Ionic strength and calcium regulate membrane interactions of myelin basic protein and the cytoplasmic domain of myelin protein zero

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ABSTRACT  
The formation of a mature myelin sheath in the vertebrate nervous system requires specific protein-membrane interactions. Several myelin-specific proteins are involved in stacking lipid membranes into multilayered structures around axons, and misregulation of these processes may contribute to chronic demyelinating diseases. Two key proteins in myelin membrane binding and stacking are the myelin basic protein (MBP) and protein zero (P0). Other factors, including Ca2+, are important for the regulation of myelination. We studied the effects of ionic strength and Ca2+ on the membrane interactions of MBP and the cytoplasmic domain of P0 (P0ct). MBP and P0ct bound and aggregated negatively charged lipid vesicles, while simultaneously folding, and both ionic strength and calcium had systematic effects on these interactions. When decreasing membrane net negative charge, the level and kinetics of vesicle aggregation were affected by both salt and Ca2+. The effects on lipid membrane surfaces by ions can directly affect myelin protein-membrane interactions, in addition to signalling effects in myelinating glia.  
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1. Introduction  
Myelinated nervous fibers are formed by the processes of myelination. This process is essential for the insulation of the axons, thus enabling their fast propagation. Myelination involves the formation of myelin sheets, which are formed by compact and non-compact myelin [11–13]. MBP is involved in multiple sclerosis (MS) [14,15], being an intrinsically disordered protein (IDP) that folds upon lipid binding [16–18]. MBP carries a positive charge and presents post-translationally citrullinated forms with differing net charge [19]. It requires negatively charged lipids for membrane adhesion, but correct myelin morphology also depends on other lipids and ions [10,18,20]. Binding of divalent cations to MBP [21–24] could, together with altered membrane compositions, influence its in vivo functions, having implications in MS and the stability of CM.  
Myelin protein zero (P0) is a transmembrane protein in PNS myelin, relevant in human peripheral neuropathies and animal models [25,26]. P0 has an extracellular immunoglobulin-like domain, a single transmembrane helix, and a cytoplasmic tail (P0ct). Like MBP, P0ct is positively charged and interacts with phospholipids through electrostatics, gaining secondary structure [27,28]. This might make it susceptible to effects caused by cationic species.  
We studied the effects of ionic strength and Ca2+ on the structural and functional aspects of MBP and P0ct binding to membranes using biophysical techniques. The results shed light on the interplay between ions and proteins in CM and support the role of Ca2+ in regulating bilayer-MBP interactions.
2. Experimental procedures

2.1. Protein expression and purification

Mouse MBP and human P0ct were expressed and purified as described [18,28]. The final size-exclusion chromatography after affinity purification and tag removal was performed using Superdex75 16/60 HiLoad and Superdex75 increase 10/300 GL columns (GE Healthcare) in 20 mM HEPES, 150 mM NaCl, pH 7.5 (HBS). When required, the buffer was exchanged to water by sequential dialysis. Purified proteins were either used fresh or snap-frozen in liquid N₂ and stored at −80 °C.

2.2. Vesicle preparation

1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC), 1,2-dimyristoyl-sn-glycero-3-phospho-(1'-rac-glycerol) (Na-salt) (DMPG), and the deuterated d₅₄-DMPC, d₅₄-DMPG, and d₇-cholesterol were from Avanti Polar Lipids (Alabaster, Alabama, USA). n-dodecylphosphocholine (DPC) was from Affymetrix.

Lipid stocks were prepared by dissolving dry lipids in chloroform or chloroform:methanol (9:1 v/v) at 10 mg/ml. Mixtures were prepared at desired molar ratios, followed by solvent evaporation under N₂ and lyophilization. The dried lipids were stored air-tight at −80 °C.

2.3. Vesicle turbidimetry

For turbidimetric measurements, samples containing 0.5 mM DMPC:DMPG (1:1) SUVs with 2.5 μM P0ct or MBP were prepared in duplicate at 150 μl final volume on a Greiner 655161 96-well plate. Optical density at 450 nm was recorded immediately after thorough mixing at ambient temperature, to ensure that no unsuspended lipids remained. Multilamellar vesicles were prepared by 7 freeze-thaw cycles using liquid N₂ and a warm water bath, with vortexing between. Small unilamellar vesicles (SUV) were prepared by sonicating fresh multilamellar vesicles using a probe tip sonicator (Materials Inc. Vibra-Cell VC-130) until clear, while avoiding overheating. All lipid preparations were immediately used for experiments.

2.4. Synchrotron radiation circular dichroism spectroscopy

SRCD data were collected from 0.1 to 0.4 mg ml⁻¹ protein samples in water or 0.5% SDS on the AU-CD beamline at ASTRID2 (ISA, Aarhus, Denmark). Samples containing lipids were prepared immediately before measurement by mixing P0ct (P/L ratio 1:200) or MBP (P/L ratio 1:300) with freshly sonicated SUVs. 100-μm and 1-mm pathlength closed circular cells (Suprasil, Hellma Analytics) were used, and SRCD spectra were recorded from 170 to 280 nm at 30 °C. Data were processed using CDtoolX [31].

2.5. Stopped-flow SRCD

Rapid kinetic SRCD data were collected using an SX-20 stopped-flow instrument (Applied Photophysics) mounted on the AU-SRCD branch line of the AU-AMO beamline at ASTRID2 (ISA, Aarhus, Denmark) at 30 °C. Acquisition of kinetic data comprised of 6–10 shots (160-μl total volume per shot) per sample. The two 1-ml syringes were loaded as follows: syringe 1 contained 0.1 mg ml⁻¹ protein and syringe 2 a lipid solution (final molar P/L ratios of 1:200 and 1:300 for P0ct and MBP, respectively). The contents of each shot were rapidly mixed (2 ms dead time) before injection into the measurement cell (2-mm pathlength). CD signal was monitored at 195 nm for 5 s. Successful overlaying shots were averaged into a single curve for each sample. All sample sets were prepared and measured in duplicate. Water baselines were subtracted, and data were analyzed in GraphPad Prism 7 using single- and two-phase exponential decay functions to obtain rate constants.

3. Results

The folding and lipid membrane aggregation of MBP and P0ct were studied. Optical density measurements were used to observe the aggregation of lipid vesicles in the presence of proteins and different ions. The studies were expanded to different ratios of DMPC:DMPG, which affects the surface charge of the SUVs. Using SRCD, protein folding was probed, and time-resolved experiments provided data on membrane aggregation kinetics.

3.1. The effect of ionic content on myelin protein-induced vesicle turbidity

As a functional assay for MBP and P0ct, turbidimetric experiments on SUVs were carried out. In our earlier studies, we conclude that in the presence of MBP, net μM, negatively charged lipid vesicles aggregate and subsequently form myelin-like bilayer stacks, resulting in sample turbidity [18]. P0ct, on the other hand, can cause SUVs to undergo fusion events to produce large aggregated lipid bodies [28], which may cause additional sample turbidity.

The turbidity induced by the proteins for 1:1, 4:1, and 9:1 DMPC:DMPG SUVs, at a 1:200 protein-to-lipid (P/L) ratio, was determined. The experiments were carried out at various concentrations of NaCl, NaF, and CaCl₂. Especially multivalent cations may induce lipid aggregation in the absence of proteins [32], control samples without proteins were included (Fig. 1). Protein-free sample turbidity was observed mostly in the presence of CaCl₂, with more CaCl₂ needed for induction, when the lipids became more negatively charged. The effect of monovalent ions as inducers of turbidity per se in protein-free samples was low.

The turbidity induced by MBP was higher than that of P0ct, with the exception of DMPC:DMPG (9:1), in which P0ct was more efficient than MBP. In the presence of either protein, the induced turbidity systematically increased with salt concentration in DMPC:DMPG (1:1) (Fig. 1A). In DMPC:DMPG (4:1) and (9:1), the turbidity decreased above a certain ionic strength (Fig. 1B and C), and the balance between ionic strength and lipid net charge plays a role in protein-induced membrane aggregation. The turbidification of DMPC:DMPG (9:1) was abolished at salt concentrations of 150 mM or above. In all lipid mixtures, both proteins in the presence of NaCl and NaF produced roughly the same levels of turbidity and concentration-dependent trends. Thus, the choice of monovalent anion matters little in the level of induced turbidity.

To mimic the physiological concentration, CaCl₂ was included at much lower concentration than NaCl and NaF, but increased turbidity was still observed. CaCl₂ induced lipid turbidity in the absence of protein; the levels varied as a function of concentration. Calcium-induced turbidity showed differences between the three DMPC:DMPG ratios: the turbidification of DMPC:DMPG (1:1) vesicles was more resistant to CaCl₂ than 4:1 and 9:1 mixtures (Fig. 1). CaCl₂ changed the turbidity induced by MBP and P0ct, depending on the membrane composition. In DMPC:DMPG (1:1), turbidity was highest, at a similar level to that induced by the presence of salt (Fig.1A). DMPC:DMPG (9:1) showed least turbidity, and high calcium concentrations brought it to baseline levels (Fig. 1C). Interestingly, in DMPC:DMPG (4:1), 0.5 mM and 1 mM CaCl₂ showed a large difference in turbidity in the absence of...
was carried out with 150 mM NaF and 1 mM CaCl₂. NaF was used to investigate the effect of ionic strength and divalent cations on folding, SRCD behaviour to NaCl in turbidimetry (Fig. 1).

The presence of lipids often suffer from turbidity-induced light scattering - here, especially at 1:1 and 4:1 DMPC:DMPG ratios with CaCl₂ (Fig. 2A and B). Under such conditions, the positions of the spectral peaks can be used to detect the presence of secondary structure. The observed changes in conformation correspond well to the turbidimetry assays under the same conditions, whereby the negative peak at 200 nm indicates unfolded structure and possibly lack of membrane binding.

MBP reached a similar secondary structure content in all lipid compositions (Fig. 2). In DMPC:DMPG (9:1), NaF and CaCl₂ decreased MBP helical content, potentially indicating less binding to lipids (Fig. 2C). Also P0ct obtained secondary structure in lipid mixtures, and the decrease of membrane negative charge (DMPG) correlated with the loss of P0ct helical structure - seen as the movement of the main minimum towards 200 nm (Fig. 2). NaF had a small unfolding effect on P0ct in all lipid mixtures.

### 3.3. The kinetics of initial vesicle nucleation

To visualize the kinetics of MBP and P0ct binding to lipid vesicles, stopped-flow SRCD measurements were carried out. The original goal was to study protein folding kinetics when encountering lipid membranes, but initial measurements and optimization revealed that the proteins gained their folding completely within the instrument dead time (~2 ms), making the kinetics practically impossible to follow.

Despite the setback with folding kinetics, a strong decay in SRCD signal was observed within a 5–10 s measurement window (before the onset of induced photodegradation), when proteins were mixed with lipids in the presence of 150 mM NaF or 1 mM CaCl₂ (Fig. 3A). This resembled exponential decay and could be described with two potentially independent rate constants (Supplementary Table 1).

Since we observed similar behaviour in the absence of proteins (Supplementary Fig. S2, Supplementary Table 1), the effect does not involve protein folding. Considering the wavelength of the beam (195 nm), in the size range of large vesicular lipid bodies, and decay strength (tens of mdeg), the effect is caused by light scattering from vesicle fusion and/or aggregation, especially since the lipid mixtures and ionic conditions correlate with the observed steady states in turbidimetry (Fig. 1).

While lipids mixed with CaCl₂ alone produced quite noisy data with individual shots and replicates not overlaying very well, all protein samples had excellent reproducibility both in terms of shot quality and between replicates, which is evident from the small standard deviations. The data could be fitted with confidence, and in most cases, two rate constants were obtained: $k_1$ which describes a fast event ($>2 \text{ s}^{-1}$), and $k_2$, for a slower event ($<1 \text{ s}^{-1}$). The rate constants in the presence of proteins vary between lipid compositions (Fig. 3B). The accuracy of $k_2$ is rather low, given the limited time window of the experiments.

In the absence of proteins, the turbidity kinetics induced by Ca²⁺ were similar in DMPC:DMPG (4:1) and (9:1), and slower in DMPC:DMPG (1:1) (Supplementary Fig. S2, Supplementary Table 1), suggesting a role for electrostatic repulsion. As the two less charged lipid compositions aggregate with roughly the same rate, the effect is likely to occur due to saturation of the surface with Ca²⁺. The concentration of CaCl₂ in these experiments was 1 mM, while the total concentration of lipid negative headgroups was either 0.25, 0.1, or 0.05 mM. It appears that the membranes with less charge are fully neutralized, while the neutralization might be partial for the 1:1 mixture.

In protein-lipid samples, in the presence of NaF and CaCl₂, DMPC:DMPG (1:1) presents a higher $k_1$ than the other two lipid compositions. The effects of the two salts on $k_1$ increase with the molar percentage of DMPG (Fig. 3B, Supplementary Table 1).

In DMPC:DMPG (9:1), the aggregation effect was dominated by proteins, but when proteins were included, the levels were almost identical, indicating that the proteins could override the effect of CaCl₂ at moderate concentrations (Fig. 1B). 1.5 mM CaCl₂ decreased the turbidity induced by MBP or P0ct in all membrane compositions. Thus, while salt can influence the interactions between proteins and lipids, and protein-decorated vesicles with each other, the effect of Ca²⁺ with phospholipid headgroups is rather specific. Even small changes in Ca²⁺ levels, in the range 0.5–1.5 mM, can alter the protein-lipid interactions required for vesicle aggregation and fusion, giving that negatively charged lipid species are not predominant.

### 3.2. Folding of lipid-bound MBP and P0ct

We earlier observed lipid-induced increase in secondary structure for MBP and P0ct in the absence of salt [18,28]. To investigate the effect of ionic strength and divalent cations on folding, SRCD was carried out with 150 mM NaF and 1 mM CaCl₂. NaF was used due to the high UV absorbance of Cl⁻. NaF had a very similar effect to CaCl₂ (Fig. 2A and B). Under such conditions, the positions of the spectral peaks can be used to detect the presence of secondary structure. The observed changes in conformation correspond well to the turbidimetry assays under the same conditions, whereby the negative peak at 200 nm indicates unfolded structure and possibly lack of membrane binding.

MBP reached a similar secondary structure content in all lipid compositions (Fig. 2). In DMPC:DMPG (9:1), NaF and CaCl₂ decreased MBP helical content, potentially indicating less binding to lipids (Fig. 2C). Also P0ct obtained secondary structure in lipid mixtures, and the decrease of membrane negative charge (DMPG) correlated with the loss of P0ct helical structure - seen as the movement of the main minimum towards 200 nm (Fig. 2). NaF had a small unfolding effect on P0ct in all lipid mixtures.
the 1 mM CaCl₂ over proteins (Supplementary Table 1); this correlates with steady-state turbidity (Fig. 1C). The rate constants are similar regardless of the presence of protein, indicating that the aggregation under this condition is governed by the cation. On the other hand, 150 mM NaF inhibits aggregation of DMPC:DMPG (9:1) vesicles (Fig. 1C, Supplementary Table 1), and in salt-free conditions, the proteins display aggregation, which was not the case for the other two vesicle compositions (Fig. 3A). Taken together, the kinetics correlate well with turbidimetry, providing insights into determinants of membrane binding and stacking.

4. Discussion

Myelination requires a balance between myelin protein expression, protein-membrane interactions, and regulatory factors, such as calcium. Both MBP and P0ct are IDPs in solution, but upon binding to membranes, they fold into helical structures, altering the physical properties of the membrane [13,17,18,28,33]. The salt sensitivity of the protein-membrane interactions supports the role of electrostatic interactions in them, and the effects of calcium on the molecular interactions indicate more specific regulation.

Phospholipid vesicle aggregation was followed to see how salt content and Ca²⁺ influence the activity and folding of MBP and P0ct. Our results show that while vesicle turbidity increases with ionic strength, both in the steady state and in initial kinetics, the effect of salt varies with membrane surface net charge. Considering the combination of ionic strength and membrane charge, high salt concentrations effectively screen the protein-membrane interactions, while lower concentrations, in fact, have an inducing effect. The results are in line with extractability of MBP from myelin

Fig. 2. Lipid-induced protein folding. Three ratios of DMPC:DMPG were used: 1:1 (A), 4:1 (B), and 9:1 (C). No additive, green; 150 mM NaF, blue; 1 mM CaCl₂, red. Protein controls in water (black) are plotted for reference. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
and myelin swelling under different conditions [34–36].

The effects of calcium are likely to be more specific than the electrostatic competition caused by near-physiological ionic strength per se. Ca\(^{2+}\) alone induces spontaneous vesicle turbidity, showing that it affects the surface properties of the membrane. Calcium influences both the steady-state levels and the kinetics of MBP- and P0ct-mediated vesicle aggregation. The binding of divalent cations by MBP has been observed in earlier studies [21–23], and calcium affects both MBP solubility in purified myelin as well as myelin membrane spacing [36,37]. It, thus, appears that calcium might be able to affect the surface of both the protein and the membrane in a manner that can regulate the direct molecular interaction.

Time-resolved SRCD was used to follow the kinetics of vesicle aggregation. While it is at the moment unclear what the rate constants represent, the observed kinetics may be relevant to myelin membrane stacking. Considering the \(k_1\) values, P0ct is faster than MBP at inducing turbidity. The difference could arise from the smaller size of P0ct, which potentially tumbles in solution faster than MBP and folds more easily onto membranes. Being larger, MBP experiences more steric and repulsive effects from proteins already on the membrane.

Several studies have highlighted calcium as a regulator of proper myelination [5,7,8,30,38], and in addition to being crucial for signalling and cellular homeostasis, it has been suggested to directly affect MBP-membrane interactions [10,30,36]. Phosphoinositides are important constituents of myelin, involved in MBP-membrane interactions regulated by Ca\(^{2+}\) [30]. Various mechanisms have been proposed for the effects of calcium on myelin structure and formation; these include activation of calpains, the involvement of calmodulin, signalling pathways, cytoskeletal events, and effects on protein-lipid interactions [5,7–10,30].

It was proposed that a “sweet spot” concentration of calcium exists during normal myelination [9]. Deviations from this range might be relevant for dys/demyelination. Similar effects were observed in early studies on MBP extraction from nerve tissue [29,36]. Here, concentrations around the physiological [Ca\(^{2+}\)] were used to follow myelin protein-membrane interactions, and concentration dependence around 1 mM [Ca\(^{2+}\)] is evident. Together with the membrane surface charge, calcium may either promote or inhibit lipid membrane stacking induced by myelin proteins. The obvious next steps involve studies on the effects of calcium on myelin protein-membrane interactions in more physiological lipid compositions corresponding to the myelin cytoplasmic leaflet; the experiments here using simplified lipid mixtures pave the way for these studies.

MBP and P0 are not related by sequence; however, the physicochemical properties of MBP and P0ct are similar [13,17,18,27,28,33]. Both bind to membrane surfaces through electrostatic interactions, get embedded and fold, and affect properties of the lipid bilayer. Functional similarity extends to the effects of ionic strength and calcium. Our observations point towards common regulatory mechanisms for myelin proteins during myelin maturation and maintenance.

**Author contributions**

Study design, original text and figures: A.R., P.K.
Prepared samples and performed experiments: A.R.
Processed and analyzed data: A.R., N.C.J., S.V.H., P.K.
Competing financial interests
The authors declare no competing financial interests.

Data availability
The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

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