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ABSTRACT

Integral membrane proteins (IMPs) constitute ~30% of all proteins encoded by the genome of any organism and Escherichia coli remains the first-choice host for recombinant production of prokaryotic IMPs. However, the expression levels of prokaryotic IMPs delivered by this bacterium are often low and overproduced targets often accumulate in inclusion bodies. The targets are therefore often discarded to avoid an additional and inconvenient refolding step in the purification protocol. Here we compared expression of five prokaryotic (bacterial and archaeal) IMP families in E. coli and Saccharomyces cerevisiae. We demonstrate that our S. cerevisiae-based production platform is superior in expression of four investigated IMPs, overall being able to deliver high quantities of active target proteins. Surprisingly, in case of the family of zinc transporters (Zrt/Irt-like proteins, ZIPs), S. cerevisiae rescued protein expression that was undetectable in E. coli. We also demonstrate the effect of localization of the fusion tag on expression yield and sample quality in detergent micelles. Lastly, we present a road map to achieve the most efficient expression of prokaryotic IMPs in our yeast platform. Our findings demonstrate the great potential of S. cerevisiae as host for high-throughput recombinant overproduction of bacterial and archaeal IMPs for downstream biophysical characterization.

1. Introduction

Integral membrane proteins (IMPs) mediate efficient and selective transport of organic and inorganic molecules across cellular membranes, and many signaling processes rely on membrane-embedded receptors (Cournia et al., 2015). IMPs are critical components of each cell and constitute ~30% of the proteome across all kingdoms of life. In humans IMPs represent the largest class of drug targets, accounting for ~60% of currently approved pharmaceuticals (Gong et al., 2019).

Access to high-resolution structures of the membrane proteome is required to comprehend the membrane biology field (Cournia et al., 2015). While more than 120,000 structures are deposited in the Protein Data Bank (PDB; http://www.rcsb.org/) (Burley et al., 2019), currently just above 1% represents IMPs (http://www.blanco.biomol.uci.edu/mpstruc/). This bias reflects the main challenges associated with membrane protein structural biology, i.e., obtaining high expression levels of prime quality IMPs, followed by development of purification procedures that preserve protein fold and activity (Pandey et al., 2016). Overall, this laborious task typically involves screening for optimal expression and purification conditions for a large number of orthologue proteins to achieve optimal yields of pure IMPs.

Structural studies of prokaryotic IMPs have almost exclusively relied on recombinant production in Escherichia coli (E. coli) (Schlegel et al., 2014), which is partly attributed to fast and cheap cultivation protocols. Furthermore, E. coli has been extensively studied with a large toolbox of biochemical methods and this host remains one of the best characterized organisms for recombinant gene expression (Gileadi, 2017; Dilworth...
Moreover, it is generally assumed that the expression host most closely related to the organism from which the desired protein originates is likely to deliver the protein sample in the highest amounts and the best quality (Bill et al., 2011). Hence, *E. coli* may intuitively provide the most suitable lipid environment for recombinant production of Gram-negative bacterial IMPs. In contrast, *E. coli* may not be optimal for IMPs from Gram-positive prokaryotes and in particular not for archaeal counterparts, as the latter are found in membranes with a lipid composition that differs significantly from those found in bacteria (Jain et al., 2014). Instead of fatty acids esterified to D-glycerol-3-phosphate, archaeal phospholipids are composed of repeating units of isopenone connected through an ether bond to L-glycerol-1-phosphate (Dowhan et al., 2008).

In the present paper we challenge the dogma that *E. coli* offers the most optimal expression platform as we hypothesized that *Saccharomyces cerevisiae* (*S. cerevisiae*) may possess a number of advantageous features beneficial for expression of bacterial and archaeal IMPs. First, we avoid accumulation of overproduced IMPs in inclusion bodies, hereby circumventing the requirement for protein refolding, a laborious and inefficient recovery step, often impossible due to the particularly complicated nature of this class of proteins, which often results in targets being discarded at this initial stage (Schlegel et al., 2014; Lyons et al., 2016). Secondly, yeast possess a more advanced machinery for IMPs biogenesis that may be superior to that found in *E. coli* (Freigassner et al., 2009).

To investigate the capacity for expression of IMPs in *E. coli* and *S. cerevisiae*, we selected prokaryotic targets belonging to five protein families of Gram-negative, Gram-positive or archaeal origin and maintaining different functions (Table 1). We performed an extensive comparison between the most commonly used *E. coli* systems (Studier and Moffatt, 1986) and our previously described *S. cerevisiae* platform (Pedersen et al., 1996). Specifically, we compared to what extent recombinant proteins could be produced in the two hosts, extracted from the cell membranes, and purified in a stable and active form in high yields. Based on the obtained results, we highlight the experimental conditions that have to be considered to achieve optimal expression of bacterial and archaeal IMPs in *S. cerevisiae*. To our knowledge, such a study comparing prokaryotic targets belonging to *S. cerevisiae* has not been conducted before. Our findings demonstrate the potential of *S. cerevisiae* for overcoming issues with poorly overproduced bacterial and archaeal IMPs for downstream studies.

## 2. Materials and Methods

### 2.1. *S. cerevisiae* and *E. coli* strains

*S. cerevisiae* recombinant protein production was carried out in the PAPI500 strain (α ura3-52 trp1: GAL10-GAL4 lys2-801 leu2Δ1 his3Δ200 pep4:His3 prb1Δ1.6R can1 GAL) (Pedersen et al., 1996). BL21(DE3) (Studier and Moffatt, 1986), BL21(DE3)-pLyS5 (Studier and Moffatt, 1986; Studier, 1991), C41 (Miroux and Walker, 1996) and C43 (Miroux and Walker, 1996) strains were used as hosts for recombinant protein production in *E. coli*.

### 2.2. Plasmid construction

For yeast-based expression, the pEMBLyex4 vector was used (Cesarini et al., 1987). The *E. coli* - *S. cerevisiae* shuttle plasmid, pPAP10286, carrying the pET52b(+) expression cassette (Supplementary Figure 1) (Dowhan et al., 2008; Studier, 1991) was used for *E. coli* expression. Two sets of codon-optimized full-length cDNAs for each target were used, i.e., optimized for expression in *E. coli* and *S. cerevisiae*, respectively (GenScript, USA).

The cDNAs were amplified by PCR using primers listed in Supplementary Tables 1 and 2, and inserted into the pEMBLyex4 or pPAP10286 expression plasmid to encode proteins N-terminally fused to His10-GFP-TEV (cleavage site for Tobacco Etch Virus protease) or C-terminally fused to TEV-GFP-His10. All PCR reactions were performed using AccuPol DNA polymerase (Amplicon, Denmark). Both *S. cerevisiae* and *E. coli* expression plasmids were generated by homologous recombination in *S. cerevisiae* using the transformation protocol developed by Gietz and Schiestl (2007). PCR primers contained 5′-extensions designed for homologous recombination, hereby ensuring correct assembly of PCR fragments, and the SalI, HindIII- and BamHI-digested plasmids (Supplementary Figure 1). Transformed yeast cells were selected on minimal medium containing 2% glucose, and supplemented with leucine (30 mg/L) and lysine (20 mg/L). Correct assembly of expression plasmids was verified by DNA sequencing (Eurofins MWG Operon, Germany). *E. coli* expression plasmids were subsequently transferred to the designated *E. coli* expression strain. Plasmids encoding histidine-tagged zinc transporters (Zrt/Itt-like proteins, ZIPs) for expression in *E. coli* were generated by insertion of NdeI/XhoI digested PCR products into similarly digested pET15b or pET52b(+) vectors (Merck, Germany) and did not contain the GFP-tag.
<table>
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<tr>
<th>Name of the target</th>
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<th>UniProt ID*</th>
<th>Biological function</th>
<th>Total length (aa)</th>
<th>Topology (TMhelices)**</th>
<th>Length of N-term (aa) and orientation**</th>
<th>Length of C-term (aa) and orientation**</th>
<th>Oligomeric state</th>
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<td>Voltage-gated potassium channel</td>
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<td>6out</td>
<td>1out</td>
<td>Dimer (putative)</td>
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2.3. Expression screening and temperature optimization of protein production

E. coli pre-cultures were inoculated from frozen stocks, grown O/N at 30 °C in 50 mL of 2 x YT or LB medium supplemented with tetracycline and subsequently diluted to OD₆₀₀ = 0.05. At OD₆₀₀ = 0.6, half of the culture was transferred to 15 °C and the other half to 30 °C. After thermal equilibration, protein production was induced with a final concentration of 0.1, 0.5 or 1 mM of isopropyl-β-D-1-thiogalactopyranoside (IPTG), respectively.

Yeast pre-cultures were inoculated from frozen stocks in 5 mL of minimal medium containing leucine (30 mg/L) and lysine (20 mg/L), and grown at 30 °C until saturation (typically for 48 h). Subsequently, 200 µL of the culture was transferred to 5 mL of glucose minimal medium containing lysine, but lacking leucine, and grown for 24 h at room temperature (RT). Then 500 µL of the pre-culture was transferred to 50 mL of glucose minimal medium containing lysine, but lacking leucine, and grown for another 24 h at RT. The culture was used to inoculate 100 mL of expression media (YP medium containing 0.5% glucose and 3% glycerol) to OD₆₀₀ = 0.05. At OD₆₀₀ = 1.0, half of the culture was transferred to 15 °C and the other half to 30 °C. After thermal equilibration, recombinant protein production was induced by adding 10 mL of YP medium containing 20% galactose and 3% glycerol.

For both E. coli- and S. cerevisiae-based expression, OD₆₀₀ was measured at different time points up to 120 h post induction. For each time point, whole cell GFP fluorescence was measured in 1 OD₆₀₀ unit in a white micro plate in a Fluoroscan Ascent fluorometer (Thermo Scientific, USA) with excitation at 485 nm and emission at 520 nm.

2.4. Production of membrane proteins

Yeast cells were pre-cultured and grown at RT as described above. 2 L of YP supplemented with 0.5% glucose and 3% glycerol was inoculated to an OD₆₀₀ of ~0.05. At OD₆₀₀ = 1, the culture was transferred to 15 °C and galactose was added to a final concentration of 2%. Cells were harvested at their expression maximum as estimated by the expression screens.

Yeast cells from 2-L cultures were harvested at 1600 g for 10 min at 4 °C, at the protein accumulation optimum established by the expression screen. Cells were lysed by glass bead homogenization in ice-cold lysis buffer (1 M NaCl, 25 mM imidazole, 1 mM EDTA, 10% glycerol; pH 7.5) containing 1 mM phenylmethylsulfonyl fluoride (PMSF) and 1 µg/mL leupeptin, pepstatin and chymostatin (L, P, C) in a 50-mL Falcon tube. Cell lysates were centrifuged at 3000 g for 10 min at 4 °C to remove cell debris. Crude membranes were pelleted from the supernatant by ultracentrifugation at 160,000 g for 90 min at 4 °C (using a Sorvall SS-34 rotor; Thermo Scientific, USA). Crude membrane pellets were resuspended and homogenized in lysis buffer containing protease inhibitors and kept at –80 °C until use.

Protein concentration in crude membranes was determined by the bicinchoninic acid (BCA) assay according to specifications of the manufacturer (Sigma Aldrich, USA).

To normalize expression yields, GFP fluorescence in 25 µg of crude membrane proteins was measured in a Fluoroscan Ascent microplate fluorometer (Thermo Scientific, USA) as described above. Fluorescence was converted to pmol of TEV-GFP-His₆.tagged protein using a standard curve generated from the fluorescent signal of purified GFP mixed with crude membranes from PAP1500 yeast as previously established (Bombolt et al., 2013) and normalized to the molecular weight of the respective target protein to calculate its percentage of expression in isolated crude membrane fractions.

2.5. Small-scale crude membrane preparation and SDS-PAGE separation for comparing expression in S. cerevisiae and E. coli

Pairs of 1-L cell cultures were grown as described above at RT. At OD₆₀₀ = 0.6, one of the cultures was transferred to 15 °C and the other one to 30 °C, and induced with IPTG (E. coli) or galactose (S. cerevisiae). Subsequently, 25 mL of cell culture samples were collected at 0, 24, 48, 72, 96, 108 and 120 h after induction.

Yeast cells from 25-mL cultures were harvested at 1000 g for 10 min at 4 °C and the cell pellet was resuspended in 400 µL of ice-cold lysis buffer (25 mM imidazole, 1 M NaCl, 1 mM EDTA, 10% glycerol; pH 7.5) containing 1 mM PMSF and 1 µg/mL L, P, C. Cells were opened with glass beads homogenization in a Bertin Precellys 24 homogenizer (4 cycles of 15 s; Bertin Instruments, France) with 2-min cooling breaks. Glass beads were subsequently washed several times with ice-cold lysis buffer and the homogenate was cleared from cell debris by centrifugation at 3000 g for 10 min at 4 °C. Crude membranes were isolated from the supernatant by ultracentrifugation at 210,000 g at 4 °C for 20 min (using a Beckman TLA 100 rotor; Beckman, USA). Crude membrane pellets were resuspended in 200 µL of lysis buffer supplemented with protease inhibitors and stored at ~80 °C.

E. coli cells from 25-mL cultures were harvested at 6000 rpm for 20 min at 4 °C (using a Sorvall SS-34 rotor). Cell pellets were resuspended in 1 mL of ice-cold lysis buffer (25 mM imidazole, 1 M NaCl, 1 mM EDTA, 10% glycerol; pH 7.6) containing 1 mM PMSF and 1 µg/mL L, P, C, and cells were disrupted by sonication on ice for 15 min using pulsation with 5-sec intervals and amplitude 50% (using Sonopuls 3100 homogenizer; Bandelin, Germany). Cell debris was removed by centrifugation at 15,000 g for 10 min at 4 °C (using a Sorvall SS-34 rotor). Crude membranes were isolated from the supernatant by ultracentrifugation at 160,000 g for 20 min at 4 °C (using a Beckman TLA 100 rotor). Crude membranes were resuspended and homogenized in 500 µL of ice-cold lysis buffer supplemented with protease inhibitors and stored at ~80 °C.

Crude membranes were analyzed by SDS-PAGE prepared according to Laemmli, U. K. (1970) (Laemmli, 1970) and stained with Coomassie blue (R-250). Prior to electrophoretic analysis, crude membrane samples were denatured with sample buffer containing 4% SDS and protease inhibitors at RT for 20 min before loading. Imaging of both in-gel GFP fluorescence and Coomassie blue stained gels was performed using the Image Quant LAS-4000 system (GE Healthcare, USA).

2.6. Detergent screens

Crude membranes were incubated in solubilization buffer (25mM Tris-HCl, 10mM imidazole, 0.5 M NaCl, 10% glycerol; pH 7.6) supplemented with protease inhibitors (1 mM PMSF and 1 µg/mL L, P, C) at a protein:detergent ratio (w/w) of 1:3. The screens included the following detergents: 5-cyclohexyl-1-pentyl-β-D-maltoside (CY-S), n-dodecyl-β-maltopyranoside (DDM), n-decyl-β-D-maltopyranoside (DM), n-dodecylphosphocholine (FC-12), n-tridecylphosphocholine (FC-13), n-hexadecylphosphocholine (FC-16) and lauryldimethylamine N-oxide (LDAO). Detergents were purchased from Affymetrix, UK (as Analagre quality) or from Glycon Biochemicals, Germany. Solubilization was performed at slow rotation for 1 h at 4 °C. Unsolubilized material was removed by ultracentrifugation at 160,000 g for 20 min at 4 °C (using a Beckman TLA 100 rotor). GFP fluorescence in the resulting supernatant was monitored as described above and solubilization efficiency was estimated by normalization to the total GFP fluorescence in the sample prior to detergent treatment.

2.7. Fluorescence-detection size-exclusion chromatography

Solubilized crude membranes were analyzed by fluorescence-detection size-exclusion chromatography (FSEC) performed using a Superose 6 Increase 200 10/300 GL column (AqpZ) or a Superose 12 10/300 GL column (KcsA, NavAb, ClCec1) attached to an Äkta Purifier system (all from GE Healthcare, USA) coupled to a fluorescence detector (Shimadzu Prominance RF-20A; Shimadzu, Japan) to visualize the elution profile of the GFP-tagged proteins. All FSEC runs were conducted in 20 mM Tris-HCl, 0.15 M NaCl, 0.03% (w/v) DDM; pH 7.5.
2.8. Small-scale affinity purification

For small-scale purification of the target IMP-TEV-GFP-His\textsubscript{10} fusions, the choice of detergent was based on the results from the respective solubilization screens and FSEC analysis. Crude membranes isolated from the 2-L yeast cultures were solubilized for 4 h at 4 °C by slow rotation in solubilization buffer (described below) containing the most optimal detergent in the final concentration of 1% (w/v). Unsolubilized material was removed by ultracentrifugation at 160,000 x g at 4 °C for 1.5 h (using a Sorvall T-865 rotor). The resulting supernatant was subsequently diluted four times in the respective solubilization buffer and subjected to immobilized metal affinity chromatography (IMAC) using nickel chelated resin (see below). Following binding, the resin was washed and bound target IMPs were eluted with the below-mentioned buffers supplemented with imidazole and containing the primary detergent at a final concentration of 1.5 x CMC. The amount of recombinant protein in each fraction was determined by GFP fluorescence as described above. Essentially, three purification protocols were applied as described below.

1. AqPZ. Solubilized protein was loaded onto 2 x 1-ml HisTrap FF columns (GE Healthcare) connected in series and eluted in IMAC buffer (25 mM Tris-HCl, 500 mM NaCl and 10% glycerol; pH 7.6) with a linear imidazole gradient (0–500 mM).

2. ClCec1, NavAb and KcsA. Following O/N binding at 4 °C to 3 ml of high-affinity Ni-NTA resin (GenScript, USA), the material was packed into an Econo-column (Biorad, USA), flow-through was collected and the resin was washed with the IMAC buffer indicated above with a step imidazole gradient (i.e., 10, 30, 100, 250 and 500 mM, respectively) to remove impurities and elute bound target IMPs.

3. TmZIP. FC-16-solubilized crude membranes containing His-tagged TmZIP were diluted in solubilization buffer (50 mM HEPES-NaOH, 500 mM NaCl, 20% glycerol, 1 mM PMSF and 1 mM sodium dodecyl sulfate) and subsequently loaded onto a 5-ml Ni-NTA HisTrap HP column (GE Healthcare, USA) connected to an Acta Pure system (GE Healthcare, USA). The column was washed with 5 x CV of IMAC buffer (20 mM HEPES-NaOH, 250 mM NaCl, 20% glycerol, 0.003% FC-16; pH 7.5) and eluted with a linear imidazole gradient (0–500 mM).

2.9. Live cell bio-imaging

Localization of heterologously-expressed GFP-tagged membrane proteins was visualized by imaging GFP fluorescence in whole cells at 1000× magnification using a Nikon Eclipse E600 microscope (Nikon Instruments Inc., USA) coupled to an Magnafire camera model S99802 (Optronics, USA).

2.10. Electrophysiology

Single channel currents from DDM-solubilized, IMAC-pure KcsA and NavAb (1 mg/mL) were recorded using the Orbit Mini with EDR3 software (Nanon Technologies, Germany). Briefly, a four-electrode recording chip (MECA4) (50–100 μm; Ionera Technologies, Germany) was mounted in the Orbit Mini apparatus and recording buffer was applied to the chip. Planar lipid bilayers consisting of 10 mM 1,2-diphytanoyl-sn-glycero-3-phosphocholine (DPhPC), Avanti Polar Lipids, USA) and 1 mM cholesterol dissolved in nonane were painted on the recording wells. 0.2 μL of KcsA or NavAb in recording buffer (10 mM HEPES-NaOH, 100 mM KCl, 100 mM NaCl; pH 7.2) was added to the cis-side of the bilayer. Subsequently, alternating voltages were applied to facilitate insertion of the channels. Recordings were carried out at a sampling rate of 1.25 kHz and afterwards a low pass filter, Bessel (8-pole) with a 3-dB cutoff of 200 Hz, was applied. Mean current amplitudes were plotted as a function of tested voltages. Single channel conductances were determined as the slope factor of a linear regression to the current-voltage (IV) plot. Clampfit 10.7 (Molecular Devices, USA) and Prism 7 (GraphPad Software, USA) were used for data analysis.

3. Results

3.1. Target integral membrane proteins (IMPs)

The target proteins selected for the present paper belong to five different families and represent a wide variety of prokaryotic IMPs of bacterial (including Gram-negative and Gram-positive) and archaeal origin (Table 1). The five target protein families exert diverse biological functions, and possess different topologies with respect to the number of transmembrane (TM) helices and lengths and orientation of N-/C-termini. They furthermore differ in molecular weight and oligomeric state, as also reflected by available structural information (Table 1, Supplementary Figures 2 and 3). Due to our selection of prokaryotic IMPs included in this paper, the results may apply to the production of α-helical prokaryotic membrane proteins in general.

3.2. The E. coli and S. cerevisiae expression platforms

To compare the capacity of E. coli with S. cerevisiae for recombinant production of prokaryotic IMPs, we selected BL21(DE3), BL21(DE3)-pLYS, C41 and C43 expression strains, and the pET vector system (Studier and Moffatt, 1986) for E. coli expression and our own PAP1500 platform for yeast expression. The BL21(DE3)/pET host-vector system is the most commonly used in recombinant protein production in E. coli, while the S. cerevisiae PAPI500 expression platform has previously been shown to successfully express many challenging classes of IMPs of eukaryotic origin (Pedersen et al., 1996; Bombolt et al., 2013; Gotfryd et al., 2018; Wang et al., 2019; Molbaek et al., 2015; Scharff-Poulsen and Pedersen, 2013; Bjorkskov et al., 2017).

To simplify construction and manipulation of E. coli expression plasmids, we used the pPAP10286 vector for engineering expression constructs with AqPZ, ClCec1, NavAb and KcsA. The pPAP10286 shuttle vector can replicate in both S. cerevisiae and E. coli (Supplementary Figure 1) and carries the E. coli expression cassette from pET52b(+) and the ability of the plasmid to replicate in yeast enables fast, cheap and accurate assembly of prokaryotic expression constructs (Supplementary Figure 1), without introducing any “contaminating” sequences, such as restriction sites or site-specific recombination sequences. Expression of ZIP transporters was attempted using an unmodified pET15b or pET52b(+) vector to produce N- or C-terminally His-tagged targets, respectively.

3.3. E. coli expression host strain and temperature influence protein accumulation

To simplify optimization of protein production and purification, all target IMPs were expressed as TEV-GFP-His\textsubscript{10} tag fusions. We used codon-optimized cDNAs, i.e., tuned for expression in an E. coli-based platform. GFP can be directly used for quantification and in vivo localization of properly folded fusion proteins (Drew et al., 2001) and the TEV protease cleavage site enables removal of the entire fusion tag following metal affinity chromatography.

Expression of KcsA, NavAb and ClCec1 was screened in the four most used E. coli strains, i.e., BL21(DE3), BL21(DE3)-pLYS, C41(DE3) and C43(DE3), at both 15 °C and 30 °C induction temperature, using a standard IPTG concentration of 0.5 mM and followed by assessment of GFP fluorescence accumulated in intact cells. The highest accumulation of ClCec1 was observed in BL21(DE3) at 30 °C and peaked 24 h after induction. NavAb accumulated to the largest extent in BL21(DE3) and C41(DE3) strains at 30 °C 8 h post induction, while the most pronounced KcsA expression was achieved in BL21(DE3) induced for 20 h at 30 °C (Fig. 1).
Applying AqpZ as a model protein, we subsequently investigated whether IPTG concentrations used for induction affect accumulation of recombinant protein in a specific E. coli host strain at a given temperature. Data in Fig. 2 show that accumulation of AqpZ in the BL21(DE3) strain was higher at 15°C, while in the C41(DE3) and C43(DE3) strains the course of expression was elevated at 30°C. The peak accumulation observed in C41(DE3) and C43(DE3) strains at 30°C after 30 and 50 h, respectively, was comparable to that observed in BL21(DE3) strain at 15°C after 120 h. However, the final OD₄₅₀ was much higher for all strains grown at 30°C. The effect of IPTG concentration tested for induction of AqpZ expression indeed appeared to be host strain- and temperature-specific, and should therefore be optimized independently for each of the strains used.

Expression originating from the pET52b(+)–based shuttle plasmid was tested by examining production of a soluble protein; the GFP-tagged tetracycline repressor protein (TetR; UniProt ID: P04483) from transposon Tn10 to ensure that the low levels of accumulation observed for the prokaryotic target IMPs in E. coli were not caused by using the pET52b(+)–derived yeast shuttle vector. As highlighted by the data presented in Supplementary Figure 4, the generally observed low accumulation of prokaryotic IMPs in E. coli, compared to the TetR, cannot be attributed to the use of the engineered pET52b(+)–pEMBLyex4 plasmid, but rather to the membrane embedded nature of the expressed targets.

3.4. Temperature-dependent accumulation of prokaryotic integral membrane proteins (IMPs) in S. cerevisiae

To determine how temperature affects the accumulation of recombinantly produced IMPs in S. cerevisiae, we investigated the time course...
Fig. 2. Effect of IPTG concentration and temperature on accumulation of AqpZ C-terminally fused to TEV-GFP-His$_{10}$ in four E. coli host strains. E. coli strains (A) BL21(DE3), (B) BL21(DE3) pLysS, (C) C41(DE3) and (D) C43(DE3) were grown at 30 °C until OD$_{450}$ = 1, and transferred to either 15 °C (left panels) or 30 °C (right panels) prior to induction with 0.1, 0.5 or 1 mM IPTG. Protein accumulation was measured by GFP fluorescence in the cell amount corresponding to 1 OD$_{450}$ unit at each time-point after induction. Fluorescence measurements are shown as solid lines and OD$_{450}$ measurements as dashed lines, respectively. Color codes for the respective concentration of IPTG used for induction of the expression are indicated below each panel.
of their accumulation at 15 °C and 30 °C. As seen in Fig. 3A–D, at 15 °C the accumulation increased over the entire production period and was much higher than that observed at 30 °C. In contrast, accumulation at 30 °C already peaked after ~24 h, followed by a rapid decline for all tested proteins.

We used live cell bio-imaging to visualize the expressed GFP fusions in the yeast expression host. Fig. 3E illustrates that all target proteins accumulated in distinct intracellular compartments and not in the plasma membrane.

3.5. Prokaryotic integral membrane proteins (IMPs) accumulate to a high density in yeast membranes

The structures of the some of the IMP targets investigated in this work have already been determined using X-ray crystallography (Table 1) and the samples for structural characterization were produced using E. coli-based expression systems. Here we investigated whether the S. cerevisiae expression platform is capable of expressing the same targets, and subsequently compared both protein quantity and quality.

To achieve optimal protein production, yeast cells were induced at 15 °C and harvested at the established expression optimum for each target protein (Fig. 3A–D). The data in Supplementary Table 3 show that many target IMPs accumulated to a high density in yeast membranes relative to their yield in E. coli membranes. Because the molecular weight of the produced targets differs significantly, we also calculated the molar amounts accumulating per mg of crude membrane protein content. This revealed that the quantity of accumulated recombinant protein varied a lot between the different targets. From the estimated content of membrane-located recombinant protein, we also calculated the expected expression levels of the respective target IMPs (Supplementary Table 3). This would give rise to theoretical yields ranging from 0.9 to 149 mg of protein per liter shake flask culture. Such promising expression levels highlight the great potential for producing prokaryotic IMPs in large quantities needed for subsequent biophysical characterization.

3.6. Homogeneity of solubilized prokaryotic membrane proteins is detergent independent

Following the expression tests, we performed a screen to compare the ability of a selection of six non-ionic and zwitter-ionic detergents to extract the produced target IMPs from crude yeast membranes. The results shown in Fig. 4 (left panels) indicate that each target protein could be solubilized in all tested detergents. We identified AqpZ to be the most refractory to extraction, with an approximate solubilization efficiency of 20% in all applied surfactants (Fig. 4A). In contrast, ClCec1 generally solubilized well, with the highest extraction efficiencies observed for FC-12, DDM and CY-5 (80–100%; Fig. 4B). KcsA could also be effectively extracted with 40–75% solubilization efficiency (Fig. 4C), with DDM being the most potent surfactant. Finally, for NavAb, solubilization was most pronounced in DM, LDAO and DDM (80–100%), whereas the other tested detergents resulted in solubilizations amounting to 50% in CY-5, and 30% in both FC-12 and FC-13 (Fig. 4D).

To maintain the activity of the extracted prokaryotic IMPs during purification, it is essential to apply a detergent that not only efficiently extracts the target from the membranes, but also preserves the sample in a homogeneous and non-aggregated state. As permitted by the GFP fusion tag, we used fluorescence-detection size-exclusion chromatography (FSEC) to assess the quality of detergent-solubilized samples. The results in Fig. 4 (right panels) show the FSEC profiles of each target protein in all six tested detergents, with color codes corresponding to those used in the detergent screen. In general, protein elution peaks were narrow and monodisperse, with only marginal amounts of aggregated protein eluting in the void volume, overall indicating a high protein quality. Only NavAb showed a double peak, revealing two populations of protein solubilized in LDAO, DM or DDM, and to some degree also in CY-5, FC-12 or FC-13.

3.7. IMAC yields pure fusion proteins from yeast expression

Based on the results from the detergent screen and FSEC analysis (Fig. 4), several detergents proved to be promising candidates for subsequent protein purification. Following solubilization in the respective detergents, i.e., AqpZ in LDAO, ClCec1 in DM, and both NavAb and KcsA in DDM, all tested IMP targets were purified by immobilized metal affinity chromatography (IMAC). Fig. 5 shows the IMAC elution profiles of the respective proteins. In all cases, the top fraction (containing also the highest amount of GFP) was separated by SDS-PAGE and the purity of the samples was visualized by in-gel GFP fluorescence and Coomassie blue staining.

AqpZ eluted as a single broad peak starting at 100 mM imidazole (Fig. 5A). SDS-PAGE separation of purified AqpZ revealed a highly pure protein band, with only minor bands of higher molecular weight visible by in-gel GFP fluorescence and Coomassie blue staining, indicating that AqpZ tetromers have not been fully dissociated and the sample also contains a population of higher oligomeric states.

ClCec1 eluted as one narrow and symmetrical peak at 500 mM imidazole (Fig. 5B). IMAC resulted in a large amount of pure protein, with only minor impurities visible in the Coomassie blue stained SDS-PAGE gel.

KcsA eluted mainly as a single peak with a shoulder at 100 mM imidazole (Fig. 5C). The IMAC profile revealed a minor amount of protein eluting at 50 mM imidazole and a second small elution peak at 250 mM imidazole, respectively. SDS-PAGE of the top fraction confirmed one band corresponding to KcsA, but a substantial amount of the fluorescent sample did not enter the gel. An impurity of higher molecular weight was only seen in the Coomassie blue stained SDS-PAGE gel.

The elution IMAC profile of NavAb resembled that of KcsA. However, the SDS-PAGE gel revealed more impurities only apparent in the Coomassie blue stained gel (Fig. 5D, lane 1). The SDS-PAGE gel also disclosed several bands of higher molecular weight visible by Coomassie blue and in-gel GFP fluorescence, hence not all of NavAb tetromers were fully dissociated after exposure to SDS (Fig. 5D).

The molecular weights of the purified fusion proteins, as determined by SDS-PAGE, correspond to the predicted molecular size of the monomeric proteins (i.e., AqpZ of 23 kDa, ClCec1 of 50 kDa, KcsA of 18 kDa and NavAb of 30 kDa, respectively), considering that correctly folded GFP contributes to the overall molecular weight of the fusion proteins only with 10–15 kDa (Geertsma et al., 2008).

3.8. The prokaryotic ion channels produced in S. cerevisiae are functional

The above-described data confirm that prokaryotic IMPs can be produced in yeast and purified to homogeneity. Both DDM-solubilized KcsA and NavAb channels were purified by IMAC and subsequently analyzed by FSEC to demonstrate that the detergent-solubilized samples maintained stability during purification (Supplementary Figure 5). While the IMAC and FSEC profiles indicate high protein quality, they do not provide information on the functionality of the purified channels. We therefore decided to investigate the activity of both KcsA and NavAb.

After reconstitution into lipid bilayers, single channel current was recorded at a series of different voltages (Supplementary Figure 6A). A representative KcsA recording at ~40 mV is shown in Fig. 6A. The reconstituted KcsA channel proved to be functional, and the corresponding recording displays channel openings and closings of at least three single channel molecules (O1–O3). Fig. 6B illustrates the mean single KcsA channel current amplitude plotted as a function of voltage (V–I curve) and reveals a conductance of 77.3 ± 5.6 pS.

The reconstituted NavAb channel also displayed functionality and representative single-channel current recordings measured at ±120 mV reflect fast transitions between the open and the closed state (Fig. 6C), with an estimated single channel conductance of 57.6 ± 4.0 pS (Fig. 6D). The recordings also revealed that the activity was apparently voltage-dependent, with a larger open probability observed at positive voltages.
Fig. 3. Temperature-dependent accumulation of AqpZ, ClCec1, KcsA and NabAb C-terminally fused to TEV-GFP-His10 in S. cerevisiae. Yeast cultures were grown at RT until OD$_{450} = 1$ before half of the culture was transferred to 15 and the other half to 30 °C. After thermal equilibration, recombinant protein production was induced by addition of galactose to the final concentration of 2%. Accumulation of (A) AqpZ, (B) ClCec1, (C) KcsA and (D) NavAb was measured by GFP fluorescence in the cell amount corresponding to 1 OD$_{450}$ at each time-point after induction at 15 (blue lines) and 30 °C (red lines). (E) Live cell bio-imaging of yeast cells expressing the target proteins as C-terminal TEV-GFP-His$_{10}$ fusions. Pairwise differential interference contrast and GFP fluorescence images are shown.
Fig. 4. High-throughput solubilization screen and fluorescence-detection size-exclusion chromatography (FSEC) of AqpZ, ClCec1, KcsA and NabAb expressed in S. cerevisiae as C-terminal TEV-GFP-His10 fusions. Crude yeast membranes overexpressing (A) AqpZ, (B) ClCec1, (C) KcsA and (D) NavAb were solubilized for 1 h at 4 °C in 5-cyclohexyl-1-pentyl-β-D-maltoside (CY-5), n-dodecyl-β-D-maltopyranoside (DDM), n-decyl-β-D-maltopyranoside (DM), n-dodecylphosphocholine (FC-12), n-tridecylphosphocholine (FC-13) or lauryldimethylamine N-oxide (LDAO) in a protein:detergent ratio (w/w) of 1:3. Normalized solubilization efficiencies (left panels) were estimated from GFP fluorescence in the supernatant after ultracentrifugation. Subsequently, detergent-solubilized material was separated by FSEC where GFP fluorescence was monitored during the elution (right panels). The colors of the chromatograms correspond to the respective colors of detergents shown in the bar diagrams (left panels). The void volume of the used columns is ~8 mL for Superose 6 Increase 200 10/300 GL and is ~9 mL for Superose 12 10/300 GL.
suggesting that the NavAb channel shows a preference for opening at positive membrane potentials. It should be noted, however, that due to random insertion into the bilayer during reconstitution, the orientation of the channels cannot be predicted, thus the polarity of channel insertion is speculative.

3.9. N-terminal tagging improves accumulation of AqpZ in E. coli and S. cerevisiae

To test whether the position of the fusion tag affects the expression level and quality of recombinant prokaryotic IMPs produced in E. coli and S. cerevisiae, we used the previously described vectors (Supplementary Figure 1) to express AqpZ fused N-terminally to His10-GFP-TEV. Data in Fig. 7 show accumulation of this fusion protein in four BL21-derived E. coli host strains. It can be seen that BL21(DE3), C41(DE3) and C43(C43) strains were able to express the N-terminally tagged AqpZ to a larger degree at 15 °C than at 30 °C. By comparing Figs. 2 and 7, it is evident that relocating the GFP-His10 tag from the C- to the N-terminus of AqpZ greatly increased accumulation of the protein. The induction curves also revealed that production levels were not significantly affected by the IPTG concentration used for induction, except for the expression in the BL21(DE3) strain, where accumulation was higher after induction with 0.1 mM IPTG, compared to 0.5 and 1.0 mM (Fig. 7A).

Subsequently, we examined whether a similar trend can be observed in S. cerevisiae. Indeed, we observed that also in yeast N-terminally tagged AqpZ accumulated to a higher membrane density as compared to the C-terminal fusion (Fig. 8). Moreover, protein accumulation was also considerably higher at 15 °C than at 30 °C.

Whole cell fluorescence does not necessarily reflect accumulation of fusion proteins in the membranes, but may also potentially result from GFP that has been proteolytically liberated from the membrane protein. Hence, using in-gel GFP fluorescence we assessed membranes isolated from E. coli and S. cerevisiae expressing either the N- or C-terminally tagged AqpZ to investigate the combined effects of tag position, temperature, expression host and induction time on protein accumulation. To unbiasedly compare the accumulation of GFP-tagged AqpZ, all gels were imaged simultaneously (Fig. 8C). In accordance with the whole cell fluorescent measurements, the N-terminally tagged AqpZ displayed a higher accumulation in the membranes at 15 °C than at 30 °C.

(Supplementary Figure 6B), suggesting that the NavAb channel shows a preference for opening at positive membrane potentials. It should be noted, however, that due to random insertion into the bilayer during reconstitution, the orientation of the channels cannot be predicted, thus the polarity of channel insertion is speculative.
were solubilized and purified in DDM prior to reconstitution into planar lipid bilayers on a 50- or 100-μm MECA 4 recording chip in the Orbit Mini system. (A) Representative single KcsA channel current recordings at −40 mV. At least three individual channels have been incorporated into the planar lipid bilayer. “C” indicates the closed state and “O1-3” indicates the open states, respectively. (B) The Current-Voltage relationship (I–V curve) for KcsA with a linear regression fitted to the data revealing a slope conductance of 77.3 ± 5.6 pS. (C) Representative single NavAb channel current recordings at +100 mV. (D) I–V curve for NavAb with a linear regression fitted to the data revealing a slope conductance of 57.6 ± 4.0 pS. Results are shown as mean current amplitudes ± SEM, where each data point was based on at least 3 single channel openings (n = 3–1300 events, from 3 independent experiments).

Fig. 6. Single-channel recordings of immobilized metal affinity chromatography-pure KcsA and NavAb C-terminal TEV-GFP-His$_{10}$ fusions. Both channels were solubilized and purified in DDM prior to reconstitution into planar lipid bilayers on a 50- or 100-μm MECA 4 recording chip in the Orbit Mini system. (A) Representative single KcsA channel current recordings at −40 mV. At least three individual channels have been incorporated into the planar lipid bilayer. “C” indicates the closed state and “O1-3” indicates the open states, respectively. (B) The Current-Voltage relationship (I–V curve) for KcsA with a linear regression fitted to the data revealing a slope conductance of 77.3 ± 5.6 pS. (C) Representative single NavAb channel current recordings at +100 mV. (D) I–V curve for NavAb with a linear regression fitted to the data revealing a slope conductance of 57.6 ± 4.0 pS. Results are shown as mean current amplitudes ± SEM, where each data point was based on at least 3 single channel openings (n = 3–1300 events, from 3 independent experiments).

3.10. N-terminal position of the fusion tag improves solubilization of AqpZ from E. coli and S. cerevisiae membranes

To investigate the influence of the GFP-His$_{10}$ tag position on solubilization and quality of produced recombinant AqpZ, we performed a detergent screen and FSEC analysis of the His$_{10}$-GFP-TEV-AqpZ fusion. Results from the screens presented in Fig. 9A revealed a significant increase in solubilization efficiency of the N-terminally tagged AqpZ compared to the C-terminal fusion (see Fig. 4 for comparison). Specifically, the N-terminally tagged AqpZ can be extracted with 80–100% efficiency, while only 20% of the C-terminal fusion could be solubilized. Although the FSEC profiles revealed a wider elution peak in LDAO, all obtained profiles were monodisperse and devoid of aggregates (Fig. 9B).

Subsequently, we attempted affinity purification of the LDAO-solubilized N-terminally tagged AqpZ produced in yeast, using identical IMAC protocol as applied for the C-terminal construct. The chromatogram in Fig. 9C revealed that the N-terminally AqpZ fusion eluted in a single narrow peak at ~60 mM imidazole, resulting in a pure protein sample as revealed by both Coomassie blue staining and in-gel GFP fluorescence.

3.11. Rescue of expression of recombinant prokaryotic ZIPs in S. cerevisiae

With the aim of producing prokaryotic ZIPs for structural studies, we constructed a library consisting of archaeal and Gram-negative bacterial members. We selected ten prokaryotic ZIPs representing the entire ZIP family, with the two types of predicted membrane topologies shown in Supplementary Figure 2. Utilizing our selection of BL21 (DE3) based E. coli host strains we attempted to produce the zinc transporters from pET vectors with both N- and C-terminal histidine tags. However, after numerous attempts none of the ten targets were detectable in either cytoplasmic or membrane fractions of E. coli even by immunoblotting (data not shown).

Based on the improved expression of AqpZ, ClCc1, KcsA and NavAb in S. cerevisiae, we hypothesized that yeast would also have the capability to express this particularly difficult family of IMPs. Initially, we performed small-scale expression screens, where accumulation of the protein in yeast was visualized by live cell bio-imaging 48 h after induction (Fig. 10A). The S. cerevisiae-based platform enabled successful expression of all selected ZIPs, in contrast to the lack of expression in E. coli. Again, based on GFP fluorescence detected in the yeast membranes and the percentage of crude membrane protein, as described previously for other target IMPs, the expected expression levels of accumulated ZIPs ranged from 0.8 to 12.9 mg of protein per liter shake flask culture (Supplementary Table 3).

Following these expression tests, we performed a detergent screen using DDM and FC-16, a mild and a harsh surfactant, respectively. Fig. 10B shows that FC-16 was most effective, however, for PfZIP and TmZIP the achieved extraction efficacy was comparable with that observed for C-terminally tagged AqpZ (~30%). Based on the highest target content in the crude membrane fraction and the highest solubilization efficiency (Fig. 10B and C, and Supplementary Table 3), TmZIP emerged as the most promising target for downstream purification. Hence, we performed IMAC-based purification of both C- and N-terminally GFP-His$_{10}$-tagged TmZIP variants from crude S. cerevisiae membranes solubilized in FC-16 (Fig. 10C and D, respectively). In case of both constructs, bound protein eluted at an imidazole concentration of ~300 mM and yielded high purity after a single purification step, as reflected by both Coomassie blue staining and in-gel GFP fluorescence. Although both TmZIP variants were purified mainly in their monomeric forms, small fraction of the GFP-fluorescent dimer (MW of ~110 kDa with electrophoretic mobility shifted to ~80 kDa) could also be detected. In addition, irrespective of tag location, the...
Fig. 7. Effect of IPTG concentration and temperature on accumulation of AqpZ N-terminally fused to His\textsubscript{10}-GFP-TEV in four *E. coli* host strains. *E. coli* strains (A) BL21(DE3), (B) BL21(DE3) pLysS, (C) C41(DE3) and (D) C43(DE3) were grown at 30 °C until OD\textsubscript{450} = 1, and transferred to either 15 (left panels) or 30 °C (right panels) prior to induction with 0.1, 0.5 or 1 mM IPTG. Protein accumulation was measured by GFP fluorescence in the cell amount corresponding to 1 OD\textsubscript{450} unit at each time-point after induction. Fluorescence measurements are shown as solid lines and OD\textsubscript{450} measurements as dashed lines, respectively. Color codes for the respective concentration of IPTG used for induction of the expression are indicated below each of the bottom panels.
two IMAC-pure TmZIP fusions contained a significant population of non-fluorescent species (with electrophoretic mobility of ~50 kDa). Mass spectrometry and immunoblotting of this non-fluorescent fraction suggested that it originated from TmZIP misfolding or aggregation (data not shown), as malfolded GFP migrates with a different moiety than correctly folded GFP, hence using GFP as a folding reporter (Geertsma et al., 2008). However, this fraction could be completely removed during the reverse-IMAC step following TEV protease cleavage of the N-terminal fusion (Supplementary Figure 7), greatly improving the final purity of the produced TmZIP sample. It is also evident from Supplementary Figure 7 that the electrophoretic mobility of the GFP tagged TmZIP and TmZIP liberated after TEV protease digestion are higher than what would be expected from their molecular weight. This is commonly observed for membrane proteins due to altered binding of SDS (Rath et al., 2009). The approximate difference between the GFP tagged and non-tagged TmZIP is around 15 kDa in accordance with previous observations, that correctly folded GFP only contributes 10–15 kDa to the molecular weight of a corrected folded membrane protein (Geertsma et al., 2008).
4. Discussion

In the present paper we selected prokaryotic IMPs from bacteria and archaea representing five different protein families to systematically investigate the production levels in *E. coli* and in *S. cerevisiae*. Our results demonstrate that *S. cerevisiae* was a more robust expression host as all selected IMPs accumulated in the yeast membranes at a higher density than in *E. coli*. Strikingly, switching to the *S. cerevisiae*-based platform proved to be the only successful experimental strategy in overproducing ZIP transporters, providing complete rescue of the expression of all tested members of this IMP family. An important observation is that all tested targets accumulated to the highest density at the same expression conditions. Time consuming optimization of expression for each target was not required.

ZIP transporters remain rather understudied, largely due to difficulties in their production. Previously, ZIPs were selected as targets for a structural genomics study (Punta et al., 2009), where out of 52 homologous targets tested, only 4 displayed detectable expression levels (Lin et al., 2010). Furthermore, the generally accepted notion that members of this IMP family are refractory to overproduction in *E. coli* was also confirmed by another study (Ma et al., 2015). Although *E. coli* remains a widely used and robust expression system for expression of prokaryotic IMPs, problems can arise when attaching a poly-histidine tag to a terminus facing the periplasm, i.e., Nout or Cout. Reportedly, difficulties with membrane insertion and folding lead to severely impaired expression compared to tag placement on a cytoplasmic terminus (Baker et al., 2017). This problem is inevitable when working with proteins displaying Nout-Cout topologies as in case of ZIPs. Protein engineering studies have been successful in adding auxiliary TM helices and thus flipping the N- or C-terminus intracellularly to allow more efficient tagging and expression (Quick and Wright, 2002; Hsieh et al., 2010).

None of the 10 ZIPs investigated in this study was expressible in *E. coli*, including PaZIP, the only target predicted to possess Nout-Cout topology (Supplementary Figure 2F). However, it cannot be excluded that the location of the N-terminus in PaZIP is predicted incorrectly, as its sequence may not harbor an additional (false-positive) TM. Here, without employing any protein engineering, we switched to our *S. cerevisiae* platform and successfully rescued expression of all the attempted ZIP. This demonstrates clearly the strengths of the *S. cerevisiae* platform and likely its superior machinery for IMPs biogenesis, at least for the studied targets.

Microbial expression systems play a vital role in producing recombinant proteins for biotechnological and medical applications in addition to basic protein science. IMPs and their interactions are mainly of interest.
Fig. 10. Recombinant expression of bacterial zinc transporters (Zrt/Irt-like proteins, ZIPs) in *S. cerevisiae*. (A) Live cell bio-imaging of yeast cells expressing ten bacterial ZIPs as C-terminal TEV-GFP-His10 fusions. Yeast cultures were grown in 2-L scale and induced for protein expression for 48 h at 15 °C. Pairwise GFP fluorescence and differential interference contrast images are shown. (B) Crude yeast membranes overexpressing ten bacterial ZIPs were solubilized for 1 hour at 4 °C in n-dodecyl-β-D-maltopyranoside (DDM) or n-hexadecylphosphocholine (FC-16) in a protein:detergent ratio (w/w) of 1:3. Normalized solubilization efficiencies were estimated from GFP fluorescence in the supernatant after ultracentrifugation. (C) and (D) Immobilized metal affinity chromatography (IMAC) purification of C- and N-terminally GFP-tagged TmZIP, respectively. IMAC profiles after solubilization in 1% (w/v) FC-16 for 1 h at 4 °C are shown. Protein elution was monitored by absorbance at 280 nm. The dashed lines indicate the used imidazole gradient. Inserts show the SDS-PAGE gels, where the top fraction from each elution was visualized by Coomassie blue staining (lane 1) and in-gel GFP fluorescence (lane 2).
to the biomedical industry and to basic science as targets for medical treatments and structure-function studies, respectively (Yin and Flynn, 2016). The structures of prokaryotic IMPs deposited in the PDB (Burley et al., 2019) almost exclusively originate from samples produced in *E. coli*, reflecting the notion that this organism is an ideal host for production of proteins of prokaryotic origin. In contrast, *S. cerevisiae* has been used as a workhorse for production of eukaryotic IMPs for structural biology (Bill et al., 2011; Routledge et al., 2016).

*E. coli* has continuously been used for historical reasons. After it was found to express some recombinant IMPs well in a relatively easy to manipulate and cultivate manner, many laboratories pursued to use this organism as their standard expression host (Schlegel et al., 2014). Another reason given for choosing *E. coli* as the preferred expression host for expression of prokaryotic IMPs is that it offers low cost for culturing. However, the prices of *E. coli* and *S. cerevisiae* growth media are comparable, and both expression systems are applicable for large-scale cultivation in bioreactors.

Parallel expression screening of many targets requires fast and efficient generation, and, potentially, manipulation of recombinant expression plasmids. As exemplified in the present paper, yeast as expression host also offers an option of using homologous recombination to generate the expression plasmids directly in the *S. cerevisiae* production strain readily and without applying expensive enzymes.

Expressing IMPs in yeast has the indisputable advantage of avoiding accumulating the protein in inclusion bodies. Although IMPs can be expressed in very high quantities in inclusion bodies and the success rate for refolding β-barrel IMPs was shown to be high (Bannwarth and Schulz, 2003), it is often inefficient or impossible to refold α-helical IMPs, properly due to the complicated folding and hydrophobic nature of this class of proteins and therefore such efforts to refold are often avoided (Lyons et al., 2016; Kiefer, 2003; Krogh et al., 2001). Secondly, the eukaryotic machinery for membrane protein biogenesis in yeast may support correct folding of heterologous membrane proteins better than the simpler prokaryotic secretory pathway of *E. coli* (Pechmann et al., 2013; Bohnsack and Schleiff, 2010; Cross et al., 2009; Stephenson, 2005). In addition, the rate of protein synthesis and folding is much slower in *S. cerevisiae* than in *E. coli*, correlating with their generation time, which is approximately five times slower for yeast (Widmann and Christen, 2000). Consequently, the rates of protein synthesis in eukaryotic cells have been shown to be an order of magnitude slower as compared with prokaryotic hosts (Bill, 2014). Lowering the rate of recombinant protein production in *E. coli* strains C41 and C43 is the well characterized reason for their success. Both strains carry mutations in the promoter responsible for transcription of the T7 RNA polymerase gene, giving rise to slower recombinant protein synthesis rates than wild-type *E. coli* (Kwon et al., 2015).

The role of the lipid environment surrounding embedded proteins has been the focus of intense research and debate, as lipids have been proposed to be crucial for structure and function of IMPs (Martens et al., 2016). This may result from direct interactions between specific lipids and the protein or from the physico-chemical properties of the membrane itself, such as its fluidity, thickness, intrinsic curvature or lateral pressure (Renne and de Kroon, 2018). At the post transcriptional level, successful production of a prokaryotic IMP in *S. cerevisiae* at least requires that the protein can be targeted to the endoplasmic reticulum, membrane inserted through the Sec61 translocon and assembled into a stable and correctly folded form (Cymer et al., 2015). In this light, the *S. cerevisiae* homologues have permitted expression of all included targets in their native higher oligomeric forms, which includes dimerization (ClCec1, ZIPs) and tetramerization (AqpZ, KcsA and NaVAb).

Based on the previous paragraphs, it may seem counterintuitive to attempt producing prokaryotic IMPs in a eukaryotic host such as *S. cerevisiae*, as the lipid compositions of the Gram negative, Gram positive, archaean and the yeast plasma membranes differ considerably. Bacterial and yeast membranes are both mainly made of phospholipids esterified to glycerol-3-phosphate, while the building blocks in archaean membranes are isoprenes connected to glycerol-1-phosphate through an ether bond (Jain et al., 2014). As seen from Tables 2 and 3, major differences in the type and content of the phospholipids include the carbon length of the fatty acids and their degree of unsaturation. Specifically, yeast membranes compared with *E. coli* contain a higher percentage of C16 and C18 fatty acids, in particular their mono-unsaturated versions (Klose et al., 2012; Kanonenberg et al., 2019). The longer fatty acids found in yeast membranes result in an overall membrane thickness (9 nm) that is 3 nm thicker than the *E. coli* membrane and corresponds to a hydrophobic width of 30 Å and 20 Å, respectively (White and Wimley, 1999). The head group composition of the phospholipids also varies significantly between the two species, as phosphatidylethanolamine (PE) and phosphatidylglycerol (PG) constitute the great majority in *E. coli*, while these are much less represented in yeast. Conversely, yeast membranes include substantially higher amounts of phosphatidylinositol (PI) and phosphatidylcholine (PC). Furthermore, cardiolipin (CL) is more enriched in the *E. coli* membranes in contrast to sphingolipids and sterols that are solely found in yeast (Klose et al., 2012; de Kroon et al., 2013). Presence of the latter component, e.g., in a form of cholesterol, has previously been shown to preferentially interact with IMPs to restrict their dynamics (Grouleff et al., 2015) and enhance stability (Fanti et al. and Barrantes, 2013). Thus, concerning that protein dynamics and function are intimately entwined, this can further highlight the role of endogenous sterols of the yeast membranes in enhancing stability of overproduced prokaryotic IMPs that, identically to their eukaryotic counterparts, may also be sensitive to the presence of certain lipids. Indeed, despite the extensive differences in composition of the host membranes, our experiments demonstrate that the lipid environment in yeast is not prohibitive for accumulation of correctly folded prokaryotic IMPs. Moreover, only yeast was able to produce the prokaryotic ZIP transporters and deposit them in the membranes to a high concentration. Furthermore, also the other target proteins accumulated to a higher level in yeast even though they were well produced in *E. coli*. Importantly, our electrophysiological characterization of the KcsA and NaVAb ion channels purified from *S. cerevisiae* showed that they display the same functional characteristics as those produced in *E. coli* (LeMasurier et al., 2001; Shaya et al., 2011; Callahan and Roux, 2018) and, to our knowledge, represent the first published single channel recordings of these proteins. Based on the portfolio of prokaryotic IMPs analyzed here, our yeast expression platform emerged to be more robust than the BL21(DE3)/pET expression system. Consequently, while our yeast system was able to produce all proteins to a high level at 15 °C, expression in *E. coli* differed significantly among the tested strains. As example, AqpZ showed the highest accumulation in C41 and C43 strains, while BL21 was superior at expressing ClCec1, KcsA and NaVAb, and none of these expressed well in BL21(DE3) pLysS. Production of the ZIP transporters was even more challenging, as they all were refractory to production in all tested *E. coli* strains. Based on these results, a drawback of using the BL21(DE3)/pET expression system may implicate that it is necessary to screen a number of BL21-derived strains and expression conditions to achieve a satisfactory production level, as the optimal temperature for accumulation in the *E. coli* strains was observed to be IMP-dependent. These factors combined render the yeast-based expression system highly attractive and competitive.

The popularity of *E. coli* as host is undoubtedly due to the large number of successful functional studies and structures solved from recombinant membrane protein samples produced in this host, but it also has historical reasons. It was discovered often to produce recombinant proteins well, to be easy and cheap to manipulate and cultivate. Associated with it is a short generation time, laboratories continue to use this organism as their standard expression host for prokaryotic membrane proteins (Schlegel et al., 2014). However, the prices of *E. coli* and *S. cerevisiae* growth media are practically identical and protocols for large-scale cultivation in bioreactors are well established for both expression systems. The longer generation time in yeast does, however, inevitable increase production time relative to *E. coli*.

The present study focused on the production of prokaryotic IMPs in the membranes of the host organism since expressing membrane proteins...
in yeast has the advantage of avoiding expressing the protein in inclusion bodies. Membrane proteins can be expressed in very high quantities in inclusion bodies, but while the success rate for refolding β-barrel IMPs was shown to be high (Bannwarth and Schulz, 2003), it is often inefficient or impossible to refold α-helical IMPs properly due to the complicated folding and hydrophobic nature of this class of proteins (Lyeons et al., 2016; Kiefer, 2003; Krogh et al., 2001). Secondly, yeast possesses eukaryotic machinery for membrane protein biogenesis that may be more compatible with correct folding of membrane proteins than the simpler prokaryotic secretory pathway of E. coli.

So what drawbacks may potentially emerge from using yeast as expression host for prokaryotic IMPs? Expression in yeast may potentially result in non-native posttranslational modifications including N- and O-glycosylation (Conde et al., 2004; Neubert et al., 2016) that may detract from the downstream biophysical applications, e.g., crystallization (Mesters and Hilgenfeld, 2007). Here, we did not assess such modifications, but the very sharp bands observed for SDS-PAGE-resolved IMAC-pure samples do not implicate heavy protein glycosylation (e.g., Figs. 5, 9 and 10).

Even though AqpZ expressed better in yeast than in E. coli and we were able to purify it as the C-terminally tagged fusion, re-localizing the tag to the N-terminus not only increased the expression level in both hosts, but also resulted in a remarkably improved solubilization in all six tested detergents. This agrees with the general strategy that target expression level and solubilization efficiency need to be tested with both C- and N-terminal tags to identify the most optimal fusion construct (Mohanty and Wiener, 2004).

Based on our data and previous experience with production of the prokaryotic IMPs, we present a roadmap generalizing strategies for enhanced production of recombinant prokaryotic IMPs in S. cerevisiae (Fig. 11). Starting with molecular biology, the following should be considered: codon optimization of the prokaryotic target proteins to the yeast host, selection of tag(s) and its/their position and choice of a suitable vector for a given expression strain. Based on these decisions, the expression plasmid can be assembled from PCR products directly in S. cerevisiae by homologous recombination. Here, we successfully used a poly-histidine tag in combination with GFP for rapid screening and quantification of expression levels. Correct construct design can then easily be verified “in vivo” by assessing live cell GFP fluorescence concomitantly with sequencing of the expression construct. The following step encompasses small-scale expression trials, where screening of the most optimal expression conditions is performed applying, e.g., assessment of GFP fluorescence in the cells and SEC-MALS analysis to confirm integrity of accumulated targets. Subsequently, upon isolation of crude yeast membranes, detergent screen has to be conducted to identify the optimal surfactant that is able to efficiently extract the protein of interest in the most stable and native form. Here FSEC-based analysis proves to be useful, as it permits rapid determination of the monodispersity of expressed GFP fusions after solubilization, an important quality check in the whole strategy. Finally, affinity chromatography can be conducted, with the overall aim to obtain a protein sample of the highest possible purity and integrity that can be assessed by, e.g., SDS-PAGE analysis (in-gel GFP fluorescence, Coomassie blue staining, immunoblotting). Finally, if a sample of satisfactory quality is produced, it can be used for the subsequent characterization, e.g., activity measurements or structural determination that may require implementation of additional purification/reconstitution steps. However, prior to structure-function studies, an additional analysis may be required to further assess the quality of produced samples. Such validation may include, e.g., SEC-multi-angle light scattering (MALS), a technique useful in analyzing the homogeneity of proteins (Some et al., 2019). If any of the stages in the above-described expression strategy do not meet the success criteria, troubleshooting can be attempted.

5. Future perspectives

Our comparative study demonstrates that S. cerevisiae is a promising starting organism for overproduction of prokaryotic IMPs, both for well-characterized and difficult to express protein classes. Although there is not so much known on how specific lipids interact with IMPs to regulate their stability and activity, the natural diversity of membrane lipid composition might be one of the important factors to consider when attempting expression of prokaryotic targets in yeast and E. coli. Further improvements of our expression platform to excel at producing challenging targets could then include fine-tuning of the native membrane environment of the S. cerevisiae strain to suit a given type of IMP, both to improve expression levels and quality of the protein.

6. Conclusions

In summary, we showed that expression of KcsA, NavAb, AqpZ and ClCe1 was achievable in both E. coli and yeast, but switching to an S. cerevisiae-based platform improved production levels. We find it particularly important that we were able to express all of the selected prokaryotic ZIP transporters in S. cerevisiae that was not possible employing E. coli, demonstrating that the yeast-based system could be useful for overproduction of other challenging families of IMPs. Overall, there are many strengths of the expression platform described herein, as reflected by robust recombinant protein overproduction of all targets. They could all be purified to high purity by a single affinity chromatography step and for two tested ion channels activity was preserved. Hence, this suggests that the powerful S. cerevisiae-based methodology presented here can be applied to a wider range of challenging prokaryotic IMP targets for subsequent biochemical and structural analysis.
Fig. 11. Workflow of recombinant production of prokaryotic integral membrane proteins (IMPs) in yeast. The chart comprises four steps critical for a successful overproduction strategy. Each step includes end-points for verification and ideas for troubleshooting; a green arrow indicates that the criteria is met, a red arrow indicates the procedure was unsuccessful and needs to be optimized further before proceeding. **Molecular biology:** This step should include codon optimization of the cDNA encoding the prokaryotic IMP of interest, choosing the fusion tag (here a detachable GFP-His10 fusion is used as the reporter/affinity tag) and its position, application of a vector suitable for the selected yeast expression strain and engineering the final expression plasmid using PCR and homologous recombination. Correct construct assembly should be verified by DNA sequencing and the initial expression can be confirmed by live cell bio-imaging. Unsuccessful construction of the target should be followed by starting over with PCR and homologous recombination. **Small-scale expression:** In this step screening of optimal induction temperature, time course and concentration of inducer in chosen expression strains should be considered. Assessment can be performed by monitoring the course of GFP accumulation, live cell bio-imaging, SDS-PAGE-based analysis employing in-gel GFP fluorescence or immunoblotting. Unsuccessful protein accumulation should be followed by optimization of growth conditions (e.g., medium, additives or growth temperature), construct redesign (e.g., removal of large cytoplasmic termini, change of fusion tag position or introduction of a linker between target and tags). **Detergent screen:** Ability of detergents to efficiently solubilize the yeast membranes to isolate prokaryotic target IMP in a stable form should be verified. Extraction efficacy can be estimated directly in solubilized membranes (after removal of insoluble fraction by ultracentrifugation) using GFP fluorescence. This should be followed by testing the integrity of detergent-solubilized sample by fluorescence-detection size-exclusion chromatography (PSEC) to assess its level of aggregation and monodispersity. Additionally, sample quality (e.g., preservation of its native state) can be analyzed using non-reducing SDS-PAGE. Unsuccessful solubilization of target protein should be followed by change of solubilization conditions (pH, buffering system, reducing agent, ligands, additives, lipids) or by using alternative extraction methods (e.g., styrene maleic acid co-polymers (SMAs) or direct reconstitution into nanodiscs). **Affinity chromatography:** A purification protocol must be established for each individual IMP target, but overall it should aim at obtaining of pure sample eluting in one main peak. Sample purity should be assessed using SDS-PAGE (Coomassie blue staining, in-gel GFP fluorescence or immunoblotting) and, if necessary, the binding conditions should be revised (e.g., pH and buffering system, concentration of salt). Following successful affinity purification, activity and stability of the protein sample should be investigated. Unsuccessful purification should be followed by optimization of binding and elution conditions (pH, buffering system, concentration of salt, type of gradient) or changing the type of affinity resin used.

**CRediT authorship contribution statement**

**Sarah Spruce Preisler:** Writing - original draft, generated the expression constructs, performed protein accumulation screens, performed protein purification, performed protein purification, designed the project, wrote the initial version of the manuscript and the final version in association, All authors contributed with comments on the final manuscript. **Anders Drabaek Wiuf:** Writing - original draft, generated the expression constructs, performed protein accumulation screens, performed protein purification, performed protein purification, All authors contributed with comments on the final manuscript. **Marc Friis:** Writing - original draft, performed protein purification. **Lasse**
Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.crbi.2021.02.001.

References


Kjaergaard: Writing - original draft, performed protein purification.

Molly Hurd: Writing - original draft, performed activity measurements.

Eva Ramos Becares: Writing - original draft, performed protein purification.

Casper Normann Nurup: Writing - original draft, performed detergent screens and FSEC analysis.

Frederik Bühring Bjoerkskov: Writing - original draft, designed the project. Kirstine Calloe: Writing - original draft, performed activity measurements. Dan A. Klaerke: Writing - original draft, performed activity measurements. Kamil Gotfryd: Writing - original draft, generated the expression constructs, performed protein accumulation screens, designed the project. Per Amstrup Pedersen: Writing - original draft, generated the expression constructs, performed protein accumulation screens, designed the project.

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