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Tracy Arakaki,<sup>a,b</sup> Isolde Le Trong,<sup>a,b</sup> Eric Phizicky,<sup>b,c</sup> Erin Quartley,<sup>b,c</sup> George DeTitta,<sup>b,d</sup> Joseph Luft,<sup>b,d</sup> Angela Lauricella,<sup>b,d</sup> Lori Anderson,<sup>a,b</sup> Oleksandr Kalyuzhniy,<sup>a,b</sup> Elizabeth Worthey,<sup>b,e</sup> Peter J. Myler,<sup>b,e</sup> David Kim,<sup>b,f</sup> David Baker,<sup>a,b,f</sup> Wim G. J. Hol<sup>a,b,f</sup> and Ethan A. Merritt<sup>a,b\*</sup>

<sup>a</sup>Department of Biochemistry, University of Washington, Seattle, WA 98195-7742, USA, <sup>b</sup>Structural Genomics of Pathogenic Protozoa (SGPP) Consortium, USA, <sup>c</sup>Department of Biochemistry and Biophysics, University of Rochester School of Medicine and Dentistry, Rochester, NY 14642, USA, <sup>d</sup>Hauptman– Woodward Institute, Buffalo, NY 14203, USA, <sup>c</sup>Seattle Biomedical Research Institute, Seattle, WA 98109, USA, and <sup>f</sup>Howard Hughes Medical Institute, University of Washington, Seattle, WA 98195 USA

Correspondence e-mail: merritt@u.washington.edu

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# Structure of Lmaj006129AAA, a hypothetical protein from *Leishmania major*

The gene product of structural genomics target Lmaj006129 from *Leishmania major* codes for a 164-residue protein of unknown function. When SeMet expression of the full-length gene product failed, several truncation variants were created with the aid of *Ginzu*, a domain-prediction method. 11 truncations were selected for expression, purification and crystallization based upon secondary-structure elements and disorder. The structure of one of these variants, Lmaj006129AAH, was solved by multiple-wavelength anomalous diffraction (MAD) using *ELVES*, an automatic protein crystal structure-determination system. This model was then successfully used as a molecular-replacement probe for the parent full-length target, Lmaj006129AAA. The final structure of Lmaj006129AAA was refined to an *R* value of 0.185 ( $R_{\rm free} = 0.229$ ) at 1.60 Å resolution. Structure and sequence comparisons based on Lmaj006129AAA suggest that proteins belonging to Pfam sequence families PF04543 and PF01878 may share a common ligand-binding motif.

# 1. Introduction

The aim of the Structural Genomics of Protozoan Parasite (SGPP) Consortium is to determine protein structures from major tropical eukaryotic pathogens, specifically the protozoa *Trypanosoma brucei* (African sleeping sickness), *T. cruzi* (Chagas disease), *Leishmania* spp. (leishmaniasis) and *Plasmodium* spp. (malaria). Targets are selected based on criteria of length, pI, predicted disorder and size of Pfam family. The Lmaj006129AAA gene product (Gene DB LMJF36.6870; Hertz-Fowler *et al.*, 2004) is a member of a large Pfam sequence family PF04543 (DUF589) present in eukaryotes and eubacteria (Bateman *et al.*, 2004). The family had no representative structure at the time of target selection. The target sequence was expressed as a protein of molecular weight 20.2 kDa (172 residues) including a hexa-His tag. Below, we describe the efforts that culminated in the determination of the three-dimensional structure of the Lmaj006129AAA at 1.6 Å, encompassing residues 7–163.

Although native crystals of Lmaj006129AAA diffracted to 1.6 Å resolution, SeMet expression failed and heavy-atom derivatives were unachievable despite considerable trials. We therefore searched for variants of the original target sequence that might yield a new crystal form or be more amenable to SeMet expression. Laboratory techniques such as limited proteolysis (Gao *et al.*, 2005) were considered, but we decided to use a computational approach suitable for structural genomics on a large scale. This approach is the generation of truncation variants through the use of *Ginzu*, a domain-prediction method (Kim *et al.*, 2005), and it proved essential for the structure determination of Lmaj006129AAH.

# 2. Materials and methods

# 2.1. Truncation variants

The sequence of Lmaj006129AAA was submitted to the *Robetta* server (Chivian *et al.*, 2003; Kim *et al.*, 2004) for analysis by *Ginzu*, a domain-prediction method (Kim *et al.*, 2005), based upon sequence similarity to known structures, regions of conserved sequence,

	10	155	164
AAA	MSRKRVRAED	LRMNE	EQQRK
AAB	MSRKRVRAED	LRMNE	EQ
AAC	MSRKRVRAED	LRMNE	
AAD	KRVRAED	LRMNE	EQQRK
AAE	KRVRAED	LRMNE	EQ
AAF	KRVRAED	LRMNE	
AAG	RAED	LRMNE	EQQRK
AAH	RAED	LRMNE	EQ
AAI	RAED	LRMNE	~
AAJ	D	LRMNE	EOORK
AAK	D	LRMNE	ΕÕ̃
AAL	D	LRMNE	~

#### Figure 1

Truncation variants of target ORF Lmaj006129. Structures were determined for the truncation variant Lmaj006129AAH and for the full-length target Lmaj006129AAA.

predicted secondary-structural elements and regions of predicted disorder. 11 truncations of the parent gene Lmaj006129AAA were selected, expressed and purified (Fig. 1). The primary guidance for truncations came from disorder prediction (Ward *et al.*, 2004) and from multiple sequence alignment of family members (Fig. 2). Of these truncated variants, identified by postfixes AAB through AAL, ten were expressed, nine were soluble and one variant (Lmaj006129AAH) yielded crystals during screening. Once crystals were obtained, work on the remaining variants was discontinued.

## 2.2. Expression and purification

The gene encoding Lmaj006129AAA was cloned into vector BG1861, a modified form of pET14b (Alexandrov *et al.*, 2004). BG1861 contains an N-terminal non-cleavable hexa-His tag. The protein was expressed in *Escherichia coli* and purified using an Ni–NTA column and by gel filtration on a HiLoad Superdex 200 26/60 column (Amersham Pharmacia Biotech).

The gene encoding Lmaj006129AAH variant was cloned into a vector AVA421, which is also based on pET14b. In contrast to

BG1861, the AVA421 vector contains a cleavable N-terminal His tag, although the crystallization success came from uncleaved protein. The protein was purified using an Ni–NTA column and the bound protein was cleaved by protease 3C overnight at 277 K. The released protein was further purified by gel filtration on a HiLoad Superdex 200 26/60 column (Amersham Pharmacia Biotech).

# 2.3. Crystallization

Lmai006129AAA was shipped on dry ice for high-throughput crystallization screening at the Hauptman-Woodward Medical Research Institute (Luft et al., 2003). The sample was thawed at 296 K. Crystallization experiments were set up using the microbatchunder-oil technique (Chayen et al., 1992) with a Robbins Scientific Tango liquid-handling system. Each of the 1536 experiments contained 200 nl crystallization cocktail solution combined with 200 nl protein solution under paraffin oil (Fluka catalog. No. 76235) in a 1536-well plate (Greiner BioOne catalog No. 79101). The plate was stored at 277 K for one week after setup and stored at 296 K for three additional weeks. The plate was imaged weekly at 296 K. Images were manually reviewed; eight of the 1536 crystallization conditions for Lmaj006129AAA produced outcomes considered marginally suitable for optimization trials. Following the same screening procedure for the Lmaj006129AAH truncation variant, 20 outcomes considered suitable for optimization trials were observed.

Initial hits were optimized and crystals grown by the sitting-drop method. For native Lmaj006129AAA crystals, the crystallization drop consisted of 1 µl protein solution (8.2 mg ml<sup>-1</sup>) mixed with 1 µl reservoir solution containing 2.8 *M* sodium malonate pH 6. For SeMet-incorporated Lmaj006129AAH crystals, the crystallization drop consisted of 0.4 µl protein solution (16.4 mg ml<sup>-1</sup>) mixed with 0.4 µl reservoir solution containing 35% PEG 8000, 0.1 *M* lithium bromide, 0.1 *M* TAPS, 5 m*M* DTT pH 9.



#### Figure 2

Multiple sequence alignment of PF04543 family members. TrEMBL sequence identifiers and residues extents are shown on the right. Secondary-structure elements and residue numbering are from the present *L. major* structure (PDB code 2ar1). Sequence alignment is based on *CLUSTALW* analysis of 106 sequence family members. Residue positions are colored if the hydropathy class is conserved in >50% of the family members. Below this is shown the sequence fingerprint of *P. horikoshii* protein PH1033 (PDB code 1wmm) from Pfam sequence family PF01878, aligned based on structural similarity to the *L. major* structure. Residues lining a binding pocket observed in the four sequences with known structure (PDB codes at far right) are indicated by brown dots. This figure was generated using  $T_EXshade$  (Beitz, 2000).

#### Table 1

Data-collection and phasing statistics.

Values in parentheses are for the last resolution shell.

	Lmaj006129AAH (truncation)			Lmaj006129AAA (full length)	
	Peak	Inflection	Remote	Native	
Wavelength	0.9795	0.9797	0.9537	0.9746	
Resolution (Å)	2.27	2.27	2.27	1.6	
Space group	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	P21212	
Unit-cell parameters (Å)	<i>a</i> = 78.4, <i>b</i> =	122.9, <i>c</i> = 77.	5	a = 31.2, b = 64.6, c = 72.6	
Total reflections	103179	103731	108733	55401	
Total unique reflections	16790	16792	17115	19933	
R <sub>svm</sub>	0.084 (0.64)	0.087 (0.73)	0.095 (0.90)	0.041 (0.69)	
Completeness (%)	100.0 (77.1)	100 (77.3)	100 (83.8)	95.3 (94.6)	
$I/\sigma(I)$	10.0 (2.0)	9.8 (1.9)	9.1 (1.7)	26.7 (1.8)	
Redundancy Phasing power† (dis; ano) Figure of merit	6.2 (3.4) 0.57; 0.66 0.496	6.4 (3.4) 0.87; 0.46	6.4 (3.9) 0; 0.42	2.9 (2.8)	

† Phasing power =  $(\sum |\Delta F_H|^2 / \sum \varepsilon^2)^{1/2}$ , where  $\Delta F_H$  is either the dispersive or anomalous difference in scattering factor attributed to Se.

# 2.4. X-ray diffraction and structure determination

Native crystals of Lmaj006129AAA were cryoprotected with malonate and glycerol before flash-freezing directly in liquid nitrogen. Data were collected to 1.6 Å resolution from a single crystal maintained at 100 K at a wavelength of 0.974609 Å on beamline 11-3 at SSRL (Stanford Synchrotron Radiation Laboratory). Data were processed and scaled using HKL2000 (Otwinowski & Minor, 1997). Crystallographic data statistics are given in Table 1.

SeMet-derivatized crystals of Lmaj006129AAH, the truncation variant of Lmaj006129AAA, were flash-frozen directly in liquid nitrogen in a cold room at 277 K. Data were collected to 2.3 Å from a single crystal maintained at 100 K at a wavelength of 0.9797 Å on beamline 8.2.2 at the Advanced Light Source, Berkeley, CA, USA. Data were processed and scaled using *ELVES*, an automatic protein crystal structure-determination system (Holton & Alber, 2004). Crystallographic data statistics are given in Table 1. All ten selenium sites belonging to two monomers in the asymmetric unit were found by *ELVES* using *SOLVE* (Terwilliger & Berendzen, 1999). *RESOLVE* (Terwilliger, 2000, 2003) built 236 of 318 residues (74%,

Tab	le	2	

Refinement	statistics.
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Resolution (Å)	1.6
R	0.185
R <sub>free</sub>	0.229
R.m.s.d. bonds (Å)	0.021
R.m.s.d. angles (°)	1.93
Residues in most favored region of $\varphi/\psi$ (%)	91
Residues in allowed region of $\varphi/\psi$ (%)	8.3
No. of protein atoms	1303
No. of non-protein atoms	104
Wilson <i>B</i> factor $(Å^2)$	33.5
TLS groups	7-37, 38-127, 128-163
Mean (e.s.d.) of $(B_{iso} + B_{TLS})$ for protein atoms $(Å^2)$	28 (11)
Mean (e.s.d.) $B_{iso}$ for non-protein atoms (Å <sup>2</sup> )	33 (7)

159 residues per monomer) and added side chains to 135 residues (42%). Additional manual building using XFIT (McRee, 1999) yielded a single polypeptide that was as complete as possible before submission to REFMAC for refinement (Murshudov et al., 1997). Model building and refinement in REFMAC continued until R and  $R_{\rm free}$  were 0.33 and 0.40, respectively, at which point the model was used for molecular replacement using MOLREP (Vagin & Teplyakov, 1997) against the full-length native 1.6 Å data. The structure of Lmaj006129AAA contains one molecule in the asymmetric unit and was refined using REFMAC (Murshudov et al., 1997), keeping 5% reflections for  $R_{\text{free}}$ . In the final cycles of refinement, the protein chain was described by three TLS groups identified by the TLSMD server (Painter & Merritt, 2006) and TLS parameters were refined for each group (Table 2). The final model consists of residues 7-163 (Table 2), one glycerol molecule and 99 waters. Waters were added using Coot. Model quality was validated using Coot and MolProbity (Emsley & Cowtan, 2004; Lovell et al., 2003).

#### 3. Results and discussion

The structure of the Lmaj006129AAA monomer is shown in Fig. 3(*a*) and exhibits an  $\alpha\beta$ -fold containing six strands and six helices. Datacollection, refinement and model statistics are summarized in Table 1. The final model contains one monomer in the asymmetric unit, with all residues and side chains built except for the hexa-His tag, the first six N-terminal residues and the C-terminal lysine residue. No electron density was observed for these residues owing to disorder. The



#### Figure 3

(a) A ribbon diagram showing the tertiary structure of Lmaj006129AAA color coded from the amino-terminus (blue) to the carboxy-terminus (red) with secondary-structure elements labeled as  $\beta 1$  (13–18),  $\alpha 1$  (24–32),  $\beta 2$  (33–35),  $\alpha 2$  (41–51),  $\beta 3$  (56–61),  $\beta 4$  (68–81),  $\alpha 3$  (83–87),  $\beta 5$  (105–120),  $\alpha 4$  (121–125),  $\alpha 5$  (126–131),  $\beta 6$  (143–146),  $\alpha 6$  (147–162). (b) Stereo figure depicted as a coil with every tenth residue numbered. It is in the same orientation and color scheme as (a).

model also contains a glycerol molecule that was likely to be incorporated from the cryoprotectant.

In an endeavor to identify and classify the fold of Lmaj006129AAA, a structural similarity search was completed using the *DALI* server (Holm & Sander, 1993). The *DALI* server found eight instances of significant structural similarity (Z > 3). Four of these are hypothetical proteins with unknown function, including two with Z scores of greater than 10. Three *DALI* hits have functional annotations, of which the highest Z score (4.7) is associated with the N-terminal domain of PDB code 2ane, Pfam sequence family PF02190. The remaining *DALI* hit, PDB code 1zbo, has no previous functional annotation but is homologous in both structure and sequence to 2ane.

## 3.1. Structural similarity within the same sequence family

The homolog with the highest DALIZ score, 19.1, is a hypothetical protein from the Gram-negative bacterial plant pathogen Agrobacterium tumefaciens (PDB code 1zce) with sequence identity of 44% over 139 residues as reported by the DALI server. This structure was deposited in the interim between data collection of SeMet crystals of Lmaj006129AAH variant and the structure refinement. Secondary-structure alignment of 1zce and Lmaj006129AAA within *Coot* (Krissinel & Henrick, 2004; Emsley & Cowtan, 2004) yields an r.m.s.d. of 1.1 Å for 134 C<sup> $\alpha$ </sup> atoms in three chain fragments.

An additional close structural homolog, a hypothetical protein from *Pseudomonas syringae* (PDB code 2eve), was found subsequent to the initial submission of this paper by a *BLAST* search against the PDB. 2eve has sequence identity of 51% and an r.m.s.d. of 0.80 Å aligned over 145 residues within *Coot*.

# 3.2. Structural similarity to members of other sequence families

Three weak structural neighbors identified by *DALI* have annotated functions. The tRNA pseudouridine synthase b fragment (PDB code 1k8w; *DALI* Z score 3.3) is a fragment of a two-domain protein that catalyzes the isomerization of uridines to pseudouridines. The N-terminal domain contains the active site; the C-terminal domain binds to the adaptor stem of the RNA (Hoang & Ferré-D'Amaré, 2001). The C-terminal domain shows weak structural similarity to the present protein, but has not been assigned to a Pfam sequence family. The residues of the C-terminal domain interacting with RNA are Arg307, Lys308 and Asn266. The equivalent Lmaj006129AAA residues are Val112, Ala113 and Lys30, respectively, as identified by a structural alignment within *Coot*. The secondary-structure alignment yields an r.m.s.d. of 1.9 Å aligned over 53 residues and a low sequence identity of 9%. Common secondary-structure elements which overlay well include  $\beta$ 1,  $\alpha$ 1 and  $\beta$ 3– $\beta$ 5.

ATP sulfate adenylyltransferase (ATPS; PDB code 1g8f; *DALI Z* score 3.2) catalyzes the formation of adenosine-5-phosphosulfate from ATP and activated inorganic sulfate (Ullrich *et al.*, 2001). ATPS has four domains, with domain II containing the active site and substrate-binding pocket. Lmaj006129AAA is structurally similar to the N-terminal domain I, which belongs to the ATP-sulfurylase Pfam family PF01747. Secondary-structure matching in *Coot* yields an r.m.s.d. of 2.7 Å aligned over 76 residues and a sequence identity of 12%. Although the structures of Lmaj006129AAA and ATPS are distinctly different, secondary-structure elements  $\beta$ 1,  $\alpha$ 1 and  $\beta$ 3– $\beta$ 5 overlay well.

There is also weak structural similarity to residues 8–117 of the N-terminal domain of *E. coli* Lon protease (PDB code 2ane; *DALI* Z score 4.7). *DALI* aligned 84 residues with an r.m.s.d. of 3.0 Å. This has been hypothesized to be a peptide-binding domain (Li *et al.*,

2005). However, this domain belongs to an unrelated sequence family, PF02190, as does a second weak structural homolog with no annotated function, PDB entry 1zbo.

Remarkably, the second highest *DALI Z* score of 10.5 corresponds to a protein from a different Pfam sequence family, PF01878 (DUF55). This is a hypothetical protein PH1033 from the hyperthermophilic archeon *Pyrococcus horikoshii* OT3 (PDB code 1wmm), which has a sequence identity of 17% and an r.m.s.d. of 2.1 Å over 143 residues. In comparison to 1wmm, the present Lmaj006129AAA structure has a shorter  $\beta 3-\beta 4$  loop and an additional insertion which includes  $\alpha 3$  and the  $\alpha 3-\beta 5$  loop consisting of 21 residues (residues 87–107).

## 3.3. Functional implications

The present structure contains one significant surface cleft, which is observed to contain a well ordered glycerol molecule. The CASTp server (Liang et al., 1998) finds a molecular surface area of 240 Å<sup>2</sup> for this cavity and a corresponding volume of 390  $Å^3$ . The structurally homologous 2eve structure contains a 3-(N-morpholino)propane sulfonic acid (MOPS) molecule in the same location. The ring system of MOPS overlays well with glycerol and both molecules form hydrophobic interactions with phenylalanine Phe23 (Phe13 in 2eve) and tryptophan residue Trp36 (Trp26 in 2eve). The hydrophobic nature of these residue positions is conserved across the PF04543 sequence family (Fig. 2). The site defined by association of the larger MOPS molecule with 2eve also includes residues (Val29, Ser133, Tyr82), which are structurally equivalent to residues (Val39, Ser142, Tyr92) in the present structure. The valine is conserved across the sequence family, but the latter two residues are not. These residues line a hydrophobic groove near the junction of strand  $\beta 2$  and helix  $\alpha 2$ at the lower right in Fig. 3. It is notable that the same structural motif and equivalent pocket-lining residues are also observed in the structure of the P. horikoshii protein PH1033 (PDB code 1wmm), although this sequence is assigned by Pfam to a different sequence family (Fig. 2). The phenyl ring of residue Tyr88 in the P. horikoshii structure is positioned equivalently to the phenyl ring of Tyr92 in the present L. major protein, although this is not a homologous pair of residues by sequence or backbone conformational alignment. For this equivalent cavity in 1wmm, the CASTp server finds a molecular surface area of 270  $\text{\AA}^2$  and a corresponding volume of 380  $\text{\AA}^3$ .

The identification of a ligand-binding site shared by these proteins may facilitate future interpretation of functional assignment made for any member of either sequence family. Furthermore, the conserved structure/sequence motif forming the binding site may also aid in binding-site recognition in more disparate sequence families.

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