A Novel Human Tectonin Protein with Multivalent β-Propeller Folds Interacts with Ficolin and Binds Bacterial LPS

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Abstract

Background: Although the human genome database has been completed a decade ago, ~50% of the proteome remains hypothetical as their functions are unknown. The elucidation of the functions of these hypothetical proteins can lead to additional protein pathways and revelation of new cascades. However, many of these inferences are limited to proteins with substantial sequence similarity. Of particular interest here is the Tectonin domain-containing family of proteins.

Methodology/Principal Findings: We have identified hTectonin, a hypothetical protein in the human genome database, as a distant ortholog of the limulus galactose binding protein (GBP). Phylogenetic analysis revealed strong evolutionary conservation of hTectonin homologues from parasite to human. By computational analysis, we showed that both the hTectonin and GBP form β-propeller structures with multiple Tectonin domains, each containing β-sheets of 4 strands per β-sheet. hTectonin is present in the human leukocyte cDNA library and immune-related cell lines. It interacts with M-ficolin, a known human complement protein whose ancient homolog, carcinolectin (CLS), is the functional protein partner of GBP during infection. Yeast 2-hybrid assay showed that only the Tectonin domains of hTectonin recognize the fibrinogen-like domain of the M-ficolin. Surface plasmon resonance analysis showed real-time interaction between the Tectonin domains 6 & 11 and bacterial LPS, indicating that despite forming 2 β-propellers with different Tectonin domains, the hTectonin molecule could precisely employ domains 6 & 11 to recognise bacteria.

Conclusions/Significance: By virtue of a recent finding of another Tectonin protein, leukolectin, in the human leukocyte, and our structure-function analysis of the hypothetical hTectonin, we propose that Tectonin domains of proteins could play a vital role in innate immune defense, and that this function has been conserved over several hundred million years, from invertebrates to vertebrates. Furthermore, the approach we have used could be employed in unraveling the characteristics and functions of other hypothetical proteins in the human proteome.

Introduction

Advances in sequence genomics have resulted in an accumulation of a large number of protein sequences derived from genome sequences. Although the human genome database has been completed a decade ago, about 50% of the human proteome still remains hypothetical as their functions are unknown [1]. The elucidation of the functions of these hypothetical proteins can lead to additional protein pathways and revelation of new cascades, thus completing our fragmentary knowledge on the proteome complex. Furthermore, information on the network of protein–protein interactions will increase logarithmically. New hypothetical proteins may serve as disease markers and pharmacological targets.

The prime targets for the discovery of functional proteins are those which show homology to counterparts in lower species by way of sequence similarities and domain conservation. An alternate approach is to examine the proteins of invertebrates that do not have homologs in the vertebrate system. One example of such a group of proteins is the Tectonin domain-containing proteins in humans. Tectonin domain containing proteins, which belong to a subclass of proteins of the larger β-propeller family, have thus far only been studied in the fish, horseshoe crab, slime mold and sponge [2–5]. Tectonin domains were first reported in the Tectonins I and II proteins of the slime mold, Physarum polycephalum. The Tectonins I and II were characterized to have repeats of Tectonin domains [2]. Because the proteins are located
at the surface of this organism, where they were postulated to scavenge food including bacteria, the Tectonin proteins have been speculated to function as bacterial sensors. Tachylectin-1 in the horseshoe crab, Tachypleus tridentatus, also has Tectonin domain classification [4,6], and was shown to be able to bind bacterial lipopolysaccharide (LPS). Study on the Tectonin protein, LEC_SUBDO, of the sponge, Suberites domuncula, also revealed a possible LPS-binding function [3]. A recent investigation on the galactose-binding protein (GBP) in the horseshoe crab (Carcinoscorpius rotundicauda), a protein consisting of only 6 Tectonin domains revealed that the Tectonin domains function to differentiate host from pathogen and simultaneously bridge a host-pathogen interactome (Low et al., unpublished).

An exhaustive search in the databases for vertebrate proteins failed to reveal any potential homologs with significant sequence similarity, indicating that perhaps these Tectonin domain-containing proteins (henceforth referred to as Tectonin proteins) have evolved through the species, although more recently, other proteins with Tectonin domains are being uncovered, for example, the human leukolectin (GenBank Accession No. ACM77812).

There are many examples of other families of meiosis-related proteins, kinetochores, cell gap contacts and nuclear pore complexes which show no homology at the primary amino-acid sequence level. However, they hint at the conservation of their domain architecture organization. Furthermore, the three-dimensional structure of functionally important domains in proteins in the budding yeast, nematode, Drosophila, Arabidopsis, and human have been conserved [7–11]. Here, we have used several databases like SCOP, CATH, SMART, which also employ domain and secondary structure classification for structure sorting and function prediction, to search for β-propeller structures and possibly distance relationships by domain conservation. This is

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**Figure 1. hTectonin is distantly related to the invertebrate Tectonins.** (A) The phylogenetic tree constructed after a PSI-Search query using the invertebrate Tectonin proteins revealed K1358 family of proteins as closely related Tectonin domain containing proteins in the mammals and also in lower species like the frog. The numbers at the nodes are an indication of the level of confidence for the branches as determined by bootstrap analysis (1000 bootstrap replicates). (B) Bioinformatics domain analysis utilizing SMART [22,23] shows existence of Tectonin domain-containing proteins both in invertebrates and vertebrates from the horseshoe crab lectins, worm, up to humans. Of interest in this study is the protein hTectonin (red asterisk) which appear to have homologues in other species as well, for example in P. troglodytes (chimpanzee), P. pygmaeus (orangutan), M. musculus (mouse), G. gallus (chicken), C. elegans (worm) and D. melanogaster (fruitfly).

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especially useful when searching for related proteins with low sequence homology or when sequences have diversified through evolution from the invertebrates to the mammals. We thus seek to identify Tectonins in the vertebrates, and compare their domain architecture and function with ancient homologs from the invertebrates in order to gain insights into their functional conservation in the vertebrates, particularly in view of host-pathogen interactions.

By using known invertebrate Tectonin proteins, we performed domain- and conserved position-specific iterated sequence searches, and identified a potential human homolog, which we dubbed the hTectonin. We also discovered that the domain architecture of hTectonin is well conserved throughout the different species, suggesting that it is an important functional protein. Sequence motif analysis, and prediction of the secondary and tertiary structures suggests that hTectonin is a β-propeller protein, in accordance to the definition of the Tectonin domain. Specifically, only the Tectonin domains of hTectonin were found to interact with the fibrinogen-like domain of M-ficolin, an important complement initiator [12]. In addition, the hTectonin domains 6 and 11 also exhibited LPS-binding properties. The specificity of recognition of LPS by certain Tectonin domains is consistent with the invertebrate Tectonins such as the limulus GBP. We suggest that hTectonin forms a β-propeller structure involved in protein-protein interaction amongst host proteins and also in pathogen-detection, thus playing a vital role in bridging the host immune defense proteins to the invading pathogen, and this phenomenon is probably conserved over a vast number of organisms.

Results

hTectonin identified from the human genome database – a hypothetical protein?

In mammals, the identity and role of proteins with Tectonin domains are unknown. Those identified or studied in the invertebrates [2–4,6,13–21] as well as the first vertebrate, fish, exhibit immune defense properties. Here, we sought to examine whether the Tectonin domains are structurally and functionally conserved in the mammals. A position-iterated search using known Tectonin domain-containing proteins in the invertebrates revealed a family of vertebrate Tectonin proteins to be distantly related (Figure 1A). This includes the human protein, Q7Z6L1, which is one of 3 human hypothetical proteins (GenBank Accession Nos: Q7Z6L1, Q15040 and O95714) that contain the Tectonin domain architecture [22,23], when a domain architectural search was done on Tectonin domain-containing proteins. Q7Z6L1 codes for a predominantly Tectonin domain-containing protein (Figure 1B), suggesting that the domains probably form an essential part of the molecular structure and play a vital role. Furthermore, the high architectural homology of Q7Z6L1, from the slime mold to the human, suggests its evolutionary conservation and functional significance (Figure 2). We thus selected Q7Z6L1 which codes for ‘hTectonin’ for molecular expression, and further structural and functional analyses.

hTectonin consists of β-propeller secondary structure

From the multiple sequence alignment (MSA) of the Tectonin domains, we confirmed a pattern of sequence repeats of 40 to 50
residues in length, which is a unique characteristic of β-propellers [9,10]. In addition, secondary structure prediction of hTectonin by PSIPRED [24,25] predicted these conserved repeats to form the β-strands of a β-sheet topology, consistent with β-propeller architecture (Figure 3).

hTectonin interacts with ficolin through its Tectonin domains
Based on our observations that a Tectonin protein, GBP (GenBank Accession No. AAV65031.1), interacts with two complement proteins, C-reactive protein (CRP) and carcirolec-
tin (CL5), and is therefore immune-related, we reasoned that the hTectonin might play a similar role in immune defense. We tested and showed that the hTectonin gene is expressed in the human T cell line (A549), monocytes (U937) and the human leukocytes (Figure 4A), corroborating its immune relevance. Based on the rationale that (i) hTectonin is an architectural homolog of GBP and (ii) as a pathogen pattern-recognition receptor, GBP interacts with CL5 [26], which is a homolog of the human ficolin, we performed yeast 2-hybrid analysis using hTectonin as bait and the three isoforms of ficolins (L-, H- and M-ficolin) as prey. Results showed that the hTectonin (clone QZ7L1) interacts specifically with M-ficolin (GenBank Accession No. O00602) (Figure 4B). M-ficolin has in turn been shown to interact with the CRP [26]. Since both the CRP and M-ficolin are key proteins of the complement classical and lectin pathways, respectively, this is the first evidence for the potential function of a human Tectonin domain-containing protein in frontline immune defense. Further delineation of hTectonin to isolate its functional domains showed that only the sub-clones expressing the predicted Tectonin domains interacted with M-ficolin. Furthermore, only the fibrinogen-like (FBG) domain of M-ficolin was shown to interact with the hTectonin, concurring with recent findings that the FBG domain is responsible for ligand-binding [12]. These results suggest that the protein-protein interaction between the hypothetical hTectonin and M-ficolin is not random, but structurally and positionally specific, and that the hTectonin is potentially involved in immune regulation, acting through its Tectonin domains.

Figure 4. hTectonin exists and interacts with immune-related genes. (A) hTectonin cDNA is found in the human T cells (A549), monocytes (U937) and leukocytes. (B) hTectonin interacts with ficolin. Yeast 2-hybrid shows that hTectonin interacts (i) with itself, suggesting the possibility of oligomerization, as observed in other beta-propeller proteins; and (ii) with ficolin, a human complement protein. Furthermore, interaction with ficolin specifically occurs through the Tectonin domains of the hTectonins. This demonstrates a possible functional conservation of Tectonin domains since the Tectonin domains of GBP (horseshoe crab Tectonin lectin) was shown to interact with carcinolectin-5, a homologue of ficolin [26].

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Tectonin domains harbor high avidity LPS-binding motifs

Gram-negative bacterial endotoxin or lipopolysaccharide (LPS) is a prominent and well-studied representative pathogen-associated molecular pattern. Proteins harboring LPS-binding motifs, with alternating basic-hydrophobic/polar residues (BHB(P)HB), have been shown to bind LPS via the lipid A moiety [27,28], which is the most conserved bioactive pathophysiological centre of the LPS molecule (Supporting Figure S1A). Based on the BHB(P)HB pattern, we identified two such motifs in the 6th and 11th Tectonin domains of the hTectonin and found that these motifs were well-conserved among the mammalian homologs of hTectonin in addition to being in a region of high sequence conservation (Figure 5). Representative Tectonin peptides were synthesized around the BHB(P)HB motifs in Tectonin domains 6 & 11, and their efficacy of binding of lipid A was compared with peptides derived from the GBP Tectonin domains 1 & 6 (Supporting Figure S1B), where similar BHB(P)HB motifs exist. Real-time biointeraction of these Tectonin peptides to lipid A immobilized on biacore HPA chip showed that indeed the hTectonin peptides bound the lipid A at affinities of KD $10^{-7}$ to $10^{-8}$ M, which are similar to the GBP peptides (Figure 6 and Table 1). We also showed that both the hTectonin and the GBP peptides exhibited similar level of binding affinity to ReLPS and LPS (Figure 6 and Supporting Figure S1A,C). Table 1 summarises and compares the binding affinities of various peptides derived from the hTectonin and GBP. This corroborates our hypothesis and demonstrates the pathogen-binding ability of Tectonin domains and its functional conservation across species, from horseshoe crab GBP to human hTectonin.

Discussion

In order to classify and complete the functional characterization of the human proteome, many of the unknown proteins are usually inferred from their counterparts in other species. This seems to be an easy option if the proteins share high sequence similarity, as they can be matched to each other by performing a simple sequence matching. However, the task is more complicated if the proteins do not show homology in their primary sequences. Nevertheless, many related proteins show conserved functionality more in terms of domain and structural conservation.

In this paper, we report our discovery of a human Tectonin protein, hitherto classified as being hypothetical. By structure-function analyses, we inferred its function as an immune-related protein. We showed that similar to its invertebrate counterparts, the hTectonin protein functions via its Tectonin domains. Furthermore, a distance PSI-BLAST sequence matching indicates that although the hTectonin shows low sequence homology, it is phylogenetically related to known proteins with Tectonin domains, functioning as immune proteins. By SMART domain comparison, we show that hTectonin contains multiple homologs widespread in the vertebrate kingdom, implying that it is not a one-off protein in the human proteome, but rather, an important one conserved throughout many species. We also discovered that the hTectonin gene is expressed in the human leukocytes. This is interesting, as a recent addition to the human database of proteins showed another human leukocyte Tectonin protein called the leukectin (GenBank Accession No. ACM77812.1) [29], which also exhibits five Tectonin domain repeats (Figure 1B). This further implicates hTectonin to be immune-related. Like its limulus counterpart, GBP, which interacts with an important complement initiator (CRP), we find that the hTectonin also interacts with a cognate complement lectin, Ficolin. Furthermore, this interaction is specific, involving only the Tectonin domains within the hTectonin protein. We also identified LPS-binding motifs within two of the Tectonin domains which are located in the highly conserved sequence of the β-propeller fold. The affinity...
of these motifs for bacterial LPS and the truncated active forms of the endotoxin molecule (ReLPS and lipid A) was verified experimentally. Thus, we propose that hTectonin is a novel human protein that forms a β-propeller structure which is involved in protein-protein interaction with immune-related proteins such as ficolin, and it simultaneously interacts with pathogens via PAMPs like LPS. Thus, the hTectonin plays a vital role in immune defense, which is conserved over a vast number of organisms.

Materials and Methods

Identification of Tectonin proteins

Tectonin domain containing proteins were identified using domain search on the SMART database [22,23]. A position-specific iterated search using the primary sequence on PSI-Search on the EMBL server was performed using GBP as the query sequence. Related sequences were chosen after 2 iterations of PSI-Search. Hits were put through the SMART prediction server to confirm their propensity to form Tectonin domains. Multiple sequence alignment was carried out on the curated list of proteins using Promals3D [30]. A phylogenetic tree was then constructed from sequences showing strong domain alignments using PHYLIP [31] with a bootstrap value of 1000.

Human hTectonin cDNA clones

The hTectonin (Q7Z6L1) cDNA was obtained from iDNA Open-Biosystems (MHS1010-9205594). The cDNA was subcloned into pGBK7T and pGADT7 vectors for the yeast 2-hybrid experiments.
Table 1. Dissociation constants of Tectonin peptides when bound to LPS, ReLPS, lipid A.

| Bacterial ligand | Peptide | Sequence (LPS-binding motif underlined) | $K_0$ (m$^{-1}$) |
|------------------|---------|-----------------------------------------|----------------|
| LPS              | GBP6-1  | KSCWLNPFLAEWTHINGKLSH                  | 2.55 $\times$ 10^{-7} |
|                  | GBP6-1  | FESVPASKAEWTHINGKLSH                   | 3.56 $\times$ 10^{-7} |
|                  | hTectonin6 | LSSLSCESKVQGPRP0PAI                 | 2.03 $\times$ 10^{-7} |
|                  | hTectonin11 | IGGWDHISVRANATRAPRS              | 9.27 $\times$ 10^{-7} |
| ReLPS            | GBP6-1  | 2.04 $\times$ 10^{-7} |
|                  | hTectonin6 | 3.79 $\times$ 10^{-8} |
|                  | hTectonin11 | 2.24 $\times$ 10^{-6} |
| Lipid A          | GBP6-1  | 9.83 $\times$ 10^{-8} |
|                  | hTectonin6 | 5.63 $\times$ 10^{-6} |
|                  | hTectonin11 | 4.17 $\times$ 10^{-8} |
|                  |                         | 1.42 $\times$ 10^{-6} |

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Yeast-2-hybrid assay for protein-protein interaction

Co-transformations of the different bait and prey plasmids into *S. cerevisiae* AH109 strain were performed in accordance to [16]. The full-length and subclones of hTectonin and the ficolin cDNAs (without their signal sequences) were each fused to the activation domain of Gal4 in the prey vector. Positive transformants were selected in SC media lacking Leu and Trp. hTectonin plasmid in pGBKT7 was then transformed into the library-positive yeast. DNA from resulting colonies from the co-transformations of the different bait and prey plasmids into SC media lacking Leu and Trp. hTectonin plasmid in pGBKT7 was then transformed into the library-positive yeast. DNA from resulting colonies from the co-transformations on QDO agar were extracted and identified through sequencing.

Peptide design and synthesis

The hTectonin protein sequence was scanned for LPS-binding motif. Two potential sites with the BHPHB pattern were found in hTectonin domains 6 (KVQGR) and 11 (HISVR). Henceforth, these peptides are referred to as hTec peptides (hTec6 and hTec11). The hTec peptide length and region surrounding the LPS-binding motif was chosen and optimized based on hydrophilicity and solubility values. The hTec was: LSSLCSERKVQGPRP0PAI and hTec11 was IGGWDHISVRANATRAPRS. For comparison, one BHPHB site was found in the limulus GBP, Tectonin domain 1 (HNGK). The GBP peptides are: GBP6-1(tail) KSCWLNPFLAEWTHINGKLSH and GBP6-1(no tail) FESVPASKAEWTHINGKLSH, which are annotated based on the amino acid residues which encompass the domains 6-to-1. Peptides were also designed from the combination of GBP Tectonin domains 1 and 6. The peptides were synthesized by Gensyn, Inc., USA, and purified to >95% under pyrogen-free conditions.

Surface plasmon resonance analysis of the peptides

Surface plasmon resonance analysis for real-time biointeraction between the Tectonin peptides and bacterial LPS was performed using a Biacore 2000 instrument (Biacore AB). LPS, ReLPS and lipid A from *Salmonella minnesota* (List Biologicals, UK) were diluted to 0.25 mg/ml in 20 mM sodium phosphate, 150 mM NaCl, pH 7.4 and immobilized on the surface of an HPA sensor chip (Biacore AB) according to the manufacturer's specifications. Binding of the Tectonin peptides to the immobilized ligands was measured at a flow rate of 20 μl/min in 10 mM Tris, 150 mM NaCl, pH 7.4. Regeneration of the chip surface was achieved by injection of 20 μl 0.1 M NaOH until steady baseline was achieved. The dissociation constant, KD was calculated using BioEvaluation software, version 3.2.

Supporting Information

Figure S1 (A) Structure of the bacterial LPS. LPS structure and the truncated forms, ReLPS and lipid A. (B) The structure of GBP, with the tail (circled) at the C-terminal end, which does not form the β-propeller structure of GBP. (C) Control Tectonin peptides which do not harbor the LPS-binding motif of BHPHB do not bind lipid A. Found at: doi:10.1371/journal.pone.0006260.s001 (0.24 MB TIF)

Author Contributions

Conceived and designed the experiments: DHPL QY VF JLD. Performed the experiments: DHPL ZA QY. Analyzed the data: DHPL ZA VF BH JC JLD. Wrote the paper: DHPL ZA VF BH JC JLD.

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