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Analytical ultracentrifugation and preliminary X-ray studies of the chloroplast envelope quinone oxidoreductase homologue from *Arabidopsis* thaliana

Sarah Mas y mas, a Cécile Giustini, b Jean-Luc Ferrer, c Norbert Rolland, b Gilles Curien and David Cobessid*

^aInstitut de Biologie Structurale, Université Grenoble Alpes, CNRS, CEA, 71 Avenue des Martyrs, 38044 Grenoble, France, ^bLaboratoire de Physiologie Cellulaire et Végétale, CNRS, Université Grenoble Alpes, CEA, INRA, 17 Rue des Martyrs, 38054 Grenoble, France, ^cInstitut de Biologie Structurale, CEA, Université Grenoble Alpes, CNRS, 71 Avenue des Martyrs, 38044 Grenoble, France, and ^dInstitut de Biologie Structurale, CNRS, Université Grenoble Alpes, CEA, 71 Avenue des Martyrs, 38044 Grenoble, France. *Correspondence e-mail: david.cobessi@ibs.fr

Quinone oxidoreductases reduce a broad range of quinones and are widely distributed among living organisms. The chloroplast envelope quinone oxidoreductase homologue (ceQORH) from *Arabidopsis thaliana* binds NADPH, lacks a classical N-terminal and cleavable chloroplast transit peptide, and is transported through the chloroplast envelope membrane by an unknown alternative pathway without cleavage of its internal chloroplast targeting sequence. To unravel the fold of this targeting sequence and its substrate specificity, ceQORH from *A. thaliana* was overexpressed in *Escherichia coli*, purified and crystallized. Crystals of apo ceQORH were obtained and a complete data set was collected at 2.34 Å resolution. The crystals belonged to space group *C*222₁, with two molecules in the asymmetric unit.

1. Introduction

The chloroplast plays central roles in plant growth and development by catalyzing a remarkable range of metabolic and physiological functions (Armbruster et al., 2009; Rolland et al., 2012). Most of the chloroplast proteins are encoded by the nuclear genome, synthesized in the cytosol and then imported into the chloroplast through the TOC/TIC (translocons at the outer and inner envelope of the chloroplast) machineries (Strittmatter et al., 2010). If most of the outer envelope proteins are excluded, these chloroplast proteins possess a predictable and cleavable transit peptide at their N-terminus (Jarvis & López-Juez, 2013; Richardson et al., 2014). Recently, some chloroplast proteins, such as the inner envelope protein IEP32, also called Tic32 (Nada & Soll, 2004), and the chloroplast envelope quinone oxidoreductase homologue (ceQORH; Miras et al., 2002, 2007), have been shown to be targeted to the chloroplast inner envelope membrane while lacking this predictable and cleavable transit peptide. Moreover, the two abovementioned chloroplast proteins were also shown to be targeted to the chloroplast by an as yet unknown alternative targeting pathway (Miras et al., 2007).

While information for the targeting of canonical chloroplast proteins is located at their N-terminus (*i.e.* within the transit peptide sequence), ceQORH displays a nonterminal hydrophilic domain (residues 59–100) that is essential for its targeting to the chloroplast (Miras *et al.*, 2007). Based on sequence alignment, the 329-residue ceQORH protein was proposed to belong to the quinone oxidoreductase (QOR)



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 Table 1

 Macromolecule-production information.

Source organism	A. thaliana
DNA source	AT4G13010 cDNA
Cloning vector	pET-30a+
Expression vector	pET-30a+
Expression host	E. coli Rosetta2 (DE3)
Complete amino-acid sequence	AGKLMHALQYNSYGGGAAGLEHVQVPVPTPKSNE-
of the construct produced	VCLKLEATSLNPVDWKIQKGMIRPFLPRKFPC-
	IPATDVAGEVVEVGSGVKNFKAGDKVVAVLSH-
	LGGGGLAEFAVATEKLTVKRPQEVGAAEAAAL-
	PVAGLTALQALTNPAGLKLDGTGKKANILVTA-
	ASGGVGHYAVQLAKLANAHVTATCGARNIEFV-
	KSLGADEVLDYKTPEGAALKSPSGKKYDAVVH-
	CANGIPFSVFEPNLSENGKVIDITPGPNAMWT-
	YAVKKITMSKKQLVPLLLIPKAENLEFMVNLV-
	KEGKVKTVIDSKHPLSKAEDAWAKSIDGHATG-
	KIIVEP

family (Miras et al., 2002). This enzyme family shares common structural motifs and oxidizes NADPH to reduce a broad range of quinones, and is widely distributed among living organisms (Pan et al., 2009). Enzymes belonging to the quinone oxidoreductase (QOR) family are often dimeric, such as those from Escherichia coli (Thorn et al., 1995) and Saccharomyces cerevisiae (Guo et al., 2011), but can also be monomeric, such as that from Coxiella burnetii, which is the only known structure of a monomeric QOR (PDB entry 3tgh; M. C. Franklin, J. Cheung, M. Rudolph, M. Cassidy, E. Gary, F. Burshteyn & J. Love, unpublished work). No significant sequence identity is observed between the region corresponding to the transit peptide of ceQORH and the QOR sequences of known structures. Recently, ceQORH has also been reported to bind calmodulin (CaM; Dell'Aglio et al., 2013), as does IEP32 (Chigri et al., 2006), a property that has never been reported for other QORs.

In order to unravel the structural characteristics of the uncleaved transit peptide and to obtain insights into the substrate specificity and binding of CaM to ceQORH, we undertook structural studies of this protein. Here, we report the cloning, purification and crystallization of ceQORH from *Arabidopsis thaliana* and data collection from crystals of the apo form. Analytical ultracentrifugation studies are also discussed.

2. Materials and methods

2.1. Macromolecule production

2.1.1. Protein expression and purification. The AT4G13010 cDNA coding for a 329-amino-acid protein was inserted into a pET-30a+ vector to transform E. coli Rosetta2 (DE3) cells. The cells were grown overnight in 20 ml LB with 30 μ g ml⁻¹ kanamycin and 34 μ g ml⁻¹ chloramphenicol at 37°C. This first culture was used to inoculate 2 l LB (with antibiotics). Growth was continued at 37°C. When the OD₆₀₀ reached 0.6, the temperature was decreased to 20°C and 0.4 mM isopropyl β -D-1-thiogalactopyranoside (IPTG) was added. After overnight induction, the bacteria were harvested by centrifugation at 5500g for 30 min at 4°C. The cell pellet was suspended in 50 ml lysis buffer [50 mM Tris–HCl pH 7.5, 1 mM EDTA, 1 mM

Table 2 Crystallization.

Method	Hanging drop
Plate type	Linbro plate
Temperature (K)	293
Protein concentration (mg ml ⁻¹)	5
Buffer composition of protein	50 mM Tris-HCl pH 7.5, 200 mM KCl,
solution	2 mM DTT, 1 mM EDTA, 10% glycerol
Composition of reservoir	0.2 M sodium acetate, 0.1 M Tris-HCl
solution	pH 8.5, 32%(v/v) PEG 4000
Volume and ratio of drop	$2 \mu l + 2 \mu l$
Volume of reservoir (ul)	500

DTT, 10%(v/v) glycerol]. A cOmplete protease-inhibitor cocktail tablet (Roche) and, after lysis, 0.2%(w/v) streptomycin sulfate were added. The lysate was centrifuged at 14 000g for 30 min at 4°C. The purification was performed at 4°C. The supernatant was applied onto a DEAE Sepharose column in 50 mM Tris-HCl pH 7.5, 5 mM DTT, 1 mM EDTA, 10%(v/v) glycerol. ceOORH displays a high theoretical isoelectric point (pI = 9.05) and thus eluted in the flow through. It was collected, concentrated and applied onto an SP Sepharose column. ceQORH was eluted with a linear gradient to 50 mM Tris-HCl pH 7.5, 1 M KCl, 5 mM DTT, 1 mM EDTA, 10%(v/v) glycerol. The fractions containing ceQORH eluted between 150 and 180 mM KCl. They were pooled and concentrated with an Amicon Ultra 4 ml centrifugal filter with a 10 kDa membrane cutoff; they were then loaded onto a HiLoad 16/60 Superdex 200 column and eluted with 50 mM Tris-HCl pH 7.5, 200 mM KCl, 2 mM DTT, 1 mM EDTA, 10%(v/v) glycerol. The pool containing pure ceQORH was concentrated and frozen in liquid nitrogen before use. Macromolecule-production information is summarized in Table 1.

2.1.2. Analytical ultracentrifugation. The analytical ultracentrifugation experiment was performed with an Proteome-Lab XL-I analytical ultracentrifuge from Beckman Coulter (Palo Alto, USA) and an An-60 Ti rotor (Beckman Coulter). The cells had a double sector (Nanolytics) with optical routes of either 12, 3 or 1.5 mm and were equipped with Sapphire windows. Absorbance and interference measurements were performed with the ProteomeLab XL-I v.6.0 software. The samples were prepared at 0.6 and 6.0 mg ml⁻¹ in 10 mM Tris-HCl pH 7.5, 150 mM NaCl (AUC buffer) supplemented with NADPH (1.45 mM) as indicated in Supplementary Table S1. The interference and absorbance were recorded for 12 h at 50 000 rev min⁻¹ at 20°C. ceQORH was considered to be a compact spherical protein (molecular weight 34 304 Da) with $\overline{v} = 0.745 \text{ ml g}^{-1}$, $f/f_{\text{min}} = 1.25$, $\rho = 1.005 \text{ g ml}^{-1}$ and $\eta = 1.02 \text{ cP}$ at a temperature of 20°C. Using these parameters, the sedimentation coefficient was estimated by SEDNTERP assuming several oligomerization states (Supplementary Table S2). Data analysis was performed with SEDFIT v.14.c.

2.2. Crystallization

ceQORH at 5 mg ml $^{-1}$ in 50 mM Tris–HCl pH 7.5, 200 mM KCl, 2 mM DTT, 1 mM EDTA, 10%(v/v) glycerol was subjected to crystallization in its apo form using the sitting-

drop vapour-diffusion technique and the high-throughput crystallization facility at EMBL, Grenoble at 4°C. Crystallization hits were optimized using Linbro plates at 20°C. Crystallization information is summarized in Table 2.

2.3. Data collection and processing

Diffraction data were collected on beamline FIP-BM30A (Roth *et al.*, 2002) at the European Synchrotron Radiation Facility (ESRF), Grenoble, France at a wavelength of 0.9800 Å at 100 K using an ADSC 315r detector. The crystal-to-detector distance was 412.20 mm. The data were processed and scaled using *XDS* (Kabsch, 2010).

3. Results and discussion

3.1. Purification and analytical ultracentrifugation

After overnight expression, ceQORH was found mainly in the soluble fraction. Most of the contaminants were eliminated using a DEAE Sepharose column (Fig. 1). Two peaks of ceQORH with different OD_{260}/OD_{280} ratios were obtained from the SP Sepharose column, suggesting that ceQORH in one of these peaks is bound to nucleic acids or other absorbing molecules. The peak with the lower OD_{260}/OD_{280} ratio was selected and subjected to gel filtration. Pure protein was obtained with a yield of 14 mg per litre of culture. Analysis of the AUC sedimentation results for apo ceQORH showed numerous peaks between 2 and 10 S (Fig. 2a) corresponding to monomer, dimer and tetramer and possible aggregation at the two protein concentrations. When NADPH was added, only one peak at 2.7 ± 0.1 S ($s_{20,w} = 2.8$ S) was observed and the frictional ratio was 1.27. The theoretical frictional ratio

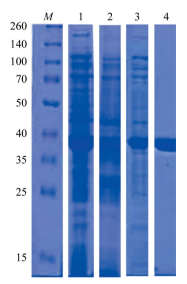


Figure 1
Purification of recombinant A. thaliana ceQORH. Proteins were separated on a 12% SDS-PAGE stained with Coomassie Brilliant Blue R-250. Lane M, molecular-weight markers (labelled in kDa); lane 1, crude soluble extract; lane 2, flow through from the DEAE Sepharose column containing ceQORH; lane 3, pool of the fractions containing ceQORH eluted from the SP Sepharose column; lane 4, fraction containing ceQORH eluted from the gel-filtration column.

a globular monomeric protein is 1.25 (Fig. 2b). Therefore, ceQORH–NADPH is monomeric (Supplementary Table S3). The results suggest that ceQORH undergoes a conformational change upon NADPH binding, leading to a transition from a dimer or a tetramer to a monomer.

3.2. Crystallization and data collection

Crystallization trials with ceQORH–NADPH failed even though this enzyme form was monodisperse in solution. These results suggest that ceQORH bound to NADPH is probably flexible and prevents crystal nucleation and growth. This is a difference from other QORs, since crystals of QOR from *E. coli* or *S. cerevisiae* bound to NADP+ or of QOR from *C. burnetii* bound to NADPH were obtained. Surprisingly, apo ceQORH crystallized in the presence of 0.2 *M* sodium acetate, 0.1 *M* Tris–HCl pH 8.5, 32%(v/v) PEG 4000, as according to the AUC result apo ceQORH has different oligomeric states in solution. A complete X-ray diffraction data set was collected to 2.34 Å resolution from a crystal of apo ceQORH (Fig. 3). The crystals belonged to the orthorhombic system, with unit-cell parameters a = 44.72, b = 142.65, c = 209.89 Å.

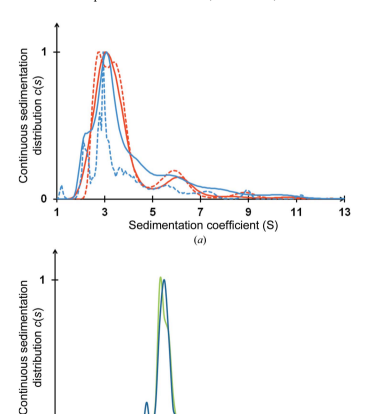


Figure 2
(a) Sedimentation curves of ceQORH calculated from absorbance (solid line) and interference (dashed lines) measurements for two ceQORH concentrations: 6 mg ml⁻¹ (red) and 0.6 mg ml⁻¹ (blue). (b) Sedimentation curves of ceQORH-NADPH calculated from absorbance (green) and interference (blue) experiments (ceQORH concentration 6 mg ml⁻¹; NADPH concentration 1.45 mM).

Sedimentation coefficient (S)

(b)

5

2

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Table 3
Data collection and processing.

Values in parentheses are for the outer shell.

Diffraction source	FIP-BM30A, ESRF
Wavelength (Å)	0.9800
Temperature (K)	100
Detector	ADSC 315r
Crystal-to-detector distance (mm)	412.20
Rotation range per image (°)	0.75
Rotation range (°)	180
Mosaicity (°)	0.13
Space group	$C222_{1}$
$a, b, c \stackrel{\circ}{(A)}$	44.72, 142.65, 209.89
α, β, γ (°)	90, 90, 90
Resolution range (Å)	41.82-2.34 (2.49-2.34)
Total No. of reflections	172766 (13536)
No. of unique reflections	28220 (4078)
Completeness (%)	97.9 (88.9)
Multiplicity	6.12 (3.31)
$\langle I/\sigma(I)\rangle$	13.03 (2.62)
$R_{\rm meas}$ (%)	13.20 (45.30)

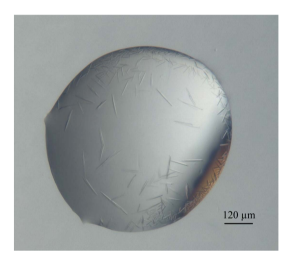


Figure 3 ceQORH crystals obtained at 4° C using 0.2 M sodium acetate, 0.1 M Tris–HCl pH 8.5, 32% PEG 4000 with the protein at 5 mg ml⁻¹.

Data statistics are summarized in Table 3. Systematic extinctions along the 00l axis showed that the crystals belonged to space group $C222_1$. Solvent-content analysis suggests that two molecules are present in the asymmetric unit, with a solvent content of 45.8%. Analysis of the self-rotation function reveals the presence of a peak corresponding to a noncrystallographic twofold axis ($\kappa = 180, \omega = 55, \varphi = 155^{\circ}$). Therefore, a dimer of apo ceQORH is present in the asymmetric unit.

Ligand screening to study the activity of the protein has been undertaken. Combined with X-ray crystallography, these studies should provide new insights into the ligand-binding specificity, protein–protein interaction (CaM binding) and the structure of the internal transit peptide sequence.

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