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Permalink
https://escholarship.org/uc/item/52f5q2p4

Journal
PLoS ONE, 9(12)

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Publication Date
2014-12-05

DOI
10.1371/journal.pone.0114685

Peer reviewed
RESEARCH ARTICLE

Structure of the Essential *Plasmodium* Host Cell Traversal Protein SPECT1

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Abstract

Host cell traversal by *Plasmodium*, the protozoan cause of malaria, is an essential part of this parasite's virulence. In this process, the parasite enters a host cell through a parasite-induced pore, traverses the host cell, and then exits the host cell. Two *P. berghei* proteins, SPECT1 and SPECT2, are required for host cell traversal by the sporozoite form of the parasite. In the absence of either, no pore formation is observed. While SPECT2 has sequence homology to pore-forming proteins, SPECT1 has no homology to proteins of known structure or function. Here we present the 2.75 Å resolution structure of a slightly truncated version of *P. berghei* SPECT1. The structure reveals that the protein forms a four-helix bundle, with the rare feature of having all of these helices in parallel or antiparallel alignment. Also notable is the presence of a large, conserved, hydrophobic internal cavity in the protein, which may constitute a ligand-binding site or be indicative of partial instability in SPECT1, or both. The structure of SPECT1 will make possible targeted mutagenesis experiments aimed at understanding its mechanism of action in host cell traversal.

Introduction

Malaria exerts a global burden on human health and welfare, being endemic in 100 countries and causing 250 million clinical illnesses and 1 million deaths each year [1, 2]. This devastating disease is caused by the infective sporozoite form of the protozoan parasite *Plasmodium*, which is introduced into the dermis of a host through the bite of an infected female Anopheline mosquito. Sporozoites are highly motile and transit from the dermis through the circulatory system to the liver, where they invade hepatocytes and mature into the red blood cell-infective form of the parasite [3, 4]. Prior to invading hepatocytes, sporozoites have the
ability to traverse host cells [5–7]. Host cell traversal involves entry into a host cell by the sporozoite, transit through the host cell cytosol, and finally exit from the host cell. Entry and presumably exit require the formation of a pore in the host cell plasma membrane [5,7,8], resulting in a membrane wound that causes necrosis in most instances, although in the some cases the wound is resealed and the cell survives [5,9,10]. Host cell traversal was first observed in macrophages [11], but has since been documented in other cell types, including hepatocytes, epithelial cells, and fibroblasts [5,7,10]. There appears to be no cell type specificity for host cell traversal, suggesting that only *Plasmodium* factors are important for this process.

Host cell traversal plays a critical role in *Plasmodium* infectivity. Loss of host cell traversal results in a loss in infectivity due to the killing of the parasite by Kupffer cells, the resident macrophage of the liver [6,8,12]. A further loss in infectivity occurs due to the entrapment of sporozoites by phagocytic leukocytes as well as nonphagocytic cells in the dermis [7]. The combined loss in infectivity due to effects in the dermis and liver appears to be ~300-fold (10-fold in the dermis and 30-fold in the liver).

Two proteins from *P. berghei*, a rodent-infecting parasite, have been identified to be essential for host cell traversal by sporozoites [8,12]. These are *Pb*SPECT1 and *Pb*SPECT2 (sporozoite microneme protein essential for cell traversal). No pore formation or host cell membrane wounding is seen in the absence of either of these proteins [7,8,12]. This suggests that *Pb*SPECT1 and *Pb*SPECT2 work together or in consecutive fashion to form a pore. Both *Pb*SPECT1 and *Pb*SPECT2 have signal sequences and localize to micronemes [8,12], secretory organelles at the apical end of sporozoites.

The sequence of *Pb*SPECT2 (also called *Plasmodium* perforin-like protein1, PPLP1) suggests that it has a direct role in pore formation. This protein of ~800 amino acids has a central domain of ~330 amino acids that has homology to pore-forming proteins, namely those belonging to the membrane attack complex/perforin (MACPF) and cholesterol-dependent cytolysin (CDC) family [12–15]. MACPF/CDC proteins are synthesized as soluble proteins and are triggered to undergo a conformational change that promotes insertion and pore formation in target membranes. Recombinant *P. falciparum* SPECT2 has been shown to cause lysis of red blood cells in a Ca\(^{2+}\)-dependent manner, as has the MACPF/CDC domain of *Pf*SPECT2 [16]. The role of *Pf*SPECT2 in host cell traversal by sporozoites, however, has not been demonstrated. Instead *Pf*SPECT2 has been implicated in the Ca\(^{2+}\)-dependent egress of *P. falciparum* merozoites from red blood cells [16]. *Pf*SPECT2 and *Pb*SPECT2 appear to have different functions, as the latter has been detected only in sporozoites and is absent from merozoites [12]. *Pb*SPECT2 has not yet been demonstrated to form a pore, and as noted above, the genetic data indicate that the presumed pore-forming activity of *Pb*SPECT2 is dependent on *Pb*SPECT1.

In contrast to *Pb*SPECT2, *Pb*SPECT1 (~25 kDa) has no detectable primary sequence homology to proteins of known structure or function [8]. *Pb*SPECT1 has no identifiable membrane-spanning or attachment regions. *Pb*SPECT1 has
been found to be expressed in salivary gland sporozoites by not in merozoites, ookinetes, or midgut sporozoites [8]. Homologs of PbSPECT1 are encoded in the genomes of a number of Plasmodium species, including P. falciparum. These proteins have ~40% sequence identity, suggesting that they share a common structure and mode of action. Homologs of SPECT1 do not exist outside of Plasmodium species. To begin to understand how PbSPECT1 contributes to host cell traversal, we recombinantly expressed, purified, and determined the crystal structure of a slightly truncated version of this protein. We found that PbSPECT1 forms a four-helix bundle, with the rare feature of having all of these helices in parallel or antiparallel alignment. This along with a large, conserved, hydrophobic internal cavity in PbSPECT1 are suggestive of conformational lability in the molecule, which would accord with a mechanism in which the protein is triggered to undergo a conformational change from soluble to membrane-associated form, as typical for many pore-forming proteins (including those belonging to the MACPF/CDC family).

**Materials and Methods**

*P. berghei* SPECT1 expression and purification

*P. berghei* SPECT1Δ24, containing amino acids 25–241, and SPECT1Δ41, containing amino acids 42–241, were cloned from a plasmid generously provided by Masao Yuda into a modified pET28 vector (Novagen) as a C-terminal fusion to *S. cerevisiae* SUMO (SMT3, NP_010798.1); these constructs included an N-terminal His-tag preceding SUMO (Table S1). The expression vectors were separately transformed into *E. coli* BL21 (DE3). Bacteria were grown with shaking at 37°C in LB media containing 50 µg/mL kanamycin to an OD600 of 0.8–1.0, whereupon expression was induced with 0.4 mM IPTG; the cultures were maintained thereafter for 16–18 h at room temperature. Bacteria were harvested by centrifugation (2246 × g, 20 min, 4°C), resuspended in lysis buffer (100 mM sodium phosphate buffer [NaPi], pH 8.0, 500 mM NaCl, and 40 mM imidazole), and lysed using an Emulsiflex (Avestin). The lysates were clarified by centrifugation (37,500 × g, 30 min, 4°C), filtered through a 0.45 µm filter, and applied to columns containing Ni²⁺-nitrilotriacetic acid (NTA, Sigma) agarose beads (10 mL per L of bacterial culture); prior to application, the columns were equilibrated in binding buffer [NaPi, pH 8.0, 500 mM NaCl] containing 40 mM imidazole. The columns were washed with 10 column volumes of binding buffer, and bound proteins were eluted in binding buffer containing 250 mM imidazole.

The His-tag and SUMO fusion partners were cleaved from PbSPECT1Δ24 and PbSPECT1Δ41 using His-tagged *S. cerevisiae* ULP-1. The cleavage reaction was carried out overnight at a 4:1 substrate:ULP-1 molar ratio at 4°C in binding buffer supplemented with 250 mM imidazole, 1 mM DTT, and 5 mM EDTA. The samples were then dialyzed (6000–8000 MWCO membrane) in binding buffer at 4°C and applied to Ni-NTA columns in the same buffer. Fractions of
PbSPECT1Δ24 or PbSPECT1Δ41 that were devoid of their fusion partners were collected from the flow-through of these columns. These fractions were then exchanged into 20 mM Tris, pH 8, 50 mM NaCl using a HiPrep 26/10 desalting column (GE Healthcare) and applied in this same buffer to a Q Sepharose anion exchange column (GE Healthcare). Protein that was bound to this column was eluted using a step-wise gradient to 20 mM Tris pH 8, 200 mM NaCl. PbSPECT1Δ24 and PbSPECT1Δ41 were further purified by gel filtration chromatography (Superdex 75 26/60, GE Healthcare) in 20 mM Tris, pH 8, 10 mM NaCl. Purified PbSPECT1Δ24 and PbSPECT1Δ41 were concentrated by binding the proteins to a 1 mL HiTrap Q anion exchange column (GE Healthcare) and eluting the protein using a step-wise gradient to 20 mM Tris, pH 8, 500 mM NaCl. The concentrated proteins were dialyzed at 4 °C using a 3500 MWCO Slide-A-Lyzer (Thermo Scientific) in 1 mM Tris, pH 8, 10 mM NaCl. The dialyzed proteins were flash frozen in liquid N2 and stored at −80 °C.

Two dual Met-substitution mutants of PbSPECT1Δ41, L57M/L75M and L57M/I133M, were constructed for the purpose of phasing. The mutagenesis was carried out using a combination of MegaPrimer [17] and QuikChange (Agilent Technologies, Inc.) techniques (Table S1). Selenomethionine (SeMet) was incorporated into PbSPECT1Δ41 L57M/I133M using methionine pathway inhibition as described previously [18], except that biotin was omitted and 20% glucose (w/v) was included in the M9 minimal salts media. SeMet-labeled SPECT1Δ41 L57M/I133M was expressed and purified as described above. PbSPECT1Δ41 L57M/L75M was produced without SeMet-labeling, and expressed and purified as described above; this protein was derivatized with thimerosal, as described below, for the purpose of phasing.

Proteinase K Digestion

PbSPECT1Δ24 (4 μg) was incubated with 3×10⁻⁵ μg, 3×10⁻⁶ μg, 3×10⁻⁷ μg, 3×10⁻⁸ μg, or 3×10⁻⁹ μg proteinase K in a 50 μL reaction volume containing 25 mM Tris, pH 8.0, 25 mM CaCl₂, 75 mM NaCl for 1 hr at 37 °C. Digestion products were visualized by Coomassie-stained reducing SDS-PAGE.

Multiangle Light Scattering

Absolute molecular masses were determined by multiangle static light scattering. Protein samples in 150 mM NaCl, 50 mM Tris or HEPES, pH 7.4 were applied to a TSK-Gel G3000 SWxl size-exclusion column (Tosoh, Bioscience) attached to a miniDAWN TREOS SLS detector and an Optilab T-rEX refractive index detector (Wyatt, Santa Barbara, CA). Data were processed using ASTRA 5 software (Wyatt, Santa Barbara, CA).

Crystallization

PbSPECT1Δ24 was crystallized by hanging drop vapor diffusion. The wells of a 24-well plate were filled with 1 mL of 0.1 M Bis-tris methane, pH 5.7, 2.5% (w/v)
PEG 400, 1.2 M (NH₄)₂SO₄. PbSPECT1Δ24 at ~13 mg/mL was mixed with the reservoir buffer at a 1:1 ratio in a 1 μL crystallization drop.

Native PbSPECT1Δ41 was crystallized using an Oryx8 Crystallization Robot (Douglas Instruments) in a 96-well sitting drop plate (CrystalClear Duo, Douglas Instruments). The reservoir contained 0.08 mL of 0.2 M MgCl₂•6H₂O, 0.1 M HEPES, pH 7.5, and 25% (w/v) PEG 3350. Crystallization drops were 1 μL and contained 70% protein at 19.3 mg/mL (determined using an ε₂₈₀ of 8300 M⁻¹cm⁻¹) and 30% reservoir solution. Plates were incubated at 18°C. Clusters of crystals formed after ~2 weeks. The clusters were mechanically broken up and large fragments were dragged using a loop through an oil mixture consisting of 3:1 paraffin:Paratone N before flash cooling in liquid N₂. X-ray diffraction data were collected at beamline 23-IDB at APS.

SeMet-labeled PbSPECT1Δ41 L57M/L133M was crystallized by a modified batch method. The wells of a 24-well dish were filled with 0.5 mL of 0.3–0.4 M MgCl₂•6H₂O, 0.1 M HEPES, pH 7.5, and 30–37% (w/v) PEG 3350. SeMet-labeled PbSPECT1Δ41 L57M/L133M at ~45 mg/mL was mixed with the reservoir buffer at a 1:1 ratio in a 2 μL crystallization drop. The reservoir buffer was diluted 1:1 with water before sealing the well with a coverslip. Crystals formed within ~4 days and were cryopreserved as described above for native crystals.

PbSPECT1Δ41 L57M/L75M was crystallized by the modified batch method described above. Mercury derivatization of crystals of this protein was achieved by picking up a single crystal of thimerosal using a loop and adding it directly into the crystallization drop containing the protein crystal; the thimerosal crystal was observed to dissolve under a microscope. The PbSPECT1Δ41 L57M/L75M crystal was incubated in thimerosal at 18°C for 2 hours before flash cooling as described above.

Structure Determination

Native diffraction data were collected at APS beamline 23-ID-D from a crystal of PbSPECT1Δ41, which belonged to space group P2₁2₁2₁ and had four PbSPECT1Δ41 molecules in the asymmetric unit. Multiple wavelength anomalous dispersion (MAD) data were collected at SSRL beamline 9.2 from a crystal of SeMet-labeled PbSPECT1Δ41 L57M/L133M, and single wavelength anomalous dispersion (SAD) data were collected at APS beamline 23-ID-B from a crystal of thimerosal-derivatized PbSPECT1Δ41 L57M/L75M. Crystals of Met-substituted PbSPECT1Δ41 belonged to space group C22₂₁, and had two PbSPECT1Δ41 molecules in the asymmetric unit. Crystallographic data were processed with HKL2000 [19]. The SeMet MAD and thimerosal SAD datasets were assigned separate phasing groups in Phenix's AutoSol program [20] and the phasing information from both datasets were combined into a single set of phases. Four selenium sites and two mercury sites were identified in each asymmetric unit, and the density modified electron density map showed clear evidence for a four-helix bundle. Four polyalanine α-helices were manually placed into electron density maps using Coot [21]. The resulting model was used in Phenix as a search model
in molecular replacement phasing of diffraction data from crystals of native PbSPECT1Δ41 [20]. Electron density for amino acid side chains became more apparent in the resulting electron density map, and side chains were assigned based on the positions of the anomalously scattering atoms. Twenty-four iterations of maximum likelihood restrained refinement using Phenix (with default parameters) followed by manual rebuilding into $\sigma_A$-weighted 2mFo-DFc and mFo-DFc maps using COOT were carried out. Non-crystallographic symmetry restraints were applied at each step of refinement using default parameters in Phenix. Electron density for the main chain was continuous for the following residues: 48–82, 88–197, and 206–240 in chain A; 50–82, 89–239 in chain B; 47–82, 86–167, 174–197, and 210–239 in chain C; and 47–82, 91–197, and 207–240 in chain D. Analysis by MolProbity [20] yielded a clashscore of 5.8 and a MolProbity score of 1.74; MolProbity reports these as the 100th percentile for both score categories.

Molecular figures were made with PyMol (). Structure-based sequence alignments were generated using 3D-Coffee [22] and displayed using ESPript [23]. Mapping of conserved residues to the molecular surface of PbSPECT1Δ41 was carried out with ConSurf [24]. Analysis of the cavity in PbSPECT1Δ41 was carried out with CASTp [25]. The crystal structure and structure factors have been deposited with the Protein Data Bank (accession code 4U5A).

Results and Discussion

We sought to crystallize and determine the structure of PbSPECT1 to gain insight into its mechanism of action in host cell traversal. A slightly truncated version of PbSPECT1 containing amino acids 25–241 (called PbSPECT1Δ24) was initially chosen for crystallization trials. The first nineteen amino acids of PbSPECT1 make up a predicted signal sequence [26] and were therefore excluded from this construct (Fig. 1a), and the first several amino acids of mature PbSPECT1 are predicted to be disordered [27] and were therefore also excluded from this construct. PbSPECT1Δ24 was expressed in E. coli, purified, and crystallized. These crystals, however, were found to yield no observable X-ray diffraction (with the use of a synchrotron source) under a variety of cryoprotectant conditions. Similarly, no diffraction was observed (using a rotating anode source of X-rays) when these crystals were held at room temperature.

To overcome this issue, a further truncation of PbSPECT1 was carried out. For this, PbSPECT1Δ25 was subjected to proteinase K digestion, resulting in the production of a stable, shorter fragment. This fragment consisted of residues 42–241, as identified by N-terminal sequencing and MALDI-TOF mass spectrometry. A construct encoding PbSPECT1 residues 42–241, called PbSPECT1Δ41 (Fig. 1a), was then expressed in E. coli and purified. PbSPECT1Δ41 was observed by size-exclusion chromatography (SEC) to exist as two separable species (Fig. 1b). These species were stable, as the isolated forms maintained their state over the course of days as assessed by SEC. Multiangle light scattering, which provides a shape-
independent measure of mass, showed that these species corresponded to monomeric (Mₙ 21,380 Da; calculated 23,184 Da) and dimeric (Mₙ 45,070 Da) forms of SPECT1Δ41 (Fig. S1). While the dimeric form of PbSPECT1Δ41 did not crystallize, the monomeric form did. It is worth noting that PbSPECT1Δ24 also formed monomeric and dimeric species, and it was the monomeric form that yielded the non-diffracting crystals mentioned above.

Crystals of PbSPECT1Δ41 diffracted to 2.7 Å resolution (Table 1), and its structure was determined by a combination of isomorphous replacement and multi-wavelength anomalous dispersion (Fig. S2). While monomeric PbSPECT1Δ41 was used in the crystallization, an association suggestive of a dimer was seen in the asymmetric unit. Two PbSPECT1Δ41 molecules came together in a “V”-shaped arrangement, with a total of ~1260 Å² being buried at the interface (Figs. 1c and S3). This is the largest intermolecular buried surface area in the crystal, with the next largest being ~1020 Å². The asymmetric unit is constituted by two of these “V”-shaped, presumptive dimers that are then related by noncrystallographic translational symmetry (Fig. S3b). In crystals of methionine-substitution mutants of PbSPECT1Δ41 (produced for phasing purposes), the same intermolecular packing was seen but the translational noncrystallographic symmetry became crystallographic (changing the space group from P2₁2₁2₁ to C22₁). Whether PbSPECT1 forms oligomers in effecting host cell traversal is not known, but the dimeric state observed by SEC and the association observed in the crystal raise the possibility that a dimer may be involved.

The structure of PbSPECT1Δ41 revealed that it forms a four α-helix bundle with a ‘hook’-like feature at one end (Figs. 2a and 2b). No electron density was visible for the first 5–9 residues or the last 1–2 residues of PbSPECT1Δ41, likely due to the flexibility of these amino acids. The helices in the four-helix bundle range between 29–39 residues in length and proceed in an up-and-down fashion. Helices 2 (α2) and 4 (α4), which pack against one another, contain prolines (P101

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Figure 1. Monomeric and Dimeric Forms of PbSPECT1Δ41. a. Schematic to scale of PbSPECT1. Yellow denotes the putative signal sequence, purple the portion of the mature protein not included in PbSPECT1Δ41, and blue PbSPECT1Δ41. b. Size exclusion chromatogram of PbSPECT1Δ41 showing the appearance of monomeric and dimeric species. c. “V”-shaped association between two PbSPECT1Δ41 molecules (red and blue) observed in the asymmetric unit of the crystal.

doi:10.1371/journal.pone.0114685.g001
in \( \alpha2 \) and P216 in \( \alpha4 \) and are kinked due to these. These prolines are not conserved in \( PbSPECT1 \) homologs (Fig. 2c), and thus the kinks are unlikely to be functionally consequential. The connections between the four helices in the bundle are between nearest neighbors, except for a diagonal connection between \( \alpha2 \) and \( \alpha3 \). These connections mainly consist of short loops. The loop connecting \( \alpha1 \) with \( \alpha2 \) appears to be highly flexible, as no density for 4–8 residues in this loop, depending on the \( PbSPECT1\Delta41 \) molecule, was evident. Analysis of crystals confirmed that the lack of density in this region was not due to inadvertent proteolysis in the loop (Fig. S4). The “hook” feature of \( PbSPECT1\Delta41 \) is composed of short antiparallel \( \alpha \)-helices (\( \alpha3a \) and \( \alpha3b \)) that are situated obliquely to the four-helix bundle and form a connection between helices \( \alpha3 \) with \( \alpha4 \). The segment leading from the “hook” to the \( \alpha4 \) helix appears also to be flexible, as

<table>
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<th>( PbSPECT1\Delta41 )</th>
<th>Thimerosal-derivated ( PbSPECT1\Delta41 ) L57M L75M</th>
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<th>SeMet ( PbSPECT1\Delta41 ) L57M/L133M (peak)</th>
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<td>50.00–3.52 (3.65–3.52)</td>
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electron density for this region (residues 198–205) was visible in only one of the four PbSPECT1Δ41 molecules in the asymmetric unit; this electron density was sufficient for modeling only the main chain. Other than this difference, the four PbSPECT1Δ41 molecules are nearly identical in structure (root-mean-square-deviation ranging between 0.49–0.87 Å for main chain atoms).

One face of PbSPECT1Δ41 has several patches of conserved residues (Fig. 3a, left). One of these patches (composed of L77, D78, E80, S82, S95, K173) is proximal to the “hook” and has an overall negatively charged electrostatic character (Fig. 3b, left). The other (K56, D60, D64, K139, C142, L146) is distal to the “hook” and has a negatively charged stripe separated from a positively charged one. These two patches are connected by a set of conserved residues that line a deep pocket (K69 and E102, among others) that is described below. The opposite face of PbSPECT1Δ41 is more variable and has no distinctive electrostatic character (Figs. 3a and 3b, right).

The four-helix bundle is a common structural motif and occurs in proteins of diverse functions [28]. For the vast majority of four-helix bundle proteins, the helices are tilted at about +25° or -35° with respect to one another [28–31]. This optimizes the packing between side chains. In contrast to this pattern, the helices in PbSPECT1Δ41 are nearly parallel or antiparallel to one another; the angles between the helices range between 1–8° from parallel/antiparallel. Parallel/antiparallel packing is rare in four-helix bundle proteins, but has been observed to occur at a frequency greater than that predicted by random chance [30, 31]. This latter point suggests that the parallel/antiparallel feature of these proteins is under positive selection.

Since the presence of parallel/antiparallel helices in four-helix bundles can be a sign of poor side chain packing, we examined the core of PbSPECT1Δ41 [25]. The portion of the structure distal to the ‘hook’ has a close-packed interior, which is composed of a large number of conserved leucines and isoleucines (Fig. 4a). In contrast, a striking and spacious cavity occurs in the interior of the structure proximal to the “hook” (Fig. 4a). The volume of this cavity is ~750 Å³, but in actuality may be just somewhat smaller as six residues in the α1-α2 loop, which is at the top of the cavity, were not modeled. This cavity is lined predominantly by hydrophobic residues (L75, F76, I79, A81, S82, D89, V90, V91, I94, Y97, N156, K160, I163, I164, Y167, G168, K169, V207, L210, I214, I217, Y218, F221), with most of these being conserved in PbSPECT1 homologs (Fig. 2c). Of further note,
this cavity is juxtaposed against an unusually deep pocket (145 Å² in area) on the surface of PbSPECT1Δ41 (Fig. 4). Most of the residues lining this deep pocket (K69, Q70, I72, K73, F76, S98, P101, E102, and L105) are also conserved in

Figure 3. Surface of PbSPECT1Δ41. a. Residues conserved among homologs of PbSPECT1 were mapped to the molecular surface of PbSPECT1Δ41. Two views of PbSPECT1 related by a 180° rotation around the indicated axis are shown. Color scale for conservation is shown at bottom. b. The electrostatic surface mapped to the van der Waals surface of PbSPECT1Δ41 is shown. As in panel a, two views and a color scale are shown.

doi:10.1371/journal.pone.0114685.g003

Figure 4. The internal cavity of PbSPECT1Δ41. a. Stereoview of the interior cavity in PbSPECT1Δ41 depicted as a surface, and the backbone of PbSPECT1Δ41 as sticks. The positions of the cavity, pocket, and hook are indicated. b. A molecular surface representation of PbSPECT1Δ41 with its deep pocket is shown. Residues at the entrance of the pocket are indicated.

doi:10.1371/journal.pone.0114685.g004
PbSPECT1 homologs (Fig. 2c). The cavity and the pocket are not continuous, but with motions in the protein, it is possible that the cavity would have access to solvent through the pocket. A second possibility for solvent access to the cavity is near the “hook”. As noted above, this area contains the disordered (and unmodeled) α1-α2 loop residues, and so it is difficult to say based on the present data whether the cavity opens to solvent at this location.

Conclusions

The structure of PbSPECT1 supplies some clues as to how this protein may function in pore formation and consequent host cell traversal. While the four-helix bundle fold of PbSPECT1 is quite widespread, the parallel/antiparallel alignment of the four helices in PbSPECT1 is quite rare [30, 31]. The great majority of four-helix bundle proteins have their helices tilted with respect to another, which enables favorable packing of side chain packs. One four-helix bundle protein, cytochrome b_{562}, has been studied in both tilted and parallel/antiparallel conformations. In its heme-bound form, cytochrome b_{562} (i.e., the holoprotein) was found to have its helices tilted at the canonical angle of ∼25° [32]. Holocytochrome b_{562} was observed to be stable [33]. However, without the heme bound, cytochrome b_{562} (i.e., the apoprotein) was found to be only marginally stable and have a poorly packed set of parallel/antiparallel helices [33, 34]. These observations bring up the possibility that the four-helix bundle in PbSPECT1Δ41 is also marginally stable and, correspondingly, conformationally labile. This would accord with a mechanism in which PbSPECT1 is triggered to undergo a conformational change from soluble to membrane-associated or inserted form.

The most distinctive structural feature of PbSPECT1Δ41 is its large interior cavity. It is intriguing to note that the marginally stable apocytochrome b_{562} has a similar feature. In cytochrome b_{562}, the hydrophobic residues that contact the heme in the stable holoprotein form become exposed to solvent and form a large exterior cavern in the unstable apoprotein form. The presence of the cavity in PbSPECT1Δ41 further suggests marginal stability, as the interiors of well folded, stable proteins are as densely packed as crystals of small organic molecules [35]. In several cases, the filling of cavities with larger side chains has been found to increase stability [36, 37]. The cavity in PbSPECT1Δ41 may be indicative of a structural defect, but may also constitute a ligand-binding pocket. In this latter regard, it is notable that the residues that line the cavity are mostly hydrophobic and conserved among SPECT1 homologs. As a point of reference for the size of ligands that could be accommodated by the ∼750 Å³ cavity of PbSPECT1Δ41, we point out that cholesterol has been measured to occupy a volume of ∼630 Å³ [38]. It is tempting to suggest that the binding of a specific ligand, which could be another protein, triggers a conformational change required for membrane activity in PbSPECT1Δ41. The structure of PbSPECT1Δ41 lays the foundation for testing this and other possibilities.
Supporting Information

Figure S1. Multiangle light scattering. Multiangle static light scattering of the PbSPECTΔ41 monomer (red) and dimer (blue) fractions applied to a gel filtration column. The fractions were run separately, but the chromatograms were superimposed for display purposes. doi:10.1371/journal.pone.0114685.s001 (TIF)

Figure S2. Electron Density. Electron density (contoured at 1σ) calculated from a 2Fo-Fc omit map shown as a mesh. The residues shown in the figure, which line the internal cavity of PbSPECTΔ41, were omitted from calculations. doi:10.1371/journal.pone.0114685.s002 (TIF)

Figure S3. “V”-shaped dimer. a. Molecular surface representation of the “V”-shaped interaction between two PbSPECT1Δ41 molecules (red and blue). Helices α1 and α2 form the greater part of the interface, with a few residues from α4 being involved as well. b. Contents of the PbSPECT1Δ41 crystal unit cell. PbSPECT1Δ41 is shown in ribbon representation, with the four molecules that constitute an asymmetric unit having the same color. There are four asymmetric units in the unit cell. doi:10.1371/journal.pone.0114685.s003 (TIF)

Figure S4. State of PbSPECT1Δ41 in crystals. Crystallization drops containing crystals of PbSPECT1Δ41 were solubilized in SDS-PAGE buffer and analyzed by Coomassie-stained SDS-PAGE. Lane 1: One-third of a PbSPECT1Δ41 crystallization drop. Lane 2: Two-thirds of a PbSPECT1Δ41 crystallization drop. Lane 3: PbSPECT1Δ41 (~4 μg). Lane 4: PbSPECT1Δ41 (~4 μg) diluted 1:1 with crystallization reservoir buffer. Lane 5: PbSPECT1Δ41 (~8 μg) diluted 1:1 with crystallization reservoir buffer. Molecular mass standards are shown between lanes 3 and 4. The crystallization reservoir contains 25% PEG 3350, which results in PbSPECT1Δ41 migrating slightly faster in the crystallization drop (lanes 1 and 2) and in samples containing PEG (lanes 4 and 5) than in the sample containing no PEG (lane 3). No proteolysis of PbSPECT1Δ41 in the crystallization drop was evident. doi:10.1371/journal.pone.0114685.s004 (TIF)

Table S1. Sequences of Primers.

doi:10.1371/journal.pone.0114685.s005 (DOC)

Acknowledgments

We thank Sumit Handa for analysis of crystals; Tomoko Ishino, Robert Ménard, Kailash Patra and Joseph Vinetz for discussions; Masao Yuda for the gift of a clone of PbSPECT1; Dirk Walther for analysis of four-helix bundle proteins; and Thu Banh for initial work on expression and purification of PbSPECT1.
Author Contributions
Conceived and designed the experiments: BYH PG. Performed the experiments: BYH. Analyzed the data: BYH PG. Wrote the paper: BYH PG.

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