SCIENTIFIC REPORTS

Received: 09 August 2016 Accepted: 25 October 2016 Published: 15 November 2016

OPEN Structure and Function of **Cyanobacterial DHDPS and DHDPR**

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Lysine biosynthesis in bacteria and plants commences with a condensation reaction catalysed by dihydrodipicolinate synthase (DHDPS) followed by a reduction reaction catalysed by dihydrodipicolinate reductase (DHDPR). Interestingly, both DHDPS and DHDPR exist as different oligomeric forms in bacteria and plants. DHDPS is primarily a homotetramer in all species, but the architecture of the tetramer differs across kingdoms. DHDPR also exists as a tetramer in bacteria, but has recently been reported to be dimeric in plants. This study aimed to characterise for the first time the structure and function of DHDPS and DHDPR from cyanobacteria, which is an evolutionary important phylum that evolved at the divergence point between bacteria and plants. We cloned, expressed and purified DHDPS and DHDPR from the cyanobacterium Anabaena variabilis. The recombinant enzymes were shown to be folded by circular dichroism spectroscopy, enzymatically active employing the quantitative DHDPS-DHDPR coupled assay, and form tetramers in solution using analytical ultracentrifugation. Crystal structures of DHDPS and DHDPR from A. variabilis were determined at 1.92Å and 2.83Å, respectively, and show that both enzymes adopt the canonical bacterial tetrameric architecture. These studies indicate that the quaternary structure of bacterial and plant DHDPS and DHDPR diverged after cyanobacteria evolved.

Lysine is synthesised *de novo* in bacteria, plants and some fungi¹⁻³. This occurs through either the α -aminoadipate pathway in fungi⁴ or the diaminopimelate pathway in bacteria and plants¹⁻³. The diaminopimelate pathway commences with the condensation of pyruvate and (*S*)-aspartate semialdehylde (ASA) to form (4*S*)-4-hydroxy-2,3,4,5-tetrahydro-(2*S*)-dipicolinic acid (HTPA) (Fig. 1A)^{1-3,5,6}. This reaction is catalysed by dihydrodipicolinate synthase (DHDPS) (EC4.3.3.7), which is the product of an essential gene in bacteria^{1,2,7-9}. HTPA is subsequently non-enzymatically dehydrated to dihydrodipicolinate (DHDP), which is then reduced by dihydrodipicolinate reductase (DHDPR) (EC1.17.1.8) to form tetrahydrodipicolinate (THDP) (Fig. 1A)^{1,10,11}. The pathway then diverges into four sub-pathways, namely the acetylase, aminotransferase, dehydrogenase and succinylase pathways, which operate across different genera and kingdoms^{2,12,13}. For example, the aminotransferase pathway is canonical to plants, but is also innate to cyanobacteria, including Anabaena variabilis (Kyoto Encyclopedia of Genes and Genomes www.genome.jp/kegg-bin/show_module?ava_M00527+Ava_3607, 2016). The structure of DHDPS has been determined from a number of bacterial species, including Agrobacterium tumefaciens¹⁴, Bacillus anthracis^{15,16}, Escherichia coli^{17,18}, Legionella pneumophila¹⁹, Mycobacterium tuberculosis²⁰, Pseudomonas aeruginosa²¹, Staphylococcus aureus^{22,23}, Streptococcus pneumoniae^{9,19} and Thermotoga maritima²⁴. The canonical bacterial DHDPS structure is a homotetramer best described as a 'head-to-head' dimer-of-dimers with four identical $(\beta/\alpha)_8$ -barrel subunits (Fig. 1B)^{3,7,25}. The structures of DHDPS from several plant species have also been determined showing that the plant orthologues also form homotetramers but in a 'back-to-back' dimer-of-dimers arrangement (Fig. 1B)²⁶⁻²⁹. Additionally, there are also reports of dimeric DHDPS enzymes from *P. aeruginosa*²¹, S. aureus^{22,23}, and Shewanella benthica³⁰.

Similarly, the structure of DHDPR has been determined from several bacterial species, including Corynebacterium glutamicum³¹, E. coli^{32,33}, M. tuberculosis^{34,35} and S. aureus³⁶. These studies show that the enzyme exists as a homotetramer with a unique quaternary architecture. Each monomer is comprised of an amino (N)-terminal NAD(P)H-binding domain that adopts a Rossman fold³⁷, and a carboxyl (C)-terminal tetramerisation domain with the substrate-binding pocket formed between the N- and C-terminal domains (Fig. 1C).

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Figure 1. Diaminopimelate biosynthesis pathway. (**A**) The pathway commences with the condensation of pyruvate and ASA catalysed by DHDPS to produce HTPA. HTPA is then non-enzymatically dehydrated to yield DHDP, which is subsequently reduced by DHDPR to form THDP. THDP is converted into the final product (*S*)-lysine in a series of enzymatic steps utilising 4 different sub-pathways. (**B**) The three dimensional structures of bacterial DHDPS (PDB ID: 1YXC) and plant DHDPS (PDB ID: 3TUU). (**C**) Crystal structure of bacterial DHDPR (PDB ID: 1DIH) and the DAMMIN model of A. thaliana DHDPR²⁸ (grey spheres) overlaying the structure of bacterial DHDPR (cyan) overlaid with the DAMMIN model of *A. thaliana* DHDPR²⁸.

However, the structure of a plant DHDPR has not yet been determined, but a recent study employing small angle X-ray scattering suggests that the enzyme from *Arabidopsis thaliana* adopts a novel dimeric structure (Fig. 1C)²⁸. Accordingly, there appears to be structural diversity between bacterial and plant orthologues of both DHDPS and DHDPR.

In this study, we aimed to determine the structure and function of the first DHDPS and DHDPR enzymes from a cyanobacterial species. Given that endosymbiotic theory suggests that the chloroplasts of plants were derived from the symbiosis of separate single bacterial cells³⁸, we were interested in characterising the structure and function of DHDPS and DHDPR from the model cyanobacterial species, *Anabaena variabilis* (Av)³⁹. Here, we present an in-depth characterisation of the structure and function of Av-DHDPS and Av-DHDPR in both solution and crystal states. We show that Av-DHDPS and Av-DHDPR both adopt the canonical bacterial structures, suggesting that the point of quaternary structural divergence between the bacterial and plants enzymes occurred after cyanobacteria evolved.

Results and Discussion

Purified recombinant Av-DHDPS and Av-DHDPR are active and folded. Av-DHDPS and Av-DHDPR were expressed in *E. coli* as His-tagged constructs and purified to >98% homogeneity using immobilised metal affinity chromatography (IMAC) (Fig. 2). The specific activity of purified Av-DHDPS and Av-DHDPR were determined to be 8.81 and 66.7 U/mg, respectively (Table 1), which correlate well to previous studies of recombinant orthologs^{10,40}. MS/MS sequencing following trypsin digestion demonstrates that both recombinant *A. variabilis* enzymes are comprised of the correct primary structure (Table 1). CD spectroscopy was subsequently employed to demonstrate that recombinant Av-DHDPS (Fig. 3, open symbols) and Av-DHDPR (Fig. 3, solid symbols) contain 45–51% α/β structure, which is consistent with previous studies of bacterial and plant orthologues^{7,9,14,16,19,22,25,27,41}.

Enzyme kinetic properties. Having determined that the recombinant cyanobacterial enzymes were homogenous, folded and enzymatically-active, we next set out to quantify their enzyme kinetic properties. Firstly, we characterised Av-DHDPS. Plots of initial rate as a function of varying pyruvate and ASA concentrations reveal typical Michaelis-Menten hyperbolic relationships (Fig. 4A). These data were globally fitted to yield a best

Enzyme	Purification step	Total activity (U)	Total protein (mg)	Specific activity (U mg ⁻¹)	Fold ¹	MS/MS coverage ²
Av-DHDPS	Crude	179	144	1.24	_	45%
	IMAC	170	19.3	8.81	7.10	
Av-DHDPR	Crude	265	186	24.8	_	100%
	IMAC	93.2	69.2	66.7	2.68	

 Table 1. Purification of recombinant Av-DHDPS and Av-DHDPR. ¹Fold of purified enzyme post-IMAC relative to crude. ²Sequence coverage from MS/MS analyses post in-gel tryptic digestion.





Figure 2. SDS-PAGE analyses of recombinant Av-DHDPS and Av-DHDPR. Lanes 1 & 6: molecular weight markers, kDa; <u>lane 2</u>: supernatant of non-ITPG treated cultures transformed with pRSETA-*dapA* (i.e. noninduced); <u>lane 3</u>: crude lysate post IPTG treatment of pRSETA-*dapA* transformed *E. coli* BL21-DE3 pLysS cells (i.e. induced); <u>lane 4</u>: post-IMAC purified recombinant Av-DHDPS; <u>lane 7</u>: supernatant of non-IPTG treated cultures transformed with pRSETA-*dapB*; <u>lane 8</u>: crude lysate post IPTG treatment of pRSETA-*dapB* transformed *E. coli* BL21-DE3 pLysS cells; <u>lane 9</u>: post-IMAC purified recombinant Av-DHDPR.



Figure 3. CD spectroscopy of recombinant Av-DHDPS and Av-DHDPR. Spectra were recorded using a protein concentration of 0.15 mg ml⁻¹ between wavelengths of 195–240 nm employing a step size of 1.0 nm with 4 s averaging time. Raw data for Av-DHDPS (\odot) and Av-DHDPR (\bullet) were fitted by nonlinear regression using the CDPro software and the CONTINLL algorithm (—), resulting in 33% α -helix, 18% β -structure, 14% β -turn and 35% unordered structure for Av-DHDPS with a RMSD of 0.070, and 18% α -helix, 27% β -structure, 13% β -turn and 42% unordered structure for Av-DHDPR with a RMSD of 0.037.

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fit to a bi-bi ping-pong mechanism without substrate inhibition ($R^2 = 0.99$), providing the kinetic parameters summarised in Table 2. The resulting kinetic parameters agree well with previous studies of bacterial orthologues^{9,14,16,20,22,24}. To establish whether recombinant Av-DHDPS is sensitive to allosteric feedback inhibition by



Figure 4. Enzyme kinetic profiles of recombinant Av-DHDPS. (A) Michaelis-Menten analyses of Av-DHDPS. The initial velocity is plotted as a function of pyruvate concentration at varying ASA concentrations of 0.0375 mM (\diamond), 0.075 mM (\Box), 0.125 mM (\blacksquare), 0.25 mM (\diamond), 0.5 mM (\diamond). The global nonlinear best-fit using the ENZFITTER software (Biosoft) was obtained to a bi-bi ping pong model without substrate inhibition and resulted in $R^2 = 0.991$. Data are presented as mean and error bars as standard deviation (n = 3). (B) Dose response curve showing initial rate plotted as a function of (S)-lysine concentration, for Av-DHDPS (\blacklozenge), Ec-DHDPS (\blacksquare), and Vv-DHDPS (\circ). The data were fitted to a ligand binding, four-parameter logistic function using ENZFITTER yielding $R^2 = 0.994$ for Av-DHDPS (\cdots), $R^2 = 0.982$ for Ec-DHDPS (-) and $R^2 = 0.982$ for Vv-DHDPS (-). Data are presented as mean and error bars as standard deviation (n = 3).

Enzyme	$K_{\rm M}^{\rm PYR}$	K _M ASA	k _{cat}	$IC_{50}^{LYS}(mM)$
Av-DHDPS	0.41	0.14	15	0.068
	$K_{\rm M}^{\rm DHDP}$	K _M ^{NADH}	k _{cat}	
Av-DHDPR	3.6	0.31	3.0	N/A

Table 2. Enzyme kinetic parameters for recombinant Av-DHDPS and Av-DHDPR.

(S)-lysine, which is the end product of the diaminopimelate pathway (Fig. 1A), enzyme assays were also performed with increasing (S)-lysine concentrations^{1-3,18,19,29}. DHDPS from *E. coli* (Ec-DHDPS) and *V. vinifera* (Vv-DHDPS) were used as controls, given that previous studies show these orthologues are allosterically inhibited by (S)-lysine^{18,29}. The dose-response curves for Av-DHDPS, Ec-DHDPS and Vv-DHDPS are shown in Fig. 4B with the nonlinear best fits to a four-parameter logistic function yielding an $IC_{50}^{LYS} = 0.068 \text{ mM}$ ($R^2 = 0.99$) for Av-DHDPS (Table 2), which is closer to the value obtained for Vv-DHDPS [$IC_{50}^{LYS} = 0.030 \text{ mM}$ ($R^2 = 0.98$)] than for Ec-DHDPS [$IC_{50}^{LYS} = 0.210 \text{ mM}$ ($R^2 = 0.98$)]. Interestingly, a recent study revealed that the amino acid at position 56 (*E. coli* numbering) determines whether DHDPS enzymes will be inhibited by (S)-lysine¹⁹. Moreover, a His or Glu at this position is a marker of allosteric inhibition, whereas DHDPS sequences that contain Lys or Arg at position 56 are insensitive to lysine-mediated allosteric inhibition. For Av-DHDPS, there is a Glu at position 58 (equivalent to position 56 in Ec-DHDPS), which is consistent with the recently established determinants of allostery for DHDPS enzymes¹⁹.

For Av-DHDPR, the enzyme kinetic parameters were determined employing *E. coli* DHDPS as the coupling enzyme using increasing DHDP and NADH concentrations (Fig. 5A). The nonlinear least squares global fit was obtained to a ternary complex model (R^2 = 0.98), yielding the kinetic values reported in Table 2. A comparison of NADH and NADPH showed that Av-DHDPR is inhibited by its substrate, DHDP, when NADPH is employed as the cofactor (Fig. 5B). Subsequent thermodynamic measurements using microscale thermophoresis^{19,42} revealed that the cyanobacterial enzyme binds the substrate analogue 2,6-pyridinedicarboxylate (2,6-PDC)³⁴ only when NADP⁺, and not NAD⁺, is present in the titration (Fig. 5C). This suggests that Av-DHDPR is inhibited by DHDP in the presence of the oxidised phosphorylated cofactor, which is consistent with *S. aureus* DHDPR¹¹.

Av-DHDPS and Av-DHDPR are tetramers in solution. To characterise the quaternary structure of Av-DHDPS and Av-DHDPR in solution, sedimentation velocity experiments were conducted in the analytical ultracentrifuge. The absorbance versus radial position profile for Av-DHDPS and Av-DHDPR at initial protein concentrations in the range of 0.1–7.0 μ M show a distinct sedimentation boundary profile consistent with the presence of a single species (data not shown). Continuous size-distribution analyses of the data at 4.0 μ M reveal that Av-DHDPS (Fig. 6A) and Av-DHDPR (Fig. 6B) have molar masses of 147 and 139 kDa with standardised sedimentation coefficients ($s_{20,w}$) of 7.1S and 6.9S, respectively (Table 3). This indicates that both enzymes exist as stable tetramers in solution^{16,24,41,43}. Additionally, the calculated f/f_0 and axial ratio values (Table 3) are consistent with the asymmetric structures previously reported for DHDPS and DHDPR enzymes^{16,24,41,43}.



Figure 5. Kinetic parameters and cofactor preference of Av-DHDPR. (A) The initial rate of Av-DHDPR is plotted against increasing NADH concentrations of 0.0375 mM (\bullet), 0.075 mM (\circ), 0.125 mM (\bullet), 0.25 mM (\circ) and 0.5 mM (\blacksquare). The (-) represent the nonlinear best fit to a ternary complex mechanism without substrate inhibition using the ENZFITTER software (Biosoft), resulting in a R^2 of 0.986. Data is presented as mean \pm SD (n=3). (**B**) The initial rate is plotted against either 0.2 mM [NADH] (\blacksquare) or [NADPH] (\circ) with trend lines shown as solid lines (-). Data is presented as mean \pm SD (n=3). (**C**) Microscale thermophoresis binding curve of 2,6-PDC to Av-DHDPR in the presence of 150 μ M NADP⁺ using the signal from Thermophoresis + T-Jump (\bullet). The solid line (-) represents the nonlinear best fit to the single binding site model to yield a K_D of 100 \pm 8.20 μ M. *Inset:* 2,6-PDC titrations in the presence of 150 μ M NAD⁺. Data are presented as mean and error bars as standard error (N=3).



Figure 6. Analytical ultracentrifugation sedimentation velocity analyses. The c(M) distribution is plotted as a function of molar mass (kDa) for (**A**) recombinant Av-DHDPS and (**B**) recombinant Av-DHDPR. Top: Residuals plotted as a function of radial position resulting from the c(M) distribution best fit using N = 200 and P = 0.95, which yielded a rmsd = 0.00951 and runs test Z = 1.94 for Av-DHDPS and a rmsd = 0.00870 and runs test Z = 2.12 for Av-DHDPR.

Enzyme	$s_{20,w}(S)^1$	Molar Mass (kDa) ²	f/f_0^{3}	Axial ratio (a/b) ⁴
Av-DHDPS	7.1	147	1.3	2.5
Av-DHDPR	6.9	139	1.3	2.6

Table 3. Hydrodynamic properties of recombinant Av-DHDPS and Av-DHDPR. ¹Standardised sedimentation coefficient calculated from SEDNTERP software. ²Determined from the ordinate maximum of the c(M) distribution best fits (Fig. 6). ³Frictional coefficient calculated from $s_{20,w}$ using the ∇ method employing SEDNTERP software. ⁴Calculated from SEDNTERP using a prolate model (∇ method).

Crystal structures of Av-DHDPS and Av-DHDPR. Av-DHDPS (7 mg/ml) was crystallised at 281 K in 16% (w/v) PEG3350, 0.4 M trisodium citrate, 0.1 M bis-Tris propane chloride, 10 mM pyruvate, pH 6.5, yielding crystals with dimensions of 0.1 mm \times 0.05 mm (Fig. 7A). These crystals diffracted to a highest resolution of 1.92 Å (Fig. 7B,C). Similarly, Av-DHDPR (6 mg/ml) was crystallised at 281 K in 21% (w/v) PEG3350, 0.2 M lithium sulphate, 0.1 M bis-Tris chloride, pH 5.5, yielding crystals with dimensions of 0.04 mm \times 0.015 mm (Fig. 7D) that



Figure 7. Crystallisation and X-ray diffraction data of recombinant Av-DHDPS and Av-DHDPR. (A) Photograph of Av-DHDPS crystal that was used to generate the diffraction patterns shown in (B) 90° rotation and (C) 180° rotation. (D) Photograph of Av-DHDPR crystal used to generate the diffraction patterns shown in (E) 90° rotation and (F) 180° rotation. Diffraction patterns were obtained using the MX2 beamline at the Australian Synchrotron as reported in the Methods section.



Figure 8. Av-DHDPS structure determined by X-ray crystallography. (A) Structure of Av-DHDPS (PDB ID: 5KTL) was solved to a resolution of 1.92 Å. (B) The active site of Av-DHDPS (red, PDB ID: 5KTL) overlayed with Ec-DHDPS (black, PDB ID: 1YXC). Shown in purple is the substrate pyruvate bonded to Lys164 in Av-DHDPS. (C) The allosteric site of Av-DHDPS (PDB ID: 5KTL) overlayed with Ec-DHDPS (black, PDB ID: 1YXC), with the allosteric inhibitor (S)-lysine shown in green (PDB: 1YXD).

diffracted to 2.83 Å (Fig. 7E,F). The diffraction data were subsequently used to determine the three-dimensional structure of Av-DHDPS and Av-DHDPR by molecular replacement.

Consistent with the analytical ultracentrifugation (AUC) studies in solution, the crystal structure of Av-DHDPS reveals a homotetramer that resembles the 'head-to-head' dimer-of-dimers canonical to bacterial orthologues (Fig. 8A). Each monomer is comprised of 297 residues that folds to form a N-terminal $(\beta/\alpha)_8$ -barrel domain and a C-terminal domain consisting of 3 α -helices. Close inspection of the active site shows that it is comprised of the key conserved residues known to be important for catalysis, namely Lys164, Thr46, Tyr109, Tyr136



Figure 9. Av-DHDPR structure determined by X-ray crystallography. (**A**) Structure of Av-DHDPR (PDB ID: 5KT0) was determined to a resolution of 2.83 Å. (**B**) Nucleotide binding site of Av-DHDPR (red, PDB ID: 5KT0) overlayed with Ec-DHDPR (black, PDB ID: 1ARZ). Shown in cyan is the substrate NADH (PDB ID: 1ARZ). (C) Substrate binding site of Av-DHDPR (red, PDB ID: 5KT0) overlayed with Ec-DHDPR (black, PDB ID: 1ARZ). Shown in yellow is the substrate analogue 2,6-PDC (PDB ID: 1ARZ). (**D**) Extended loop in Av-DHDPR (red) between residues Val107 and Gly116 located between β-sheets B4 and B5 overlayed with Ec-DHDPR (grey, PDB ID: 1ARZ) and *M. tuberculosis* DHDPR (black, PDB ID: 1YL5).

and Arg141, which are equivalent to Lys161, Thr44, Tyr107, Tyr133 and Arg138 in *E. coli* DHDPS (Fig. 8B)^{17,18}. Likewise, inspection of the allosteric site confirms the presence of Glu at position 58 (His56 in Ec-DHDPS), but also reveals that Trp occupies position 55 (His53 in Ec-DHDPS), which is common in plants but not bacteria (Fig. 8C)²⁶⁻²⁹. The presence of a Trp at this positon is likely to explain the plant-like lysine IC_{50} for Av-DHDPS (Fig. 4B, Table 2).

The Av-DHDPR crystal structure also reveals a homotetramer (Fig. 9A), which is consistent with the in-solution AUC studies, and agrees well with previously characterised bacterial DHDPR structures^{31–36}. Each monomeric unit consists of 287 residues with an N-terminal nucleotide and a C-terminal tetramerisation domain connected via a hinge region. Although residue variation is observed at the nucleotide binding site, the physic-ochemical properties of the residues are still conserved (Fig. 9B) as observed for other bacterial DHDPR species^{31–36}. By contrast, the substrate binding cleft of Av-DHDPR is predominantly conserved (Fig. 9C). However, Av-DHDPR has a unique prolonged solvent-exposed loop located between β -sheets B4 and B5, consisting of residues Val107 to Gly116 (Fig. 9D). Overall, the loop residues have a neutral, slightly hydrophobic nature, and the function of the loop is unclear. This prolonged loop is absent in all other published DHDPR structures suggesting this is a unique feature of cyanobacterial DHDPR^{31–36}.

Bioinformatics analysis. This study reveals that both Av-DHDPS and Av-DHDPR adopt the canonical bacterial tetrameric architecture. This was unexpected given the endosymbiosis theory³⁸ and the shared aminotransferase pathway found in both cyanobacteria and plants. Consequently, bioinformatics sequence analyses of DHDPS and DHDPR from several bacterial and plant species were performed to predict when the plant structures first evolved. For DHDPS, the dataset employed consisted of sequences from 150 bacteria, 85 cyanobacteria, 84 plants, 12 green algae and 2 red algae from the NCBI protein database (www.ncbi.nlm.nih.gov/protein, 2016). A representative subset of these sequences are aligned in Fig. 10A. It was noted that the motifs Arg43 to Asp45, Arg108 to Gln116 and Gly325 to Tyr/His327 (*V. vinifera* numbering) are conserved in plants and form an interacting network at the tetramerisation interface in plant structures^{26–29}. These motifs were also found in

•			43	108	325	Phylum	NCBI ID	
A	v.	vinifera	RFD	REAIHATEQ	GRY	Plant	400261070	
•••	N.	sylvestris	RFD	REAIHATEQ	GRY	Plant	698557988	
	z.	mays	RFD	REAVHATEQ	SRY	Plant	162462912	
	ο.	sativa	RFD	REAIHATEQ	SRY	Plant	563440366	
	м.	notabilis	RFD	REAIHATEQ	DRY	Plant	703085863	
	c.	reinhardtii	KFD	REALHATEQ	GRH	Green algae	158281680	
	v.	carteri	KFD	REALHATEQ	GRH	Green algae	300267543	
	c.	variabilis	KFD	REALHATEQ	SSY	Green algae	552813123	
	М.	pusilla	KID	REAVHASSQ	SEW	Green algae	226462645	
	в.	prasinos	KID	REAVHATSQ	DEW	Green algae	612387409	
	G.	sulphuraria	EVD	EEAVEATHK		Red algae	452822686	
	с.	merolae	EVD	EEAIEATHK		Red algae	544210540	
	A.	variabilis	SVN	KEAIAATQK		Cyanobacteria	75703537	
	ο.	nigro	SVN	SEAVAATKK		Cyanobacteria	428244203	
	s.	hofmanni	SVN	KEAIAATQK		Cyanobacteria	657931409	
	G.	violaceus	NVH	REAVVATRK		Cyanobacteria	37520587	
	s.	subsalsa	TVN	HEAIARSOK		Cyanobacteria	703194018	
	s.	aureus	KVN	EKSIQASIQ		Bacteria	473009079	
	E.	coli	NVC	TAEAISLTQ		Bacteria	190906811	
	s.	enterica	NVS	AEAISLTOR		Bacteria	261247688	
	L.	monocytogenes	KVC	AETIAFTRE		Bacteria	984410294	
	н.	salinarum	SLD	ROTVAHAEH		Bacteria	10580053	
				183		208	Phylum	NCBI ID
D	G.	max		GV-SY-DMDQI	KLIRDP	ROQLEMVGPEE	Plant	255671801
D	P.	mume		GV-CF-DMEOI	QIIRDT	QQQIEMVGPEE	Plant	645276001
	м.	notabilis		GV-SF-DMDQI	OMIRDP	KQQIEMVGPEE	Plant	703069664
	м.	truncatula		GA-SF-DMDOI	OLIRDP	ROOVEMVGPEE	Plant	922333438
	G.	soia		GV-SY-DMDOI	RLIRDP	ROOLEMVGPEE	Plant	734362028
	т.	ap. GSL018		GC-PF-KETDI	EKLRDD	DASHORLGVPE	Green algae	654118802
	к.	flaccidum		GV-KF-NVDEI	ERVRDP	SEOVERMHVPE	Green algae	971513782
	А.	protothecoides		GL-EF-DLSNI	ELVREA	AEOIGRMOVPE	Green algae	675353396
	с.	subellinsoidea		GT-PT-BEADT	FRVRDA	STOVERMOVER	Green algae	384253568
	o.	tauri		GC-DE-DVSOI	FLVRDV	ENOTGRMGVPE	Green algae	116055387
	c.	merolae		GITGEGGEODI	OKTRDP	ALOVSEMGVEE	Red algae	544214461
	G.	sulphuraria		GA-HFEYDEDL	PMTRDP	ALCOOFLSTPE	Red algae	452824477
	<u>5</u> -	variabilia		GE-TENSATVEET	FRIDGA	P======G9	Cvanobacteria	123610708
	~	nigro-wiridia		CE-DENDDEVEED	FRIDCA	P68	Cyanobacteria	428244018
	č.	eninsammum		GK-SFNPPTVTFT	FRIAGA	R63	Cyanobacteria	428246863
	č.	eprosentium		CY-DEMODIVEE	DE TOCA	R00	Cyanobacteria	420240003
	p.	ap. PCC 0303		CD-FFNIFINDDFF	PULMCA	R	Cyanobacteria	427999930
	£.	flavnari		DE-DIEDCAUVOD	EGUM	RGR	Cyanobacteria Rectaria	24111477
	5.	ilexner1		DK-DLKDGAVYSR	EGHT	GE	Bacceria	241114//
	E.	COLL		DK-DLKDCAVYSR	EGHT	GE	Bacteria	3091903
	5.	enterica		DK-NLKDCAVYSR	EGIT	GE	Bacteria	320084300
	к.	pneumoniae		NK-DLKDCAVYSR	EGYT	GE	Bacteria	6/3508/59
	v.	parahaemolyticu	18	DK-DLKECAVYSR	EGYT	GE	Bacteria	821090286

Figure 10. Bioinformatics sequence analyses of DHDPS and DHDPR in multiple phyla. (A) Multiple sequence alignment of the DHDPS tetramerisation interface residues (*V. vinifera* DHDPS numbering). The sequence motifs 43–45, 108–116 and 325–327 are conserved in plants and green algae but absent in bacteria and cyanobacteria. (B) Multiple sequence alignment of the flexible loop motif (183–204, *E. coli* numbering) in DHDPR. The motif length is significantly shorter in bacteria and cyanobacteria compared to red algae, green algae and plants. The long dashed line in panels A and B indicates the point of structural divergence based on sequence similarity, whereas individual dashes (–) represent a gap in the sequence. For simplicity, this alignment only shows a fraction of the sequences analysed.

green algae but not in bacteria, cyanobacteria or red algae species (Fig. 10A). This finding suggests that the point of divergence of the DHDPS quaternary structures occurred between red and green algae. This remains to be verified experimentally.

To examine whether the same pattern is observed for DHDPR, sequences from 25 bacteria, 57 cyanobacteria, 38 plants, 12 green algae and 2 red algae from the NCBI protein database were obtained (July 2016). Figure 10B shows a representative multiple sequence alignment of a subset of these species. The length of the loop motif in cyanobacteria (Gly183-Ser203, *E. coli* DHDPR numbering) is similar in length to other bacterial species (Fig. 10B). However, this loop is significantly longer in plant, red algae and green algae sequences (Fig. 10B). This suggests that DHDPR from red and green algae may adopt a similar dimeric quaternary architecture to plant orthologues. This also remains to be verified experimentally.

Conclusions

In this study, we determined for the first time the enzyme kinetic parameters, solution properties and three-dimensional structures of DHDPS and DHDPR from a cyanobacterial species. We show both enzymes exist as homotetramers in solution and in the crystal state, and that they adopt the canonical bacterial quaternary architecture. Our results suggest that the point of structural divergence differentiating bacterial and plant DHDPS and DHDPR and DHDPR enzymes occurred between cyanobacteria and lower plants.

Parameter	Av-DHDPS	Av-DHDPR
Wavelength (Å)	0.9537	0.9537
No. of images	360	360
Oscillation angle per frame	0.5	1.0
Space group	P21212	I222
Unit-cell parameters		
a, b, c (Å)	75.73, 154.35, 55.79	72.73, 89.36, 95.92
α, β, γ (°)	90, 90, 90	90, 90, 90
Resolution (Å)	1.92	2.83
Observed reflections	335888 (47075)	110053 (17182)
Unique reflections	49053 (7452)	7762 (1225)
Completeness (%)	99.1 (94.5)	99.7 (98.7)
R _{merge}	10.3 (78.0)	10.0 (36.6)
Mean $I/\sigma(I)$	14.50 (2.68)	25.08 (8.36)
Redundancy	6.85 (6.32)	14.18 (14.03)
Molecules per asymmetric unit	2	1
Wilson-B	31.86	40.86
R _{work} /R _{free} [%]	18.0/22.4	18.8/25.3
R _{free} test set count	1016	452
Protein/solvent/metal atoms	4413/228/5	2037/1/0
Average <i>B</i> factor [Å ²]	50.6	43.8
Favored/additionally allowed/generously allowed/ disallowed residues in Ramachandran plot [%]	91.3/8.3/0.0/0.4	89.5/8.8/1.3/0.4

 Table 4. X-ray data collection and refinement statistics for recombinant Av-DHDPS (PDB ID: 5KTL) and Av-DHDPR (PDB ID: 5KT0). Values in parentheses are for the highest resolution bin.

Methods

Cloning, expression and purification of Av-DHDPS and Av-DHDPR. Synthetic codon-optimised Av-DHDPS (i.e. *dapA*) and Av-DHDPR (i.e. *dapB*) genes cloned into pRSET-A expression vectors were purchased from GeneArt. The plasmids were subsequently transformed into *E. coli* BL21-DE3 pLysS cells for the overexpression of the recombinant enzymes. Recombinant protein was produced by treating *E. coli* BL21-DE3 pLys cells with 1.0 mM IPTG at 25 °C for 8 h. Cells were harvested by centrifugation ($5000 \times g$) and resuspended in 20 mM Tris-HCl, pH 8.0, 500 mM NaCl, 20 mM imidazole, 5% (ν/ν) glycerol, which included 10 mM pyruvate for Av-DHDPS, given that pyruvate is known to stabilise DHDPS enzymes^{16,22}. The cell suspension was lysed on ice by sonication using a Vibra Cell VC40 (Sonics & Materials) at 40 micron using 6 cycles of 10 sec on followed by 2 min off. Recombinant His-tagged enzymes were isolated from the cell lysate using IMAC employing a 5 ml His-Trap column (GE Healthcare) and a 0–500 mM imidazole, respectively. The purified protein was dialysed overnight against 20 mM Tris-HCl, pH 8.0, 100 mM NaCl, 5% (ν/ν) glycerol, which included 10 mM pyruvate for Av-DHDPS and Av-DHDPR eluted at 195 mM and 140 mM imidazole, respectively. The purified protein was dialysed overnight against 20 mM Tris-HCl, pH 8.0, 100 mM NaCl, 5% (ν/ν) glycerol, which included 10 mM pyruvate for Av-DHDPS.

Tandem mass spectrometry. Purified recombinant Av-DHDPS and Av-DHDPR were subjected to trypsin digestion and tandem mass spectrometry (MS/MS) sequencing using a Thermo Scientific LTQ Orbitrap Elite ETD Mass Spectrometer as previously reported^{44,45}.

Circular dichroism spectroscopy. Circular dichroism (CD) spectra of Av-DHDPS and Av-DHDPR were obtained using an Aviv Model 420 CD spectrometer using similar methods reported previously^{14,16,22,27,41,46}. Briefly, wavelength scans were performed between 195 and 240 nm with a 4.0 sec averaging time in 20 mM Tris, pH 8.0, 150 mM NaCl (also containing 1 mM pyruvate for Av-DHDPS) in a 1.0 mm quartz cuvette. Data were analysed using the CDPro software package incorporating the SP22X database^{47,48}.

DHDPS-DHDPR coupled enzyme kinetic assay. Kinetic analyses of Av-DHDPS and Av-DHDPR were performed employing the DHDPS-DHDPR coupled assay as previously described^{7,9,11,14,16,19,22,24,25,27-29}. Briefly, assays were performed in triplicate at 30 °C in a 1 cm acrylic cuvette using *E. coli* DHDPR and *E. coli* DHDPS as the coupling enzymes for Av-DHDPS and Av-DHDPR, respectively. Mixtures were allowed to equilibrate in a temperate-controlled Cary 4000 UV-Vis spectrophotometer for 12 min before initiating the reaction with ASA. The initial reaction rate data were analysed using the ENZFITTER software (Biosoft). Data were fitted to various models, including the bi-bi ping-pong and ternary complex models with and without substrate inhibition, with best fits determined from the highest R^2 value.

Microscale thermophoresis. Affinity measurements using microscale thermophoresis (MST) were carried out with a Monolith NT. LabelFree instrument (NanoTemper Technologies)^{19,42}. 2,6-PDC diluted in water

 $(5.0 \text{ mM to } 2.4 \mu\text{M})$ was mixed 1:1 with the enzyme pre-incubated with $150 \mu\text{M}$ NAD⁺ or NADP⁺, yielding a final DHDPR concentration of $10 \mu\text{M}$ and a dilution series of 2.5 mM to $1.2 \mu\text{M}$ of 2,6-PDC. Controls were performed in the absence of NAD⁺/NADP⁺, with water added instead, but with the same dilution series for 2,6-PDC. All experiments were incubated for 30 min at $30 \,^{\circ}$ C, before applying samples to Monolith NT Standard Treated Capillaries (NanoTemper Technologies). Thermophoresis was measured at $30 \,^{\circ}$ C with $5 \, \text{s}/30 \, \text{s}/5 \, \text{s}$ laser off/on/ off times. Experiments were conducted at 30% LED power and 40% MST IR-laser power. Data from three independently performed experiments were fitted to the single binding site model (NT. Analysis software version 1.5.41, NanoTemper Technologies) using the signal from Thermophoresis + T-Jump.

Analytical ultracentrifugation. Sedimentation velocity experiments were performed at 20 °C in a Beckman Coulter Model XL-A analytical ultracentrifuge using double sector quartz cells and An50-Ti rotor^{49,50}. 400 μ l of buffer and 380 μ l of sample at an initial concentration ranging from 0.1 μ M to 7.0 μ M were loaded into the reference and sample sectors of the cells, respectively. The rotor was accelerated to 40,000 rpm and data were collected continuously at 230 nm using a step size of 0.003 cm without averaging. Initial scans were carried out at 3,000 rpm to determine the optimal wavelength and radial positions for the high speed experiment. Solvent density, solvent viscosity, and estimates of the partial specific volume of Av-DHDPS (0.736 ml/g) and Av-DHDPR (0.736 ml/g) at 20 °C were calculated using SEDNTERP⁵¹. Data were fitted using the SEDFIT software (www. analyticalultracentrifugation.com) to a continuous size-distribution model^{52–55}.

Crystallisation and X-ray diffraction. Av-DHDPS and Av-DHDPR were crystallised using the hanging-drop vapour diffusion method as described previously^{10,30,40,56}. For X-ray data collection, crystals were transferred to reservoir solution containing 20% (*v*/*v*) glycerol and 12.5 mM MnCl₂, and directly flash frozen in liquid nitrogen. Intensity data were collected at the Australian Synchrotron using the MX2 beamline. Diffraction data were processed using MOSFLM⁵⁷ and scaled using SCALA⁵⁸. Molecular replacement was performed using the MR protocol of Auto-Rickshaw⁵⁹ with *B. anthracis* DHDPS (PDB ID: 1XKY) as the search model for Av-DHDPS and *M. tuberculosis* DHDPR (PDB ID: 1YL5) as the search model for Av-DHDPR. Structural refinement was performed using REFMAC5⁶⁰ with iterative model building using COOT⁶¹. The refinement statistics are provided in Table 4. For Av-DHDPS, Ramachandran statistics showed 91.3% in the preferred region, 8.3% in the additionally allowed region and 0.4% in the disallowed region consistent with previous structural reports^{15,21,62}. For Av-DHDPR, Ramachandran statistics showed 89.5% in the preferred region, 8.8% in the additionally allowed region and 1.3% in the generously allowed region consistent with previous structural .4% of residues (i.e. Ser89 and Gln112) were in the disallowed region due to poor electron density.

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Acknowledgements

We would firstly like to acknowledge the support and assistance of the friendly staff at the CSIRO Collaborative Crystallisation Centre (www.csiro.au/C3), Melbourne, Australia and the beamline scientists at the Australian Synchrotron, Victoria, Australia. We would also like to thank the *New York Structural Genomics Research Consortium* for providing purified protein for preliminary crystallisation studies. M.A.P. and S.P. acknowledge the Australian Research Council for funding support (DP150103313) and T.P.S.C. the National Health and Medical Research Council of Australia for fellowship support (APP1091976). We would also like to acknowledge the *La Trobe University-Comprehensive Proteomics Platform* for providing infrastructure and expertise. Finally, we thank all members of the Perugini laboratory for helpful discussions during the preparation of this manuscript.

Author Contributions

J.B.C. performed experiments, analysed the experimental data, and co-wrote the manuscript. T.P.S.C. obtained and analysed the MST experimental data and revised the manuscript. P.F. performed the trypsin digestion and MS/MS sequencing. F.G.P. provided essential reagents and revised the manuscript. S.P. determined the crystal structures of Av-DHDPS and Av-DHDPR and revised the manuscript. M.A.P. devised the project, planned the experiments and co-wrote the manuscript.

Additional Information

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Christensen, J. B. *et al.* Structure and Function of Cyanobacterial DHDPS and DHDPR. *Sci. Rep.* **6**, 37111; doi: 10.1038/srep37111 (2016).

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