Three-dimensional structure and ligand binding properties of trichosurin, a metatherian lipocalin from the milk whey of the common brushtail possum

Trichosurus vulpecula

Randall P. WATSON*,†, Jerome DEMMER‡, Edward N. BAKER* and Vickery L. ARCUS†,1,2

*Laboratory of Structural Biology, School of Biological Sciences, University of Auckland, Private Bag 92-019, Auckland, New Zealand, †AgResearch Structural Biology Laboratory, School of Biological Sciences, University of Auckland, Private Bag 92-019, Auckland, New Zealand, and ‡Halcyon Bioconsulting Limited, P.O. Box 89-106, Torbay, Auckland, New Zealand

INTRODUCTION

The metatherians have historically been regarded by science as curiosities, the name quite literally means ‘other beasts’. Among the metatherians, the biology of the opossum has recently come to the fore as it occupies a unique position on the evolutionary tree, sharing a common ancestor with humans approx. 130 million years ago. This is the midpoint between the divergence of placental mammals such as humans and mice which have a common ancestor approx. 80–90 million years ago, and non-mammalian animals which diverged from humans 300–350 million years ago. The unique biology of the early development of opossum young (and other metatherians) also provides impetus for research. Research into the milk proteins of Trichosurus vulpecula, the common brushtail possum, stems from a concerted effort to study the physiological differences between metatherians and other mammals. T. vulpecula, the common brushtail possum, stems from an academic evolutionary point of view as well as an applied environmental perspective. It was introduced to New Zealand from its native Australia in the middle of the 19th century to establish a fur trade [1]. It has since become widely established, damaging large areas of forest every year and exerting considerable resource pressure upon New Zealand’s endemic fauna [2–4]. Analogous to the badger in the U.K., the possum in New Zealand also acts as an efficient vector for bovine tuberculosis. These factors make the brushtail possum one of the most important vertebrate pests in New Zealand.

Marsupials have invested in lactation, as opposed to placenta tion, as a way of rearing their young. Reproduction in the Australian brushtail possum (T. vulpecula) is typified by a short gestation (17 days) followed by a prolonged period of lactation (> 200 days) [5]. Lactation is divided into at least three distinct phases between which milk composition changes dramatically [1,6,7]. Possum milk contains a large number of whey proteins, most of which are homologues of eutherian milk proteins [8–10]. Trichosurin is one of three predominant lipocalins found in the milk of T. vulpecula [10] (Figure 1). It bears significant similarity (34% amino acid identity) to the rodent MUPs (major urinary proteins) and to the major horse dander allergen esu_c1 (28% amino acid identity). More recently, trichosurin homologues have been identified in both the tammar wallaby (87% amino acid identity) and the South American opossum (77% amino acid identity). This high level of conservation in species that diverged approx. 80 million years ago due to continental separation suggests that the function of trichosurin is critical in metatherian lactation. Currently, its biochemical and biological functions are not understood.

The lipocalins are a large family of proteins (diverse in sequence but structurally homologous) that bind and transport small hydrophobic molecules in a central hydrophobic pocket [11,12]. The binding specificity exhibited by lipocalins varies widely across the family. For example, BLG (β-lactoglobulin) is known to bind a wide array of molecules from retinal to fatty acids [13], whereas other lipocalins such as the MUPs from mice and rats bind a smaller range of compounds in a manner more in keeping with their role in scent-based territory marking [14]. The internal topology of the hydrophobic pocket is a key factor in the type and selectivity of binding, as is access to the binding pocket which is usually controlled by residues forming a loop between β-strands at one end of the lipocalin β-barrel.

Abbreviations used: BLG, β-lactoglobulin; MUP, major urinary protein; NGAL, neutrophil gelatinase-associated lipocalin; rmsd, root mean square deviation.

1 Present address: Department of Biological Sciences, University of Waikato, Private Bag 3105, Hamilton, New Zealand.

2 To whom correspondence should be addressed (email varcus@waikato.ac.nz).
The lipocalin superfamily is defined by a common fold, an eight-stranded anti-parallel β-barrel, with variable loop regions flanking and enclosing the top and bottom of the barrel to form a hydrophobic pocket in the interior [15]. A conserved disulfide bond links one of the C-terminal β-strands with the C-terminus. Identification of lipocalins is generally made via the presence of three small conserved sequence motifs which are also used to demarcate family membership between ‘core’ lipocalins and ‘outliers’, based on whether all three, or just the first two, sequence motifs are present.

In the present study, we provide the first three-dimensional structure of a metatherian lipocalin, trichosurin. This structure was solved by X-ray crystallography in two different crystal forms at two disparate values of pH (pH = 4.6 and pH = 8.2). To investigate the binding of ligands by trichosurin, six small molecules known to bind within the barrel of the closely related rodent MUPs have been used as molecular probes. Two of these small molecules were shown by fluorescence to bind to trichosurin, and we present the three-dimensional structures of trichosurin with 2-naphthol and 4-ethylphenol bound in the central pocket of the β-barrel.

2-Naphthol binds in the cavity of trichosurin with micromolar affinity and a preference for small phenolic compounds suggests a possible biological role for trichosurin.

**EXPERIMENTAL**

**Protein expression and refolding**

DNA encoding the mature trichosurin protein was amplified by PCR from a possum cDNA library and cloned into the vector pGEMTeasy. The trichosurin gene was then subcloned into the expression vector pET23a which was transformed into *E. coli* cells [BL21 (DE3)] carrying the pRI rare codon plasmid. Small overnight cultures (5 ml) were used to inoculate 1 litre cultures and these were grown at 37°C and protein expression was subsequently induced at an *D*₅₉₀ of 0.5–0.7 using 1 mM IPTG (isopropyl β-D-thiogalactoside). At 3 h post-induction, cells were harvested by centrifugation (6000 g at 4°C for 10 min) and cell pellets were washed with 50 mM Tris/HCl (pH 8.2) and 50 mM NaCl, then repelleted and frozen at −20°C. Frozen cell pellets were thawed and re-suspended in buffer (50 mM Tris/HCl, pH 8.2, and 50 mM NaCl), then lysed by passing once through a Constant Systems cell disruptor at 25 kPa. The cell lysate was then centrifuged at 10000 g for 10 min to pellet the insoluble inclusion bodies. The inclusion bodies were washed by resuspending them in buffered 0.1% (v/v) Triton X-100, incubated on ice for 10 min, centrifuged (10000 g at 4°C for 10 min) and then suspended again in buffer. This was then treated with 50 ml of 8 M urea and 50 mM Tris/HCl (pH 8.2), and 50 mM NaCl, then lyzed by passing once through a Constant Systems cell disruptor at 25 kPa. The cell lysate was then centrifuged at 10000 g for 10 min to pellet the insoluble inclusion bodies. The inclusion bodies were washed by resuspending them in buffered 0.1% (v/v) Triton X-100, incubated on ice for 10 min, centrifuged (10000 g at 4°C for 10 min) and then suspended again in buffer. This was then treated with 50 ml of 8 M urea and 50 mM Tris/HCl (pH 8.2), and incubated for 1 h on ice to denature and solubilize the inclusion bodies. The denatured protein was refolded by dialysis at 18°C against 1 litre of 50 mM Tris/HCl (pH 8.2), 50 mM NaCl, 100 mM sucrose, 1 mM EDTA, 1 mM GSH and 0.1 mM GSSG. After 8 h the buffer was changed to 1 litre of 50 mM Tris/HCl (pH 8.2) and 50 mM NaCl.

**Protein purification**

The refolded trichosurin was bound to a HiTrap Q column (Pharmacia Biotech) and then eluted using a gradient of 50–500 mM NaCl. Fractions containing trichosurin were pooled and
Crystallization

Initial crystallization screening was carried out by the hanging drop vapour diffusion method in 24-well format and using the Hampton I crystallization screen [16]. Drops of 2 µl were set up by mixing 1 µl of trichosurin (10 mg/ml) with 1 µl of reservoir solution. Successful crystallization conditions from the initial screen were refined and diffraction quality crystals were grown from drops consisting of 3 µl of protein/3 µl of reservoir solution [100 mM Tris/HCl (pH 8.2), 200 mM Li₂SO₄ and 10–14 % poly(ethylene glycol) 4000]. Crystallization was also refined at low pH values, with diffraction-quality crystals being obtained using a precipitant solution of 100 mM sodium acetate (pH 4.6), 200 mM Li₂SO₄ and 10–14 % (v/v) 2-propanol.

Ligand complexes of trichosurin were obtained by co-crystallization. Each ligand was dissolved in 2-propanol at a concentration of 1 M, and this solution was added to the reservoir solution immediately prior to crystallization in sufficient quantity to give a final ligand concentration in each case of 3–

Table 1. Data collection and refinement statistics

<table>
<thead>
<tr>
<th></th>
<th>Apo-trich8,6</th>
<th>Apo-trich8,2</th>
<th>2Nap-trich8,6</th>
<th>4Eth-trich8,6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data collection</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Space group</td>
<td>P2₁</td>
<td>P4₁2,2</td>
<td>P2₁</td>
<td>P2₁</td>
</tr>
<tr>
<td>Unit cell (Å)</td>
<td>57.85, 44.58, 70.41</td>
<td>100.16, 100.16, 140.55</td>
<td>44.3, 100.4, 83.69</td>
<td>44.3, 100.4, 84.9</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>1.54</td>
<td>1.54</td>
<td>1.54</td>
<td>1.54</td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>26–1.9 (1.97–1.90)</td>
<td>20–2.5 (2.29–2.50)</td>
<td>15–1.5 (1.6–1.5)</td>
<td>30–1.5 (1.6–1.5)</td>
</tr>
<tr>
<td>Rmerge (%)</td>
<td>0.043 (0.192)</td>
<td>0.085 (0.516)</td>
<td>0.086 (0.231)</td>
<td>0.072 (0.289)</td>
</tr>
<tr>
<td>completeness (%)</td>
<td>95.1 (88.9)</td>
<td>100 (100)</td>
<td>96.9 (88)</td>
<td>98.6 (96)</td>
</tr>
<tr>
<td>Nref</td>
<td>108260</td>
<td>534113</td>
<td>284253</td>
<td>520556</td>
</tr>
<tr>
<td>Nrefine</td>
<td>26646</td>
<td>25922</td>
<td>92603</td>
<td>116743</td>
</tr>
<tr>
<td>Refinement</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rwork (%)</td>
<td>21.6 (31.1)</td>
<td>22.0 (27.3)</td>
<td>22.6 (30.9)</td>
<td>22.7 (33.3)</td>
</tr>
<tr>
<td>Rfree (%)</td>
<td>24.9 (35.3)</td>
<td>28.6 (36.9)</td>
<td>24.0 (33.9)</td>
<td>24.5 (33.9)</td>
</tr>
<tr>
<td>B (Å²) protein</td>
<td>28.4</td>
<td>33.7</td>
<td>24.2</td>
<td>32.7</td>
</tr>
<tr>
<td>B (Å²) solvent</td>
<td>33.7</td>
<td>32.0</td>
<td>33.7</td>
<td>34.0</td>
</tr>
<tr>
<td>B (Å²) ligand</td>
<td>23.8</td>
<td></td>
<td>23.8</td>
<td></td>
</tr>
<tr>
<td>Nrefine</td>
<td>2373</td>
<td>4883</td>
<td>5124</td>
<td>5248</td>
</tr>
<tr>
<td>Bond length (Å)</td>
<td>0.0052</td>
<td>0.0065</td>
<td>0.006</td>
<td>0.005</td>
</tr>
<tr>
<td>Bond angle (°)</td>
<td>1.26</td>
<td>1.2</td>
<td>1.24</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Structure solution, model building and model refinement

Diffraction images were collected using a MAR345 detector with Cu-Kα X-rays from a Rigaku H3R rotating anode generator. Data for the trichosurin–ligand complex crystals were collected at the ESRF (European Synchrotron Radiation Facility, Grenoble, France). These synchrotron data were collected at 113 K on beamline ID 14-4 using an ADSC Quantum4 Q4R CCD (charge-coupled-device) camera. For data collection statistics see Table 1.

The structure of apo-trichosurin (crystals grown at pH 4.6) was solved by molecular replacement with AMoRe [18], using the rat α2C-globulin structure (PDB entry: 2A2U) as a search model. Two solutions were found, corresponding to two molecules in the asymmetric unit and a Matthews coefficient (V_M) of 2.3 Å³/Da. The two molecules were related by a local 2-fold rotation axis and positioned such that a dimer interface is clearly defined. Initial automated model building was with wARP [19] which built a model comprising 214 of the expected 332 residues (side chains were automatically added by wARP for 210 residues). A further 72 residues were added during iterations of building and refinement using O [20] and CNS [21] respectively. The structure of high-pH apo-trichosurin was solved using the low-pH apo-trichosurin coordinates and molecular replacement. Similarly, the structures of trichosurin with bound substrate were solved by molecular replacement (AMoRe) using the low-pH trichosurin co-ordinates as a search model. In all cases refinement and model-building were carried out in CNS and O [20,21].

Concentrated. The protein was then purified in a second step by gel-filtration chromatography (Superdex75 16/60; Pharmacia Biotech) in 50 mM Tris/HCl (pH 8.2) and 150 mM NaCl. The protein quality in each fraction was assessed by dynamic light scattering, and those fractions with the highest degree of monodispersity were pooled and concentrated to ~10 mg/ml for use in crystallization screening. N-terminal sequencing confirmed that the N-terminus of the protein was intact.
The small molecules for ligand-binding experiments were thymol, trans-2-nonenal, 2-naphthol, 4-ethylphenol, 2-isobutyl-3-methoxy pyrazine and 2-sec-butyl thiazole and were purchased from Sigma–Aldrich.

Ligand binding using fluorescence

Fluorescence binding experiments were carried out using a Hitachi F-4500 fluorescence spectrophotometer and a square quartz cuvette with a 5 mm path length. Trichosurin was used at a concentration of 20 μM in 10 mM sodium phosphate buffer (pH 7.96). Stock solutions of substrates were made up to 1 mM in ethanol and added to 400 μl aliquots of protein to a final concentration of 20 μM. Changes in fluorescence maxima were observed using an excitation wavelength of 282 nm by comparison against appropriate reference spectra of trichosurin and trichosurin with 2% ethanol.

RESULTS AND DISCUSSION

Structure determination

Trichosurin was expressed in E. coli as insoluble inclusion bodies, denatured in 8 M urea and refolded by dialysis against a suitable refolding buffer. Washing the inclusion bodies and then refolding had the advantage of producing very pure protein in the absence of an affinity purification tag. Following the final size-exclusion purification step, the protein showed excellent monodispersity as determined by dynamic light scattering.

Molecular replacement using the co-ordinates for α2v-globulin (PDB code: 2A2U) as a search model allowed the structure of trichosurin to be solved using the data collected from a single crystal grown at pH 4.6. This model, termed apo-trich8, consisted of residues 24–166 of trichosurin, the first 23 residues were determined by dynamic light scattering. For apo-trich8 the first 20 amino acids of the peptide chain are absent. The two structures are very similar, with an rmsd (root mean square difference) in Ca atom positions of 0.54 Å over residues Leu34 to Cys186.

As expected, the overall structural fold of trichosurin is an eight-stranded, anti-parallel β-barrel with a helix near the C-terminus which is common to all lipocalins (see Figure 2A). An extended loop region between β-straands A and B forms a short helix and an open turn which, together with residues on the loop between β-strands E and F, close one end of the barrel. Residues which are N-terminal to β-strand A pack against residues in the bottom of the barrel, effectively sealing the other end of the barrel. A disulfide bond, the last major conserved feature of the lipocalin fold, is formed between Cys73 and Cys186 and ties the C-terminus to β-strand D.

Trichosurin forms an unusual dimer which has not previously been seen among structures from the lipocalin superfamily. The mode of dimerization is conserved between structures determined at pH 4.6 and pH 8.2 and additional biophysical evidence from size-exclusion chromatography and DLS (dynamic light scattering) indicate that trichosurin is a dimer in solution (protein concentration = 0.5 mg/ml; molecular mass from DLS ~38.6 kDa; molecular mass from size-exclusion chromatography ~38 kDa). The surface area buried per monomer, by dimerization, is 905 Å² (11.8% of the monomer surface, calculated from the pH 4.6 structure). At the core of the dimerization interface is the ring-to-ring packing of Tyr100 with Tyr115. These aromatic interactions are complemented further by hydrophobic burial of the side-chains of Val212 and Val213. Two salt bridge interactions between Arg30 and Asp98 also contribute to the stability of the dimer along with numerous inter-molecular hydrogen bonds. At least fifteen water molecules are present within the dimer interface in the apo-trich8 structure. The lower resolution of the apo-trich8 structure precluded the modelling of as many water molecules and, as a consequence, only four bridging water molecules are seen in the dimer interface for the higher pH structure.

A review of lipocalin dimerization modes reveals six variations in orientation. The most closely related dimer is that of porcine BLG [22], which uses part of the same face to dimerize as trichosurin (Figure 3). However, the porcine BLG dimer also has an N-terminal crossover between the monomers in which hydrogen bonding and other interactions occur, taking the effective buried surface area to a reported 1757.7 Å² per monomer [22]. The dimerization of trichosurin is sufficiently different to that of porcine BLG that it represents a seventh variation on the theme (Figure 3).

Structural comparison with lipocalins

Bovine BLG is regarded as the prototypic milk lipocalin. A structural alignment of apo-trich8 with the BLG structure 1BEF, solved at pH 6.5, gives an rmsd of 2.76 Å over 139 equivalent residues (Figure 4, and Supplementary Table S1 and Figure S1 at http://www.BiochemJ.org/bj/408/bj4080029add.htm) and a sequence identity of 21% after structure-based sequence
Figure 2  Trichosurin structure, topology and ligand binding

(A) Stereo diagram showing the trichosurin monomer, with \( \beta \)-strands in blue (labelled \( \beta \)A–H) and \( \alpha \)-helices yellow. The residues whose side chains define the binding pocket are rendered as grey sticks with oxygen atoms in red and nitrogen in blue. The transparent orange surface represents the approximate size of the internal cavity as a molecular contact surface described by a sphere of 1.4 Å radius.  

(B) A 2(\( F_o \)) – (\( F_c \)) electron density map contoured at 1\( \sigma \) depicted in blue mesh showing the 2-naphthol ligand bound within the trichosurin barrel and the hydrogen bonds made to water W3 and Asn\(^{49} \) ND1.  

(C) A 2(\( F_o \)) – (\( F_c \)) electron density map contoured at 1\( \sigma \) showing 4-ethylphenol (surrounded by electron density mesh) donating a hydrogen bond to Leu\(^{78} \) O and occupying the lower part of the trichosurin-binding pocket. A 2-propanol molecule (orange and red) is still bound in the upper part of the binding pocket along with the water molecules W2 and W3 (red spheres).  

(D) The second binding site for 2-naphthol between the helix, the outside of the barrel and the dimer interface. The blue mesh is from a 2(\( F_o \)) – (\( F_c \)) electron density map contoured at 1\( \sigma \). The thin blue lines and the purple residues are from the second molecule of the dimer.
alignment. In general the two structures align very well, although
differences are seen in the N-terminus (due to this region being
undefined in the trichosurin structure) and for the GH loop (in
BLG a short kinked helical structure is formed, whereas in tri-
chosurin the same loop forms a textbook $\beta$-turn). Interestingly
the EF loop is also similar between the two proteins. This loop
is responsible for the pH-dependent opening and closing of the
barrel in BLG; a conformational change known as the Tanford transition [23]. The determination of the structure of trichosurin
at two different pHs affords an opportunity to assess the presence
of an equivalent conformational change over the pH range studied.
Trichosurin lacks a similar ionizable group when compared with
BLG (Glu$^{89}$) and does not appear to exhibit a similar opening
and closing action. The orientation of the Tyr$^{94}$ side chain does
however change position in the higher pH structure, leading
to a tighter closing of the barrel indicated by generally lower
B-factors in this area of the pH 8.2 structure when compared
with the main barrel. A mechanism responsible for access to
the barrel interior is not immediately apparent from the current
structures.

A structure–comparison survey of the SCOP [24] database
places trichosurin in the ‘retinol-binding-like’ protein family.
This family encompasses a total of 141 structures which include
retinol-binding proteins, odourant-binding proteins, BLGs, MUPs
from mice and rats, human NGAL (neutrophil gelatinase-
associated lipocalin) and C8$\gamma$ proteins, nitrophorins and $\alpha$-crust-
acyanin. A multiple structure alignment carried out using
MAMMOTH [25] showed that trichosurin clusters with a subset of

---

Figure 3  The trichosurin dimer and its relationship with other lipocalin dimers

(A) Trichosurin dimer (blue and yellow) looking down the 2-fold dimerization axis. Buried surface area upon dimerization is 905 $\AA^2$. For reference, in all cases a monomer of trichosurin is depicted in blue and yellow in the same orientation to that in (A). (B) Major horse allergen dimer (red with trichosurin monomer for reference) (PDB code 1EW3); buried surface area is 1023.3 $\AA^2$. (C) Porcine BLG dimer (yellow) (PDB code 1EXS); buried surface area is 1757.7 $\AA^2$. (D) Bovine odourant-binding protein dimer (magenta) (PDB code 1OBP); buried surface area is 2399.7 $\AA^2$. (E) Bovine BLG dimer (green) (PDB code 1B0O); buried surface area is 484.1 $\AA^2$. (F) Porcine OBP dimer (slate) (PDB code 1E06); buried surface area is 848.2 $\AA^2$; and (G) Nitrophorin 4 dimer (gold) (PDB code 1EQD); buried surface area is 789.1 $\AA^2$. 

© The Authors Journal compilation © 2007 Biochemical Society
the retinol-binding-like proteins which include porcine odourant-binding protein, the bovine d2 and equine c2 allergens, BLGs and the MUPs from mice and rats (Figure 4, and Supplementary Table S1 and Figure S1 at http://www.BiochemJ.org/bj/408/bj4080029add.htm). The majority of the structural features are well conserved among this subset with rmsd values of 1.3–3.4 Å. In contrast, the sequence identity is relatively low (16%–34%); see Supplementary Figure S2 at http://www.BiochemJ.org/bj/408/bj4080029add.htm). This points towards a general function for trichosurin as a protein which binds small ligands in the central cavity but, in the absence of tight sequence conservation, there are few clues about the precise nature of these ligands.

Ligand-binding cavity

From examination of the trichosurin structure, and from knowledge of the binding properties of other lipocalins, the central pocket inside the trichosurin barrel is expected to be the major ligand-binding site. The trichosurin-binding cavity is a hydrophobic pocket with approximate dimensions of 11 × 6 × 6 Å. The internal molecular surface, described by a probe sphere of radius 1.4 Å and using a triangulation algorithm that places six grid points every 1.4 Å [26], is 425 Å² and encloses a total volume of ~464 Å³. It has a bulbous top and bottom but is constricted in the middle by the side chains of Phe65 and Phe114 which protrude into the pocket (Figure 2A).

There are three asparagine residues, Asn49, Asn97 and Asn127, which lie in the cavity, and offer hydrogen-bonding sites for potential ligands. Two water molecules are bound inside the apo-trich8 barrel. Both are near the top of the hydrophobic pocket, with one (labelled W3 in Figure 2B) hydrogen-bonded to the carbonyl oxygen of Phe47 and the Asn127 OD1 oxygen, and the other (labelled W2 in Figure 2B) held by hydrogen bonds to Asn49 O, Asn127 OD1 and Thr10 OG1. Additionally, two regions of electron density in the cavity of apo-trich8 have been modelled as 2-propanol which is a constituent of the crystallization conditions. One is held by two hydrogen bonds to Asn49 OD1 and the water molecule W3 (see Figure 2C), whereas the other is buried further down in the cavity, hydrogen-bonded with Leu59 O. These 2-propanol molecules are not present in the apo-trichh3 structure as they originate from the crystallization mother liquor.

In the apo-trichh3 structure three water molecules are bound near the top of the pocket. Two of these water molecules occupy equivalent positions to W2 and W3 in the apo-trich8 structure (Figure 2B), whereas the third occupies a position in place of one of the 2-propanol molecules, bonded to Asn49 OD1 and W3. The position occupied by the 2-propanol molecule hydrogen-bonded to Leu59 O in the apo-trichh3 structure is left unoccupied in the apo-trichh2 structure. Water molecules found within the predominantly non-polar pocket are not unique to trichosurin. MUP has been shown to utilize internalized water molecules in the formation of water mediated ligand-to-protein hydrogen bonds [14].

There are subtle differences between the binding pockets in the apo-trichh6 and apo-trichh2 structures. The side chains of the 22 residues that line the binding pocket align closely. The volumes of the binding pocket at pH 4.6 and at pH 8.2 vary only slightly. This is in direct contrast with bovine BLG where significant conformational changes accompany pH changes across quite a narrow range (pH 6.2–8.2) [27].

The N-terminus

The N-terminal region, residues 1–23, is not observed in the structure determined at pH 4.6, whereas residues 1–20 are absent in the pH 8.6 structure. To ensure that the N-terminus was intact, N-terminal sequencing was performed prior to crystallization. In other lipocalins this region of the amino acid chain helps to close the bottom end of the barrel. In trichosurin, the bottom of the barrel is effectively closed by Phe89, Leu24, Phe112 and Ile46. However, in the apo-trichh3 structure Trp21, Glu23, Gin23 and Leu24 form a classic lipocalin-like 3_10 helix which further protects the hydrophobic residues from the aqueous environment and no doubt plays a stabilizing role in this region of the structure.

In structures of MUP, part of the N-terminus is often disordered, but this only accounts for the first five residues, with the rest of the N-terminal region hydrogen bonding to the kinked end of β-strand F in pseudo β-sheet fashion before turning to pass under the bottom of the barrel and form the 3_10 helix. The trichosurin N-terminal residues do not follow the same path. At this stage it is not possible to assign structure or a role to these 20 residues; however, they may perform a specific function such as a receptor-binding interaction whereupon they might assume a definite structure upon complexation with a suitable receptor. Such an interaction is known for the C-terminus of plasma retinol-binding protein, a lipocalin, upon formation of its complex with transthyretin [28].

Bound ligands

There have been many studies on ligand binding by various lipocalin family members, such as BLG [29], retinol-binding protein [30], MUP [14,31] and various human lipocalins [32]. In some cases, for example MUP, these studies have proven useful in the determination of probable function, but, in others, such as BLG, the information has failed to offer clear insights into biological function.

Given the close structural similarity of trichosurin to MUP, and initial hypotheses surrounding pouch young retention and possible pheromone communication [10], six ligands known to bind to MUP were selected to investigate the pocket binding specificity in trichosurin (Figure 5). The six potential ligands thymol, 2-naphthol, 4-ethylphenol, 2-sec-butyl-thiazole and 2-isobutyl-3-methoxyxpyrazine were used as molecular probes. Ligand screening of six potential small molecules was carried out by co-crystallization of trichosurin in the presence of 3–5 mM ligand. With the exception of 2-sec-butyl-thiazole, structures were solved for each of the co-crystallizations and this showed that, of the five ligands tested, two bound in the pocket of trichosurin: 2-naphthol and 4-ethylphenol. For the remaining three ligands, no electron density could be seen in the trichosurin-binding pocket. The 2-naphthol ligand binds in the top and centre of the pocket and makes a hydrogen bond to an internal water molecule, which
ligands binding to trichosurin: 2-napthol, 4-ethylphenol, trans-P were calculated using a simple model for consecutive binding; change in fluorescence at 326 nm (Figure 6) and binding constants. Curves were fitted to the data for 2-naphthol following the nonenal, thymol and 2-isobutyl-3-methoxy pyrazine (Figures 5a–g). The chemical structures for six ligands which were assessed for binding to trichosurin in vitro using fluorescence and crystallography (a–f) and two ligands assessed using in silico docking (g and h).

In turn, is hydrogen-bonded by Phe O, Asn OD1 and Asn ND1 (see Figure 2B). Despite the presence of the ligand, the barrel remains completely closed. The binding of this ligand has resulted in a minor side chain rearrangement inside the binding pocket. The plane of the aromatic ring of Phe is rotated such that a small cavity is formed between the ring and the amide side chain of Asn and this is occupied by a water molecule. The water molecule (W232 in Figure 2B) accepts a hydrogen bond from the amide nitrogen but is also close enough to the aromatic ring to form an unconventional hydrogen bond by donation to the delocalized electron density of the aromatic ring plane. This phenomenon has been previously observed and described.

In contrast with the binding of 2-naphthol, 4-ethylphenol binds in the lower of the two lobes of the binding pocket forming a hydrogen bond to Leu O. The axis of the ligand is oriented perpendicular to the barrel axis (see Figure 2C), which is also in contrast with the binding of 2-naphthol. With the ligand in this position the trichosurin-binding pocket is only partially filled and in this structure a 2-propanol solvent molecule is able to occupy the upper lobe of the binding pocket. The trichosurin barrel is again completely closed.

In addition to the ligand electron density within the trichosurin barrel, the 2-naphthol structure also shows some electron density in a region bounded by the external barrel wall, the N-terminal helix and the dimer interface (see Figure 2D). This region of electron density lies in a trough formed by these three elements and is enclosed on a fourth side by the side chain of Arg. It was designated as a second, externally bound 2-naphthol molecule, making mainly hydrophobic interactions and also a favourable aromatic interaction with the delocalized electrons of the guanidine head group of Arg. In this position the 2-naphthol molecule remains partially solvent exposed with 6–12 Å accessible to the bulk solvent. The occupancy for this second binding site has been refined at 0.7 and is lower than that found for the main binding site of 0.9.

X-ray crystallographic studies on crystals grown in the presence of three of the four remaining possible ligands (trans-2-nonenal, thymol and 2-isobutyl-3-methoxy pyrazine) indicated that these molecules did not bind to trichosurin in the main pocket or externally. No suitable crystals were grown in the presence of 2-sec-butyl thiazole.

Fluorescence titration data were collected for five potential ligands binding to trichosurin: 2-naphthol, 4-ethylphenol, trans-2-nonenal, thymol and 2-isobutyl-3-methoxy pyrazine (Figures 5a–5e). Curves were fitted to the data for 2-naphthol following the change in fluorescence at 326 nm (Figure 6) and binding constants were calculated using a simple model for consecutive binding: P + L ↔ PL + L ↔ PL2 (where P is protein and L is ligand).

This ignores complicating factors such as cooperativity between binding sites both within individual binding sites and between monomers of the dimer, as well as the dimerization .

The change in fluorescence at 326 nm (ΔF) for ligand binding to trichosurin, ΔF on the y-axis is in arbitrary units. The upper line is the best fit of two exponentials to the data (C) with K values of 14.4 (middle line) and 24.3 μM (lower line).

For the change in fluorescence at 326 nm the curve fitting resulted in a K of 14 μM for the stronger site and of 24 μM for the weaker site. These binding data make sense in the context of the structure where binding into the cavity is expected to be favoured over binding in a secondary solvent-exposed site. Changes in fluorescence for 4-ethylphenol binding to trichosurin were followed at 320 nm where the intrinsic fluorescence of the ligand is at a minimum and the fluorescence of the protein is near its maximum. The amplitude of the changes in fluorescence were moderately small and are practically linear for ligand
concentrations up to 60 µM. This makes the data difficult to interpret; however, it sets a lower bound for the $K_d$ for 4-ethylphenol of 30 µM with the possibility that this $K_d$ is much weaker.

Fluorescence spectra for titrations with trans-2-nonenal, thymol and 2-isobutyl-3-methoxy pyrazine showed no changes in protein fluorescence for these compounds. This indicates that either these compounds are binding to trichosurin and not affecting the fluorescence of the protein or that they are not binding. The latter case is supported by the crystallographic data which shows no binding to trichosurin even at concentrations of 3–5 mM ligand. These data indicate that trichosurin preferentially binds small phenolic compounds. Furthermore, the preference for 2-naphthol and 4-ethylphenol over thymol (Figures 5a, 5b and 5f) indicates a specificity for phenolic compounds with particular substituents around the aromatic ring.

In comparison with other lipocalins, the $K_d$ for the binding of 2-naphthol to MUP is 0.38 µM [35] and $K_d$ values for a group of four ligands [retinol, retinoic acid, DAUDA (11-[5-(dimethylamino)-1-naphthalenesulfonyl]amino]undecanoic acid] and ANS (8-anilino-1-naphthalenesulfonic acid)] binding to ten different human lipocalins range from 0.1–10 nM ligand [32]. The Xenopus laevis coroid plexus lipocalin Cpl11 binds retinal in the nanomolar range and retinoic acid in the micromolar range [36,37].

The specificity of trichosurin for 2-naphthol and 4-ethylphenol with relatively high $K_d$ values suggests that small phenolic compounds of this type may have higher affinities and constitute the natural ligand for trichosurin. This is the subject of an ongoing screen and some possible suggestions are made in the subsequent discussion.

In summary, the structure of trichosurin is that of an eight-stranded β-barrel typical of the lipocalins. The binding of two solvent-derived isopropanol molecules within the putative trichosurin-binding pocket is suggestive of the manner in which the natural substrate or substrates may be bound. The shape and charge distribution within the pocket indicate that a small hydrophobic molecule perhaps with substrate groups may easily be accommodated within the closed pocket, with sites provided by the protein to satisfy potential hydrogen bonding requirements. This is confirmed by the specific binding of 4-ethylphenol and 2-naphthol over other possible ligands. It is unclear at this point what, if any, mechanism is required to enable access to the binding pocket for molecules larger than 2-propanol. They may well enter and exit through the N-terminal end of the barrel given the disorder seen in the trichosurin structure in this region, although this would be contrary to established theories about lipocalin ligand binding. There are few differences between the structures of trichosurin solved from crystals grown at two different pH values. The overall mode of dimerization is consistent between the two structures and does not appear to show a dependence on pH as is the case for bovine BLG. Furthermore, at the two pH values investigated, trichosurin did not show a major conformational difference consistent with the pH-induced barrel opening and closing seen for BLG [38]. It is possible that such an opening and closing mechanism could occur outside of the pH range studied. The biological purpose, if any, of a bound Zn$^{2+}$ ion on the barrel exterior remains unknown, although it could be involved in orientation and binding during a receptor interaction.

Trichosurin is a highly conserved protein found in the milk of metatherians but its role in the milk is not understood. It is most closely related to the MUPs (from mice and rats) whose name is somewhat historical as the primary site of expression of these proteins is the liver and they have also been shown to be expressed in many other tissues such as the submaxillary, lachrymal, parotid and mammary glands where they play a very diverse set of functions. From the similarity between trichosurin and the MUPs, it is tempting to speculate that the role of trichosurin might be in delivering attractants to guide the precarious path taken by the foetus from uterus to nipple in the early stages of marsupial development. However, trichosurin is a significant constituent of milk throughout postumous lactation [39] and this argues against a chemical attractant hypothesis.

A second hypothesis regarding the putative function of trichosurin is suggested by the demonstrated preference that we see for small phenolic ligands. Phenolic compounds are a significant constituent in plants and they are utilized by the plants as toxic feeding deterrents. These compounds have a very significant effect on feeding behaviour among mammalian herbivores as the animals ability to metabolize (detoxify) these compounds directly limits dietary intake. This has been demonstrated to be the case in possums [40]. Furthermore, female possums are able to concentrate phenolic compounds in their milk to levels appreciably higher than those found in the blood plasma [41]. This provides the young with a source of phenolics to enable priming of the neonate liver in the production of detoxifying enzymes which possums are known to utilize to survive on otherwise toxic foliage. The specificity for small phenolics that we observe in the present study is consistent with the hypothesis that trichosurin may facilitate the process of carrying secondary plant phenolics through the milk and help in transfer across the neonate gut. We have used the program GOLD [42] to dock common plant phenolics such as cinnemon and gallic acid (Figures 5g and 5h) into the ligand-binding pocket of trichosurin and this shows that these ligands are of a size and have hydrogen-bonding groups which make them compatible ligands for trichosurin. As all the marsupials are generalist herbivores, this hypothesis of phenolic priming of detoxification pathways is consistent with the high level of sequence conservation of trichosurin among the metatherians (from the South American opossum to the tammar wallaby). We have also used GOLD to survey other possible ligands and we note that planar-fused benzene rings analogous to 2-naphthol, such as 2-phenanthrenol and 2-hydroxyanthracene, can also be accommodated sterically and chemically (with good GOLD scores) by the ligand-binding pocket. This may point to a more promiscuous binding function for trichosurin.

We have determined the structures of the first metatherian lipocalin which is present at high levels throughout lactation in the possum, pointing towards an important role in the unusual development of the marsupial foetus. Marsupials occupy a unique position on the evolutionary tree as they diverged from the mammals midway between the divergence of placental mammals and the divergence of mammals from other animals such as birds. We have shown a preference for small phenolic ligands by trichosurin which bind with micromolar affinity and provides a possible functional hypothesis. However, the crucial next step is to discover the natural ligands for trichosurin and this is the subject of ongoing work.

We would like to thank M. Grigor (Faculty of Science, University of Auckland, Auckland, New Zealand) for the donation of the original trichosurin cDNA clone and for discussions and critical reading of the manuscript, and Dr Didier Nutrizio for help with synchrotron data collection at the ESRF, and Dr Armin Widmer for custom curve-fitting algorithms. We acknowledge financial support from the Marsden Fund of New Zealand.

**REFERENCES**


