Research Article
Cell-Free Expression of Unnatural Amino Acid Incorporated Aquaporin SS9 with Improved Separation Performance in Biomimetic Membranes

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Aquaporins (AQPs) are widely applied in biomimetic membranes for water recycling and desalination. In this study, a novel aquaporin was isolated from Photobacterium profundum SS9 (AQP SS9), which showed high water permeability and potential for practical water purification applications. To improve the stability of the AQP SS9 embedded biomimetic membranes, a modified AQP SS9 was obtained by incorporation of an unnatural amino acid (p-propargyloxyphenylalanine, pPpa) (P-AQP SS9) in vitro using a mutated Methanocaldococcus jannaschii tyrosyl-tRNAsynthetase (TyrRS) and the cell-free expression system. The modified AQP SS9 can covalently link with phospholipids and hence significantly improve the stability of biomimetic membranes. The concentration of Mg2+ and fusion expression with signal peptides were evaluated to enhance the expression level of P-AQP SS9, resulting in a highest yield of 49 mg/L. The modified AQP SS9 was then reconstituted into DOPC liposomes and analyzed by a stopped-flow spectrophotometer. The obtained water permeability coefficient (Pf) of 7.46×10−4 m/s was 5.7 times higher than that of proteoliposomes with the wild-type AQP SS9 (Pf=1.31×10−4 m/s) and 12.1 times higher than that of the DOPC liposomes (Pf=6.15×10−5 m/s). This study demonstrates the development of a cell-free system for the expression of membrane proteins with much higher stability and the potential application of the modified aquaporins for water filtration.

1. Introduction
Aquaporins belong to a large family of water-channel proteins, and the so-called orthodox aquaporins (AQPs) possess highly defined nanoscale pores to allow water molecules to rapidly pass through while retaining the dissolved solutes effectively [1]. The incorporation of AQPs into the lipid bilayer could enhance the permeability of reconstituted bilayer by an order of magnitude, while the high retention of solutes was still maintained [2]. Such ideal separation properties of aquaporins have led to an intensive interest in synthesizing aquaporin based high-performance biomimetic membranes, especially for desalination applications [3, 4].

Considering the high osmotic pressure and salinity conditions in practical applications, AQPs from the barophilic bacteria could be suitable and promising candidates for the fabrication of biomimetic membranes. Photobacterium profundum SS9 was isolated from Sulu Trough amphipod living in 2550 meters undersea [5, 6] and has been classified as a moderate barophilic bacterium [7]. Therefore, the aquaporin from P. profundum SS9 (AQP SS9) should have certain advantages in resisting high hydrostatic pressure, high salinity, and some other extreme conditions in water filtration and desalination applications.

The tyrosyl-tRNA synthetase (TyrRS) from Methanocaldococcus jannaschii and its corresponding tRNA (MjTyrRS/MjtRNA Tyr) have been found to be able to identify the amber codon (TAG) and transport tyrosine [8]. Further study [9] showed that the mutated MjTyrRS/MjtRNA Tyr could transport certain unnatural amino acids to the TAG...
2. Materials and Methods

2.1. Strains and Plasmids. E. coli DH5α was used for plasmid construction and E. coli BL21 (DE3) (Novangen, USA) as the host for making cell-free extract. pIVEX2.4c (Roche, Grenzacherstrasse, Switzerland) was used to construct cell-free expression vectors. pIVEX2.4c-AQP SS9, pETDuet-CK-T7, pUC-MjtRNA, p15a-MjpPpaRS, and pIVEX2.4c-MjpPpaRS were constructed in our previous studies.

2.2. Reagents. Restriction endonucleases, LA Taq DNA polymerase, and T4 DNA ligase were purchased from Takara Company (Dalian, China). All primers for PCR amplification were synthesized by Sangon Biotech (Shanghai, China) Co. Ltd. and listed in Table 1. The unnatural amino acid p-propargyloxyphenylalanine (pPpa) was provided by Amatek Chemical Company. Western-blot reagents were purchased from Beyotime Biotechnology (Shanghai, China).

2.3. Vector Construction. The amber mutations were introduced into the sites of F35TAG and L39TAG of AQP SS9 using the primer set AQP SS9TAG-F/AQP SS9TAG-R with pIVEX2.4c-AqpSS9 as the template, resulting in the "FIGURE 1: Schematic diagram of genetic incorporation of pPpa into AQP SS9."
the construction of pIVEX2.4c-AQP SS9TAG. As signal peptides (sp) were found to enhance the expression level of target proteins in cell-free system [10], six fusion expression plasmids including pIVEX2.4c-pelBsp-AQP SS9TAG, pIVEX2.4c-ompEsp-AQP SS9TAG, pIVEX2.4c-ompAsp-AQP SS9TAG, pIVEX2.4c-ompCsp-AQP SS9TAG, and pIVEX2.4c-phoAsp-AQP SS9TAG were constructed. The expression of the mutated AQP SS9 with different signal peptides including dsbEsp (GenBank ID BAL40591), ompEsp (GenBank ID BAL38095), ompCsp (GenBank ID BAL39006), pelBsp (GenBank ID AAA24848), and phoEsp (GenBank ID BAL37591) was carried out according to the previously reported method [11]. The signal peptide sequences were predicted using the Signal Peptide Database (http://proline.bic.nus.edu.sg/spdb/index.html) (Box 1).

Box 1: Signal peptides used in this study to enhance the expression of AQP SS9 in cell-free system. The peptidase cleavage sites Ala-X-Ala were underlined.

2.4. Expression and Purification of Proteins Using E. coli Derived Cell-Free Expression System. Two plasmids pUC-MjtRNA and p15a-MjtPPaRS were cotransformed into E. coli BL21 (DE3), which was used for the preparation of cell-free extracts following a previously reported protocol [12]. The cell-free extracts following a previously reported protocol [12]. The cell-free extracts were stored at -80 °C for future use. The E. coli cell-free expression system was set up accordingly with minor modifications [11, 12]. The unnatural amino acid pPpa was prepared as a stock solution of 50 mM and supplemented to the cell-free system at a final concentration of ~0.45 mM, unless specifically mentioned. 1% Brij58 was supplemented to the reaction mixture for soluble expression of the membrane protein. The cell-free reaction mixture was shaken at 400 rpm and 37°C (thermostat metal bath) for 4 h and then centrifuged at 12000 g for 10 min. The resulting supernatant was dialyzed 5 times with column buffer (20 mM Tris-HCl, pH 7.8, 300 mM NaCl, 10 mM imidazole, 0.05% Brij-78) and mixed with 1 mL Ni^{2+}-NTA resin and then shaken at 4°C and 120 rpm for additional 1 h. After rinsed with 5 mL column buffer, the target protein was eluted with 1 mL wash buffer (MOPS-KOH, pH 7.5, 300 mM imidazole, 0.05% Brij-78).

2.5. Preparation of AQP SS9 and P-AQP SS9 Proteoliposomes. To prepare AQP derived proteoliposome, 100 μg/mL AQP SS9 or P-AQP SS9, 4 mg/mL DOPC liposome and 0.25% Triton X-100 (m/v) [13] were mixed and shaken at 25°C, 120 rpm for 30 min. Then 0.2 g/mL SM-2 Bio-Beads (Sigma, St. Louis, MO, USA) were added to the mixture and shaken for additional 2 h. The reaction mixture was ultracentrifuged at 300000 g and 4°C for 15 min, and the precipitates were washed and resuspended with MOPS-KOH buffer (100 mM, pH 7.5) to get the AQP SS9 or P-AQP SS9 proteoliposomes.

2.6. Water Permeability Assay of Reconstituted P-AQP SS9 Proteoliposomes. The water permeability of DOPC liposome, AQP SS9 proteoliposome, and P-AQP SS9 proteoliposome was measured by light scattering at 436 nm, 500 V, and 25°C using a stopped-flow spectrophotometer (SFM-300, BioLogic) [14]. An equal volume of liposome or proteoliposome solution (100 μg/mL) and hypertonic solution (100 mM MOPS-KOH, pH 7.5, 400 mM NaCl) was mixed and measured under the reported conditions [15].

3. Results

3.1. Phylogenetic Analysis of AQP SS9. Due to the lack of experimental data, a phylogenetic analysis was performed to study the evolutionary origins and the potential solute transport properties [16]. Some known bacterial orthodox aquaporins and aquaglyceroporins, as well as an animal (AQPI) and a plant (TIP) aquaporin, were aligned using Clustal Omega [17] and included in the phylogenetic tree. As shown in Figure 2, AQP SS9 showed the highest homology with the E. coli AqpZ [18], which has been characterized as an orthodox aquaporin and was not included in the aquaglyceroporin branch of the phylogenetic tree. Therefore, based on the phylogenetic analysis, AQP SS9 belongs to the orthodox aquaporin subfamily of water-channel proteins, which should transport water molecules efficiently and specifically.

3.2. Design of Unnatural Amino Acid Incorporation in AQP SS9. As the crystal structure of AQP SS9 has not been reported yet, the E. coli AqpZ with similar structure and high homology was selected for evaluation. Based on multiple sequence alignment (MSA) and analysis of the transmembrane region of AqpZ by TMHMM, as well as the
Figure 2: Phylogenetic analysis of AQP SS9. The amino acid sequences of aquaporin homologs were aligned and the phylogenetic tree was constructed. TIP, a plant aquaporin (NCBI Accession Number P25818.1); MgaGlpF, Mycoplasma gallisepticum aquaglyceroporin (NCBI Accession Number ADC30962.1); MgeGlpF, Mycoplasma genitalium aquaglyceroporin (NCBI Accession Number AAC71249.1); BsGlpF, Bacillus subtilis aquaglyceroporin (NCBI Accession Number AOT51422.1); EcGlpF, E. coli aquaglyceroporin (NCBI Accession Number NP_418362.1); AQP1, an animal aquaporin (NCBI Accession Number P29972.3); 6083AqpZ, Synechocystis sp. PCC6803 aquaporin Z (NCBI Accession Number P73809.1); EcAqpZ, E. coli aquaporin Z (NCBI Accession Number NP_415396.1).

Figure 3: Western-blot analysis of pPpa incorporated AQP SS9. Lane 1, pellet without pPpa; Lane 2, supernatant without pPpa; Lane 3, pellet with 2 μL pPpa; Lane 4, supernatant with 2 μL pPpa; Lane 5, pellet with 4 μL pPpa; Lane 6, supernatant with 4 μL pPpa; Lane 7, pellet with pPpa and pIVEX2.4c-MjpPpaRS; Lane 8, supernatant with pPpa and pIVEX2.4c-MjpPpaRS.

3.3. Optimization of the Expression of pPpa Incorporated P-AQP SS9. The pIVEX2.4c-AqpSS9TAG carrying amber mutations was expressed in E. coli cell-free expression system under different conditions. P-AQP SS9 expression could not be detected neither in the soluble nor in the insoluble fractions without the addition of pPpa (Figure 3, Lane 1 and Lane 2), while the supplementation of 2 μL pPpa (~0.45 mM) led to the expression of P-AQP SS9 (Figure 3, Lane 3 and Lane 4). These results indicated that MjpPpaRS/MjtRNA in the cell-free expression system could recognize the stop codon TAG and specifically incorporate pPpa into AQP SS9. Since further increment of pPpa up to 4 μL could not improve the amount of the recombinant protein (Figure 3, Lane 5 and Lane 6), 2 μL pPpa was set as the optimal amount for subsequent studies. Although MjpPpaRS has already been overexpressed in the cell-free extracts, supplementation of pIVEX2.4c-MjpPpaRS further increased the production of P-AQP SS9 (Figure 3, Lane 7 and Lane 8), indicating that the amount of MjpPpaRS was the bottleneck for efficient incorporation of pPpa into the recombinant protein. Notably, in agreement with previous studies, all the membrane proteins were expressed as the insoluble form in the cell-free system (Figure 3). To enable soluble expression of P-AQP SS9, 1% Brij58 was supplemented into the cell-free system in the following studies.

In addition, the effect of Mg$^{2+}$ on the expression of P-AQP SS9 was also investigated. Varied concentration of Mg$^{2+}$ (11.0, 13.8, 16.5, 19.2, 22.0, and 24.8 mM) was added to the cell-free expression system. Western-blot analysis showed that the highest expression level of P-AQP SS9 was achieved with 22.0 mM of Mg$^{2+}$, while very low level of protein expression was observed when Mg$^{2+}$ concentration was beyond the range from 11.0 mM to 22.0 mM (Figure 4).

3.4. Fusion Expression of P-AQP SS9 with Different Signal Peptides. The fusion of signal peptide at the N-terminus has been found to increase the expression level of recombinant proteins in cell-free system [11]. Therefore, multiple signal peptides including dsbAsp, malEsp, ompAsp, ompCsp, pelBsp, and phoAsp were chosen to further enhance the expression of P-AQP SS9. Among six signal peptides tested, dsbAsp enhanced the expression of P-AQP SS9 up to 49 mg/L, which was two times higher than that without any signal peptide.
peptide (Figure 5). Interestingly, signal peptides pelBsp and phoAsp decreased the expression level of P-AQP SS9 to 11 mg/L.

3.5. Water Permeability of P-AQP SS9 Reconstituted Proteoliposomes. The two types of proteoliposomes with AQP SS9 and P-AQP SS9 were individually prepared and analyzed for its water permeability. The $P_f$ of P-AQP SS9 proteoliposomes, AQP SS9 proteoliposomes, and control empty liposomes were $7.46 \times 10^{-4}$ m/s, $1.31 \times 10^{-4}$ m/s, and $6.15 \times 10^{-5}$ m/s, respectively (Figure 6). The water permeable rate of P-AQP SS9 proteoliposomes was 5.7 times higher than that of AQP SS9 proteoliposomes, both of which were much higher than that of the empty liposomes. Thus, the modified P-AQP SS9 presented better water filtration activity compared with the wild-type AQP SS9.

4. Discussion

The AQPs incorporated biomimetic membranes have demonstrated better water permeability and salt rejection ability [19], but the low strength and poor integrity of the membranes are the biggest challenges for practical applications. To solve these defects, we have successfully incorporated $pPpa$ into AQP SS9 to achieve the covalent-link between the membrane proteins and the phospholipids bilayer. In this case, the strength and integrity of AQP embedded biomimetic membrane could be enhanced.

Considering the impact of unnatural amino acids on the 3D structure and water permeability of AQPs as well as the cross linking efficiency between AQPs and phospholipids, the type of unnatural amino acid and its incorporation sites in AQP SS9 should be rationally designed. Based on the
modeling studies of AqpZ, which is highly homologous to AQP S9, the active sites and phospholipids-linking regions of AQP S9 were avoided for amino acid substitution, and finally F35TAG and L39TAG were selected as the mutation sites. On the other hand, since the covalent cross-link of acetylene group and phospholipids may improve the protein stability in the membrane, pPPa was chosen as the target unnatural amino acid to be incorporated into AQP S9.

To achieve our engineering goal, the M. jannaschii TyrRS mutant constructed in our previous studies was used for the synthesis of corresponding aminoacyl-tRNA, and the stop codon TAG was specifically introduced into the AQP S9 gene by site-directed mutagenesis. Then E. coli cell-free expression system containing the engineered TyrRS could specifically incorporate pPPa into the F35TAG and L39TAG sites of AQP S9. The expression level of P-AQP S9 in cell-free system was further improved through the optimization of the Mg²⁺ concentration (22.0 mM) and the fusion of signal peptide (dsbAsp), resulting in the production of 49 mg/L P-AQP S9.

To characterize the obtained P-AQP S9 and explore its possible applications, P-AQP S9 was reconstituted into liposomes and biomimetic membranes. The water permeability factor (P₁) of P-AQP S9 liposomes was 7.46×10⁻⁴ m/s, 5.7 and 12.1 times higher than that of natural AQP S9 and DOPC liposomes, respectively.

5. Conclusion
In this study, the M. jannaschii TyrRS mutant was used to incorporate the unnatural amino acid pPPa into AQP S9 (P-AQP S9) via the E. coli cell-free expression system. The expression level of P-AQP S9 in cell-free system was further improved by optimizing Mg²⁺ concentration and fusing with a signal peptide. Subsequent analyses showed that the water permeability of P-AQP S9 had been significantly improved compared with that of AQP S9.

Data Availability
The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest
The authors declare that they have no conflicts of interest.

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