong to any of these groups; they are mainly from green algae and can all be confirmed by reverse PSI-Blast searches to contain a CULT domain.

Multiple alignment of the sequences identified in the search (Fig. 3A) show that several residues are highly conserved in the CULT domain. Conservation of these residues is highest in the CULT core group (cereblon, secreted eukaryotic sequences, bacterial sequences) and declines towards the periphery. Particularly conspicuous are three tryptophan residues, marked by arrows in Fig. 3A, which are seen to form the binding site for thalidomide and cellular ligands in the crystal structure of the bacterial CULT protein MGR_0879 from Magnetospirillum gryphiswaldense ([18]; PDB ID:4V2Y; Figs. 3B,C). The crystal structure, which we determined after this bioinformatic study, shows that the other highly conserved residues group around this binding site (S1 Fig.), their conservation being rationalized by an influence on substrate recognition and discrimination. The one exception to this are two CxxC cysteine motifs, which we took from the beginning of this project to be indicative of a zinc binding site and thus present for structural reasons.

We developed the bacterial model system to study the CULT domain because we found eukaryotic cereblon proteins very difficult to express in useful amounts and even more difficult to purify in a soluble state. In contrast, the bacterial protein could be produced and purified in a straightforward way [18]. We reasoned that, at 36% sequence identity and with almost all well-conserved positions similar or the same between the CULT domains of humans and Magnetospirillum, the bacterial system should represent an accurate model for the eukaryotic domain. The structures of eukaryotic CULT domains from human, mouse and chicken [15,16] now show that the expectation is true to an astonishing extent, with a root-mean-square deviation (r.m.s.d.) of around 0.9 Å over 100 Cx positions between the bacterial and human proteins (Fig. 3B). The bacterial domain is thus structurally almost as similar to the eukaryotic domains as these are to each other (Table. S1).
The β-tent fold

We searched for remote homologs of the CULT domain using profile Hidden Markov Model (HMM) comparisons in HHpred and obtained matches at probabilities better than 90% (E values < 1e-6) for multiple protein families, several of which have members of known structure (Fig. 4). The best matches were to a protein family found throughout eukaryotes, yippee [19]. Two of the five yippee paralogs in mammals have been implicated in signal transduction [20] and tumor suppression [21], respectively, but the actual mechanism of these proteins remains unknown.

Fig. 3. CULT domain sequence and structure. (a) Multiple alignment of CULT domains from representative members of the groups in Fig. 2. The alignment is based on the results of the PSI-Blast search with the CULT domain of human cereblon (first sequence in the alignment). Invariant residues of the three core groups (cereblon, secreted eukaryotic, bacterial) are underscored in black, residues conserved in at least two thirds of the sequences in the alignment are highlighted in dark grey and residues in at least one third of the sequences in light grey. The three tryptophan residues forming the thalidomide-binding site are marked by arrowheads and the two cysteine motifs coordinating the Zn ion, as well as a highly conserved motif at the tip of the inserted β-hairpin, are written out. The secondary structure above the alignment (S = β-strand) is the experimentally determined structure of the CULT domain from MGR_0879 of Magnetospirillum gryphiswaldense (first bacterial sequence in the alignment) [18]. The β-strands of the two main β-sheets are numbered according to the consensus structure of the β-tent fold and colored by whether they belong to the N-terminal β-sheet (purple) or the C-terminal one (gold); β3 is shown in brackets as it has lost its β-strand character in the CULT domain. The two β-strands of the inserted hairpin (teal) are labeled βI1 and βI2. The sequences are: (cereblon) HS - Homo sapiens NP_057386.2, DR - Danio rerio NP_001003996.1, DM - Drosophila melanogaster XP_001999319.1, CE - Caenorhabditis elegans NP_502300.1, AT - Arabidopsis thaliana NP_850069.1, EH - Entamoeba histolitica XP_657530.1; (secreted eukaryotic) DR - Danio rerio NP_001121712.1, CB - Caenorhabditis briggsae XP_000570325.1, CE - Caenorhabditis elegans NP_502300.1, AT - Arabidopsis thaliana NP_850069.1, EH - Entamoeba histolitica XP_657530.1; (bacterial) MG - Magnetospirillum gryphiswaldense CAM74667.1, GP - gamma proteobacterium WP_008249149.1, HO - Haliangium ochraceum WP_012831591.1, DT - Desulfonioanospirina sp. BDW918 XP_000887065.1, LS - Leptospira sp. BS-022 WP_020769190.1; (oomycete) PI - Phytophthora infestans XP_002999235.1, AL - Albugo laibachii CCA16326.1; (kinetoplastid) LM - Leishmania major strain Friedlin XP_001681231.1, TC - Trypanosoma cruzi EKG02463.1; (other) BP - Bathycoccus prasinos XP_007508760.1. (b) Superimposition of bacterial and eukaryotic CULT domain structures from M. gryphiswaldense (red), H. sapiens (blue; PDB ID: 4TZ4), H. mediterranei (orange; PDB ID: 4TZC), and G. gallus (green; PDB ID: 4Q12). The r.m.s. deviations in Cs positions for the pairwise comparisons ranged between 0.4 and 0.9 Å and are listed in S1 Table. (c) Superimposition of the thalidomide binding site for the structures in panel (b). The ligand and the residues of the aromatic cage are shown in stick representation and colored as in panel (b). Residue numbering is for the human protein.

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searching against Pfam we noticed that, in this database, the profile for yippee (PF03226) was generated jointly with another protein, Mis18, which in our analyses is not particularly close to yippee and indeed seems about as remote from yippee as cereblon is (Fig. 5). Mis18 proteins are broadly represented in eukaryotes, except plants, and appear to be involved in centromere assembly [22,23], although their actual mechanism remains unknown. The reason for merging Mis18 with yippee in Pfam is unclear to us.

Two protein families of known structure related at a similar level to the CULT domain as yippee and Mis18 are methionine sulfoxide reductase B (MsrB or SelR) and the regulatory domain of retinoic acid-induced gene-1 (RIG-I). MsrB is the most widely distributed protein in this study and is universal to all cellular life. It protects cells from oxidative stress by reducing methionine-R-sulfoxide residues (for a review see e.g. [24]). RIG-I has a more limited phylogenetic spectrum, being detectable only in animals. It is an RNA helicase that, upon binding viral RNA, activates the host innate immune system (for a review see e.g. [25]). The regulatory domain is the RNA 5′-triphosphate sensor of RIG-I, activating the ATPase activity of the protein by RNA-dependent dimerization [26]. These four proteins, yippee, Mis18, MsrB, and RIG-I, are sufficiently close to the CULT domain in sequence space that they usually show up in the non-significant part of sequence similarity searches, between E values of 0.005 and 10, and are occasionally included in the significant part as well. Thus, for example, PSI-Blast searches of the nr database with our bacterial model protein, Magnetospirillum MGR_0879, include the first yippee and RIG-I sequences in the second iteration and the first Mis18 and MsrB sequences in the fifth. These proteins appear roughly equidistant from cereblon in sequence space (Fig. 5). More distantly related, but showing up with fair regularity in our searches is glutathione-dependent formaldehyde-activating enzyme (GFA), a protein found in bacteria and most eukaryotes, except plants. GFA catalyzes the first step in the detoxification of formaldehyde [27].

All these proteins share a common fold, formed by two four-stranded, antiparallel β-sheets that are oriented at approximately a right angle and pinned together at their tip by a zinc ion (Fig. 6). The two sheets are connected covalently across the top on both
sides by loops, due to circular permutation. Thus, the last strand of the domain is topologically the first strand of the first sheet, yielding the strand order $b_8^{-}b_1^{-}b_2^{-}(b_3)$ for the first sheet and $(b_4)^{-}b_5^{-}b_6^{-}b_7$ for the second ($b_3$ and $b_4$ are shown in brackets as, in some structures, they have lost their $b$-strand character). Because of the striking arrangement of these $b$-sheets we have named this fold the $b$-tent.

A conserved feature of all proteins with a $b$-tent fold is an insertion between strands $b_2$ and $b_3$, which usually has a $b$-hairpin stem and reaches across the bottom of the tent to extend the second $b$-sheet at its C-terminal edge. Due to the curvature of the $b$-sheet and the sizable nature of the loops connecting $b_4$ to $b_5$ on one side and the strands of the insertion on the other, all $b$-tent proteins contain a cradle-shaped groove at this location, which hosts the binding site (Fig. 7).

The residues giving the binding site its specificity in the individual proteins are frequently found in equivalent positions. This is particularly conspicuous when comparing the binding sites of CULT and MsrB (Fig. 8). Of the four residues forming the thalidomide-binding site in *Magnetospirillum* CULT (4V2Y: W79, W85, W99, and Y101), the last three have equivalents in homologous positions in the methionine sulfoxide-binding site of MsrB (3HCJ: R97, H111, F113); the first, W79, is also a tryptophan in the MsrB binding site, but from an analogous position in the insert loop (W73), due to a shift in the position of the site caused by the shape difference between W85 of CULT and R97 of MsrB. This shift places the ligand above $b_7$ in MsrB, rather than above $b_6$, allowing the positioning of a further residue into the active site, which is the catalytic cysteine; conversely, there appears to be no need for a catalytic residue in CULT. We note that the homology of the conserved aromatic residues in CULT to the residues of the binding site in MsrB can be readily seen from the HHpred alignment.

Extending these observations to yippee, which has a similar distribution of conserved residues as CULT (S1 Fig.), we predict that the binding site of this protein is also an aromatic cage, comprising the highly conserved Y43, F45, W82, and Y84, as numbered in *D. melanogaster* yippee isoform B, ABC67182.1 (Fig. 7). Of these, W82 and Y84 are in homologous positions to W99 and Y101 of CULT and H111 and F113 of MsrB, whereas Y43 and F45 are at the same position as W73 in MsrB, but not recognizably homologous.

A striking property of the $b$-tent fold is that, in several of the proteins, the two sheets have considerable structural symmetry, such as for example in the MsrB structure 3HCJ, where superposition of the two 43-residue halves yields an r.m.s.d. of 1 Å over the $C\alpha$ positions of the core 30 residues (Fig. 6). This raises the possibility that the fold originated by duplication of a...
subdomain-sized fragment, but we note that no similarity is detectable between the two halves by sequence comparisons.

Searches in structure space for other proteins with the β-tent fold yielded three more proteins of known structure (Figs. 6, S2), which share the fold with the same topology of secondary structure elements, including the β-hairpin extension between strands β2 and β3, but have no significant sequence similarity to the other proteins in this study, or to each other (Fig. 4). These are MSS4, a guanine exchange factor and nucleotide-free chaperone for the Rab GTPase [28,29], TCTP, a pleiotropic protein involved in malignant transformation and regulation of apoptosis [30–32], and DUF427, a domain of unknown function. Whereas MSS4 and TCTP are eukaryotic proteins, TCTP being present universally and MSS4 broadly, but not in plants, DUF427 is seen mainly in bacteria and fungi, with a small number of archaea presumably having acquired this domain by lateral transfer. Of these proteins, only MSS4 has the zinc binding site (Fig. 6). In the SCOP database, MSS4 and TCTP are grouped together with MsrB and GFA as families within the MSS4-like superfamily, which is the sole representative of the MSS4-like fold (b.88).

What are the physiological ligands of CULT?

A fundamental issue in understanding the biological role of CULT domains, not directly illuminated by their homology to other proteins, is the identification of their physiological ligand(s). The only ligands known today, thalidomide and its derivatives, are clearly non-physiological. Given that the clustering of the invariant tryptophans into a cage-like arrangement was already suggested at the modeling stage (see above), we searched PDB for ligands.
bound in aromatic cages, loosely defined. For this we allowed the aromatic residues to be Phe and Tyr, as well as Trp, and provided only a very general requirement for cage-like geometry, in order to gain as broad a view as possible (see Methods). We obtained 1098 distinct ligands, which could be grouped approximately into five classes, corresponding to heterocyclic rings, hydrocarbon rings, hydrocarbon chains with and without heteroatoms, and ammonium-based cations (Fig. 9, Table S2). Upon inspection, many of the “cages” identified indeed turned out to be only approximately cage-like and for 46 ligands, all binding sites turned out to be geometrically too divergent to be considered further.

Half of the identified ligands belonged to the largest class, comprising heterocyclic rings. Many of these were enzyme inhibitors, both of natural and synthetic origin, such as indole-2,3-diones (4KWG), aryl hydrazines (4MQQ), or non-nucleoside reverse transcriptase inhibitors (1SGH). Thalidomide, which is bound in the aromatic cage of CULT domains via its glutarimide ring, belongs to this class. Among the natural compounds, we found pyrimidines and their nucleosides of particular interest, as these resemble the glutarimide ring of thalidomide [19] and are bound in similar cages. For example, the transcription factor RutR (3L0C; Fig. 9B) can bind both uracil and thymine in its aromatic cage and acts as the master regulator of genes involved in the synthesis and degradation of pyrimidines [33]. An experimental screen against Magnetospirillum MGR_0879 found that, of the nucleobases, uracil and its nucleoside (uridine) were indeed bound and their relevance for eukaryotic cereblon could be established in vivo in zebrafish. It is attractive to consider that they might also be the physiologically relevant ligand, given that DDB1 is an integrator of cellular information on DNA damage and incorporation of uracil into DNA represents a mutagenic lesion [18].

Another type of ligand that we found to be of particular interest in this analysis comprises amino acid sidechains, modified and unmodified. These include the heterocyclic rings of His, Pro, and Trp, the hydrocarbon rings of Phe (Fig. 9C) and Tyr, the hydrocarbon chains of Ile, Leu (Fig. 9D), Met, and Val, and the cationic sidechains of methylated and unmethylated Lys (Fig. 9G) and Arg (Figs. 9H, S3). Particularly the latter occur prominently in the tails of histones and are recognized by aromatic cages in a range of different domains, including bromodomains, chromodomains (Fig. 9G), and Tudor domains (Fig. 9H). Given that the DDB1-Cul4A E3 ubiquitin ligase complex is known to bind and ubiquitinate histones [see e.g. [34]], an activity of cereblon in recognizing histone tail modifications within a linear sequence motif and providing the target specificity for the ligase complex appears fully plausible [18].

Other sidechain interactions, particularly in the context of linear sequence motifs, also appear entirely possible. Thus, the homeobox transcription factor MEIS2, which is implicated in various aspects of human development, was recently identified as a cereblon interactor [15]. Its binding was exclusive with thalidomide and its derivatives, suggesting that it is recognized via the same binding site. We note that MEIS2 and its paralogs contain two folded domains, one being the homeobox domain and the other uncharacterized at present, flanked by extended regions predicted to be unstructured and with low sequence conservation. The N-terminal approximately 10 residues are however very highly conserved and contain sidechains [Arg, Tyr, His] that could easily be envisaged as the ligands of an aromatic cage. We therefore consider this region to be the most attractive first candidate for exploring the MEIS2-cereblon interaction. This said, the very high similarity between the bacterial and eukaryotic CULT domains, particularly in the area of the aromatic cage, points to a wide-spread ligand, present also outside the cell, rather than to a linear sequence motif. By similarity to methyllysine, one might envisage choline, carnitine, betaine, and related compounds, but none of these could so far be seen to interact with the CULT domain in our model system.

**Discussion**

In this article we have presented evidence that the thalidomide-binding region of cereblon is a conserved domain, CULT, present.
in several other proteins of eukaryotes and bacteria. The CULT domain is recognizably homologous to at least five other domain families, which share - where known - a common fold and a shared mechanism of ligand binding. The fold is also recognizable in three further domain families, which however do not have detectable sequence similarity to any of the other proteins, or to each other, and whose evolutionary relationship thus remains unclear (the SCOP database, however, clearly considers them homologous, as it groups them into the same superfamily). We have named the common fold of these proteins the β-tent, due to the orientation of its two constituent β-sheets.

The widely differing activities of proteins with a β-tent fold, as well as the absence of invariant residues across the domains, suggest that the β-tent is a structural scaffold, which mounts a binding site at a specific location. The binding site is formed by a cradle-shaped groove, whose sides are provided by loops connecting strands and whose bottom is formed by strands and β6 and β7. The elaboration of this site in the individual families is tailored to their specific function, but appears to follow common principles, particularly in families binding small-molecule ligands. Here, binding residues are mainly located on the two loops and strand β6, while catalytic residues appear to be located on strand β7. For families whose binding site is at present unknown, this can therefore be reasonably predicted by mapping their conservation pattern onto homology models of the relevant region.

Members of the β-tent fold show, to varying degrees, a twofold rotational symmetry around a central axis passing through the apical zinc ion (where present). The symmetry is most pronounced in MsrB and this domain also has the broadest phylogenetic spectrum, being the only one with a universal representation in all cellular life forms. It therefore seems attractive to surmise that it is the ancestral representative of this fold, from which the others evolved by duplication and differentiation, and that it itself originated by duplication of a four-stranded β-meander. We have previously argued for an origin of folded proteins from subdomain-sized peptides [35,36]. But for the apparent lack of internal sequence symmetry to support this inference, the β-tent would seem an attractive candidate for such a scenario.

The absence of statistically significant sequence similarity between MSS4, TCTP, DUF427 and the other proteins of this fold raises the possibility of a convergent origin. We note however that MsrB and RIG-I also do not share statistically significant sequence similarity between each other (Fig. 4) and are only connected conclusively in sequence space via CULT and yippee (Fig. 5). The homology of all proteins with a β-tent fold thus remains a clear possibility, which may become substantiated by new domain families found in hitherto poorly explored parts of the tree of life.

Methods

Sequence similarity searches were carried out at the National Institute for Biotechnology Information (NCBI; http://blast.ncbi.nlm.nih.gov/) and in the MPI Bioinformatics Toolkit (http://toolkit.tuebingen.mpg.de; [37]). PSI-Blast [38] at NCBI was run on the non-redundant protein sequence database (nr) with an E-value threshold of 0.005. CS-Blast [39] in the MPI Toolkit was run on a version of nr clustered at 70% sequence identity (nr70), also with a threshold of E = 0.005. The sequence relationships of proteins identified in these searches were explored by clustering them according pairwise Blast P-values [40] in CLANS [41]. Clustering was done in default settings (attract = 10, repulse = 5, exponents = 1), with other settings as given in the figure legends.

Searches for more distant homologues were made with HHpred [42] and HHsenser [43] on the databases pdb70 (sequences of protein databank structures, as available in April 2014, clustered at 70% sequence identity), CDD (conserved domain database from NCBI, as of February 2014), pfamA release 27.0, SCOP release 1.75, and profile HMM databases of all human and all Drosophila proteins built locally and available through the MPI Toolkit.

Secondary structure was predicted in the MPI Toolkit, using the meta-tool Quick2D.

Structure similarity searches were carried out on the Dali server (ekhidna.bioinformatics.rutgers.edu/dali_server; [44]). Molecular models were built using Modeller [45], and manipulated in Swiss PDB Viewer [46] and PyMol (Schrodinger, LLC). Sequence conservation patterns were visualized with ProtSkin (http://www.mcgumr.mcgill.ca/ProtSkin/; [47]).

Aromatic cage-like conformations containing ligands in PDB structures were detected by applying a set of geometric criteria (see Figure 9A). First, in each PDB structure, all non-water molecules within 6.0 Å distance to these ligands were considered in further analysis. Then, we defined a set of at least three aromatic residues from the same polypeptide chain to form a cage-like conformation interacting with a ligand if: a) all pairwise distances between their side chain mass centers were less than 10.0 Å; b) the angle between and was less than 60° for at least three of the aromatic residues, where is the normal vector of the aromatic ring, is the vector connecting and is the vector connecting and and c) at least two ligand atoms were within 3.0 Å distance to . The program was implemented in Python using Biopython [48], SciPy [49] and NetworkX [50] libraries.

We applied these geometric rules to scan 102,886 PDB files downloaded from the PDB (24 Aug 2014). In total, 6,144 putative aromatic cage-like conformations were detected with 1,098 different ligands binding to them. We grouped the cages according to the ligands they interact with. In each group, redundant cage-like conformations were removed (two cage-like conformations were considered identical if the composite residue names and numbers were the same). Subsequently, we manually examined at least one cage-like conformation in each of the 1,098 groups. Based on the ligand moiety within the aromatic cage, we further classified the 1,098 groups into different categories (S2 Table).

Supporting Information

S1 Fig  Comparative models of the β-tent domains in (a) human cereblon and (b) Drosophila yippee. Models are colored according to the templates used (black - CULT domain of M. glyphiswaldense MGR_0879, 4VZY; red - X. campestris MsrB, 3HJC; green - human RIG-I, 3EQT). The two central β-hairpins that mount the zinc binding site are shown bold. For each protein, the two following panels show the sequence conservation mapped onto the surface of the model built on the CULT domain of MGR_0879. The red-to-white scale follows highest-to-lowest conservation. Mapping was done using ProtSkin and a multiple sequence alignment derived from two iterations of PSI-Blast and filtered to 70% identity.

S2 Fig  Structure gallery of proteins of the β-tent fold superimposed on the crystal structure of M. glyphiswaldense MGR_0879 (grey). The superimposition mainly covers the two four-stranded β-sheets; the β-hairpin insertion that
reaches across the bottom of the fold to the C-terminal edge of the second β-sheet is present in all folds, but two structurally variable to superimpose well. A superimposition of the best-conserved structural core of all structures in the figure is shown in the center. The r.m.s. deviations for the Cα carbons for the pairwise superimpositions were, in clockwise order: 3HGC: 0.98 Å over 63 residues, 1XA8: 1.72 Å over 30 residues, 3EQT: 1.38 Å over 63 residues, 2FU5: 1.40 Å over 57 residues, 1YZ1: 1.31 Å over 50 residues, 3DJM: 1.83 Å over 27 residues.

S3 Fig Protein-protein association mediated by an aromatic cage. The image shows the main specific contact in the interaction between neuropilin-1 (cyan) and vascular endothelial growth factor-A (VEGF-A) (green), mediated by the docking of the C-terminal arginine of VEGF-A into an aromatic cage formed by neuropilin-1 (PDB: 4DEQ; Parker MW, Xu P, Li X, Vander Kooi CW). Structural basis for selective vascular endothelial growth factor-A (VEGF-A) binding to neuropilin-1 (2012) J Biol Chem 287: 11082–11089).

References

1. Lenz W (1988) A short history of thalidomide embryopathy. Teratology 38: 203–215.
2. World Health Organization (2014) Use of thalidomide in leprosy. Available: http://www.who.int/lep/research/thalidomide/en/. Accessed 9 May 2014.
3. Shekini J (1965) Thalidomide in the treatment of lepra reactions. Clin Pharmacol Ther 6: 303–306.
4. Teo S, Rassat KE, Schellfer MA, Kook KA, Zeldis JB, et al. (2002) Thalidomide in the treatment of leprosy. Microbes Infect 4: 1193–1202.
5. Franks ME, Macpherson GR, Fig WD (2004) Thalidomide. Lancet 363: 1802–1811.
6. D’Amato RJ, Loughman MS, Flynn E, Folkman J (1994) Thalidomide is an inhibitor of angiogenesis. Proc Natl Acad Sci U S A 91: 4082–4085.
7. Singhyl S, Mehta J, Desikan R, Ayers D, Roberson P, et al. (1999) Antimicrobial activity of thalidomide in refractory multiple myeloma. N Engl J Med 341: 1503–1507.
8. Morgan DJ, Davies FE (2013) Role of thalidomide in the treatment of patients with multiple myeloma. Crit Rev Oncol Hematol 88 Suppl 1: S14–22.
9. Crawford A (2013) Brazil’s new generation of Thalidomide babies. BBC News. http://www.bbc.com/news/magazine-23418102. Accessed 9 May 2014.
10. Higgins JJ, Paulcikova J, Lombardi RQ, Tuttle R, et al. (2010) YPEL3, a p53-regulated gene that induces cellular senescence. Cancer Res 70: 3566–3575.
11. Hayashi T, Fujita Y, Iwasaki Y, Takahashi K, et al. (2004) Mi16 and Mi18 are required for CENP-A loading and histone deacetylation at centromeres. Cell 118: 713–729.
12. Fujita Y, Hayashi T, Kiyosu M, Toso Y, Kobukou A, et al. (2007) Priming of centromere for CENP-A recruitment by human Mi18alpha, Mi18beta, and Mi18BPI. Dev Cell 12: 17–30.
13. Lee BG, Dikiy A, Kim HY, Gladyshev VN (2009) Functions and evolution of salivary transcriptomes of the sand fly Phlebotomus argentipes. J Mol Biol 390: 692–707.
14. Teo S, Resztak KE, Scheffler MA, Kook KA, Zeldis JB, et al. (2002) A thalidomide analog as an inhibitor of angiogenesis and tumor cell invasion. Proc Natl Acad Sci U S A 99: 5358–5363.
15. Groenick M, Bartoschek S, Hagermeier GH, Griesinger C, Verheuf JA (2002) A glutathione-dependent formaldehyde-activating enzyme (GfA) from Paraxoxococcus denitrificans detected and purified via two-dimensional proton exchange NMR spectroscopy. J Biol Chem 277: 3069–3072.
16. Burton J, Roberts D, Montaldi M, Novick P, De Camilli P (1993) A mammalian guanine-nucleotide-releasing protein enhances function of yeast secretory protein Sec4. Nature 361: 464–467.
17. Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, et al. (1997) Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res 25: 3389–3402.
18. Bieger A, Soding J (2009) Sequence context-specific profiles for homology searching. Proc Natl Acad Sci U S A 106: 3770–3775.

S1 Table Structural similarity of the CULT domains from Magnetospirillum MGR_00789 and human, mouse and chicken cerebrib. The numbers show the r.m.s. deviation in Å over the number of superimposed Cα carbons.

S2 Table Classes of ligands detected in cage-like binding sites in PDB (see also Methods and Fig. 9).

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Author Contributions

Conceived and designed the experiments: ANL. Performed the experiments: ANL. Analyzed the data: ANL. Wrote the paper: ANL.
40. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool. J Mol Biol 215: 403–410.
41. Frickey T, Lupas A (2004) CLANS: a Java application for visualizing protein families based on pairwise similarity. Bioinformatics 20: 3702–3704.
42. Soding J, Biegert A, Lupas AN (2003) The HHpred interactive server for protein homology detection and structure prediction. Nucleic Acids Res 33(Web Server issue): W244–248.
43. Soding J, Remmert M, Biegert A, Lupas AN (2005) HHpred: exhaustive transitive profile search using HMM-HMM comparison. Nucleic Acids Res 34(Web Server issue): W374–378.
44. Holm L, Rosenstro¨m P (2010) Dali server: conservation mapping in 3D. Nucleic Acids Res 38(Web Server issue): W545–549.
45. Wohli B, Sali A (2010) Protein structure modeling with MODELLER. Methods Mol Biol 1137: 1–15.
46. Guex N, Peitsch MC (1997) SWISS-MODEL and the Swiss-PdbViewer: an environment for comparative protein modeling. Electrophoresis 18: 2714–2723.
47. Deprez C, Lloubes R, Gavioli M, Marion D, Guerlesquin F, et al. (2005) Solution structure of the E.coli TolA C-terminal domain reveals conformational changes upon binding to the phage g3p N-terminal domain. J Mol Biol 346: 1047–1057.
48. Cock PJ, Antao T, Chang JT, Chapman BA, Cox CJ, Dalke A, Friedberg I, Hamelryck T, Kauff F, Wilczynski B, de Hoon MJ (2009) Biopython: freely available Python tools for computational molecular biology and bioinformatics. Bioinformatics 25: 1422–1423.
49. Jones E, Oliphant E, Peterson P, et al. (2001) SciPy: Open Source Scientific Tools for Python. Available: http://www.scipy.org/. Accessed 5 Oct 2014.
50. Hagberg AA, Schult DA, Swart PJ (2008) Exploring network structure, dynamics, and function using NetworkX, in Proceedings of the 7th Python in Science Conference (SciPy2008), Gael Varoquaux, Travis Vaught, and Jarrod Millman (Eds), (Pasadena, CA USA), pp. 11–3.