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Melamine-cored glucosides for membrane protein solubilization and stabilization: importance of water-mediated intermolecular hydrogen bonding in detergent performance†

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Membrane proteins play essential roles in a number of biological processes, and their structures are important in elucidating such processes at the molecular level and also for rational drug design and development. Membrane protein structure determination is notoriously challenging compared to that of soluble proteins, due largely to the inherent instability of their structures in non-lipid environments. Micelles formed by conventional detergents have been widely used for membrane protein manipulation, but they are suboptimal for long-term stability of membrane proteins, making downstream characterization difficult. Hence, there is an unmet need for the development of new amphipathic agents with enhanced efficacy for membrane protein stabilization. In this study, we designed and synthesized a set of glucoside amphiphiles with a melamine core, denoted melamine-cored glucosides (MGs). When evaluated with four membrane proteins (two transporters and two G protein-coupled receptors), MG-C11 conferred notably enhanced stability compared to the commonly used detergents, DDM and LMNG. These promising findings are mainly attributed to a unique feature of the MGs, *i.e.*, the ability to form dynamic water-mediated hydrogen-bond networks between detergent molecules, as supported by molecular dynamics simulations. Thus, MG-C11 is the first example of a non-peptide amphiphile capable of forming intermolecular hydrogen bonds within a protein–detergent complex environment. Detergent micelles formed *via* a hydrogen-bond network could represent the next generation of highly effective membrane-mimetic systems useful for membrane protein structural studies.

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Introduction

Membrane proteins are ubiquitous bio-macromolecules that reside within multifaceted cellular membranes. Approximately a quarter of the human proteome corresponds to membrane proteins.¹ These bio-macromolecules play essential roles in a variety of cellular functions and are implicated in a number of

diseases such as asthma, cancers, viral infections and neurodegenerative diseases.² Owing to their central roles in a number of physiological processes, membrane proteins constitute around 60% of approved drug targets and, therefore, their three dimensional structures are eagerly sought to assist in structure-based drug design. Despite significant and considerable recent improvements, isolation of functional membrane proteins in sufficient amounts and of the requisite quality necessary for

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their functional and structural studies is still challenging.^{3,4} Structure determination of membrane proteins has historically been and, to a certain extent, remains technically difficult, as demonstrated by the fact that membrane proteins with a known structure comprise a low fraction of all protein structures deposited in the Protein Data Bank (PDB). These challenges can arise at any point in the workflow from gene expression to structure determination, as exemplified by inadequate protein expression, poor protein extraction from native membranes, limited long-term protein stability, and recalcitrant protein crystallization. While these challenges have severely hindered membrane protein structure determination, technological advances in expression systems, solubilisation and purification techniques, and structure determination methods such as X-ray crystallography, single-particle cryogenic-electron microscopy (cryo-EM) and nuclear magnetic resonance (NMR) spectroscopy over the last decade have led to an ever-growing number of deposited membrane protein structures.^{5–7}

Detergents serve as essential tools for membrane protein solubilization, purification and structural study. These amphipathic compounds are widely used not only to extract membrane proteins from the membranes, but also to maintain protein integrity during purification and downstream characterization.^{8–10} OG (*n*-octyl- β -D-glucoside), DM (*n*-decyl- β -D-maltoside), and DDM (*n*-dodecyl- β -D-maltoside) are among the most widely used detergents for structural studies of membrane proteins.⁹ These conventional detergents have been successfully used for structural studies of relatively stable membrane proteins, but they tend to be inadequate to handle more challenging eukaryotic membrane proteins and multisubunit membrane protein complexes. Due to their canonical architecture of single head and tail groups, micelles formed by these detergents are markedly more dynamic than cell membranes.¹¹ In addition, detergent micelles, particularly those formed by harsh detergents, can remove closely associated lipid molecules important for membrane protein structure and function during protein extraction. Therefore, it is necessary to develop novel amphiphiles able to both form micelles with reduced dynamics and retain lipid molecules essential for protein function.

Over the past few decades, several membrane-mimetic systems have been developed including bicelles,¹² membrane scaffold protein (MSP)-based nanodiscs (NDs),¹³ polymeric amphiphiles [amphipols (Apol)s]^{14,15} and styrene-maleic acid copolymers (SMAs)],^{16,17} and peptide-based amphiphiles [lipopeptide detergents (LPDs),¹⁸ β -peptides (BPs),¹⁹ and Saposin A].²⁰ Among these membrane-mimetic systems, nano-assemblies formed by MSP, SMA and Saposin A, namely NDs, SMA lipo-particles (SMALPs) and Salipro, respectively, are particularly interesting as they include lipids that are either added to the isolated protein or directly extracted from the native membrane along with the protein, providing a more physiological environment. While successfully used for structural studies of membrane proteins, particularly *via* cryo-EM, many of these systems are ineffective at membrane protein extraction from the membranes and are not suitable for membrane protein crystallization. As additional alternatives to conventional detergents, small amphiphiles have been developed, as

exemplified by neopentyl glycol-based amphiphiles [*e.g.*, maltose neopentyl glycols (MNGs), glucose neopentyl glycols (GNGs), and neopentyl glycol-derived triglucosides (NDTs)],^{21–23} rigid hydrophobic group-bearing amphiphiles [*e.g.*, chobimalt and glyco-diosgenin (GDN)],^{24,25} carbohydrate-cored amphiphiles [*e.g.*, mannitol-based amphiphiles (MNAs)²⁶ and scyllo-inositol glycosides (SIGs)]²⁷ and facial amphiphiles (FAs).^{28,29} Among these small detergent molecules, two amphiphiles (LMNG and GDN) are particularly notable as they have been used for structural studies of more than 450 membrane proteins during the past 10 years.³⁰ These biochemical tools are particularly effective for structural elucidation of biologically and pharmaceutically important bio-macromolecules such as G protein-coupled receptors (GPCRs) and ATP-binding cassette (ABC) transporters, highlighting significant contributions of novel detergents to membrane protein structure determination. Very recently, we reported a couple of classes of 1,3,5-triazine-based detergents including triazine-based di-maltosides (TEMs),³¹ tris(hydroxymethyl)methanamine (TRIS)-bearing triazine-based glucosides (TTGs),³² and triazine-based tetra-maltosides (TZMs)³³ for membrane protein stability. In this study, we report another class of 1,3,5-triazine-based amphiphiles containing a melamine unit (2,4,6-triamino-1,3,5-triazine) in the central region, denoted as melamine-cored glucosides (MGs). Due to the presence of multiple hydrogen-bonding donors and acceptors in the central melamine unit, we hypothesized that these amphiphiles form a hydrogen-bond network within the micellar structure when assembled around membrane protein surfaces and that this network increases detergent–detergent interactions which in turn enhances membrane protein stability. When tested with four model membrane proteins [two transporters (leucine transporter (LeuT) and melibiose permease (MelB)) and two G protein-coupled receptors (β_2 adrenergic receptor (β_2 AR) and μ -opioid receptor (MOR))], MG-C11 proved highly effective at stabilizing the membrane proteins compared to two gold standard detergents (DDM and LMNG). Molecular dynamics (MD) simulations show that MG-C11 tends to form (dynamic) water-mediated hydrogen-bonding with other detergent molecules in the protein–detergent complexes (PDCs), which explains its remarkable protein stabilizing efficacy. Thus, MG-C11 not only has potential as a useful tool for membrane protein manipulation, but also provides a new strategy for the design of novel detergents optimal for membrane protein stability.

Results and discussion

Detergent design, synthesis and physical characterization studies

New amphiphiles containing two alkyl chains and three glucose units as the hydrophobic and hydrophilic groups were synthesised (Fig. 1). The alkyl chains were conjugated into the triazine ring *via* an amino linkage, while the three glucose head groups were introduced into the same ring *via* a tris(hydroxymethyl)aminomethane (TRIS) linker. Due to the presence of strongly nucleophilic amine functional groups, we could efficiently attach both the alkyl chain and TRIS linker into the



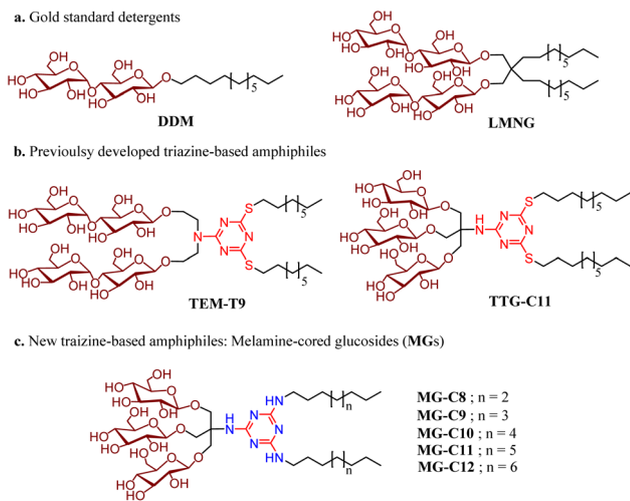


Fig. 1 Chemical structures of (a) two gold standard detergents (DDM and LMNG) and (b) previously developed triazine-based amphiphiles (TEM-T9 and TTG-C11), and newly prepared melamine-cored glucosides (MGs). The MGs differ from TEM-T9 in terms of head group identity (glucoside vs. maltoside) and from TTG-C11 in terms of the incorporation of a functional group used to connect the alkyl chains with the triazine unit (thioether vs. amino). Due to the connection of the two alkyl chains *via* the amino linkage and use of a TRIS linker for the introduction of the three glucose units, the new detergents contain a melamine unit (2,4,6-triamino-1,3,5-triazine) in the central region. The atoms/groups of the melamine unit potentially acting as hydrogen bond donors or acceptors are indicated in blue.

triazine ring *via* nucleophilic aromatic substitution reactions. The resulting triazine-based glucoside detergents contain a core structure of 2,4,6-triamino-1,3,5-triazine (A.K.A, melamine), and are thus denoted as melamine-cored glucosides (MGs). This core unit has a unique structure distinct from that of other detergent scaffolds since it contains multiple hydrogen bond donors and acceptors. The secondary amino (NH) and tertiary amino (N) groups presented in an alternating pattern around/on the heterocyclic ring are typical hydrogen-bonding donors and acceptors, respectively (Fig. 1). This distinctive feature has facilitated the wide use of melamine-bearing compounds in supramolecular assemblies.^{34–37} Of note, the MGs differ from previously developed triazine-based amphiphiles (TEMs and TTGs). Both MGs and TTGs have a glucoside head group, while the TEMs have a maltoside head group. The MGs differ from the TTGs in terms of the functional group used to introduce the two alkyl chains into the triazine ring (thioether (TTGs) vs. amino (MGs)). The MGs are structurally similar to the TTGs, but greatly differ in terms of the way they interact in micelles given the presence of hydrogen bond networks formed between the amino groups in the melamine core. The alkyl chains of the MGs varied from C8 to C12 to allow identification of the optimal alkyl chain length for membrane protein study. Detergent alkyl chain length is a critical factor determining balance between detergent hydrophobicity and hydrophilicity (*i.e.*, hydrophilic-lipophilic balance (HLB)). HLB values of the individual MGs were obtained using Griffin's method and are summarized in Table S1†. All MGs were found to give high HLB values (13.6–

15.2) compared to DDM, LMNG and previously developed TTG-C11. For example, the HLB value of MG-C11 (13.9) is higher than those of its thioether version (TTG-C11; 12.7), DDM (13.4) and LMNG (13.6), reflecting the rather hydrophilic nature of the new detergents.

The MGs were prepared in four efficient synthetic steps starting from 2,4,6-trichloro-1,3,5-triazine, comprising dialkylation, TRIS coupling, glycosylation and global deprotection (Fig. 2a). Briefly, an alkyl amine (R-NH₂) and TRIS were successively reacted with 2,4,6-trichloro-1,3,5-triazine in the presence of diisopropylethylamine as a base to produce the dialkylated triazine-based triol derivative (B). The resulting compound is subjected to β -selective glycosylation to give the corresponding glycosylated product in ~85% yield, followed by complete removal of the benzoyl protecting group using sodium methoxide (NaOMe) (see the ESI† for details). The ease of synthesis along with high synthetic yields made it possible to synthesize the designed amphiphiles in multi-gram quantities.

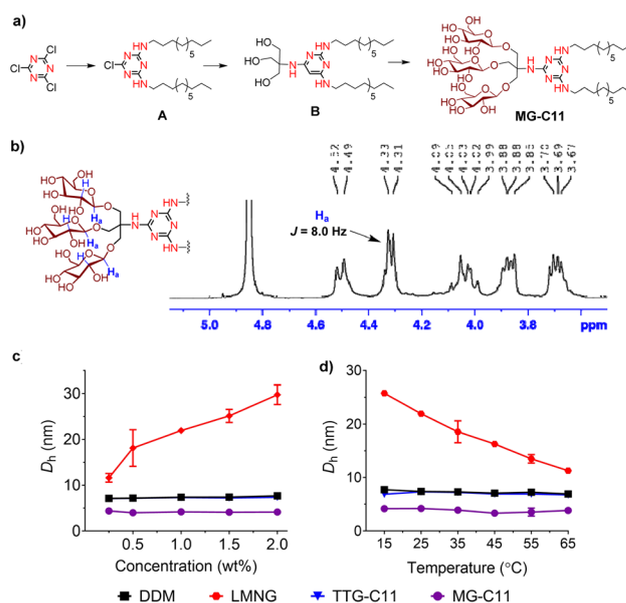


Fig. 2 (a) Synthetic scheme for the preparation of MG-C11, (b) partial ¹H NMR spectrum of MG-C11 focusing on the anomeric region (3.5–5.1 ppm), and (c and d) concentration and temperature dependency of micelle sizes formed by representative detergents (DDM, LMNG, TTG-C11 and MG-C11). (a) MG-C11 was synthesized from cyanuric chloride *via* four synthetic steps: dialkylation, TRIS coupling, glycosylation and deprotection. Dialkylation and TRIS coupling generated compounds A and B, respectively. (b) ¹H NMR spectrum of MG-C11 obtained using CD₃OD as an NMR solvent. The chemical shift (~4.32 ppm) and the coupling constant (8.0 Hz) of the NMR peak assigned to the β -anomeric protons (H_a) indicate the formation of a β -glycosidic bond during glycosylation, as expected from the mechanism of neighbouring group participation. The β -anomeric protons (H_a) are indicated in blue in the chemical structure of the detergent head group (the inset). (c and d) Hydrodynamic diameters (D_h) of self-assemblies formed by DDM, LMNG, TTG-C11 and MG-C11 were measured over a range of detergent concentrations, (0.3 to 2.0) wt% or temperatures (15 to 65 °C). Solution temperature and detergent concentration were kept at 25 °C and 1.0 wt%, respectively, in the individual studies for detergent comparison. Error bars (S.D.; n = 5) are shown only when they are larger than the symbols.



NMR spectra corroborate the β -stereochemistry of the glycosidic bonds of the MGs. The NMR spectrum of MG-C11 is shown in Fig. 2b, as an example. The anomeric proton peak appears at 4.32 ppm with a vicinal coupling constant ($^3J_{\text{aa}}$) of 8.0 Hz, which is in good agreement with the formation of the β -glycosidic bond (Fig. 2b). An anomeric proton with an α -glycosidic linkage gives higher chemical shift (5.15–5.20 ppm) and a smaller coupling constant (4.0 Hz) than the β -anomeric proton.

The individual MGs were soluble to at least 10%, with the exception of MG-A12 which had a lower water solubility of \sim 5%. The solutions containing these detergents were stable at room temperature for a month. The tendency of the MGs to form self-assemblies was investigated by measuring their critical micelle concentrations (CMCs), and the size of the self-assemblies formed by these detergents was estimated *via* hydrodynamic radii (R_{h}). Detergent CMC and R_{h} were obtained by utilizing encapsulation of the water-insoluble dye, diphenylhexatriene (DPH), in the micelle interiors and dynamic light scattering (DLS) experiments, respectively.³⁹ The results are summarized in Table 1. The CMCs of the MGs follow a typical trend; their CMCs decrease with increasing alkyl chain length. This trend is likely due to the fact that detergent hydrophobicity is proportional to the alkyl chain length. The MG CMCs were estimated to be in the range of 40 to 600 μM , significantly higher than those of other two alkyl chain-bearing detergents such as LMNG (\sim 10 μM) and TTG-C11 (\sim 4 μM).

For example, the CMC of MG-C11 is roughly 40 times higher than that of TTG-C11 (150 *vs.* 4 μM) although these two detergents have the same head and tail groups. Such a large CMC difference likely originates from the polarity difference between the two functional groups (amino *vs.* thioether) used to incorporate the alkyl chain into the detergent scaffold; the amino group is highly polar, while the thioether group is slightly polar. As a result, double alkyl-chained MG-C11 gave a surprisingly high CMC comparable to single alkyl-chained DDM (150 *vs.* 170 μM). Notably, most novel detergents developed to date including LMNG (10 μM) and NDT-C11 (6 μM) have lower CMCs than DDM. The high CMCs suggest that the MGs have a reduced tendency to self-assemble which may facilitate detergent exchange, often

necessary for membrane protein structural study.⁴⁰ The MGs are also different from the TTGs and LMNG in terms of micelle sizes. The R_{h} of MG-C11 was estimated to be 2.4 nm, substantially smaller than those of TTG-C11 (3.7 nm) and LMNG (9.8 nm) (Table 1). The micelle sizes of the MGs are comparable to conventional detergents known to form small micelles, as exemplified by OG (2.6 nm) and LDAO (1.9 nm).^{30,41} It is worth mentioning that small micelle-forming detergents are beneficial for membrane protein research, as excess detergent micelles can be efficiently reduced from protein samples through molecular weight cut-off (MWCO) filters. This property is also considered beneficial for membrane protein structural study *via* X-ray crystallography, cryo-EM and NMR spectroscopy.^{42,43} Interestingly, the sizes of the micelles formed by the MGs only slightly varied with increasing detergent alkyl chain length. The R_{h} of detergent micelles increases by only 0.1 nm with every additional methylene unit to the alkyl chain, smaller than in other classes of amphiphiles.

Micellar structures formed by MG-C11 and MG-C12 were calculated *via* MD simulations. The calculations were carried out assuming that the aggregation number (AN) of detergent micelles is either 20, 25, 30 or 40. These ANs were selected based on the experimentally determined R_{h} of micelles formed by MG-C11 or MG-C12. The calculation results are visualized in Fig. 3 and their micellar properties are summarized in Tables S2 and S3.† Interestingly, we observed little difference in the distances of individual detergent components (glucoside (Glu), TRIS-triazine (Tt) and alkyl tail (R)) from the micellar centre when the detergent alkyl chain length increases from C11 (MG-C11) to C12 (MG-C12) (Table S2†). In addition, the radii of gyration (R_{g}) of micelles increase by 0.2–0.3 Å with the same variation in the alkyl chain length of the MGs (Table S3†). The small increase in R_{h} of the MGs with increasing the alkyl chain length is consistent with this trend (Table 1). This result indicates that the micelle size is mainly determined by the hydrophilic group (*i.e.*, glucoside-TRIS-triazine) rather than the alkyl chain of the MGs. Favourable interactions between the detergent hydrophilic groups may be responsible for this observation. Of note, the calculated R_{g} values of both MG-C11 and MG-C12 micelles were smaller than the experimentally determined R_{h} of these detergents (2.4 and 2.5 nm, respectively), typical for spherical micelles. With detergent AN increasing from 20 to 40, the R_{g} values of detergent micelles increase only 1.2 times for both MGs, indicating that new detergent molecules incorporated into MG micelles are mainly positioned to decrease the empty spaces or micelle defects in the micelle interiors rather than to increase micelle volumes (Table S3†). This indication was further supported by the substantial decreases in the solvent accessible surface areas (SASAs) of MG-C11 and MG-C12 micelles with increasing detergent AN. Interestingly, the SASA of MG-C11 micelles is substantially smaller than that of MG-C12 micelles with the same AN, indicating that MG-C11 could form more tightly packed micelles than MG-C12. Based on the distances of the glucoside head group from the micelle centre (Table S2 and Fig. S2†), the ANs of MG-C11 and MG-C12 micelles are estimated to be in the range of 25 to 30,

Table 1 Molecular weights (MWs), critical micelle concentrations (CMCs), hydrodynamic radii (R_{h} ; mean \pm S.D.; $n = 5$) and water-solubility of MGs, TTG-C11, LMNG and DDM

Detergent	M.W. ^a	CMC (mM)	CMC (wt%)	R_{h} (nm) ^b	Solubility
MG-C8	941.0	\sim 0.6	\sim 0.06	2.1 \pm 0.5	\sim 10
MG-C9	969.1	\sim 0.5	\sim 0.05	2.2 \pm 0.2	\sim 10
MG-C10	997.1	\sim 0.3	\sim 0.03	2.3 \pm 0.1	\sim 10
MG-C11	1025.2	\sim 0.15	\sim 0.015	2.4 \pm 0.1	\sim 10
MG-C12	1053.3	\sim 0.04	\sim 0.004	2.5 \pm 0.1	\sim 5
TTG-C11	1059.3	\sim 0.004	\sim 0.0004	3.7 \pm 0.1	\sim 10
LMNG	1005.2	\sim 0.01	\sim 0.001	9.8 \pm 0.2	\sim 10
DDM	510.6	0.17	0.0087	3.4 \pm 0.1	\sim 10

^a Molecular weight of detergents. ^b Hydrodynamic radius of detergents measured at 1.0 wt% by dynamic light scattering.



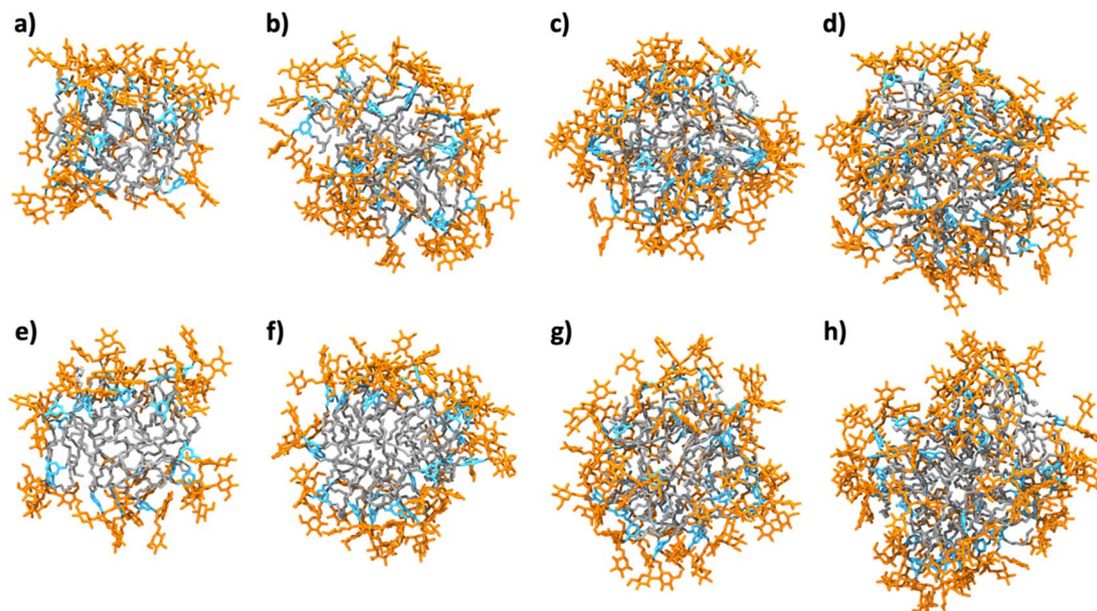


Fig. 3 Snapshots of MG-C11 micelles with an aggregation number (AN) of (a) 20, (b) 25, (c) 30, and (d) 40, and MG-C12 with an AN of (e) 20, (f) 25, (g) 30, and (h) 40 obtained after 500 ns-long MD simulations. The glucoside head group, TRIS-triazine and alkyl chain components are shown in orange, sky-blue and grey, respectively (water and ions are not shown for clarity). The ANs were chosen based on the experimentally determined R_h of MG-C11 micelles. The detergent alkyl chains within the micelle interiors are effectively covered by the hydrophilic groups on increasing the AN, thereby decreasing the solvent accessible surface areas (SASAs) of the detergent alkyl chains in the micelle interiors (see the text).

significantly smaller than those of DDM (80–150) and LMNG (~400).⁴⁴ The dynamics of the detergents in their micelles were estimated by calculating root mean square deviation (RMSD) of detergent alkyl chains. With the detergent AN increasing from 20 to 40, the RMSDs of MG-C11 and MG-C12 were reduced from 10.3 to 7.3 Å and 10.1 to 8.0 Å, respectively (Table S4†). These RMSD values are significantly smaller (14.1 Å) than that obtained from micelles formed by 130 DDM molecules. This result indicates that MG-C11 forms micelles that are less dynamic than those formed by DDM, partly due to dynamic hydrogen-bond formation between MG-C11 molecules (*vide infra*).

The small micelle size of MG-C11 compared to those of DDM, LMNG, and TTT-C11 is effectively maintained with variation in detergent concentration or solution temperature (Fig. 2c and d). The small micelle size formed by MG-C11 varied little with increasing detergent concentration from 0.3 to 2.0 wt% or increasing solution temperature from 15 to 65 °C. In contrast, self-assemblies formed by LMNG increased or decreased with the same variations of detergent concentration and solution temperature. This result indicates that MG-C11 forms small micelles with a globular/elliptical shape, while micelles formed by LMNG are cylindrical.⁴⁵ The MGs were further investigated in terms of their micelle size distributions. All the MGs showed unimodal size distributions of their micelles in the number- or volume-weighted DLS profiles, suggesting homogenous populations of their micelles (Fig. S3a and S3b†). Large aggregates shown in the intensity-weighted DLS profiles of the MGs are not due to size heterogeneity of detergent micelles, but result from the highly sensitive nature of light scattering depending on the aggregate size (Fig. S3c†).

Detergent evaluation with membrane proteins

Detergent evaluation started with melibiose permease from *Salmonella typhimurium* (MelB_{St}).^{46a-e} The permease overexpressed in *E. coli* membranes was extracted using 1.5 wt% of the individual MGs at 0 °C for 90 min. The extracts were further incubated at an elevated temperature (45, 55, or 65 °C) for another 90 min. The amounts of soluble MelB_{St} were determined using Western blot analysis following ultracentrifugation and represented as percentages (%) of the total MelB_{St} present in the untreated membranes. The amount of soluble MelB_{St} obtained from the low temperature experiment (0 °C) gives information about detergent efficiency for protein extraction, while those from the temperature variation experiments (45, 55, or 65 °C) are directly correlated with detergent efficacy for protein stabilization. Thus, this protocol provides information about both protein extraction efficiency and protein stabilization efficacy of the tested detergents. When MelB_{St} was extracted using the individual MGs or DDM (as a control) at 0 °C, DDM was most efficient at extracting and solubilizing MelB_{St} (Fig. 4). The MGs were inferior to DDM in this context, but MG-C11, the best of the five MGs, extracted MelB_{St} from the membranes in ~75% yield. When the 0 °C-extracts were additionally incubated at 45 °C, the amounts of soluble MelB_{St} increased substantially in the case of the MG-treated samples. For instance, MG-C11 and MG-C12 gave amounts of soluble MelB_{St} increasing from ~75% to 100% and from ~40% to ~65%, respectively. Increased detergent water-solubility and/or enhanced membrane dynamics induced by the elevated temperature are likely responsible for increased MelB_{St} solubilization. Detergent behaviour between the MGs and DDM



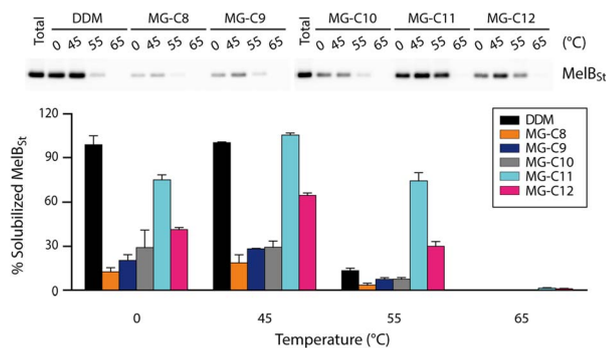


Fig. 4 Thermo-solubility of MG-extracted MelB_{St}. A conventional detergent (DDM) was used as the control. MelB_{St} produced in *E. coli* membranes was extracted using 1.5 wt% of each detergent at 0 °C. The resulting extracts were subjected to further incubation at an elevated temperature (45, 55, or 65 °C). Following ultracentrifugation to remove insoluble debris, the samples were analyzed by SDS-PAGE and Western blotting (top panel). The quantities of soluble MelB_{St} were measured from bands in the Western blot and expressed as percentages of the total MelB_{St} present in the membranes prior to detergent treatment in the histogram (bottom panel). Error bars, SEM, *n* = 3.

was differentiated in the thermo-solubility experiment at 55 °C. At this high temperature, a small amount of the transporter (~10%) remains in a soluble state following thermal treatment, indicating that most of the DDM-solubilized MelB_{St} undergoes protein denaturation and/or aggregation. In contrast, MG-C11 effectively retained protein solubility at this high temperature, resulting in ~75% soluble MelB_{St}. The other MGs, except MG-C12, were ineffective at preserving MelB_{St} in a soluble state. The same experiment at 65 °C resulted in a complete loss in soluble MelB_{St} in all tested detergents. This result indicates that MG-C11 is moderately efficient at MelB_{St} extraction and notably effective at stabilizing the transporter.

The MGs were further evaluated with another transporter, the bacterial leucine transporter (LeuT) *Aquifex aeolicus*.⁴⁷ Protein stability was assessed by measuring the ability to bind a radioactive substrate ([³H]-leucine (Leu)) *via* scintillation proximity assay (SPA).⁴⁸ LeuT purified in DDM was subjected to detergent exchange *via* sample dilution and the final concentrations of the MGs in the sample solutions were CMCs + 0.04/0.2 wt%. [³H]-Leu binding ability of the transporter in the individual MGs was monitored over a 12 day incubation period at room temperature (Fig. 5a and b). LeuT in DDM initially showed a good ability to bind Leu, but rapidly lost binding ability over the course of the incubation. After 6 days of incubation, LeuT in DDM almost completely lost its substrate binding ability. LeuT in the individual MGs, despite variation in their initial abilities to bind the substrate, was more effective at maintaining its stability than in DDM, with the best performance observed for MG-C11. The overall trend of detergent efficacy for LeuT stabilization was similar when the MGs were used at either CMCs + 0.04 wt% or 0.2 wt% (Fig. 5a and b). Along with the MelB_{St} results (Fig. 4), these findings indicate that MG-C11 is highly effective at stabilizing membrane transporters.

We selected MG-C11 for the next evaluation as this detergent showed the most promising results for solubilisation and/or

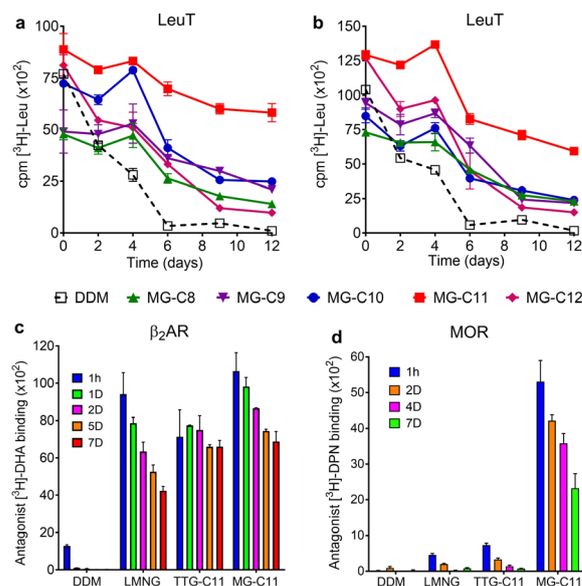


Fig. 5 Stability of LeuT (a and b), β_2 AR (c) and MOR (d) solubilized in the MGs over time. DDM was used as a control for LeuT stability analysis, while LMNG and TTG-C11 (previously developed TTG) were used as additional controls for β_2 AR and MOR stability assays. As for β_2 AR and MOR stability, MG-C11 was selected as a representative MG for detergent comparison. The detergents were tested at CMC + 0.04 wt% (a), 0.2 wt% (b), or 0.1 wt% (c and d). Stability of LeuT, β_2 AR, or MOR was assessed by measuring the ability of the protein to bind the radio-labelled substrate ([³H]-leucine (Leu)), the antagonist ([³H]-dihydroalprenolol (DHA) or [³H]-diprenorphine (DPN)) as indicated in the y-axis title. Protein stability was monitored at regular intervals during the designated time period of incubation at 4 °C (MOR) or room temperature (LeuT and β_2 AR). Error bars: SEM, *n* = 3–6 (LeuT), *n* = 2 (β_2 AR) or 3 (MOR).

long-term stabilization of two transporters (MelB_{St} and LeuT). As a model membrane protein for this evaluation, we turned to a G protein-coupled receptor (GPCR), the human β_2 adrenergic receptor (β_2 AR).⁴⁹ First, we investigated detergent efficiency for receptor solubilisation. For this purpose, we used 1.0 wt% MG-C11 to extract the receptor from the membranes at two different temperatures (4 or 25 °C) (Fig. S4†). DDM and LMNG were used as controls as these two detergents are widely used for GPCR extraction.⁵⁰ When the receptor was extracted from the membranes at 4 °C, LMNG of the tested detergents was most efficient at receptor extraction, with MG-C11 yielding a similar amount of soluble β_2 AR to DDM. When the solubilisation experiment was carried out at 25 °C, all the tested detergents yielded comparable amounts of the soluble receptor. These results indicate that MG-C11 can be used as efficiently as DDM for β_2 AR extraction. Interestingly, in marked distinction from DDM and LMNG, the receptor solubilized by MG-C11 yielded two bands (Fig. S4†). These bands are likely to correspond to the monomeric and dimeric forms of the receptor with molecular weights of ~46 and ~92 kDa, respectively. It is known that there is a substantial amount of dimeric β_2 AR in physiological cell membranes that plays a critical role in receptor and cellular function.⁵¹ The marked ability of MG-C11 to stabilize membrane protein complexes may allow effective isolation of dimeric β_2 AR. This result is important as structural study of



dimeric/oligomeric GPCRs is limited by detergent-mediated dissociation into monomers. Thus, MG-C11 may open the window to readily access dimeric GPCRs. Next, we turned to study MG-C11 efficacy for receptor stabilization. In this evaluation, we included TTG-C11 as a positive control, in addition to DDM and LMNG. LMNG is a widely used novel detergent particularly for GPCR structural study, while TTG-C11 is a previously developed triazine-based glucoside. Notably, LMNG has been used for successful determination of more than 145 GPCR structures over the past 10 years.³⁰ β_2 AR purified in LMNG was diluted in a buffer solution containing LMNG, TTG-C11, or MG-C11 to reach a final detergent concentration of 0.1 wt%. The resulting protein samples were then incubated for 7 days at room temperature and receptor stability was monitored at regular intervals during the incubation (Fig. 5c). The ability of the receptor to bind a radioactive antagonist ($[^3\text{H}]$ -dihydroalprenolol (DHA)) was used to assess β_2 AR stability.^{52–54} As expected, β_2 AR in DDM showed low affinity for the radiolabelled ligand upon detergent exchange and lost its ligand binding ability rapidly. β_2 AR in LMNG showed markedly enhanced DHA binding ability upon detergent exchange, as expected from the wide use of this NG detergent for GPCR structural studies. However, this initial binding ability of LMNG gradually decreased during the 7 day incubation. TTG-C11 was overall comparable to LMNG in this regard. When MG-C11 was used to encapsulate the receptor, this detergent showed higher initial ligand binding of the receptor than LMNG and TTG-C11, and this initial capability was effectively preserved over the course of the 7 day incubation.

Based on the encouraging results of MG-C11 for β_2 AR stability, we further evaluated this detergent for another GPCR, the mouse μ -opioid receptor (MOR).⁵⁵ This GPCR is particularly challenging to stabilize in detergent micelles. The LMNG-extracted receptor was purified in the same detergent and the resulting LMNG-purified MOR was exchanged from LMNG to the respective test detergent (DDM, LMNG, TTG-C11, or MG-C11) *via* sample dilution. The final detergent concentration for detergent efficacy comparison was 0.1 wt%. MOR stability was assessed *via* a similar method used for β_2 AR stability using a receptor-specific antagonist (*i.e.*, $[^3\text{H}]$ -diprenorphine (DPN)) instead of $[^3\text{H}]$ -DHA. MOR in DDM completely lost ligand binding ability upon detergent exchange. LMNG and TTG-C11 were better than DDM, consistent with the results of β_2 AR with these detergents, but the receptor in these novel detergents still suffered from rapid loss of DPN binding over the 7 day incubation at 4 °C (Fig. 5d). In contrast, upon detergent exchange from LMNG to MG-C11, MOR showed markedly enhanced ability to bind DPN compared to the receptor in LMNG or TTG-C11; the DPN binding increased by 11 and 7 times, respectively, when MG-C11 was used compared to LMNG and TTG-C11. Even after the 7 day incubation at 4 °C, the DPN binding ability of the receptor in MG-C11 was five/three times higher than the initial ability of the receptor in LMNG and TTG-C11. Indeed, this detergent is the most effective of recently developed novel detergents at stabilizing MOR.^{56–59}

The remarkable efficacy of MG-C11 for MOR stabilization compared to LMNG prompted us to explore molecular

interactions in the MOR-detergent complex *via* MD simulations. 500 ns-long MD simulations were performed using MOR (PDB: 6DDE) complexed with 96 molecules of MG-C11 or LMNG.⁶⁰ The simulation results for complex formation of LMNG/MG-C11 with the receptor show effective encapsulation of the hydrophobic surface of the receptor by LMNG or MG-C11 molecules (Fig. S5†). When protein–detergent interactions were analysed according to the amino-acid residues of the receptor, we found that two transmembrane helices (α_5 and α_6) interact in a higher frequency with the alkyl chains of MG-C11 than those of LMNG (Fig. S6†). The other helices showed little noticeable difference between these two detergents. In addition, the numbers of LMNG and MG-C11 molecules in contact with the receptor surfaces were calculated from the MD simulations (Fig. S7†). When using the last 200 ns trajectory data, MG-C11 gave an average contact number of ~ 79 , substantially larger than that of LMNG (~ 72). A similar result was obtained for comparison of the atom numbers of the detergent alkyl chains interacting with the receptor. The average number of detergent contact atoms increases from 677 (LMNG) to 723 (MG-C11). This increased frequency/contact number means a stronger interaction of MG-C11 with the receptor surface compared to that of LMNG, a feature favourably associated with enhanced receptor stability. Next, we investigated detergent–detergent interactions by calculating the numbers of intermolecular hydrogen-bonding within detergent micelles surrounding the receptor. Due to the presence of hydrogen bond donors and acceptors in the core structure, MG-C11 is likely to form hydrogen bonds with nearby molecules. Rather surprisingly, hydrogen-bonding that persists for more than 1 ns is absent in the simulated complex between MG-C11 and MOR. Thus, we next focused on dynamic hydrogen-bonding in the complex that constantly undergoes bond formation and breakage during the simulation trajectory (Fig. 6a and b). The extent of such dynamic intermolecular interactions was quantified by counting the average number of hydrogen bonds present in the complex at a given time in the last 200 ns trajectory. Direct hydrogen-bonding between detergent molecules in the protein–detergent complexes (PDCs), denoted as triamino (NH)–triamino (NH) or triamino (NH)–triazine (N) interaction, is found to be negligible. This is likely due to the fact that the presence of the bulky head group (*i.e.*, TRIS-triglucoside) hinders these hydrogen-bonding pairs from approaching each other (Fig. 6c). When mediated by a water molecule, however, substantially larger numbers are obtained for the two types of hydrogen-bonding interactions of this MG, denoted as triamino (NH)–water–triamino (NH) and triamino (NH)–water–triazine (N) interactions, respectively (Fig. 6d). The average number of triamino (NH)–water–triamino (NH) hydrogen-bonding is ~ 18 during the last 200 ns trajectory (Fig. 6e). This number dramatically increases to ~ 83 for the hydrogen-bonding between the triamino and triazine units of MG-C11 (triamino (NH)–water–triazine (N)). Thus, at any given instant, the total number of water-mediated hydrogen bonds between detergent molecules in the complex reaches 101, higher than the number of detergent molecules (96) employed to build the PDCs in the simulations. This result indicates that every MG-C11 molecule, at least, is competent to form a single



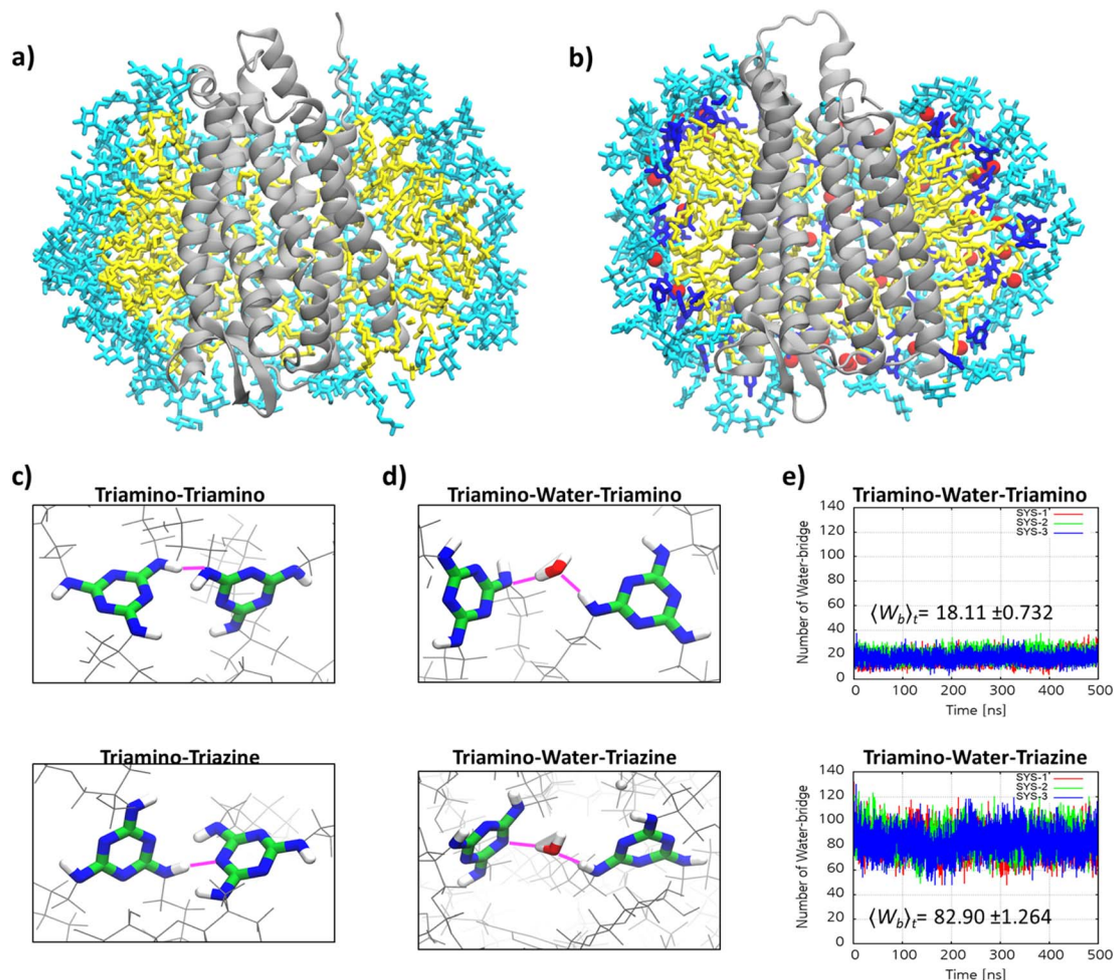


Fig. 6 MOR complex with LMNG (a) or MG-C11 micelles (b) obtained from MD simulations and hydrogen bonding analysis between MG-C11 molecules in the complex (c–e). Red spheres in (b) indicate oxygen atoms of water molecules participating in the hydrogen bonding network with the melamine unit. The number of these water molecules was found to be 87 within a single PDC. Of note, only a fraction of these water molecules are shown for clarity. (c and d) Representation of four categories of intermolecular hydrogen bonding of MG-C11. Intermolecular hydrogen bonding can form directly between the two alkyl amine (NH) groups (triamino (NH)-triamino (NH) or (c) between the alkyl amine (NH) and triazine amine (N) groups (triamino (NH)-triazine (N)). (d) Alternatively, these interactions can be mediated by a water molecule, giving two types of water-mediated hydrogen bonding (triamino (NH)-water-triamino (NH) and triamino (NH)-water-triazine (N)). (e) The numbers of water-mediated hydrogen bonding interactions between MG-C11 molecules surrounding MOR as a function of simulation time. The number of triamino (NH)-water-triamino (NH) interactions in the complexes is ~ 18 , while the number of triamino-water-triazine interactions is ~ 83 , giving a total of 101 hydrogen bonds mediated by water in the protein–detergent complex (PDC). The numbers were obtained from the simulation data of the last 200 ns trajectory.

dynamic hydrogen bond with another detergent molecule in the PDC environment, with the assistance of water molecules that penetrate into the triazine layer of the detergent micelles. It is likely that this hydrogen-bonding network in micelles formed by MG-C11, although dynamic and water-mediated, contributes to enhanced MOR stability *via* increased detergent–detergent interactions. It is noteworthy that hydrogen-bonding is much stronger than van der Waals interactions and thus hydrogen-bonding between detergent molecules, even if present in a small number, could make a big difference with respect to detergent behaviour toward membrane protein stability. Almost every detergent (*e.g.*, DDM or LMNG) can only harness relatively weak van der Waals interactions for assembly or micelle formations.

Discussion

Detergent efficacy for protein stabilization is dependent on the detergent alkyl chain length, which is closely associated with the detergent HLB and hydrophobic length. This is a reason why we typically find an optimal alkyl chain length in detergent studies for membrane protein stability. The exact alkyl chain length most effective for protein stability differs from one class of detergents to another and also varies depending on the identity of the target membrane proteins, but typically falls in the range of C10 to C13. This narrow range of optimal alkyl chain length is mainly due to small variations in the hydrophobic width of membrane proteins (28 to 32 Å).⁶¹ The compatibility of the hydrophobic dimensions of proteins and detergents is necessary for energetically favourable protein–



detergent interactions. As for the MGs, the optimal alkyl chain length was observed to be C11, the same alkyl chain length found in other detergent studies.^{23,32,59,62,63} MG-C11 was more effective than DDM at stabilizing all tested membrane proteins. Furthermore, this detergent conferred enhanced stability to two GPCRs (β_2 AR and MOR) compared to LMNG and TTG-C11. In addition, this detergent showed a reasonable extraction efficiency with both MelB_{St} and β_2 AR. Small micelle formation, with an R_h of 2.4 nm, and the rather high CMC (0.15 mM) are additional aspects of MG-C11 that are favourable for membrane protein study, distinctive from those of other novel detergents reported to date. Taken together, these results indicate that MG-C11 holds significant potential for extraction, purification and structural study of GPCRs. Notably, this detergent allowed solubilisation of the dimeric form of β_2 AR when the receptor was extracted from the membranes, implying a profound impact of this detergent on structural study of GPCR dimers.

It is difficult to pinpoint which structural feature of MG-C11 is mainly responsible for enhanced membrane protein stability observed here since protein stability is determined by a combination of multiple factors. One important factor determining detergent efficacy for protein stabilization is the hydrophobicity of the detergent hydrophobic group.³² The presence of a small number of polar groups (*e.g.*, amino and hydroxyl groups) in the detergent lipophilic region could be detrimental for membrane protein stability, as these weaken detergent interactions with membrane protein surfaces (*i.e.*, protein–detergent interactions). As a result, detergents including amino functional group(s) in the detergent core or lipophilic region are rarely used for membrane protein study. The presence of relatively nonpolar groups, as exemplified by ether and thioether groups, represent a slight compromise in detergent efficacy for membrane protein stabilization. In the MGs, each alkyl chain was conjugated to the triazine core *via* a polar amino linkage. Thus, the lipophilic groups of the MGs are relatively polar compared to those of LMNG and TTG-C11 where the corresponding alkyl chains are introduced into the detergent cores directly (LMNG) or *via* a thioether linkage (TTG-C11). The high CMCs and large HLBs of the MGs relative to those of LMNG and TTG-C11 support the relatively high polarity of the MG lipophilic groups. Given the possession of high polarity, it is remarkable that MG-C11 was superior to LMNG and TTG-C11 in membrane protein stability. In order to explore the reasons for this surprisingly enhanced stability, we investigated the molecular interactions within the MOR-MG-C11 complex *via* MD simulations and compared them with the simulation results obtained for the MOR-LMNG complex. We found that MG-C11 interacts with the receptor more frequently and extensively than LMNG, thereby increasing protein–detergent interactions. More importantly, MG-C11 micelles around MOR form rather extensive water-mediated hydrogen-bond networks between the melamine units. Therefore, the increased detergent–detergent interaction is likely to significantly contribute to the enhanced protein stabilization efficacy of MG-C11 compared to LMNG.

Development of detergents capable of forming a hydrogen-bonding network is remarkable for a couple of reasons. There is

only one class of detergents (*i.e.*, β -peptides (BPs)) reported to form hydrogen-bonding networks in detergent micelles.¹⁹ While these peptide-based detergents showed promising results in the initial evaluation with multiple membrane proteins, they exhibit poor water-solubility and are difficult to synthesize on a large scale. In contrast, MG-C11, the first example of a non-peptide-based detergent with the ability to form hydrogen-bond networks in PDCs, is convenient for large scale synthesis (requiring only four synthetic steps) and is highly water-soluble (>10%). Detergent design capable of forming intermolecular hydrogen-bonding is challenging as hydrogen-bonding can only be attained under strict restrictions in terms of orientation and distance between the hydrogen bond donor and acceptor. This challenge is particularly difficult to achieve in micellar architecture with a large curvature. Consequently, a simple insertion of a hydrogen-bonding motif into a detergent scaffold is unlikely to induce the realization of hydrogen-bond formation between detergent molecules. For instance, there is no plausible evidence that tripod amphiphiles (TPAs) and pre-assembled detergents (PADs) containing an amide linkage and triazole unit in the detergent core, respectively, form hydrogen bonds in a micellar or PDC state.^{64,65} It is remarkable that MG-C11, a small micelle-forming detergent, is able to form intermolecular hydrogen bonding mediated by water molecules in micelle interiors. Due to their strong yet reversible nature, various bio-macromolecules such as ribonucleic acids, proteins and carbohydrates have evolved to widely utilize hydrogen-bonding interactions for their individual functions. Thanks to the ability to form dynamic intermolecular hydrogen-bonds, MG-C11 can form stable assemblies around membrane proteins, resulting in the markedly enhanced protein stability demonstrated by this study. The hydrogen-bond network achieved by MG-C11 may not be optimal as the bonds are both dynamic and water-mediated. Thus, the introduction of a more extensive hydrogen-bonding network into detergent micelles may further enhance protein stability. We will pursue these directions to further improve detergent efficacy for protein stabilization. It will be also interesting to investigate the utility of this detergent for *in vitro* reconstitution of membrane proteins into proteoliposomes as only a few detergents were shown to be effective for this application.⁶⁶ Thus, new detergents capable of forming extensive and strong hydrogen-bonding could represent the next generation of novel detergents and hold significant potential for future membrane protein structural studies.

Conclusions

We have developed a set of glucoside amphiphiles with a melamine core. Of these MGs, MG-C11 showed the most effective protein stabilization and was superior to three gold standards (DDM, LMNG and TTG-C11). The MOR stability achieved by MG-C11 is particularly remarkable, indicating that this MG may be suited for structural studies of particularly unstable GPCRs. In addition, β_2 AR extraction by using this detergent yielded a dimeric receptor, a quaternary arrangement challenging to access using other detergents. MG-C11 possesses



several favorable features for membrane protein study including synthetic accessibility, small protein–detergent complex formation, reasonable efficiency for protein extraction and marked protein stabilization efficacy. Thus, the new detergent is likely to find wide use in membrane protein structural study. As indicated by our MD simulation study, MG-C11 is capable of forming dynamic hydrogen-bonding between detergent molecules when associated with membrane proteins, distinct from other detergents. The resulting increase in detergent–detergent interactions is likely to be mainly responsible for the enhanced membrane protein stability observed in this study. Therefore, the current study not only provides a detergent tool potentially useful for membrane protein study, but also a new strategy for the design of novel detergents that facilitate membrane protein research.

Data availability

The data that support the findings of this study are available in the ESI† of this article.

Author contributions

P. S. C. designed the new amphiphiles. L. G. and M. E. synthesized the amphiphiles. L. G., S. K., M. E., B. L., I. H. P., C. D. and S. K. designed and performed the research and interpreted the data. P. S. C., W. I., X. L., C. J. L., L. G. and B. B. contributed to experimental design and data interpretation. P. S. C. and L. G. wrote the manuscript, with oversight from all the other authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare the following competing financial interest(s): P. S. C. and L. G. are inventors on a patent application that covers the MGs.

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