Development of a proteoliposome model to probe transmembrane electron-transfer reactions

Gaye F. White*, Zhi Shit, Liang Shit, Alice C. Dohnalkova†, James K. Fredrickson†, John M. Zachara†, Julea N. Butt*, David J. Richardson* and Thomas A. Clarke††

*Centre for Molecular and Structural Biochemistry, School of Biological Sciences and School of Chemistry, University of East Anglia, Norwich NR4 7TJ, U.K., and †Pacific Northwest National Laboratory, P.O. Box 999, 902 Battelle Boulevard, Richland, WA 99352, U.S.A.

Abstract
The mineral-respiring bacterium Shewanella oneidensis uses a protein complex, MtrCAB, composed of two decahem cyttochromes brought together inside a transmembrane porin to transport electrons across the outer membrane to a variety of mineral-based electron acceptors. A proteoliposome system has been developed that contains Methyl Viologen as an internalized electron carrier and valinomycin as a membrane-associated cation exchanger. These proteoliposomes can be used as a model system to investigate MtrCAB function.

MtrCAB is an outer membrane electron transporter
Bacterial mineral respiration requires electrons to be transported to the cell exterior in order to reduce extracellular insoluble electron acceptors. The proteins essential for these processes have been identified and studied in model mineral-respiring organisms such as Shewanella oneidensis [1,2]. These studies indicate that multihem proteins co-operate to conduct electrons from the cell interior across otherwise impermeable cell membranes, allowing them to react with external electron acceptors such as Fe(III) minerals [3–5]. The MtrCAB complex is responsible for electron transfer across the outer membrane of S. oneidensis. This complex contains the periplasmic decahem MtrA and membrane-associated decahem MtrC that is exposed on the cell surface. These two cytochromes meet and exchange electrons inside a transmembrane sheath, MtrB [6]. Structural studies of the MtrC homologue MtrF show that the ten haems are arranged in a staggered cross which would allow for bidirectional electron transport with one membrane-facing entry and three exit sites on the protein surface [7]. However, there is much debate over how such systems deliver charge to external redox partners with a number of potential electron-transfer mechanisms being proposed [7].

Reconstitution of transport proteins into phospholipid bilayers is a recognized way of studying biochemical and biophysical parameters in a model membrane environment [8,9]. In a previous study of MtrCAB, proteoliposomes incorporating the redox dye MV (Methyl Viologen) were reduced with dithionite [6]. The interior could only be reduced if the protein complex were present, demonstrating that MtrCAB has the ability to move electrons across an otherwise impermeable lipid bilayer. Since these earlier studies, this methodology has been tested and optimized with a view to wider applications that can facilitate detailed kinetic and mechanistic studies of interactions that involve electron transport through MtrCAB and, ultimately, be used to study other protein systems connected with transmembrane electron-transfer processes.

Efficient electron transfer through MtrCAB in proteoliposomes
Proteoliposomes offer the opportunity to model the electron-transfer pathway that occurs in a bacterial cell, where the encapsulated interior has a negative potential compared with the exterior environment and the only exit pathway for electrons is through the electron transporting protein complex. Thus, when a suitable electron acceptor is introduced into the exterior environment, electrons will flow out through the protein complex as they would in vivo. The MV inside the MtrCAB proteoliposomes is designed to act both as a means of storing charge in its reduced state that will readily be released to external electron acceptors and also as an integral redox indicator to follow electron-transfer reactions. The redox potential for the one-electron reduction of MV is $E_{\text{MV}^{2+}/\text{MV}^{+}} = -0.45$ V compared with the SHE (standard hydrogen electrode) at pH 7.0 and the colour changes from colourless to blue [10,11]. The redox potentials of Fe(III) oxides are typically between $-0.3$ V and $+0.06$ V compared with the SHE at pH 7.0 [12,13]. If there is a suitable pathway through the lipid bilayer, electrons should readily transfer from the encapsulated reduced MV to externally located electron acceptors such as soluble Fe(III) chelates or insoluble Fe(III) minerals.

However, to ensure that electron transfer occurs solely via the protein complex, it is imperative to ensure that the lipid bilayer remains impermeable to charged solutes for the duration of the experiment. The reduced proteoliposomes were centrifuged to sediment them and the supernatant...
Figure 1 | Improving the stability of proteoliposomes for electron-transfer investigations

(A) Schematic representation of the MtrCAB proteoliposomes. The MtrCAB complex is shown mediating electron transfer between oxidized and reduced forms of MV (MV$^{2+}$/MV$^{•+}$) and external electron acceptors or donors (X$^+$/X). Valinomycin (V) functions as a K$^+$-specific transporter. (B) MtrCAB proteoliposomes prepared with and without valinomycin were incubated with 100 μM sodium dithionite for 20 min, then centrifuged for 1 h at 280 000 g to sediment the proteoliposomes. Any MV released remained in the supernatant. From the absorbance profile, addition of 100 μM sodium dithionite to the supernatant from the valinomycin-free MtrCAB proteoliposomes showed that 10 μM MV had been released (broken line). Absorbance after addition of 100 μM sodium dithionite to the supernatant from MtrCAB proteoliposomes containing 10 nM valinomycin showed that less than 1 μM MV was in the supernatant (continuous line).

was analysed to determine whether any MV had been released. After 20 min of anaerobic incubation with dithionite followed by 1 h of centrifugation, approximately 30% of the MV was found in the supernatant. Thus the encapsulation of the MV was not maintained after the electron transfer, as may be expected when bulk charge imbalances build up across the lipid bilayer and different redox states of MV, including MV$^{2+}$, may be formed [14]. Valinomycin is known to form channels that selectively transport K$^+$ ions through lipid bilayers [15]. KCl and valinomycin were added to the system so that charge differentials caused by electron transport could be balanced by the migration of positively charged ions. This scheme (Figure 1A) led to greatly enhanced retention of MV inside the proteoliposomes with less than 3% leakage into the surrounding medium (Figure 1B).

Biophysical characterization of MtrCAB proteoliposomes

Proteoliposomes were prepared by detergent-mediated reconstitution of Triton X-100-solubilized MtrCAB into blank phosphatidylcholine liposomes. Freeze-dried phosphatidylcholine was resuspended in MV solution and subjected to extrusion, sonication and freeze–thaw cycles to produce separate regularly sized spheroids from unilamellar lipid bilayers impermeable to charged species. Various amounts of protein solution were added to aliquots of blank liposome suspension to give 0–50 nM MtrCAB. These were incubated with Biobeads for 1 h to remove detergent. Repeated wash steps were performed using centrifugation at 280 000 g to remove MV external to blank liposomes and proteoliposomes.

The physical attributes of the proteoliposomes were characterized using dynamic light scattering and transmission electron microscopy, showing they were separate regularly sized spheroids with an average diameter of approximately 500 nm (Figure 2). After purging with O$_2$-free N$_2$ and maintaining anaerobicity, a 10-fold dilution of blank liposomes was treated with sodium dithionite. The characteristic absorbance spectrum of reduced MV, MV$^{•+}$, was not observed until 1% Triton X-100 was added to disrupt the lipid bilayer and release the contents of the liposomes (Figure 3A). This shows that the liposomal membrane is impermeable to electron transfer and there was no diffusion of MV or sodium dithionite across the lipid bilayer during the time course of the experiment. When MtrCAB proteoliposomes were subjected to the same treatment, the absorbance spectrum of reduced MV was observed after sodium dithionite was added, showing that electrons were able to cross to the interior of the liposomes solely by conduction through the MtrCAB complex (Figure 3B). The absorbance at 600 nm increased and reached a steady state after approximately 10 min.

The MV content of the proteoliposome suspensions was determined using MV$^{•+}$ $\varepsilon$$_{600}$$^\lambda$ = 13 700 M$^{-1}$·cm$^{-1}$ [10]. The efficiency of MV encapsulation and reduction was inversely proportional to the amount of protein incorporated, with optimal efficiency at 1 nM MtrCAB. Estimates of the size and volume encompassed by the optimized proteoliposomes were used to estimate their number in relation to the amount of protein added. Protein incorporation was estimated by
Figure 2 | Physical characterization of proteoliposomes

(A) Transmission electron microscopy image of an MtrCAB proteoliposome. The inset shows the lipid bilayer of a prepared liposome. (B) Size distribution (by diameter in nm) of particles in a suspension of MtrCAB proteoliposomes measured by dynamic light scattering.

Figure 3 | Reduction of MV encapsulated in liposomes and proteoliposomes

(A) Absorbance profiles of liposomes containing no protein (continuous grey line), 10 min after addition of 100 μM sodium dithionite (continuous black line), followed by addition of 1% Triton X-100 (broken line). Absorbance with maxima at 400 and 600 nm corresponds to reduced MV. Reduction of MV was only observed after the liposome contents had been released by detergent. (B) Absorbance profiles of proteoliposomes prepared with 1 nM MtrCAB (continuous grey line) 10 min after addition of 100 μM sodium dithionite (continuous black line) followed by 1% Triton X-100 (broken line). Reduction of MV (27 μM) trapped within the proteoliposomes was observed on addition of sodium dithionite. After the trapped contents were released by the addition of 1% Triton X-100, further reduction was observed (32 μM reduced MV in total).

Western blots with MtrC-specific antibodies. It was found the most efficient reduction of the proteoliposome interior was observed when the molar ratio of protein to liposomes was approximately 10–200 MtrCAB molecules per liposome. In the optimized experiments, a total of 32 μM MV was released from the proteoliposomes by detergent, of which approximately 80% could be reduced directly by dithionite via MtrCAB. In the earlier published work, additions of 0.2 μM MtrCAB were used [6]. In this case, less than 1 μM MV was trapped inside the proteoliposomes, of which less than 50% was reduced directly by dithionite via MtrCAB [6]. By incorporating a lower ratio of MtrCAB than used in the previous work and using K⁺ and valinomycin to balance charge differentials, the proteoliposome methodology has been greatly improved.

Applications for proteoliposome methodology

The viability of the scheme shown in Figure 1(A) has been tested with MtrCAB as the electron transporter, MV on the interior and sodium dithionite on the exterior of the proteoliposomes. Under strictly anaerobic conditions, this produces stable proteoliposomes that encapsulate an interior with a negative potential that, apart from the MtrCAB electron-transport pathway, is isolated from the exterior.
environment. Ultimately, the aim is to introduce an electron acceptor into the medium surrounding the reduced proteoliposomes and spectroscopically follow electron transfer from inside to outside as MV$^{3+}$ reoxidizes to MV$^{2+}$. In the first instance, it will be important to ensure that there are no other redox-active chemicals or mediators present. Thus anaerobicity must be maintained and there must be no excess dithionite remaining in the medium surrounding the proteoliposomes. Once the system can be established under these conditions, the external electron acceptor can be varied to investigate the kinetics of MtrCAB-mediated electron transfer to a range of soluble and insoluble electron acceptors. Indeed, many other factors can be varied, such as having interior redox partners with different potentials, use of different methods to reduce the interior, addition of mediators to the external environment, and incorporation of different quantities and types of proteins within the system. This methodology promises to be generally applicable to investigations of electron transfer processes in S. oneidensis and other organisms that partake in extracellular electron transfer.

**Funding**

This work was supported by the Biotechnology and Biological Sciences Research Council [grant number H007288/1] and the Subsurface Biogeochemical Research program (SBR)/Office of Biological and Environmental Research (BER), U.S. Department of Energy (DOE), and is a contribution of the Pacific Northwest National Laboratory (PNNL) Scientific Focus Area.

**References**


Received 1 May 2012
doi:10.1042/BST20120116