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# The Effect of Detergent, Temperature, and Lipid on the Oligomeric State of MscL Constructs: Insights from Mass Spectrometry

### **Graphical Abstract**



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# In Brief

Mechanosensitive channels act as emergency solute release valves, their oligomeric state being inherently linked with their function. Using mass spectrometry, Reading et al. explored the influence of construct, temperature, detergent, and lipid on the oligomeric diversity of this membrane complex.

# **Highlights**

- Mass spectrometry reveals different oligomeric forms of mechanosensitive channels
- Oligomeric state is exquisitely sensitive to temperature, detergent, and lipid
- MscL and other membrane proteins can bind to LPS when overexpressed in *E. coli*





# The Effect of Detergent, Temperature, and Lipid on the Oligomeric State of MscL Constructs: Insights from Mass Spectrometry

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http://dx.doi.org/10.1016/j.chembiol.2015.04.016

#### **SUMMARY**

The mechanosensitive channel of large conductance (MscL) acts as an emergency release valve for osmotic shock of bacteria preventing cell lysis. The large pore size, essential for function, requires the formation of oligomers with tetramers, pentamers, or hexamers observed depending on the species and experimental approach. We applied non-denaturing (native) mass spectrometry to five different homologs of MscL to determine the oligomeric state under more than 50 different experimental conditions elucidating lipid binding and subunit stoichiometry. We found equilibrium between pentameric and tetrameric species, which can be altered by detergent, disrupted by binding specific lipids, and perturbed by increasing temperature (37°C). We also established the presence of lipopolysaccharide bound to MscL and other membrane proteins expressed in Escherichia coli, revealing a potential source of heterogeneity. More generally, we highlight the use of mass spectrometry in probing membrane proteins under a variety of detergent-lipid environments relevant to structural biology.

#### INTRODUCTION

Mechanosensitive channels of large conductance (MscL) act as emergency non-selective release valves for bacteria when under high internal cellular pressures caused by hypo-osmotic shock (Booth et al., 2007; Haswell et al., 2011; Iscla and Blount, 2012; Kung et al., 2010). MscL responds to the tension of the bilayer by opening and closing its channel through a progression of subconductance states until sufficient tension is experienced to stabilize the fully open state. The monomeric subunits of MscL are relatively small (*Staphylococcus aureus* [Sa] MscL, *Escherichia coli* [Ec] MscL, and *Mycobacterium tuberculosis*  [Mt] MscL have 120, 136, and 151 residues, respectively) and possess only two transmembrane helices per subunit (Figure 1). It has been appreciated from the initial discovery of MscL (Sukharev et al., 1994) that the channel must be oligomeric; since the dimensions of the pore and hence conductance reflect the number of subunits, determination of the oligomeric state of MscL is crucial to understand the physiology.

The oligomeric state of MscL has been studied by a range of experimental approaches and has been observed to be tetrameric, pentameric, or hexameric depending on the ortholog, solution conditions, and experimental technique (Walton et al., 2015). The protein sequence is a key determinant because changes to the protein construct may perturb the oligomeric state. Particular emphasis has been placed on the role of the cytoplasmic C-terminal domain (CTD) in this process (Anishkin et al., 2003; Blount et al., 1996; Walton and Rees, 2013; Yoshimura et al., 2008). The CTD consists of a helical bundle composed of the C-terminal soluble helix from each subunit. Recently, the crystal structure of the EcMscL CTD was solved and biophysical characterization demonstrated that it formed stable pentamers devoid of the remaining EcMscL sequence (Walton and Rees, 2013). This suggests that the CTD may influence the oligomerization and stability of the full-length protein. Detergent is another important parameter and, for a given construct, the observed oligomeric state can vary with changes in detergent. Moreover, the physiological function of MscL reflects an intimate coupling to the properties of the lipid bilayer (Iscla and Blount, 2012; Laganowsky et al., 2014) and changes in the membrane environment can influence the gating properties of MscL (Moe et al., 2000). A specific interaction between MtMscL and phosphatidylinositol is crucial for the mechanosensitivity of this channel (Zhong and Blount, 2013). Indeed, since MscL experiences a detergent environment only when purified for in vitro studies, the physiological relevance of the detergent dependence of the oligomeric state of the channel has been questioned (Dorwart et al., 2010; Iscla et al., 2011).

To investigate the effects of these parameters on oligomeric state, we studied the MscL from five different bacterial sources, ranging from 120 to 151 residues in length (Figure 1C; Table 1). We used non-denaturing (native) mass spectrometry (MS) to





#### Figure 1. Structures and Sequence Alignment of Mechanosensitive Channels of Large Conductance

(A and B) Side (top) and top (bottom) views of the crystal structures of pentameric MtMscL (A, PDB: 2OAR) and tetrameric SaMscL C $\Delta$ 26 (B, PDB: 3HZO). (C) Amino acid sequence alignment for MscL homologs produced using the CLUSTAL Omega program (Sievers et al., 2011). Amino acids are colored according to the Clustal X color scheme. Accession numbers of the sequences used for alignments are: *E. coli*, UniProt: V8K223; *M. tuberculosis*, UniProt: A5U127; *S. aureus*, UniProt: P68805; *A. ferrooxidans*, UniProt: B5ERJ9; and *T. thermophilus*, UniProt: Q72L04.

reveal the oligomeric states of full-length and CTD truncation constructs in a variety of detergents, lipids, and solution temperatures. MS has been shown previously to enable determination of the mass, subunit stoichiometry, and lipid binding properties of membrane protein complexes (Barrera et al., 2009; Konijnenberg et al., 2014). The results presented here show that purification of EcMscL following established protocols (Chang et al., 1998; Liu et al., 2009) results in significant co-extraction with lipopolysaccharide (LPS) and show how co-extraction of this outer membrane lipid is common for a number of *E. coli* membrane proteins; using appropriate buffer conditions and/or detergents, however, LPS can be removed.

We then probe the interrelationships between protein subunit stoichiometry and the effects of different detergent-lipid environments. We find that while detergent can perturb the overall subunit stoichiometry, the majority of lipids investigated here did not influence the oligomeric state significantly. The effects of C-terminal truncations, however, showed marked preference for tetrameric forms of SaMscL, while the position of the tag, whether N- or C-terminal, affected the oligomeric stability in some cases with more homogenous populations of oligomers observed for C-terminally green fluorescent protein histidine (C-GFP) tagged constructs.

Overall, using this experimental dataset consisting of five homologs and various constructs, we explore the effects of lipid, detergent, and temperature on the population of different oligomeric forms. From more than 50 different experimental conditions, we provide the most comprehensive view to date of the interrelated factors that affect the conversion and stability of MscL oligomers.

#### RESULTS

#### LPS Binds to MscL

Starting with the expression and purification of MscL from E. coli with a GFP and histidine tag at the C-terminus (EcMscLC-GFP), a mass spectrum was recorded from *n*-dodecyl-β-D-maltoside (DDM) under conditions optimized for the study of membrane protein complexes (Figure 2A) (Laganowsky et al., 2013). The mass spectrum reveals two overlapping charge state series at m/z 8,000 corresponding to oligomeric species with molecular masses of 235.2 and 238.8 kDa. Overlapping charge state series at m/z 2,000 are also observed, corresponding to monomeric subunits with molecular masses of 44.0 and 14.8 kDa. The former closely matches the expected mass for this construct, while the latter surprisingly corresponds to that for the native EcMscL (the subsequent use of an E. coli strain knocking out the native gene [Laganowsky et al., 2014] eliminated this species). The oligomer peaks must correspond to EcMscL pentamers (expected molecular mass 220.4 kDa) but there is a significant discrepancy of an additional 14.8 and 18.5 kDa, respectively (Figure 2A). Performing this experiment on MtMscL C-GFP also revealed bound adducts with a clear additional mass in the range of 3.4-3.7 kDa from the apo form of MtMscL C-GFP, particularly evident in the stripped oligomer region of the mass spectrum (m/z 10,000– 22,000) (Figure 2B). This suggests that MscL membrane proteins co-purify with a varying amount of adducts possessing masses in the range of 3.4-3.7 kDa. Although the mass of the adduct does not define its chemical identity, the absence of charge on the bound adduct narrows down the possible candidates to either a small peptide or lipid-like molecule.

| Table 1. Summary and Abbreviations for MscL Constructs                   |                      |   |
|--|----------------------|---|
| Vector Construct   | Abbreviation         | Theoretical<br>Molecular<br>Weight (Da) |
| pet15b-SaMscL-S-TEV-GFP-6xHis  | SaMscL C-GFP         | 42,751.9                                |
| pet15b-SaMscL C∆(96-120)-S-<br>TEV-GFP-6xHis                             | SaMscL C∆25<br>C-GFP | 39,844.5                                |
| pet15b-10xHis-Thrombin-SaMscL  | N-His SaMscL         | 15,779.6                                |
| pet15b-10xHis-Thrombin-SaMscL<br>C∆(95-120)                              | N-His SaMscL<br>C∆26 | 12,743.2                                |
| pet15b-MtMscL-S-TEV-GFP-6xHis  | MtMscL C-GFP         | 45,296.3                                |
| pet15b-MtMscL C∆(103-151)-S-TEV-<br>GFP-6xHis                            | MtMscL C∆48<br>C-GFP | 40,077.6                                |
| pet15b-10xHis-Thrombin-MtMscL-PG   | N-His MtMscL         | 18,440.1                                |
| pet15b-10xHis-Thrombin-MtMscL<br>CΔ(102-151)                             | N-His MtMscL<br>C∆49 | 13,075.4                                |
| pet15b-EcMscL-TEV-GFP-6xHis  | EcMscL C-GFP         | 44,076.4                                |
| pet15b-AfMscL-TEV-GFP-6xHis  | AfMscL C-GFP         | 45,354.0                                |
| pet15b-TtMscL-TEV-GFP-6xHis  | TtMscL C-GFP         | 42,778.8                                |
| Af. Acidithiobacillus ferrooxidans: Ec. Escherichia coli: Mt. Mycobacte- |                      |   |

COII: IVIT. IVIVCC rium tuberculosis; Sa, Staphylococcus aureus; Tt, Thermus thermophilus.

A series of experiments were performed to identify the noncovalently bound adduct with a mass ranging from 3.4 to 3.7 kDa. First, we did not observe a lower molecular weight band in SDS-PAGE stained with Coomassie blue dye (Figure S1), suggesting this adduct was a non-protein species. We suspected that the adduct could be the main outer membrane lipid, LPS. LPS has a complex and heterogeneous structure; in laboratory E. coli K-12 strains, LPS consists of lipid A with two Kdo sugars and an R-core oligosaccharide domain consisting of six to ten sugars (hexoses and heptoses, along with side-chain modifications) corresponding to a molecular mass of  ${\sim}3.4-$ 4.1 kDa (Raetz, 1996). Since LPS may be detected by silver staining (Tsai and Frasch, 1982), we repeated the SDS-PAGE electrophoresis but this time followed by silver staining. A dark band appeared for a molecular species consistent with our observed adduct mass and migrating similarly to the purified LPS from E. coli (Figure S1). The ligand identity as LPS was further validated by dot blot analysis of purified MscL protein with an anti-lipid A antibody (Figure S1). In addition, we expressed and purified MtMscL C-GFP from ClearColi E. coli (Lucigen), a strain that harbors several gene knockouts that reduce the biochemical pathway of LPS to a precursor, lipid IV<sub>A</sub>. Recording a mass spectrum of adducts bound to the MscL complex isolated from this strain reveals adducts of 1.4 kDa consistent with the mass of lipid IV<sub>A</sub> (1,409 Da) (Figure 2C). These experiments identify the bound adduct as LPS and provide an explanation for the range in mass observed for the bound adduct, due to the heterogeneity of the core oligosaccharide of LPS.

To determine if LPS co-purifies with other membrane proteins or is enhanced with MscL, we expressed and purified three additional inner membrane proteins in E. coli (Figure S2). Following extensive purification, we found LPS bound to  $\sim$ 50% of the chloride channel and the multidrug resistance protein, and up to ~20% of the acriflavine resistance protein B. We conclude that LPS binding is not specific to MscL but can occur readily to a variety of membrane proteins expressed in E. coli and likely goes unnoticed in typical evaluation for purity and heterogeneity (Drew et al., 2008; Gutmann et al., 2007).

To assess the stability and stoichiometry of the apo protein and the effects of other lipids using non-denaturing MS, we developed purification methods for the removal of LPS. Inspired by the use of Triton X-100 in removing LPS from outer membrane proteins (de Cock et al., 1996; Magalhaes et al., 2007), mass spectra of MtMscL-GFP extracted with 1% Triton X-100 were obtained, revealing small fractions of LPS bound to the protein complex and oligomer stripped species. In the presence of 1% Triton X-100, but this time with the addition of 20% glycerol, a dramatic increase in the proportion of LPS bound to the complex was observed. Interestingly, substitution of Triton X-100 with octyl glucose neopentyl glycol (OGNG) detergent in glycerol-free buffer resulted in no detectable LPS bound to the complex (Figure S3). We conclude therefore that LPS removal is dependent on the physical properties of the detergent and that, in this case, OGNG is more efficient in removing this adduct from MtMscL than other detergents that we tested.

#### **MscL Stoichiometry in Different Detergents**

To probe the dependence of the oligomeric states of MscL within different detergents, we compared two structurally characterized homologs, SaMscL and MtMscL. When in a DDM environment, we found that the N-terminally histidine tagged (N-His) SaMscL was tetrameric (47%) with smaller oligomers present (trimers, dimers, and monomers), while N-His MtMscL showed a significant population of pentamers (38%) and tetramers (63%) (Figure 3). In the same DDM detergent environment but with a C-terminal GFP histidine tag (C-GFP), SaMscL and MtMscL both formed stable pentamers (100%). Furthermore, the C-GFP constructs of EcMscL, A. ferrooxidans (Af) MscL and T. thermophilus (Tt) MscL, all produced exclusively pentameric complexes in DDM (Figure 3).

To determine if there was a dependence on detergent and the oligomeric state, we then compared SaMscL and MtMscL constructs in three further detergents: octyl-β-D-glucopyranoside (OG), tetraethylene glycol monooctyl ether (C8E4), and lauryldimethyl amine oxide (LDAO). Interestingly, we found that N-His SaMscL, N-His MtMscL, SaMscL C-GFP, and MtMscL C-GFP in LDAO, OG, and C8E4 were exclusively pentameric, with small populations of GFP truncated species observed in some cases. This demonstrates that the protein construct can influence the observed oligomer distributions, with full-length SaMscL and MtMscL C-GFP constructs producing pentameric states predominantly, while the N-terminal histidine modifications can destabilize pentameric forms in one of the detergents investigated here.

#### The Effects of the C-Terminal Domain on the Oligomeric State

Given previous observations that the CTD of MtMscL and EcMscL have extensive intersubunit interactions (Chang et al., 1998; Walton and Rees, 2013) and the proposal that these interactions may control the oligomeric state, we generated SaMscL and MtMscL constructs (C-GFP and N-His) in which we deleted these C-terminal amino acid residues (C $\Delta$ ) (Table 1). We found





#### Figure 2. Non-Denaturing Mass Spectrometry of EcMscL C-GFP and MtMscL C-GFP Extracted and Purified in the Detergent DDM

(A) Mass spectrum of EcMscL C-GFP collisionally activated with argon gas measures pentameric complexes bound to several adducts. Two different monomeric masses are observed in the low m/z region (1,500–3,000 m/z) as a result of collisional activation; one corresponding to the expected MtMscL C-GFP fusion protein and the second corresponding to endogenous EcMscL with the initiating methionine removed. Reported in the inset of each mass spectrum is the measured mass and SD. Use of an MscL knockout strain (Laganowsky et al., 2014) eradicated endogenous MscL contamination.

(B) Mass spectrum of MtMscL C-GFP collisionally activated with sulfur hexafluoride gas reveals up to three bound adducts, with an average mass of  $\sim$ 3.6 kDa, bound to the pentamer and the collisionally dissociated tetramer.

that these truncations had little or no effect on MtMscL in the detergents DDM and LDAO, with MtMscL C $\Delta$ 49 C-GFP remaining predominantly pentameric; however, the presence of C8E4, and OG especially, disrupted the MtMscL C $\Delta$ 49 C-GFP pentamer into smaller oligomers (Figure 3).

By contrast, SaMscL CA25 C-GFP revealed almost exclusive populations of the tetrameric species in the same conditions (Figure 3). To assess the possible effects of the C-GFP on the stability of SaMscL oligomers, we investigated the N-His SaMscL CA26 in addition to the C-GFP. We found that this construct produced a mixture of oligomeric states in DDM with no pentameric channel present, similar to that observed for the full-length construct with the N-His tag. Interestingly, N-His SaMscL CA26 produced an almost 80:20 population of pentamer/tetramer in LDAO consistent with OCAM and crystallographic results but in contrast to the full-length protein, in which only pentamer was observed. In addition, we observed that the CTD truncated species of SaMscL and MtMscL in the presence of particular detergents were not compatible with MS analysis, producing unresolvable mass spectra through aggregation, insolubility, or degradation, whether N- or C-terminally tagged (pink boxes, Figure 3). This was indeed the case for the N-His MtMscL C∆49 construct; data were unobtainable in all detergents examined here.

Tetramers were observed, however, almost exclusively for the SaMscL C $\Delta$ 25 C-GFP construct in LDAO, DDM, and C8E4. Interestingly, only pentamers were seen for the full-length protein from equivalent conditions. We can conclude therefore that truncation of these C-terminal residues caused all SaMscL C $\Delta$  constructs to form populations of tetramers in contrast to the pentameric full-length constructs. This is in accordance with the crystal structure of the SaMscL tetramer purified and crystallized in LDAO (Liu et al., 2009) as well as analytical ultra-centrifugation data from the same solution conditions (Dorwart et al., 2010).

#### Influence of Temperature on MscL Oligomeric State

Our observation that SaMscL populates various oligomeric states according to the construct and the detergent environment implies that different oligomeric forms can interconvert. Reasoning that an increase in temperature could promote this interconversion, we incubated and equilibrated solutions of SaMscL C-GFP in both LDAO and C8E4 for periods of up to 1 hr at 4°C or 37°C prior to MS analysis (Figure 4). Following incubation at 4°C, SaMscL C-GFP was solely pentameric in both LDAO and C8E4. Interestingly, when equilibrated at 37°C, for 1 hr, SaMscL C-GFP remained exclusively pentameric in C8E4. By contrast, the same construct underwent complete conversion to a tetramer in LDAO following incubation at the same elevated temperature. When the LDAO solution at 37°C was then re-equilibrated to 4°C, SaMscL C-GFP precipitated. This would imply that the oligomeric switch from pentamer to tetramer at elevated temperature is irreversible, at least in LDAO.

These results reflect a relationship between temperature and oligomeric state in one of the two detergent environments

Removal of the C-GFP fusion with the TEV protease did not result in reduction of bound LPS.

<sup>(</sup>C) MtMscL C-GFP was overexpressed in ClearColi *E. coli* strain (Lucigen) and analyzed by non-denaturing mass spectrometry.



studied here. The fact that the pentameric complex was perturbed in LDAO, but not in C8E4, likely reflects the different oligomeric stabilities of SaMscL in these two detergent environments.

# Influence of Lipids on the Interconversion between SaMscL Oligomers

Given the observation that SaMscL constructs are able to form stable tetramers and pentamers in solution and can convert from pentamers to tetramers following thermal activation, the potential influence of lipids on this interconversion was investigated. Different lipids were added to SaMscL constructs solubi-

#### Figure 3. Stoichiometry and Relative Abundance of MscL Constructs Purified in Different Detergents

All constructs were kept at 4°C before analysis. All MscL constructs were solubilized, purified, and analyzed by non-denaturing mass spectrometry in the same detergent, where fractions from a single peak from size exclusion chromatography were used for non-denaturing mass spectrometry analysis (see Experimental Procedures). Cases in which the mass spectrum was unresolved or could not be acquired are denoted with a pink box. The relative abundances and fitting errors for the oligomers were calculated using an in-house deconvolution software package using an algorithm described in Marty et al. (2015). Relative abundances of the different oligomers were calculated by adjusting the mass spectra intensity for detector efficiency (Ebong et al., 2011; Fraser, 2002; Stengel et al., 2012) and assuming that the ionization efficiencies of the various oligomers are similar. All mass spectra and spectral deconvolution fits are shown in Table S1 and Figures S4-S6.

lized in either LDAO or C8E4 (Figure 5). The lipid composition of the Sa membrane mainly consists of cardiolipin (CDL), phosphatidylglycerol (PG), and lysyl-PG in relative populations of approximately 24%, 10%, and 50%, respectively (Koch et al., 1984). These lipids and the lipid phosphatidylethanolamine (PE), which is non-natural to the Sa membrane, were therefore added to full-length SaMscL C-GFP in LDAO and C8E4, and incubated for 6 hr at 4°C, to investigate the effect of natural and non-natural lipids on MscL oligomeric state.

In the LDAO-lipid environments, the predominant pentameric oligomer was retained, with a small amount of conversion to tetrameric forms observed for each lipid. The highest population of tetrameric SaMscL C-GFP was observed with CDL and lysyl-PG (23% and 21%, respectively). For the C-terminal truncation SaMscL C $\Delta$ 25 C-GFP, the addition of lipid had little influence on the oligo-

meric state, the tetramer forming exclusively in LDAO as seen previously in the absence of lipids (Figure 3). When assessing the effects of the four different lipids in C8E4 on the stability of full-length SaMscL C-GFP, we found that it retained its pentameric form, as observed in the absence of added lipids (Figure 3). However, the SaMscL C $\Delta$ 25 C-GFP in C8E4 generated predominantly tetrameric forms of the protein with a small population of pentamer observed. For the SaMscL C $\Delta$ 25 C-GFP construct, the population of pentamer in the presence of CDL, PG, and PE lipids (16%, 19%, and 19%, respectively) is greater than in C8E4 alone (10%), implying that lipids can stabilize the pentameric state of SaMscL within the C8E4 detergent.



**Figure 4. Oligomeric State of SaMscL C-GFP Is Dependent on Both Temperature and Solubilizing Detergent** Mass spectra of SaMscL C-GFP purified at 4°C in either LDAO or C8E4 (top) prior to incubation at 37°C for 1 hr (bottom).

Given that lipids may play a role in stabilizing these different oligomers, we then investigated the effects of lipid on the temperature-induced (37°C) interconversion probed above (Figure 6). Full-length SaMscL, cleaved of its C-terminal GFP tag, in LDAO was doped with lipids at 4°C for 12 hr and revealed mainly pentameric and tetrameric complexes in accordance with the observations for a detergent-only environment. However, substantially less pentamer was observed for detergentlipid solutions (detergent-only = 87% and lipid-detergent = 19%-58%). Similar populations of pentamer and tetramer were observed with PG, lysyl-PG, and PE. CDL did not follow the same trend, however, producing substantially less pentamer (19%) and forming moderate populations of trimer (27%). Incubating at 37°C for 12 hr, however, caused SaMscL to form tetramers and trimers but no pentamers irrespective of the lipid added. Surprisingly, all lipids caused an increase in lower oligomeric species in comparison with the LDAO-only environment. This suggests that lipids can affect populations of oligomers for SaMscL in the presence of LDAO detergent. Since conversion from pentamers to tetramers occurs more readily in LDAO than in C8E4, and given that CDL was the most effective in causing this transition, but was less effective than the detergent environment, we conclude that the dominant effect on oligomeric stability is therefore the detergent environment.

#### DISCUSSION

We have shown using non-denaturing MS that the detergent or detergent-lipid environment can influence the oligomeric states of MscL channels. However, to assess the effects of lipids and detergents on subunit stoichiometry, it was critical that protein preparations are lipid free initially. In this regard, knowledge that MscL co-purifies with LPS, an outer membrane lipid and endotoxin known to contaminate recombinant protein preparations from gram-negative bacteria, (Morrison and Ulevitch, 1978) is critical. We also found that, rather than binding specifically to MscL, LPS also co-purifies with a number of other membrane proteins isolated from *E. coli*, adding a potential complication to biophysical/biochemical analyses. The presence of LPS if undetected could therefore complicate the analysis of the effects of exogenously added lipids or reconstitution, especially in the case of patch clamp experiments for MscL or crystallization experiments.

Although pentameric channels are observed under many conditions for all five full-length homologs (Mt, Sa, Ec, Af, and Tt) populations of lower-order oligomers could be seen, depending on the detergent environments and tag placement. These different populations were assessed using a mass spectral deconvolution algorithm (Marty et al., 2015) revealing the relative abundance and diversity of MscL oligomeric states in vitro. C-GFP provided more homogenous oligomeric states compared with the N-His constructs. Truncation of the CTD of MscL channels was also found to be critical, leading to enhanced dissociation, aggregation, and interconversion to lower oligomeric states, with CTD truncated SaMscL forming tetramers within all detergent environments. This suggests that the CTD has a role in oligomeric stability. However, it has previously been shown that CTD truncated and full-length SaMscL forms



в

Stoichiometry (Relative abundance, %)



#### Figure 5. Stoichiometry of SaMscL Constructs in Different Detergent-Lipid Environments

(A) After addition of lipid to purified SaMscL constructs, the solution was left at 4°C for 6 hr before analysis by non-denaturing mass spectrometry (equipped with altered accelerating voltages for increased collisional activation, which enabled detergent and lipid removal while keeping oligomeric interaction intact; Hopper et al., 2013). Protein degradation (loss of C-GFP tagged unit) varied for different mixtures.

(B) The relative abundances and fitting errors for the oligomers were calculated using an in-house deconvolution software package using an algorithm described in Marty et al. (2015). Relative abundances of the different oligomers were calculated by adjusting the mass spectra intensity for detector efficiency (Ebong et al., 2011; Fraser, 2002; Stengel et al., 2012) and assuming that the ionization efficiencies of the various oligomers are similar. All mass spectra and spectral deconvolution fits are shown in Tables S2 and Figures S7 and S8.

predominantly pentamers in native membranes (Dorwart et al., 2010; Iscla et al., 2011). Interestingly, we found that the interconversion of full-length SaMscL from a pentamer to a tetramer could be stimulated by temperature, in a detergent-dependent manner. Therefore, the solubilizing detergent environment has a major influence on the oligomeric state of MscL.

Understanding the formation and interconversion between oligomeric states of membrane proteins enables the mechanisms for their function to be deciphered and understood. Comparing our results for MscL with the  $\alpha$ -helical trimeric membrane protein diacylglycerol kinase (DGK), recently shown to form trimers and monomers in vitro, we note that DGK has high kinetic stability in OG (up to several weeks) and in liposomes at 37°C (Jefferson et al., 2013). By contrast, SaMscL converts from pentamers to tetramers within 1 hr, at 37°C in LDAO, and can form different oligomers in DDM but does not convert when in C8E4. This further supports that MscL oligomeric state is highly sensitive to its detergent environment. Interestingly, the presence of lipids affected the oligomer conversion of SaMscL. CDL was found to have a more destabilizing effect on

the SaMscL pentameric state than other smaller lipids (PG, lysyl-PG, and PE). It is noteworthy, however, that CDL, PG, and lysyl-PG exist in the natural Sa membrane (Koch et al., 1984). We conclude therefore that while membrane proteins can exhibit vastly different thermodynamic and kinetic stabilities (e.g. DGK and MscL), MscL is exquisitely sensitive to its detergent-lipid environment.

Multimeric membrane proteins may possess a number of oligomeric states that are possible within the bilayer environment. It is likely that for MscL, quaternary and tertiary structural stability are dictated by the combination of physical properties of bilayers (i.e. membrane curvature, tension, charge, pressure [Booth, 2005; Phillips et al., 2009]) as well as direct lipid interactions (Bogdanov et al., 2014; Laganowsky et al., 2014; Lee, 2011). Here, we have demonstrated that the solubilizing detergent environment has a dominant and profound impact on the quaternary structure (oligomeric state) of MscL channels, especially the SaMscL channel in the presence of LDAO. Therefore, the solubilization of MscL from its native membrane environment can influence the oligomeric state detected. However,



this destabilization could arise either from the removal of stabilizing interactions with lipids or from the loss of the physical presence of the bilayer.

For the MtMscL and SaMscL channels, it has been observed that when maintained in a pentameric form, in the appropriate detergent, specific lipid interactions stabilize its tertiary structure. Interestingly, however, a range of lipids were found to stabilize MscL to a similar degree, indicating little selectivity for particular lipid classes (Laganowsky et al., 2014). However, while lipids stabilized the pentameric structure of SaMscL within C8E4, lipids caused destabilization when in an LDAO environment. We propose therefore that although lipids stabilize the

#### Figure 6. Detergent-Lipid Environments Influence the Temperature-Induced Interconversion of SaMscL

(A) Summary of the effect of lipid addition on temperature-induced oligomer dissociation within LDAO detergent. SaMscL was expressed and purified in LDAO as SaMscL C-GFP and the tag removed by TEV protease cleavage. The protein was analyzed after 12 hr at 4°C or 37°C in the presence of 2× the CMC of LDAO detergent under equivalent non-denaturing mass spectrometry conditions (equipped with altered accelerating voltages for increased collisional activation, which enabled detergent and lipid removal, while keeping oligomeric interaction intact; Hopper et al., 2013). <1 = negligible amount observed in spectrum. The relative abundances and fitting errors for the oligomers were calculated using an in-house deconvolution software package using an algorithm described in Marty et al. (2015). Relative abundances of the different oligomers were calculated by adjusting the mass spectra intensity for detector efficiency (Ebong et al., 2011; Fraser, 2002; Stengel et al., 2012) and assuming that the ionization efficiencies of the various oligomers are similar. All mass spectra and spectral deconvolution fits are shown in Table S3 and Figure S9.

(B) Representative mass spectrum for PE addition to SaMscL in LDAO for two different temperature regimes. Monomer (purple), trimer (red), tetramer (green), and pentamer (blue) charge series are represented by filled circles.

tertiary structure of MscL, they cannot prevent the dominant quaternary destabilization caused, within certain detergent environments, and by removal from their natural bilaver environment. The extent of quaternary destabilization is dependent on the homolog, construct, and detergent environment and, most importantly, how well this environment can substitute for the physical properties of native membranes. The natural bilayer possesses many different biophysical properties in comparison with a micelle, such as the greater exclusion of water as well as differences in both the profile and extent of lateral pressure. Recapitu-

lating these attributes, in a homogeneous detergent micelle, is consequently a formidable task.

The dominant factor that determines the diversity of states observed for MscL channels is therefore likely due to the kinetic stability of the various oligomers within different detergent environments. Since we can separate the effects of individual lipid binding events (Laganowsky et al., 2014) from the physical effects afforded by the physical presence of the lipid bilayer, we conclude that quaternary destabilization likely arises when the detergent micelle replaces the native membrane environment rather than as a result of delipidation. This is in accord with the evolutionary function of MscL channels, their intimate relationship with tension and pressures in the bilayers acting to open and close the non-selective pore accordingly (Martinac, 2011; Perozo et al., 2002; Phillips et al., 2009; Sawada et al., 2012), and, as proposed here, to maintain (or change) its oligomeric state after it has been formed.

More generally, the ability of non-denaturing MS to characterize these interconverting states and compositional heterogeneity as a function of protein construct, purification conditions, detergent, lipid, and applied temperature, is likely to lead to powerful insights into membrane proteins; not only to guide structural biologists but also for the many biotechnology applications of protein channels that are rapidly coming to the fore (Charalambous et al., 2012; Iscla et al., 2013; Koçer et al., 2005, 2006; Yang et al., 2013).

#### SIGNIFICANCE

Mechanosensitive channels are membrane proteins that are present within all organisms and are of great significance since they control mechanical sensation. The bacterial mechanosensitive channel of large conductance (MscL) is important for the maintenance of cell turgor, acting as an emergency solute release valve when under hypo-osmotic stress. In addition, MscL has been tailored for use in nanotechnology, for example, as a nanovalve or in building protein-protein communication networks. MscL acts in concert with its lipid environment during channel opening (and closing). The size of its pore, and therefore its function, is determined by the oligomeric state of the channel. However, the oligomeric state has been highly controversial since it has been observed in tetrameric, pentameric, and hexameric forms. Critical in the field, therefore, is the question as to whether or not these oligomers can interconvert in vitro in response to different lipid and/or detergent environments. To address this important question, we have investigated more than 50 different experimental conditions, in which we varied both the protein construct and detergent/lipid environments. We then exploited the unrivalled ability of nondenaturing MS to monitor both protein subunit stoichiometry and lipid binding simultaneously to address the question of their oligomeric states. Here, we demonstrate how this biophysical tool, coupled with de-convolution algorithms, can be used to probe the equilibria between different oligomeric states of membrane protein complexes and demonstrate an exquisite sensitivity of MscL oligomers to temperature and the detergent/lipid environment.

#### **EXPERIMENTAL PROCEDURES**

#### **MscL Expression and Purification**

MscL constructs with a C-terminal fusion to GFP and 6× histidine tag were constructed as previously described (Laganowsky et al., 2014). Constructs containing an N-terminal 6× histidine tag were constructed as previously described (Chang et al., 1998; Liu et al., 2009). Plasmids were transformed into a modified *E. coli* strain (Laganowsky et al., 2014), BL21(DE3)  $\Delta mscL::Kan^R$  unless otherwise stated. Proteins were expressed and purified as previously described (Chang et al., 1998; Laganowsky et al., 2014; Liu et al., 2009). To investigate the effect of detergent on MscL oligomeric state, an MS membrane protein detergent screen was performed on the constructs as previously described (Laganowsky et al., 2013).

Briefly, the protein was purified from the membranes with 2% TX-100, 1% DDM, 1% LDAO, 1.5% OG, or 5% C8E4 detergent in 100 mM sodium chloride, 20% glycerol, 5 mM  $\beta$ -mercaptoethanol, and 20 mM Tris (pH 7.4) at room temperature supplemented with a complete protease inhibitor tablet and incubated overnight at 4°C with gentle agitation. Insoluble material was pelleted by centrifugation at 20,000 g for 25 min at 4°C and the clarified supernatant was purified using Ni-NTA agarose (Qiagen) drip columns (Bio-spin Chromatography columns, Bio-Rad). Membrane protein samples were injected onto a Superdex 200 5/150 (GE Healthcare) column equilibrated in 130 mM sodium chloride, 10% glycerol, and 50 mM Tris (pH 7.4) at room temperature supplemented with of 2× critical micelle concentration (CMC) of the detergent of interest. Peak fractions were concentrated and buffer exchanged into MS buffer (2× CMC detergent of interest and 200 mM ammonium acetate [pH 8.0] with ammonium hydroxide) using a centrifugal buffer exchange device (Micro Bio-Spin 6, Bio-Rad).

#### **Lipid Preparation**

Lipids (Avanti Polar Lipids) were prepared and added as previously described (Laganowsky et al., 2013). Different lipids were added to MscL, solubilized in LDAO, at an empirically determined protein/lipid/detergent molar ratio in order to sustain electrospray and achieve quality mass spectra: *E. coli* L- $\alpha$ -PG (1:9:548), *E. coli* CDL (1:4.8:548), 1,2-dioleoyl-*sn*-glycero-3-[phosphor-rac-(3-lysyl(1-glycerol))] (lysyl-PG) (1:7:548), and *E. coli* L- $\alpha$ -PE (1:9.5:548).

#### **Dot Blot Analysis**

LPS standard purchased from Sigma (L2880 from *E. coli* O55:B5) was prepared at 2.3 mg/ml in 0.5 M sodium chloride. Dot blot was performed on polyvinylidene fluoride membranes spotted with 1  $\mu$ l of protein in buffer (20 mM Tris [pH 7.5], 150 mM NaCl, 0.02% DDM). The blot was probed with primary antilipid A antibody (Pierce PA73178) at a 1:1,000 dilution in TBST (Tris-buffered saline with Tween 20) plus 5% non-fat milk (probing/blocking were concurrent). After washing with TBST, the blot was probed with secondary antigoat-horseradish peroxidase conjugate at a 1:10,000 dilution in TBST plus 5% non-fat milk. After washing with TBST, blot was developed with Western Lightening Plus ECL substrate (Perkin Elmer) for 3 min and film exposed for 2 min before development. All constructs were solubilized and purified at 4°C and then flash frozen before immunodot blot analysis.

#### **Non-Denaturing Mass Spectrometry**

A modified Q-ToF2 mass spectrometer (Micromass) with a Z-spray source (Sobott et al., 2002) was used. Typical instrument settings were 5-7 µbar source pressure, 1.5-1.8 kV capillary voltage, 150-190 V cone voltage, 1-10 V for extraction voltage, 180-200 V for collision voltage, either SF6 or argon for collision gas, and 0.2-0.3 MPa for collision gas pressure. In addition, another modified Q-ToF2 was used; this instrument was modified to facilitate high-mass molecule transmission and allow for higher collision activation capabilities, as previously described (Hopper et al., 2013). The modifications were conducted by Mass Spec Service Solutions. Both mass spectrometers used a multi-channel plate detector. The accelerating voltages used were optimized so as to maintain the oligometric state of the membrane proteins while removing the maximal amount of detergent or detergent lipid. Because detergent removal is dictated by the physical properties of the detergent micelle (Reading et al., 2015), it is not possible to compare accelerating voltage conditions for constructs in different detergents. The relative abundances of oligomers were calculated using the UniDec deconvolution software program (Marty et al., 2015) with detector efficiency taken into account (Ebong et al., 2011; Fraser, 2002; Stengel et al., 2012).

#### **Non-Denaturing Mass Spectrometry Deconvolution**

Deconvolution was performed using a Bayesian deconvolution algorithm described previously (Marty et al., 2015; unidec.chem.ox.ac.uk). Mass spectra and the detail for each fit (and the fitting parameters used) are provided in Tables S1–S3 and Figures S4–S10. Discriminating between native oligomers and potential collisional induced dissociation (CID) products was based on the charge states of the protein species. CID for homo-oligomers characteristically leads to the loss of highly charged monomer subunits producing charge-stripped oligomers that possess substantially lower average charge states, resulting in series at high *m/z* values (Benesch et al., 2006). In addition,

a detergent-dependent relationship exists between membrane protein mass and the ion charge state produced by nano-electrospray ionization (Reading et al., 2015). This evidence enables a distinction between native oligomeric forms and those produced by CID.

Briefly, spectral smoothing and background subtraction were applied to the data prior to deconvolution. Peak widths were determined by fitting peaks to either a Gaussian or a split Gaussian/Lorentzian model and were between 15 and 80 *m/z*, generally ~35, full width at half maximum (mzsig). Charge was allowed to range from 1 to 100, and mass (massub and masslb; ub, upper bound; lb, lower bound) and *m/z* range (minmz and maxmz) were limited for each piece of data. Errors were estimated using k-fold cross validation, where k is 2, 3, 4, or 5. The unprocessed data were sampled by deleting every kth data point in k different reading frames. Each of these 14 sub-spectra was processed and fit with the same parameters. Abundances are reported as the mean and SD of the peak height from these 14 fits. The pseudointensities, intensity error derived from the deconvolution algorithm, relative oligomer abundances, and propagated errors are summarized in Tables S1–S3.

#### SUPPLEMENTAL INFORMATION

Supplemental Information includes three tables and ten figures and can be found with this article online at http://dx.doi.org/10.1016/j.chembiol.2015. 04.016.

#### **AUTHOR CONTRIBUTIONS**

E.R., T.A.W., I.L., and A.L. performed the experiments and analyzed the data. E.R., T.A.W., A.L., D.C.R, and C.V.R designed the study. M.T.M. developed the deconvolution software and provided analysis support for this study. E.R. and C.V.R. wrote the article with contributions from all other authors.

#### ACKNOWLEDGMENTS

This work was supported by an ERC Advanced Investigator Award (26851) (IMPRESS), a Royal Society Research Professorship, a Medical Research Council Program Grant (98101) and an NIH grant GM084211.

Received: March 11, 2015 Revised: April 23, 2015 Accepted: April 27, 2015 Published: May 21, 2015

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