A bacterial chloroform reductive dehalogenase: purification and biochemical characterization

Bat-Erdene Jugder,1* Susanne Bohl,1,2 Helene Lebhar,1 Robert D. Healey,3 Mike Mane,1 Christopher P. Marquis1,4* and Matthew Lee1

1 School of Biotechnology and Biomolecular Sciences, University of New South Wales, Sydney, NSW 2052, Australia.
2 Department of Biotechnology, Mannheim University of Applied Sciences, 68163 Mannheim, Germany.
3 School of Chemistry, University of New South Wales, Sydney, NSW 2052, Australia.

Introduction

Environmental pollution with organohalides mainly caused by unregulated use and discharge from industrial sites is a serious public health problem due to their recalcitrance and hazardous impact on all ecosystems. Organohalide-respiring bacteria (ORB) and their reductive dehalogenase (RDase; EC 1.97.1.8) enzymes represent promising bioremediation solutions to these pollutants, as organohalides are used as terminal electron acceptors for growth of ORB. During the reductive dehalogenation reaction catalysed by the RDase, one halogen atom is replaced by one hydrogen atom, yielding a lesser halogenated product which is in many cases less toxic and less recalcitrant to further dehalogenation. RDases are membrane-associated redox metalloproteins with a corrinoid and two iron-sulphur clusters as cofactors (Holliger et al., 1998; Jugder et al., 2015). Several RDases have been studied from obligate (such as, Dehalobacter and Dehalococcoides strains) and non-obligate ORB (such as, Desulfitobacterium and Sulfitospirillum strains) (Jugder et al., 2015, 2016a). Most RDases derived from Dehalococcoides members, with the exception of CbrA from D. mccartyi CBDB1, were reported to dechlorinate chlorinated ethenes (Magnuson et al., 1998, 2000; Muller et al., 2004; Jugder et al., 2016a), whereas Dehalobacter RDases exhibited dechlorination activity towards chlorinated methanes, ethanes and ethenes (Schumacher et al., 1997; Tang and Edwards, 2013; Jugder et al., 2016a). The RDases reported from Desulfitobacterium strains mostly have substrate specificity for chlorinated ethanes and ethenes as well as ortho-chlorophenols (van de Pas et al., 1999, 2001; Boyer et al., 2003; Thibodeau et al., 2004; Bisailon et al., 2010).

Trichloromethane, commonly referred to by its trivial name chloroform (CF), is a hydrophobic (partition coefficient LogP_{ow} of 1.97 at 20°C) and volatile (vapour pressure of 197 mm Hg at 25°C) organohalide used as an industrial solvent for organic materials and as a chemical intermediate for the production of PTFE and the refrigerant mono-chloro-difluoro-methane (HCFC-22) (Verschueren, 1985; Jugder et al., 2016a). The low water solubility (< 8 g l^{-1} at 25°C), high density (ρ = 1.48 g cm^{-3}) and long half-life (3400 years at pH 7) contribute to its recalcitrance (Mabey and Mill, 1978; Mackay et al., 1980). The US-EPA has classified CF as a probable human
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The period the cultures in 2-l bottles were closely monitored for their CF consumption and DCM production activity by gas chromatography (GC). The total amount of CF respired was approximately 60 mmol over the 5 month period. During the final days prior to harvest, CF dechlorination rate was $6.5 \times 10^{-11}$ mol cell$^{-1}$ day$^{-1}$ and the cell density was $(9.8 \pm 4.3) \times 10^7$ cells ml$^{-1}$. Before harvesting, we examined the expression of tmrA gene, as it was previously shown that the same gene is expressed upon CF respiration (Jugder et al., 2016b). To confirm the induction of tmrA at the gene level, transcription of tmrA gene was investigated. Qualitative gene-expression analyses by RT–PCR demonstrated that the tmrA amplified from the cDNA synthesized from the cells actively respiring with CF was highly expressed, as the band intensities appear to reflect on the template abundance (Fig. S1). The amplicon obtained with the tmrA_F and tmrA_R primers was purified and sequenced. A nucleotide sequence of about 190 bp was obtained for the PCR product, showing a 100% sequence identity to the tmrA gene of Dhb sp. UNSWDHB.

Purification of chloroform reductive dehalogenase, TmrA

To purify TmrA, the cells grown with acetate, H$_2$ and CF were anoxically harvested and disrupted mechanically. CF-reductive dehalogenating activity was predominantly observed in the membrane fraction after ultracentrifugation of cell lysate. TmrA was solubilized from the membrane in the presence of 1% Triton X-100 and purified 23-fold by anion exchange chromatography using Mono Q HR as stationary phase (Table 1). The eluted fractions were tested instantly for dechlorination activity and CF-reductase activity was recovered at approximately 200 mM NaCl as a single peak. The enzyme was purified to apparent homogeneity with an average specific activity of $(1.27 \pm 0.04) \times 10^9$ units per mg protein in the methyl viologen-dependent assay (Table 1).

The MALDI-TOF analysis of purified native TmrA revealed an intact mass detection of $44,511$ Da, corresponding to the apparent molecular mass from the

| Purification | Total activity (U$^a$) | Yield (%) | Total protein (mg$^b$) | Specific activity (10$^9$ U mg$^{-1}$) | Purification factor |
|--------------|------------------------|-----------|------------------------|----------------------------------------|--------------------|
| Cell extract | 458                    | 100       | 8.40                   | 0.05                                   | 1                  |
| Membrane fraction | 376 | 82        | 3.03                   | 0.12                                   | 2                  |
| Solubilized membrane fraction | 348 | 76        | 0.629                  | 0.55                                   | 10                 |
| Mono Q HR | 172                    | 38        | 0.135                  | 1.27                                   | 23                 |

a. Amount (nmol) DCM produced per min with CF as substrate.
b. Total protein was determined with Pierce 660 nm protein assay using bovine serum albumin as a standard.
isolated band found in SDS-PAGE (Fig. 1). After purification and SDS-PAGE separation, the mass band at 45 kDa was excised from the polyacrylamide gel and sequenced by proteolytic degradation and LC-MS/MS peptide identification. The sequence of a full-length TmrA contains a Tat-signal peptide at its N-terminus; however, in this analysis the N-terminal detected peptide sequence started at amino acid number 58, suggesting N-terminal cleavage prior to translocation to the periplasmic membrane. Using the PRED-TAT program (Bagos et al., 2010) suggests that the most likely cleavage site is between residue 53 (Ala) and residue 54 (Gly). As such, the expected mass of the intact protein without the signal peptide is 44 522.8 Da (calculated using ProtParam-ExPaSy), which very slightly differs to the mass detected by MALDI-TOF.

Biochemical properties of chloroform reductive dehalogenase

The pH dependency of TmrA activity was tested in the standard assay reaction mixtures (30°C) containing 4 μg of the purified enzyme, 2 mM titanium (III) citrate and 2 mM methyl viologen in 100 mM Tris–HCl buffer at various pH values (5.5–8.3). The pH optimum of TmrA enzyme was found to be 7.2 (Fig. 3A). The optimum temperature for dechlorinating activity was evaluated by testing activity between 25 and 70°C (pH 7.5), which resulted in 45°C being found to be the optimum value (Fig. 3B). A substantial decrease in enzyme activity was observed at temperatures higher than 55°C.

The substrate range of the purified TmrA is shown in Table 2. The highest specific activity was observed with CF, followed by 1,1,2-TCA, 1,1,1-TCA and 1,1-DCA. The substrate specificity of the purified enzyme is in agreement with that reported for the Dhb. sp UNSWDHB cell extracts (Wong et al., 2016). This confirms that a single enzyme is responsible for the dechlorination of these organohalides by this bacterium under the experimental condition used.

According to the Michaelis–Menten model, the apparent $K_m$ value of the purified enzyme for CF was $154 \pm 41 \text{ μM}$ and the apparent $V_{max}$ value was $(1.25 \pm 0.09) \times 10^3 \text{ nmol min}^{-1} \text{ mg of protein}^{-1}$ for CF.
at a methyl viologen concentration of 2 mM. The highest initial activity of \((1.28 \pm 0.04) \times 10^3 \text{ nmol min}^{-1} \text{ mg of protein}^{-1}\) was measured at 480 \(\mu\text{M}\) of CF. A decrease in the dechlorination rate was observed when substrate concentrations exceeding this value were used. The reaction kinetics modelled to include substrate inhibition (with an inhibition constant \((K_i)\) of 354 \(\mu\text{M}\)) fitted experimental data better \((R^2 = 0.998)\) than the Michaelis–Menten model \((R^2 = 0.971)\) over the whole substrate concentration range examined (Fig. 4).

The native protein presents a broad absorption maximum between 400 and 500 nm and a shoulder at 310 nm (Fig. 5). The shape and location of these bands are typical of iron–sulphur proteins, including previously isolated reductive dehalogenases (Neumann et al., 1996; Schumacher et al., 1997; Christiansen et al., 1998). This is further substantiated by the absorption maximum at around 420 nm, which is assigned to the \([4\text{Fe-4S}]\) clusters. The absorbance in the 450–475 nm region as well as the shoulder at 310 nm could indicate the presence of a cobalamin in

Table 2. Substrate specificity profile of purified TmrA.

| Substrate                  | Product(s)                  | Mean specific activity \((10^3 \text{ nmol min}^{-1} \text{ mg of protein}^{-1})\) |
|---------------------------|-----------------------------|--------------------------------------------------------------------------------|
| Chloroform                | Dichloromethane             | 1.27 ± 0.04                                                                    |
| 1,1,2-trichloroethane     | 1,2-dichloroethane, vinyl chloride | 1.15 ± 0.01                                                                    |
| 1,1,1-trichloroethane     | 1,1-dichloroethane, chloroethane | 0.12 ± 0.01                                                                    |
| 1,1-dichloroethane        | Chloroethane                | 0.03 ± 0.01                                                                    |

The mean values of duplicates are presented with standard deviations.

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the 2+ oxidation state (Neumann et al., 1996; Christiansen et al., 1998; Miles et al., 2015). Moreover, the lack of peaks at 385 nm or at 360 nm, characteristic for cob(I)alamin or cob(III)alamin, respectively, suggests the presence of cobal(II)amin in the dehalogenase, which is indeed a characteristic of many corrinoid proteins (Banerjee, 1999).

Discussion

*Dehalobacter* sp. strain UNSWDHB is able to use CF as a terminal electron acceptor for growth. It has previously been demonstrated that the enzyme directly responsible for reductive dechlorination of CF is the membrane-associated TmrA, which is specifically overproduced in response to CF (Jugder et al., 2016b; Wong et al., 2016). In this study, we for the first time report the production, purification and biochemical characterization of a chloroform-reducing enzyme. The TmrA dehalogenase was purified to near homogeneity from *Dehalobacter* sp strain UNSWDHB cells in the presence of CF. Most of the dechlorinating activity was found in the membrane fraction, as observed in most RDases purified previously, due to the translocation of the enzymes to the periplasmic membrane via TAT signal pathway (Jugder et al., 2015, 2016a). A simple purification protocol was developed, involving membrane fractionation and solubilization in the presence of Triton X-100 followed by a step gradient elution in anion exchange chromatography. The estimated specific activity of $1.27 \times 10^3$ nmol min$^{-1}$ mg protein$^{-1}$ is in the middle range of other native RDases reported previously (Jugder et al., 2016a), although it should be noted that there were differences in the activity assay methods employed among the studies (e.g., gas chromatography versus spectrophotometry, measurement of substrate dechlorination versus product formation, reaction conditions).

The purification of native RDases from wild-type ORB, especially from obligate respirers like *Dehalobacter*, is challenging due to the difficulty in obtaining sufficient biomass, oxygen intolerance of ORB and low water solubility of organohalides. Nevertheless, several RDases have been isolated from facultative and obligate ORB; some of them followed similar strategies used in this study. The PCE-degrading PceA from *Dehalobacter restrictus* was

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**Fig. 4.** CF dechlorinating kinetics of purified TmrA. Each point represents the initial rate of dechlorination determined by the concentration of DCM formed after 2 h incubation. Curves are nonlinear regression fitted to the Substrate Inhibition model (solid line) and the Michaelis–Menten model (dashed line), resulting from duplicates using GraphPad Prism v. 6.07. Error bars represent standard deviation.

**Fig. 5.** UV-visible absorption spectra of the purified TmrA. The purified enzyme (0.05 mg ml$^{-1}$) exhibits features consistent with inclusion of both [4Fe-4S] clusters (~420 nm) and cob(II)alamin (310 nm and 450–475 nm) as cofactors.
purified using membrane solubilization with Triton X-100, ultracentrifugation and anion exchanger (Schumacher et al., 1997). Van de Pas and co-workers purified the 3-chloro-4-hydroxyphenylacetate-dechlorinating CprA from Desulfitobacterium sp. sp. PCE-1 and the ortho-chlorophenol-degrading CprA from Desulfitobacterium dehalogenans, the PCE-degrading PceA from Desulfitobacterium sp. PCE-1 and from D. hafniense TCE-1 (van de Pas et al., 1999, 2001) using the same solubilization technique, followed by consecutive anion exchange steps. Other native RDases, such as the PCE, TCE, VC-degrading RDases from Dehalococcoides and Sulfurospirillum members, have been purified in a functional form employing similar approaches (Magnuson et al., 1998, 2000; Muller et al., 2004).

The molecular mass of the enzyme detected by MALDI-TOF analysis (44 511 Da) was in agreement with the 45 kDa-protein band on SDS-PAGE. This is in the same range of molecular mass of biochemically characterized RDases to date (30–64 kDa) (Jugder et al., 2016a). Although the predicted mass of the full-length protein devoid of the region containing the signal peptide (44 522 Da) differs by 11 Da, such difference can be attributed to the analytical techniques, as MALDI-TOF detects the intact protein including post-translational modifications, whereas LC-MS/MS detects digested peptide fragments and post-translational modifications were not included in the search. Previously, in our laboratory, TmrA was identified as putatively dimeric, with an approximate mass of 100 kDa using blue native polyacrylamide gel electrophoresis (BN-PAGE) coupled with LC-MS/MS (Wong et al., 2016). The purification of the native TmrA under reducing conditions may suggest dimer formation arises via intermolecular disulphide bonds.

The pH optimum at 7.2 (30°C) and temperature optimum at 45°C (pH 7.5) were in the range of values from previously reported similar enzymes (Boyer et al., 2003; Thibodeau et al., 2004; Bisaillon et al., 2010). It is worth mentioning that determined pH and temperature optima have to be regarded as solely local maxima. In case of process development, it would be justified determining the optimal parameters as pivoting function of both. The apparent K_m value of the purified enzyme for CF (149 μM) differs from the K_m value of the crude cell extract (30 μM), reported previously in our laboratory (Wong et al., 2016). The difference in K_m values between crude cell extract and the pure protein can be attributed to the surrounding environment of the protein (e.g., buffers, detergents, presence of other proteins and membrane components) that can affect the conformation of the protein. Higher concentrations of CF appear to inhibit the enzyme activity (K_i = 354 μM). Similar inhibition patterns caused by high substrate concentrations were observed with tetrachloroethene (PCE) and trichloroethene (TCE) dechlorinating enzymes (Neumann et al., 1996; Miller et al., 1998).

This study presents the first report on the isolation and biochemical characterization of a highly purified reductive dehalogenase enzyme that can catalyse the dechlorination of CF. Previous studies mainly reported multi-step purification of RDases, whereas in this study the TmrA enzyme was purified to apparent homogeneity using membrane separation and solubilization followed by a single-step liquid chromatographic purification. Kinetic parameters are reported for reduction in not only CF but three other substrates 1,1,1-trichloroethane (TCA), 1,1,2-TCA and 1,1 dichloroethene (DCA).

**Experimental procedures**

**Growth, maintenance and harvest of cells**

Dhb sp. UNSWDHB cells, previously isolated from a CF-respiring enrichment culture (Lee et al., 2012), were grown at 30°C anaerobically in 2-litre flasks containing defined mineral medium with hydrogen as the electron donor (Lee et al., 2012; Wong et al., 2016) and CF as the electron acceptor. The growth of the cells was monitored using cell counting under fluorescence microscopy. Glutaraldehyde and SYBR Green fluorescent stain (Thermo Fisher, USA) were used as the fixing and staining solution respectively. The stained cell suspensions were fixed onto a 1.5% agarose-coated microscope slide and were imaged on a fluorescent microscope OLYMPUS BX51WI (Olympus-Lifescience, USA). The medium and all buffers used were prepared using anaerobic conditions.

**Preparation of membrane fraction and purification of chloroform reductive dehalogenase**

Approximately, 1 g of wet biomass was thawed and resuspended in anoxic 100 mM Tris buffer, pH 7.5, with 5% (w/w) glycerol, 1 mM DTT and 150 mM NaCl and subjected to another round of centrifugation under the same condition. The cell pellets were stored anoxically at −80°C until further use.
were then allocated into 2 ml O-ring capped FastPrep® Lysing Matrix Tubes (MP Biomedicals, Solon, OH, USA) for cell disruption by bead beating at 50 Hz for 15 min using TissueLyser LT (QIAGEN, Hilden, Germany). Following a brief centrifugation to separate the beads, the crude cell lysate was transferred to a 12PA seal tube (Hitachi, Tokyo, Japan) and heat sealed using Tube Topper (Beckman, Palo Alto, CA, USA). The membrane fraction was separated at 140 000 g for 90 min at 4°C using CP100WX ultracentrifuge with rotor P90AT (Hitachi). Back in the anaerobic chamber, the supernatant was removed and the pellet was resuspended in 20 ml solubilization buffer (100 mM Tris, 10% (w/v) glycerol, 1 mM DTT and 1% (v/v) Triton X-100, pH 8.0) and agitated for 1 h at 4°C for solubilizing the membrane fraction. After a second ultracentrifugation under the same condition, the supernatant containing the solubilized membrane fraction was immediately applied to a Mono Q HR (1 ml) column equilibrated with anaerobic buffer A (50 mM Tris, pH 8.0, 0.1% Triton X-100, 10% glycerol and 1 mM DTT). Following washing with 20 column volumes (CVs) of buffer A, protein was eluted at a flowrate of 0.6 ml min⁻¹ with a linear gradient of 20 CVs from 0 to 300 mM NaCl in buffer A. Fractions with chloroform-reducing activity eluted in a peak at approximately 200 mM NaCl.

**Measurement of chlorinated methanes by gas chromatography**

To monitor optimal growth of *Dhb* sp. UNSWDHB cells, the chlorinated methanes in the cultures (reduction of CF to DCM) were monitored by headspace analysis using an Agilent Technologies gas chromatograph equipped with a flame ionization detector (GC-FID) and a GS-GasPro capillary column (60 × 0.32 mm), as described elsewhere (Lee et al., 2012). Inlet and detector temperatures were set at 250°C, the oven temperature was programmed as follows: 1 min at 100°C followed by a gradient of 25°C min⁻¹ increasing to 250°C where held for 5 min. The retention times of DCM and CF were 5.3 and 6 min, respectively, at a flowrate of 3 ml min⁻¹.

**Dehalogenase activity assay**

Standard enzyme assays were performed anoxically in the anaerobic chamber, as previously described with the following modifications (Grostern et al., 2009). Briefly, in 2 ml glass screw cap glass vials 1.5 ml of 100 mM Tris–HCl buffer (pH 7.5) containing 2 mM titanium (III) citrate and 2 mM methyl viologen was added. To each vial was then added 100 µl of crude protein extract or purified protein sample. The vials were then capped and CF with a final concentration of 0.5 mM was injected through the septum, leaving no headspace. The vials were incubated at 30°C inside the chamber for 5 h. After incubation, the enzymatic reaction was quenched by transferring 1 ml of each reaction mixture to a 10 ml headspace flask containing anhydrous sodium sulphate (0.5 g) and 1 M sulphuric acid (1 ml). The headspace vials were then analysed by a Shimadzu 2010 GC-FID equipped with a PAL Systems headspace auto-sampler (PAL LHS2-xt-Shim) with GS-Q capillary column (30 m × 0.320 mm) as stationary phase. Separation was carried out at 100°C for 1 min followed by 5 min ramping with 20°C min⁻¹ to reach a final temperature of 250°C. Helium with a flowrate of 30 ml min⁻¹ was used as the carrier gas. CF and DCM standards (0 mM, 0.5 mM, 1.0 mM, 1.5 mM and 2.0 mM) were prepared on the same day from 8 mM stock solutions in the same buffer and treated the same way as the samples. Enzyme specific activity was defined as the formation of 1 nmol of DCM per mg of protein per minute. As negative controls, heat-deactivated crude protein extracts (prepared by incubating at 80°C for 15 min) as well as no CF mixtures were assayed in preliminary tests. Standard activity assays were performed in triplicate.

**Analytical methods**

The total protein concentration was measured using a Pierce 660 nm protein assay kit (Thermo Fisher Scientific, Rockford, IL, USA). Proteins were separated on Bolt™ 4–12% Bis-Tris Plus polyacrylamide precast gels (Thermo Fisher Scientific) with the SeeBlue Plus2 Pre-stained Standard (Thermo Fisher Scientific) as a molecular weight marker and stained with GelCode Blue Stain Reagent (Thermo Fisher Scientific). Protein bands on SDS-PAGE were cut from the gel and digested in-gel with trypsin as previously described (Hellman et al., 1995) for mass spectrometry analysis. The tryptic peptides were separated by nano-LC using an Ultimate 3000 HPLC and autosampler system (Dionex, Amsterdam, Netherlands) using a micro C18 pre-column (500 µm × 2 mm; Michrom Bioresources, Auburn, CA, USA). Mass spectra were recorded on an Orbitrap Velos (Thermo Electron, Germany) mass spectrometer equipped with a nanoelectrospray ionization source. Peak lists were generated using Mascot Daemon/extract_msn (Matrix Science, London, UK) using the default parameters and submitted to the database search program Mascot (version 2.2; Matrix Science). Mass spectra were searched against a custom database consisting of all proteins in the genome of *Dhb* sp. UNSWDHB.

The molecular mass of the purified TmrA was determined using MALDI-TOF/TOF. The purified detergent solubilized TmrA protein solution was subjected to reverse-phase C-18 ZipTip clean up prior to analysis as
per the manufacturer’s instructions. Briefly, a C18 ZipTip was equilibrated with acetonitrile/water/trifluoroacetic acid (2/98/0.1%, v/v/v). 10 µL of detergent solubilized TmrA was aspirated over the ZipTip several times. The bound analyte was washed with equilibration solution and then eluted with a saturated solution of sinapic acid in acetonitrile/water/trifluoroacetic acid (80/20/0.1%, v/v/v). The eluted analyte was spotted directly onto a polished steel MALDI-TOF target plate (Bruker Daltonics, Leipzig, Germany) and subject to MALDI-TOF using a Bruker UltraflameXtreme MALDI-TOF/TOF.

UV-visible spectra of the purified dehalogenase were obtained with a Cary 60 UV-Vis spectrophotometer. Absorbance of the dehalogenase sample in 50 mM Tris, pH 8, containing 0.1% Triton X-100 and 10% glycerol was measured at between 300 and 600 nm in 1 cm quartz cuvettes.

The enzyme kinetic parameters, such as the apparent \( K_m \) and \( V_{\text{max}} \) values for CF, were determined under the standard assay conditions with a range of substrate concentration varying from 8 to 1500 µM. The parameters were calculated using GraphPad Prism v. 6.07, ‘Enzyme kinetics – Michaelis-Menten’ module (GraphPad Software, Inc, La Jolla, CA, USA). Inhibition constant (\( K_i \)) was determined by a nonlinear fit of experimental data to a substrate inhibition equation using the same software (‘Enzyme kinetics – Inhibition: Substrate inhibition’ module).

The optimum pH of the enzyme was determined by carrying out standard enzyme assays at different pH values between 5.5 and 8.3 in 100 mM Tris–HCl buffer at 30°C. The temperature dependence was tested by varying the incubation temperature between 25°C and 70, with 5°C increments, at pH 7.5. The average values of triplicates were used to create the graphs by GraphPad Prism v. 6.07.

The substrate range was tested on 1,1,2-TCA, 1,1,1-TCA, and 1,1-DCA under the standard activity assay conditions, with 1.5 mM of respective substrates.

**Semi-quantitative reverse transcription RT-PCR analyses**

Total RNA for RT-PCR from *Dhb* sp. UNSWDHB grown in the absence and presence of CF was isolated using RNeasy Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions, including on-column DNase digestion with the QIAGEN RNase-Free DNase Set. The cDNA was synthesized from total RNA using the SuperScript VILO MasterMix kit (Life Technologies, Carlsbad, CA, USA), according to the manufacturer’s protocol. Equivalent amounts of RNA (100 ng) from each sample were subjected to RT reaction. A No-RT control was run with DEPC-treated water instead of template RNA. The primers specifically designed for *tmrA* gene (NCBI GI number: 530294411), *tmrA_F* (5’-TTTGGCCCA-GATTGGATA TG-3’) and *tmrA_R* (5’-CTTCACAGAGTCTAACATTGT-3’), were used in PCR reactions where annealing temperature of 57.6°C was applied. A no template control (NTC), replacing the template cDNA with DEPC-treated water and a no-RT were included as negative controls. For DNA amplification, 2× PCR Master Mix (Promega, USA) was used. The PCR products were separated via electrophoresis in a 1.8% (w/v) agarose gel and were subsequently cleaned up using DNA Clean & Concentra- tion and characterization of a novel CprA reductive dehalogenase specific to highly chlorinated phenols from *Desulfitobacterium hantiense* strain PCP-1, *Appl Environ Microbiol* 76: 7536–7540.

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**Conflict of interest**

None declared.

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Supporting information
Additional Supporting Information may be found online in the supporting information tab for this article:

**Fig. S1.** Transcriptional profiling of the *tmrA* gene in *Dehalobacter* sp. UNSWDHB by qualitative RT–PCR.

**Table S1.** Genetic characteristics of chloroform reductive dehalogenase proteins of *Dehalobacter* sp. UNSWDHB.

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