

A biochemical analysis of the constraints of tail-anchored protein biogenesis

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TA (tail-anchored) proteins utilize distinct biosynthetic pathways, including TRC40 (transmembrane domain recognition complex of 40 kDa)-mediated, chaperone-dependent and/or unassisted routes to the ER (endoplasmic reticulum) membrane. We have addressed the flexibility of cytosolic components participating in these pathways, and explored the thermodynamic constraints of their membrane insertion, by exploiting recombinant forms of Sec61 β and Cytb5 (cytochrome b_5) bearing covalent modifications within their TA region. In both cases, efficient membrane insertion relied on cytosolic factors capable of accommodating a surprising range of covalent modifications to the TA region. For Sec61 β , we found that both SGTA (small glutamine-rich tetratricopeptide repeat-containing protein α) and TRC40 can bind this substrate with a singly PEGylated TA region. However, by introducing two PEG [poly(ethylene glycol)] moieties, TRC40 binding can be prevented, resulting in a block

of subsequent membrane integration. Although TRC40 can bind Sec61 β polypeptides singly PEGylated at different locations, membrane insertion is more sensitive to the precise location of PEG attachment. Modelling and experimentation indicate that this post-TRC40 effect results from an increased energetic cost of inserting different PEGylated TA regions into the lipid bilayer. We therefore propose that the membrane integration of TA proteins delivered via TRC40 is strongly dependent upon underlying thermodynamics, and speculate that their insertion is via a phospholipid-mediated process.

Key words: cytochrome b_5 (Cytb5), PEGylation, Sec61 β , small glutamine-rich tetratricopeptide repeat-containing protein α (SGTA), tail-anchored protein (TA protein), transmembrane domain recognition complex of 40 kDa (TRC40).

INTRODUCTION

TA (tail-anchored) proteins constitute a group of integral membrane proteins characterized by the presence of a single C-terminally localized stretch of hydrophobic amino acids that acts as both the subcellular targeting signal and membrane anchor [1]. Although TA proteins are found in most, if not all, intracellular membranes, the ER (endoplasmic reticulum) acts as the entry site for TA proteins destined for the various compartments of the secretory pathway [2–5]. Importantly, the C-terminal location of the membrane-spanning region precludes its co-translational recognition by the SRP (signal recognition particle), ensuring that TA protein targeting to, and insertion into, the ER membrane is post-translational [4].

The precise route of TA protein delivery to the ER membrane is precursor-dependent and correlates with the relative hydrophobicity of the TMS (transmembrane segment) of the tail-anchor region [6–8]. Thus, comparatively hydrophilic tail anchors, such as those of Cytb5 (cytochrome b_5) and protein tyrosine phosphatase 1B, define a pathway(s) that is either mediated by Hsp70 (heat-shock protein 70)/Hsp40 chaperones [8,9] and/or does not utilize any cytosolic components [10]. This pathway(s) does not rely on a specific receptor at the ER membrane; hence protease-treated microsomes are still fully capable of accepting the TA protein substrate [6,11].

The majority of TA proteins, such as RAMP4 (ribosome-associated membrane protein 4) and Sec61 β , have a TA region predicted to be more hydrophobic than that of Cytb5 [12]. These precursors use a specialized targeting factor, known as TRC40

(transmembrane domain recognition complex of 40 kDa; Asna1 in mammalian cells [13,14] and Get3 (guided entry of tail-anchored proteins 3) in *Saccharomyces cerevisiae* [15,16], for their delivery to the ER membrane. The TRC40/Get3 pathway requires additional cytosolic factors, such as Get4/Get5 and Sgt2 in yeast [17–20], and a membrane-bound receptor that has been defined as the Get1–Get2 complex in *S. cerevisiae* [16]. Recent structural analyses of the Get3 protein [21–25] and associated components [26,27] have provided new insights into the relationship between its ATPase cycle and TA protein delivery to the ER membrane and identified the likely substrate-binding site. However, since none of the current Get3 structures were obtained as a complex with a TA protein substrate, several aspects of substrate recognition by Get3 remain to be elucidated. Moreover, the precise details of TA protein recognition by upstream components, such as Sgt2 and its mammalian equivalent SGTA (small glutamine-rich tetratricopeptide repeat-containing protein α), are also currently unclear (see [20,28]).

To date, many biochemical studies addressing the biogenesis of TA proteins at the ER membrane have relied upon substrates generated by *in vitro* translation using cell lysates that contain essential cytosolic factors (see [7]). However, a bacterial system has recently been exploited to co-express model TA proteins with TRC40/Get3 [21,29], and the resulting complexes were shown to be sufficient for facilitating membrane insertion, indicating that TRC40/Get3 acts at a very late stage of TA protein delivery to the ER. Studies using purified recombinant TA proteins expressed in the absence of known cytosolic interacting partners are limited, and have largely focused on Cytb5 [10,30]. The use

Abbreviations used: BMH, bis(maleimido)hexane; BODIPY[®], boron dipyrromethene (4,4-difluoro-4-bora-3a,4a-diaza-s-indacene); Cytb5, cytochrome b_5 ; EndoH, endoglycosidase H; ER, endoplasmic reticulum; Get, guided entry of tail-anchored proteins; HisTrx, histidine–thioredoxin; Hsp, heat-shock protein; IASD, 4-acetamido-4'-[(iodoacetyl)amino]stilbene-2,2'-disulfonate; mPEG, methoxypoly(ethylene glycol); OPG, opsin glycosylation tag; PDI, protein disulfide-isomerase; PEG, poly(ethylene glycol); SGTA, small glutamine-rich tetratricopeptide repeat-containing protein α ; SRP, signal recognition particle; TA, tail-anchored; TMS, transmembrane segment; TRC40, transmembrane domain recognition complex of 40 kDa.

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of recombinant TA proteins allows polypeptide synthesis to be physically and temporally separated from the ER delivery and membrane insertion steps, facilitating studies of the components involved in these processes. Indeed, we have recently taken this approach to study Sec61 β biogenesis, and identified a novel role for Bat3 during TRC40-dependent ER delivery [28]. Notwithstanding the potential advantages of using recombinant forms of TA proteins [31], this approach has not been exploited extensively to date.

In the present study, we describe the successful production of recombinant TA protein models for each of the two major biosynthetic pathways, namely Sec61 β and Cytb5. By chemically modifying these recombinant TA protein substrates, we show that both routes are surprisingly unconstrained, and can accommodate substrates with significantly modified transmembrane domains. Remarkably, PEGylated forms of both Sec61 β and Cytb5 can also be efficiently integrated into the ER membrane, albeit via different routes. We find that the cytosolic factors SGTA and TRC40 can both accommodate substrates with a PEGylated TA region, and provide a model for the ER delivery and membrane integration of PEGylated Sec61 β . By analysing additional variants of Sec61 β , we show that the location of the PEG [poly(ethylene glycol)] moiety within the TA region strongly influences membrane integration in a manner that correlates with the predicted thermodynamic cost of this process. On this basis, we speculate that the membrane insertion step of the TRC40-dependent pathway for TA protein biogenesis may be a purely lipid-mediated event that does not require a dedicated integrase.

EXPERIMENTAL

Materials

Bacterial expression vector pHisTrx was a gift from Richard Kammerer (Paul Scherrer Institute, Villigen, Switzerland). Rabbit polyclonal antiserum recognizing TRC40 was a gift from Bernhard Dobberstein [ZMBH (Zentrum für Molekulare Biologie der Universität Heidelberg), Heidelberg, Germany]. All detergents were from Anatrace, except Triton X-100, which was from Sigma. Cross-linking reagents and EZ-link-biotin were from Pierce, whereas BODIPY[®] [boron dipyrromethene (4,4-difluoro-4-bora-3a,4a-diaza-*s*-indacene)] and IASD {4-acetamido-4'-[(iodoacetyl)amino]stilbene-2,2'-disulfonate} were from Molecular Probes. mPEG [methoxypoly(ethylene glycol)]-5000 and mPEG-20000 maleimides were from Nektar. Nuclease-treated rabbit reticulocyte lysate was supplied by Promega.

Chemical modification of recombinant TA proteins

Recombinant TA proteins were expressed and purified as described previously [28], with the various cysteine residue mutants generated by site-directed mutagenesis and confirmed by DNA sequencing. Thiol-reactive probes, as indicated, were mixed with solutions of single cysteine residue mutants of Sec61 β OPG^{S77C} (where OPG is opsin glycosylation tag) (77 μ M) in buffer B [50 mM Tris/HCl, pH 7.4, 300 mM NaCl, 10 mM imidazole, 10% (v/v) glycerol, 0.75% (w/v) OG (octyl β -D-glucopyranoside)] and Cytb5OPG^{S119C} (25 μ M) in buffer Q [50 mM Hepes/KOH, pH 7.9, 50 mM potassium acetate, 200 mM NaCl, 10% (v/v) glycerol, 0.1% LDAO (dodecanoyldimethylamine-*N*-oxide)] (see Supplementary Table S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm> for details of the mutants) using the reagents at a final concentration of 1–3 mM and incubating for 2 h at room temperature (21 °C) in the dark. Reactions were quenched by addition of 20 mM 2-mercaptoethanol and further incubation

for 10 min. Modified proteins were resolved by SDS/PAGE and then stained with Coomassie Brilliant Blue or immunoblotted with the anti-opsin tag antibody, or they were used for membrane integration reactions. All other Sec61 β OPG cysteine mutants (see Supplementary Table S1) were purified and labelled as described using 1.5 mM mPEG-5000 maleimide, pre-quenched with glycine, for 1 h at 25 °C. After quenching any unchanged maleimides, labelled proteins were analysed as above or used for integration assays. Membrane integration assays of recombinant TA proteins were performed using sheep pancreatic microsomes as described previously [28].

Binding of PEGylated Sec61 β OPG to recombinant SGTA and endogenous TRC40

Human SGTA was expressed in *Escherichia coli* BL21 Gold (DE3) pLysS cells and purified essentially as described above for TA proteins. HisTrx (histidine–thioredoxin)–SGTA, or BSA control, was then immobilized on UltraLink Biosupport beads (ThermoScientific) according to the manufacturer's instructions. Recombinant PEGylated Sec61 β OPG variants were mixed with beads in buffer S [50 mM Hepes/KOH, pH 7.5, 40 mM potassium acetate, 5 mM MgCl₂ and 0.05% pMAL-C12 amphiphile (Anatrace)], to give a final concentration of 3.3 μ M and incubated for 1 h at 4 °C. Beads were washed with buffer S plus 0.5 M NaCl and any bound material eluted first with buffer S supplemented with 0.5% (v/v) Triton X-100, and then mixed with Laemmli sample buffer.

Association of PEGylated Sec61 β OPG variants with endogenous TRC40 was tested by incubating 1–1.9 μ M of recombinant modified Sec61 β OPG with 140 μ l of rabbit reticulocyte lysate for 30 min at 30 °C. Reaction mixtures were then split into two and each half was added to 20 μ l [50% (v/v) suspension] of Protein A–Sepharose coated with 7.5 μ l of rabbit anti-TRC40 or anti-PDI (protein disulfide-isomerase) antisera. The lysate was incubated with anti-Protein A–Sepharose antibody beads for 1 h at 4 °C, the resin was then washed extensively with buffer R (50 mM Hepes/KOH, pH 7.5, 40 mM potassium acetate and 5 mM MgCl₂) supplemented with 0.5 M NaCl, and bound proteins were eluted with buffer R containing 0.5% (v/v) Triton X-100 (cf. [13]). The eluate was mixed with Laemmli sample buffer, and samples were resolved on SDS/PAGE and immunoblotted with the anti-opsin tag antibody. Quantitative analysis was carried out using a LiCor Biosciences system (described below). The non-specific binding of the indicated Sec61 β OPG variants to anti-PDI antibody-coated Protein A–Sepharose beads was subtracted from the results obtained using anti-TRC40 antibody-coated Protein A–Sepharose, and the resulting values were normalized to the amount of each Sec61 β OPG species present in the starting material (see Supplementary Figure S1 for a full worked example).

Quantitative immunoblotting

Following transfer of the samples on to a low-fluor PVDF membrane and incubation with a primary anti-opsin epitope tag antibody, a secondary anti-mouse fluorescent-dye-conjugated antibody was added at 1:10000 dilution. The membrane was incubated for 1 h at room temperature, scanned using a LiCor Biosciences system and the results were quantified with Odyssey 2.1 software.

Protease protection of membrane-integrated Sec61 β OPG variants

Two membrane integration reaction mixtures were combined. Microsomes were isolated as described previously [28] and

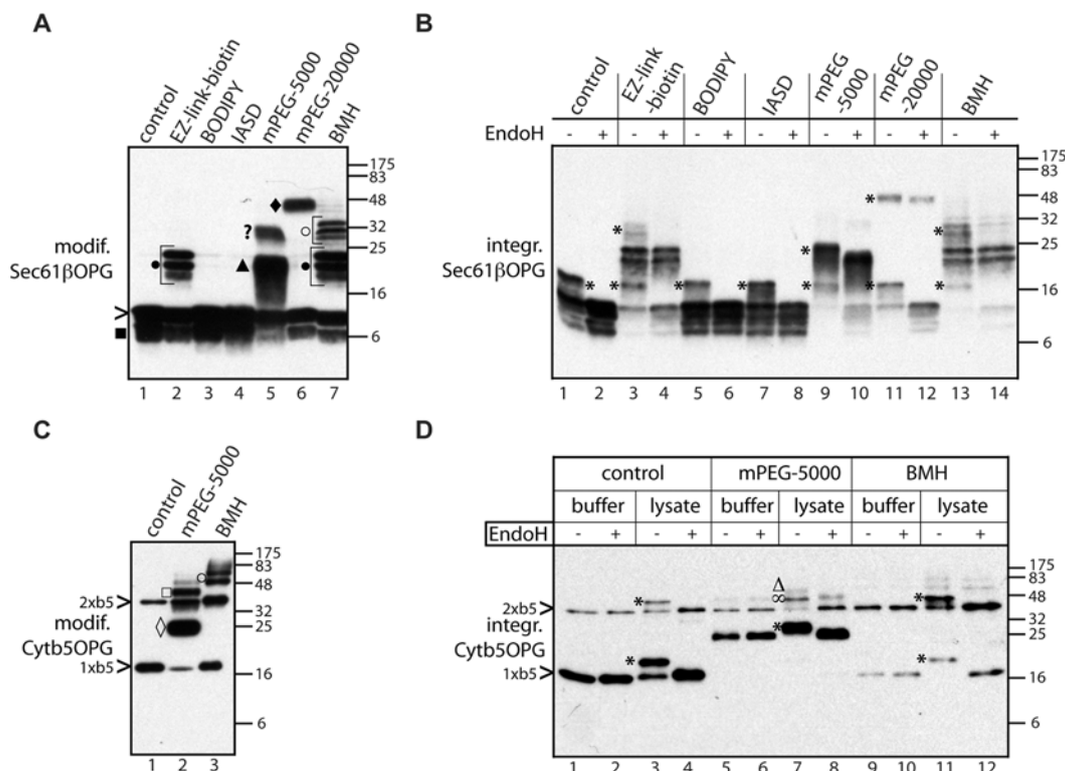


Figure 1 Chemically modified TA proteins are integrated into ER-derived membranes

Sec61βOPG^{S77C} (A) and Cytb5OPG (C) with single cysteine residues located in their TMSs (see Supplementary Table S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>) were modified (modif.) with thiol-reactive probes or the homobifunctional crosslinking reagent BMH, as indicated, and the products were analysed by immunoblotting. Full-length Sec61βOPG (>), a truncated form (■) and cross-linked dimers (●) are indicated, as are the monomer and dimer of Cytb5OPG (1xb5 and 2xb5 respectively). PEG-5000- and PEG-20000-modified Sec61βOPG (▲ and ◆ respectively), and PEGylated Cytb5OPG monomer (◇) and dimer (□) are indicated. Other products assumed to represent aberrant cross-linked species (? and ○) were not competent for membrane integration (integr.) and their precise identity was not pursued. (B and D) N-glycosylated species are indicated with an asterisk (*), except the N-glycosylated, PEG-modified Cytb5OPG dimer (Δ) (D, lane 7). A product corresponding to the PEGylated Cytb5OPG dimer, and an N-glycosylated, non-PEGylated dimer that co-migrate is also indicated (∞). Molecular mass markers in kDa are also indicated on the right-hand side of the Western blots.

resuspended in 68 μl of LSS (low-salt sucrose) buffer [250 mM sucrose, 100 mM potassium acetate, 5 mM magnesium acetate and 50 mM HEPES/KOH, pH 7.9]. The suspension was then split into three and each 20 μl aliquot received 2 μl of water, 2 μl of proteinase K (2.5 mg/ml) or 2 μl of proteinase K (2.5 mg/ml) and 2.2 μl of 10% (v/v) Triton X-100. Digestion was carried out on ice for 30 min and proteinase K was inhibited by the addition of 2.5 mM PMSF and further incubation on ice for 10 min. Laemmli buffer was then added and the samples were immediately heated to 95 °C for 10 min. Samples were resolved on a 16% (v/v) Tris/Tricine polyacrylamide gel and immunoblotted with the anti-opsin epitope tag antibody.

RESULTS

Recombinant TA proteins are inserted into the ER membrane

Preliminary analysis indicated that a HisTrx-based fusion was superior to a glutathione transferase tag for the high-level expression and purification of the recombinant TA proteins used in the present study (results not shown). We used this approach to purify and characterize two model recombinant TA proteins, Sec61β and Cytb5, each of which favours a distinct pathway for their biogenesis at the ER membrane (see the Introduction; also [7,8,13,14,32]). Cells expressing recombinant TA proteins were lysed in the presence of detergent, soluble fractions were incubated with Ni-NTA (Ni²⁺-nitrilotriacetate)-agarose and, after washing the beads with buffer containing a low

concentration of detergent, the various proteins were released from the tag by thrombin-mediated cleavage ([10,28,30]; see also the Experimental section and Supplementary Table S1). Coomassie Blue staining showed an acceptable level of protein purity, with only a fraction of Sec61βOPG and Cytb5OPG present in truncated forms (see Supplementary Figure S2A at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>).

The membrane integration capacity of the recombinant proteins was investigated using a well-established N-glycosylation assay that relies on a short C-terminal opsin-derived epitope tag [4,8,11,13,33]. Crucially, this modification can only take place when the protein spans the membrane and the tag has entered the ER lumen, consistent with authentic membrane integration. Substantial levels of EndoH (endoglycosidase H)-sensitive protein species for both Sec61βOPG and Cytb5OPG were observed (Supplementary Figure S2B, cf. lanes 1 and 2, 3 and 4), clearly indicating that these two precursors are efficiently integrated into the ER membrane in the presence of cytosol.

Chemically modified TA proteins are efficiently inserted into the ER membrane

Purified Sec61βOPG and Cytb5OPG, both bearing single cysteine residues within their TA regions (Sec61βOPG^{S77C} and Cytb5OPG^{S119C}; see Supplementary Table S1), were treated with a variety of thiol-reactive probes differing in size and physicochemical properties (Figures 1A and 1C). In the case of PEGylation, the modification was confirmed by a shift in

electrophoretic mobility (see Figures 1A and 1C), whereas for BODIPY[®], the fluorescently tagged protein could be visualized after membrane integration (see Supplementary Figure S3 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>). In the case of IASD and biotinylation, we have not directly confirmed labelling, and cannot rule out the possibility that little if any protein was modified. However, the biotinylation reagent also generated covalently linked dimers of Sec61 β OPG, most likely resulting from trace amounts of bifunctional cross-linkers present in the commercial reagent (Figure 1A, lane 2, ●). This conclusion is strongly supported by the fact that we can generate apparently identical species upon treatment with a bifunctional maleimide (Figure 1A, lane 7, ●). Treatment with mPEG-5000 maleimide and BMH [bis(maleimido)hexane] also generated even larger species (Figure 1A, lanes 5 and 7, ? and ○ respectively), but since these derivatives were not recovered in the membrane fraction after further analysis (cf. Figure 1B, lanes 9 and 10, 13 and 14), they were not further characterized. In the case of Cytb5OPG, a small amount of SDS-stable dimer was observed in the absence of any additional treatment (Figure 1C, lane 1). A similar species became more prominent after incubation with both mPEG-5000 maleimide and BMH, presumably due to covalent cross-linking of the dimer (Figure 1C). Both the monomeric and dimeric forms of Cytb5OPG appeared to be modified with mPEG-5000 (Figure 1C, lane 2, ◇ and □ respectively). Additional high-molecular-mass species were also observed with BMH (Figure 1C, lane 3, ○), but as for Sec61 β OPG, little if any of these unassigned species were recovered in association with ER-derived membranes (cf. Figure 1D). Hence our analysis showed that cysteine residues introduced into the TA regions of both recombinant proteins can be covalently modified by PEGylation and cross-linking reagents, whereas Sec61 β OPG can also accept a number of other thiol-reactive probes.

A role for cytosolic components during the membrane integration of Cytb5 is controversial [7,8,10]. Our previous analysis of recombinant Cytb5OPG found that immunodepletion of TRC40 or Bat3 from cytosol had no effect on its insertion, clearly distinguishing its integration from that of Sec61 β OPG [28]. This difference is further underlined when the kinetics of membrane insertion for recombinant Cytb5OPG and Sec61 β OPG are compared (see Supplementary Figures S2C and S2D), with Cytb5OPG integration reaching completion substantially more rapidly (cf. [34]). We therefore established whether the membrane integration of recombinant Cytb5OPG is cytosol-dependent in our assay. We can only detect the N-glycosylation of Cytb5OPG in the presence of both cytosol and ER-derived membranes (Figure 1D, lanes 1 to 4), using experimental conditions where the efficient N-glycosylation of recombinant TA proteins has been observed in the presence of buffer alone [21,29]. We therefore conclude that the efficient membrane integration of the recombinant Cytb5OPG used in our experimental system is cytosol-dependent.

To examine the constraints on the cellular components involved in the biogenesis of Sec61 β OPG and Cytb5OPG, we tested the cytosol-dependent integration of the two modified recombinant polypeptides into ER-derived membranes. Remarkably, none of the treatments with thiol-reactive reagents tested resulted in a complete block of membrane integration, despite the location of the target cysteine residue within the TA region of both proteins (Figure 1B, lanes 1–12 and Figure 1D, cf. lanes 3 and 4, 7 and 8). In fact, even homodimers of both Sec61 β OPG and Cytb5OPG were still measurably inserted into the lipid bilayer (Figure 1B, cf. lanes 3, 4, 13 and 14 and Figure 1D, cf. lanes 3, 4, 11 and 12). Membrane integration of a Cytb5OPG dimer occurred both with and without prior BMH treatment, suggesting that the

close proximity of the two TA regions of the protein monomers does not preclude membrane integration of a Cytb5OPG dimer (Figure 1D, cf. lanes 3 and 4, 11 and 12). A more detailed analysis of Sec61 β OPG dimers, generated by using cross-linking reagents of different length spacers, revealed that membrane integration was not qualitatively affected by the distance between individual subunits (see Supplementary Figure S4A at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>). Moreover, partial EndoH digestion of the membrane-associated material showed that both subunits of Sec61 β OPG and Cytb5OPG dimers are N-glycosylated, indicating that both polypeptides are fully inserted into the ER membrane (Supplementary Figures S4B and S4C). Remarkably, even the addition of large hydrophilic ER membrane-impermeable probes [35], including PEG-5000 (Figure 1B, cf. lanes 9 and 10, *, and Figure 1D, lanes 7 and 8, ∞) and, for Sec61 β OPG, PEG-20000 (Figure 1B, cf. lanes 11 and 12), to the TA-regions did not prevent authentic membrane insertion of the recombinant TA proteins. Perhaps most strikingly, a Cytb5OPG dimer modified with mPEG-5000 was still capable of being efficiently inserted into the ER membrane (Figure 1C, lane 2, and Figure 1D, cf. lanes 7 and 8, Δ). It is conceivable that the introduction of a PEG moiety into the TA region of a protein might alter its choice of biosynthetic route. In particular, we wished to confirm that the requirements for PEGylated Sec61 β OPG were the same as those previously defined for the unmodified protein. Sec61 β is known to exploit the TRC40-dependent pathway for ER integration, a route that is characterized by a protease-sensitive membrane component [11,14]. We therefore investigated the effect of trypsin treatment upon the membrane integration of Sec61 β OPG and found that the membrane association and N-glycosylation of both PEGylated and non-PEGylated forms of the protein were identically sensitive to protease treatment (see Supplementary Figure S5A at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>). In contrast, the TRC40-independent membrane integration of Cytb5 was unaffected by trypsin treatment, confirming both the resilience of the N-glycosylation machinery and the specificity of the assay (see Supplementary Figure S5A, and also [11,14]). We also discovered that prequenching of mPEG-5000 maleimide with glycine prevented the formation of aberrant products (cf. Figure 1A, lane 5 and Supplementary Figure S5B), and used this approach for all subsequent experiments.

PEGylation of Sec61 β OPG affects distinct stages of its membrane integration

Delivery of Sec61 β to the ER membrane is mediated by the TRC40 protein [14], and if this cytosolic factor is removed, the TA protein fails to reach its target destination efficiently [10,28]. Several studies also suggest a role for yeast Sgt2p [18,26], and its mammalian equivalent SGTA [28,36], acting on the same pathway but upstream of Get3/TRC40 [17,20]. To address the potential role of mammalian SGTA in TA protein biogenesis, we analysed the interaction of our recombinant TRC40 substrate, Sec61 β OPG, with immobilized recombinant SGTA. We find that Sec61 β OPG binds efficiently to SGTA, but not to a BSA control, in a Triton X-100-sensitive manner (Figure 2A, lanes 2 and 3). PEGylated Sec61 β OPG appears to bind at least as efficiently as the unmodified protein (Figure 2A, lanes 2 and 3), and is also released by Triton X-100, consistent with a hydrophobic interaction via the TA region as previously observed for TRC40 binding (cf. [13]). In contrast, an alternative version of Sec61 β that lacks the hydrophobic TA region but also bears a single PEG residue shows no association with immobilized

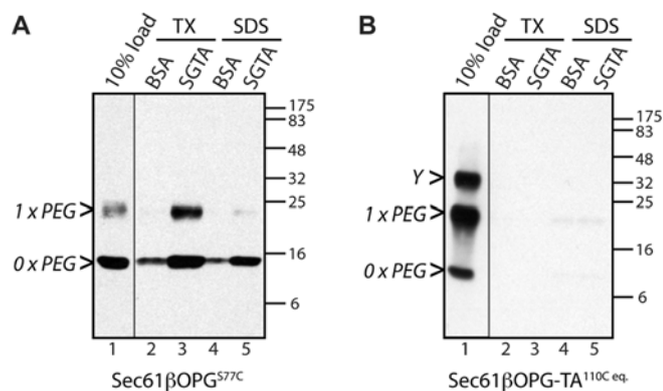


Figure 2 Recombinant Sec61 β OPG binds to immobilized SGTA

Sec61 β OPG^{S77C} (A) or Sec61 β OPG-TA^{110C eq.} (see Supplementary Table S1) (B) were PEGylated as described in the Experimental section and mixed with immobilized recombinant SGTA or a BSA control. The Triton X-100 (TX)- or Laemmli buffer (SDS)-eluted fractions were resolved by SDS/PAGE, and Sec61 β OPG variants were then detected by immunoblotting. Each panel shows samples resolved on a single blot and exposed for the same time with irrelevant lanes removed. An uncharacterized, but PEGylation-dependent, product of Sec61 β OPG-TA^{110C eq.} is indicated (Y). Molecular-mass markers in kDa are also indicated on the right-hand side of the Western blots.

SGTA, confirming that the association of a PEG moiety alone is not capable of mediating a Triton-sensitive interaction with SGTA (Figure 2B, lanes 2 and 3).

Crystal structures of Get3 [21,23,24], the yeast homologue of TRC40, reveal that it has a deep groove at the dimer interface that most likely acts as a TA protein recognition site. On the basis of our results (cf. Figure 1B), similar to SGTA, TRC40 must presumably accommodate a Sec61 β polypeptide modified with a single mPEG-5000 moiety. To investigate further the apparent flexibility of substrate binding by TRC40, we mutated both Ser⁷⁷ and Leu⁷⁹ of the Sec61 β OPG TA-region into cysteine residues (Sec61 β OPG^{S77C.L79C}) and modified this polypeptide with mPEG-5000. Assuming that the transmembrane region adopts an α -helix-like conformation, this double modification would yield a TA protein with two PEG-5000 molecules extending from opposite sides of the polypeptide chain (Figure 3A). Immunoblotting analysis confirmed the efficient labelling of Sec61 β OPG^{S77C.L79C}, yielding populations bearing both one and two mPEG-5000 molecules (Figure 3B, cf. lanes 1 and 2). When membrane insertion was tested, both unmodified and singly PEGylated forms of Sec61 β OPG^{S77C.L79C} were visibly N-glycosylated, in agreement with our previous results (Figure 3C, cf. lanes 3 and 4; cf. Figure 1B, lanes 1 and 2, 9 and 10). However, no N-glycosylation of Sec61 β OPG^{S77C.L79C} labelled with two mPEG-5000 molecules was seen, even after prolonged exposure of the immunoblot (Figure 3C, cf. lanes 3 and 4; and results not shown).

The lack of any membrane-inserted Sec61 β OPG^{S77C.L79C} that had been modified with two PEG-5000 species could result from either its inefficient delivery to the ER membrane or an inability to be integrated into the phospholipid bilayer. To address this issue, we tested the association of the PEG-labelled Sec61 β OPG^{S77C.L79C} with TRC40 by immunoprecipitation and quantitative immunoblotting. We found that the addition of two PEG probes to a single TMS of Sec61 β OPG^{S77C.L79C} almost completely abolished TA protein binding to TRC40 (Figure 3D, cf. lanes 3 and 4, see also Supplementary Figure S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>). Also, we revealed that the attachment of even a single PEG probe to Sec61 β OPG^{S77C.L79C} reduced its binding to TRC40 to approximately half the level observed with the unmodified protein

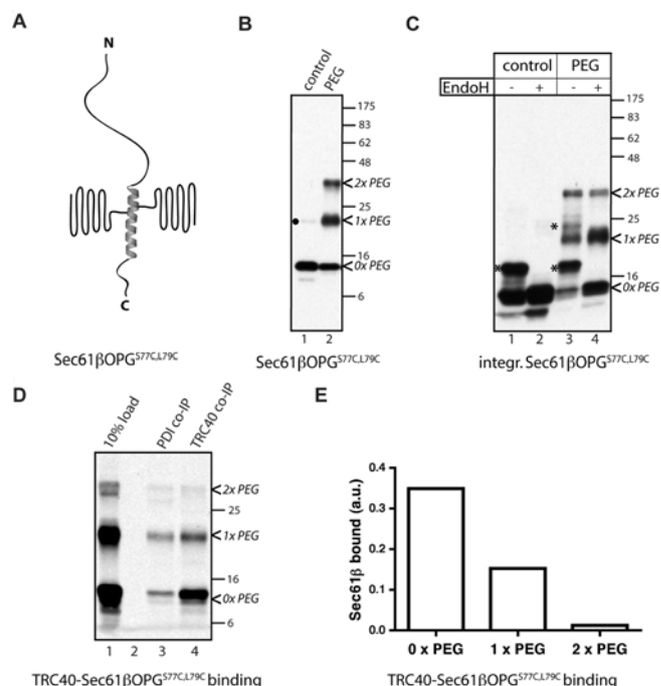


Figure 3 PEGylation of Sec61 β OPG^{S77C.L79C} prevents TRC40 binding

Sec61 β OPG^{S77C.L79C} was generated (see Supplementary Table S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>) and the purified protein was modified with mPEG-5000. A schematic diagram of the doubly labelled polypeptide is presented assuming an α -helix-like conformation of the transmembrane region (A). Covalent attachment of PEG-5000 was confirmed by immunoblotting (B) and membrane integration by EndoH sensitivity (C). Small amounts of Sec61 β OPG species most likely corresponding to a dimer formed during protein purification were observed (B, ●). Binding of PEG-modified Sec61 β OPG^{S77C.L79C} to TRC40 was addressed by pre-incubating the protein with rabbit reticulocyte lysate followed by co-immunoprecipitation with anti-TRC40 or control anti-PDI antibodies. Bound Sec61 β OPG was eluted and detected by quantitative immunoblotting (D), allowing relative binding to TRC40 to be compared (E, see Supplementary Figure S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm> for a full worked example). The values shown are the average from two independent experiments. * indicates N-glycosylated species. Molecular-mass markers in kDa are also indicated on the right-hand side of the Western blots.

(Figure 3E). This is consistent with a qualitative reduction in the membrane integration of the singly modified chain when compared with the unmodified version of Sec61 β OPG (Figure 3C, cf. lanes 3 and 4, marked 0x PEG and 1x PEG).

We next established whether a protein with a TA region modified with two PEG-5000 molecules predicted to be located on the same side of an α -helix-like conformation can be bound by TRC40 and delivered to the ER membrane. To this end, a third derivative of Sec61 β OPG with both Ser⁷⁷ and Val⁸⁴ mutated to cysteine residues (Sec61 β OPG^{S77C.V84C}) (Figure 4A) was purified and modified with mPEG-5000 (Figure 4B, cf. lanes 1 and 2). When the membrane integration of the two double cysteine residue variants of Sec61 β OPG was compared, surprisingly little if any N-glycosylation of even singly PEGylated Sec61 β OPG^{S77C.V84C} could be detected under conditions where the singly modified Sec61 β OPG^{S77C.L79C} was reproducibly integrated (Figure 4C). When the binding of Sec61 β OPG^{S77C.V84C} to TRC40 was analysed, the interaction of the singly PEGylated-Sec61 β OPG^{S77C.V84C} species was unperturbed, although association of the doubly PEGylated species was clearly diminished (Figures 4D and 4E). Hence, although TRC40 binding is relatively tolerant of the addition of a single PEG-5000 moiety to the Sec61 β TA-region, a subsequent step of the membrane insertion process appears much more sensitive to the precise

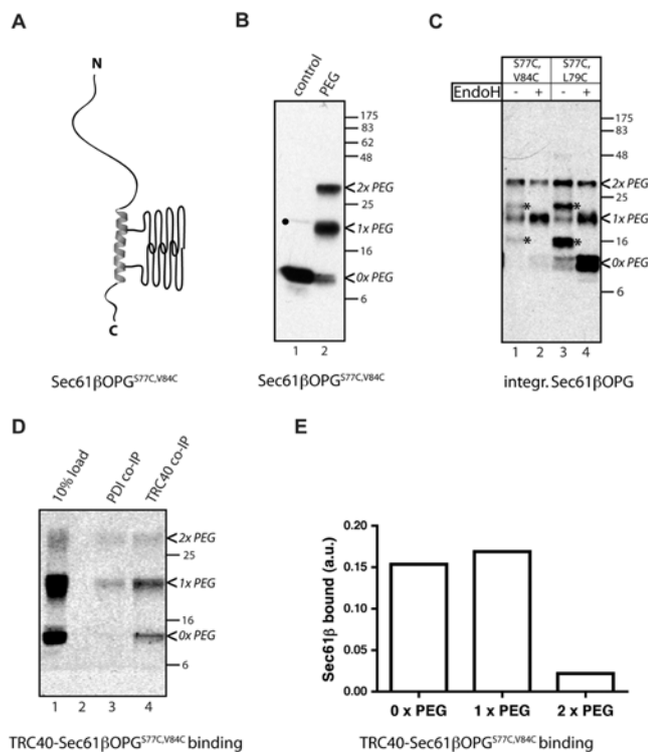


Figure 4 PEGylation of Sec61 β OPG^{S77C,V84C} inhibits membrane integration

(A) Purified recombinant Sec61 β OPG^{S77C,V84C} (see Supplementary Table S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>) was modified with mPEG-5000 and is shown schematically as described for Figure 3(A). (B–D) PEGylation was confirmed by immunoblotting (B) with a presumptive Sec61 β OPG dimer indicated (●). Membrane integration was determined by EndoH sensitivity (C), and TRC40 binding (D and E) was analysed as detailed in the legend to Figure 3 (see also Supplementary Figure S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>). * indicates N-glycosylated species. Molecular-mass markers in kDa are also indicated on the right-hand side of the Western blots.

location of such a probe and PEGylation of Cys⁸⁴ appears not to be tolerated.

In order to test this hypothesis further, we generated a single cysteine residue mutant, Sec61 β OPG^{V84C}, and studied its membrane integration compared with the Sec61 β OPG^{S77C} mutant. Cys⁸⁴ was more efficiently PEGylated than Cys⁷⁷ (Figure 5A and Supplementary Figure S1A), but the resulting Sec61 β derivative showed little if any membrane-dependent N-glycosylation, whereas Sec61 β OPG^{S77C} was again efficiently modified (Figure 5A, lanes 3 to 6), as previously observed (Figure 1B, lanes 9 and 10). Furthermore, the effect of PEGylation of residue 84 could not be correlated to a reduction in TRC40 binding (Supplementary Figure S1). Hence the PEGylation of a single cysteine residue located at amino acid 84 of Sec61 β OPG effectively prevents membrane integration as judged by N-glycosylation. One potential explanation for this location-specific effect of PEGylation is the thermodynamic cost of integrating the modified TA region into a phospholipid bilayer (see also Figure 6B, the Supplementary online data and Supplementary Table S2 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm> and the Discussion section). We therefore explored the consistency of this positional effect by analysing the membrane integration of PEGylated forms of three further single cysteine residue mutants, L74C, L87C and a novel cysteine residue, added to the C-terminus of Sec61 β OPG, denoted 110C. We observe that the membrane integration of PEGylated Sec61 β OPG^{L74C} and Sec61 β OPG^{S77C} is clearly more efficient

than that of PEGylated Sec61 β OPG^{V84C}, Sec61 β OPG^{L87C} and Sec61 β OPG^{110C} (Figures 5A, 5B and 5C). To confirm that the location-specific effect of PEGylation was due to a perturbation of membrane integration, and not N-glycosylation itself, we also carried out a protease protection analysis (cf. [10]). We could identify a discrete membrane-dependent protease-protected fragment of PEGylated Sec61 β OPG^{S77C} (Figure 5D, lane 5, filled triangle). In contrast, no such product was observed with Sec61 β OPG^{V84C} or Sec61 β OPG^{110C} (Figure 5D, lane 8 and Figure 5E, lane 2, predicted location indicated by ▲). On this basis, we conclude that the PEGylation of amino acid residues located towards the C-terminus of Sec61 β OPG preferentially inhibits membrane integration.

DISCUSSION

We have exploited recombinant TA proteins as a tool for better understanding their biogenesis at the ER. Both Sec61 β OPG [28] and Cytb5OPG (the present study) are efficiently integrated into ER-derived membranes, and in both cases this process is dependent upon the presence of cytosol. The role of cytosolic factors in the membrane integration of Cytb5 remains controversial [7,8,10]. Nevertheless, in the present study we can only detect an N-glycosylated membrane integrated form of the protein in the presence of lysate (cf. [7]). In the case of Sec61 β , the requirement for cytosol correlates to a need for TRC40/Asn-1 [13,14], and other cytosolic factors including Bat3 [28] and the mammalian homologues of Get4 and Get5 [37]. We now provide the first evidence for a direct interaction between Sec61 β and SGTA, the mammalian equivalent of yeast Sgt2. Sgt2 has recently been implicated in the transfer of TA protein substrates to Get3 [20]. These studies support our previous observation of a specific association between SGTA and TA regions [28], and we now conclude that this represents a direct interaction of SGTA with a hydrophobic substrate [38], and speculate that SGTA normally plays a role in TA protein transfer to TRC40.

Sec61 β can tolerate the covalent attachment of a variety of reagents to a single cysteine residue located within its TA region, and retain its ability to be integrated into the ER membrane. Likewise, the membrane integration of Cytb5 is not blocked by the addition of a PEG-5000 moiety close to the middle of its TA region or by the formation of a protein dimer. In the case of Sec61 β , polypeptides with various attachments including PEG-5000, and even PEG-20 000, can all be post-translationally integrated. This suggests that the cellular components that mediate its delivery to, and integration into, the ER membrane can accommodate a range of non-physiological modifications without a substantial perturbation of their function. Our analysis of the binding of Sec61 β OPG to cytosolic factors implicated in its delivery to the ER showed that the direct binding of recombinant Sec61 β OPG^{S77C} to SGTA was unaffected by PEGylation. Since the PEG moiety is attached to the hydrophobic TA region, this would suggest that SGTA has a hydrophobic groove-like binding site capable of accommodating this unnatural substrate. Interestingly, both TRC40 and the 54 kDa subunit of the SRP each have flexible hydrophobic clefts capable of accommodating a range of hydrophobic polypeptide regions [21,23,39], further supporting this notion (see also [20]).

To analyse the effects of PEGylation upon TRC40 binding, we engineered double cysteine residue mutants of Sec61 β OPG that generated a mixture of singly and doubly modified species upon PEGylation and determined their association by co-immunoprecipitation. This analysis showed that the singly PEGylated form of Sec61 β OPG^{S77C,L79C} can still bind to TRC40

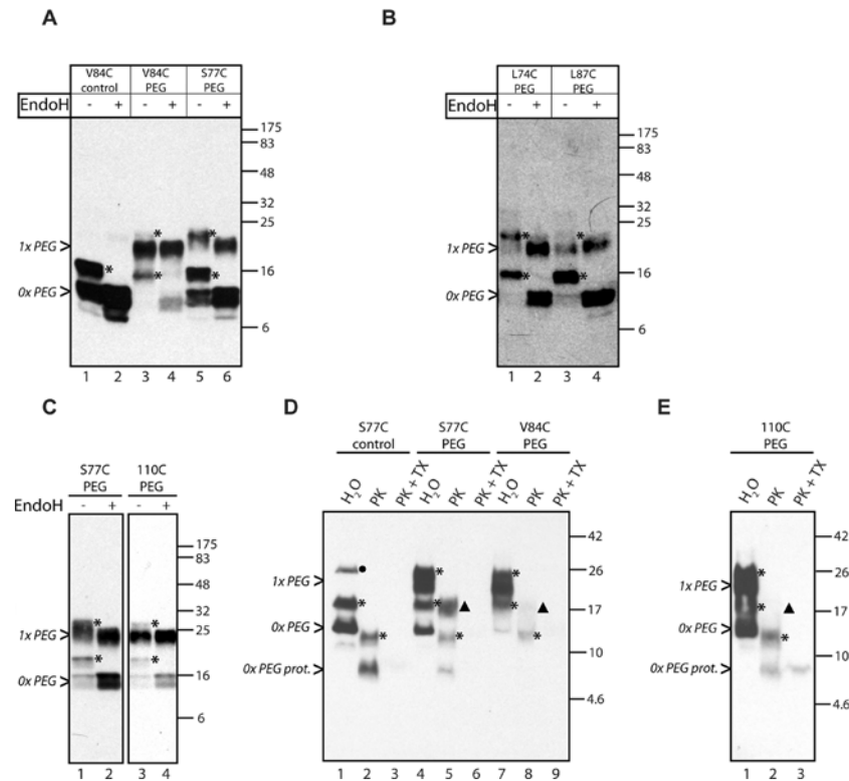


Figure 5 Distinct effects of PEGylation at different locations within the Sec61 β TA-region

Additional single cysteine Sec61 β OPG variants were generated (see Supplementary Table S1 at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>) and PEGylated. The membrane integration of PEGylated Sec61 β OPG^{S77C} and Sec61 β OPG^{V84C} (A), Sec61 β OPG^{L74C} and Sec61 β OPG^{L87C} (B) and Sec61 β OPG^{S77C} and Sec61 β OPG^{I10C} (C) were determined by EndoH sensitivity. In addition, the membrane integration status of Sec61 β OPG^{S77C} and Sec61 β OPG^{V84C} (D) and Sec61 β OPG^{I10C} (E) was determined by protease protection (see the Experimental section). The filled triangle indicates the actual (D, lane 5) or predicted (D, lane 8 and E, lane 2) location of a protease protected, PEGylated, fragment of Sec61 β OPG. In (C), the Sec61 β OPG^{I10C} samples were overexposed in order to best visualize any N-glycosylated PEGylated products. * indicates N-glycosylated species; \blacktriangle indicates predicted location of membrane-dependent protease-protected fragment of PEGylated Sec61 β OPG. Molecular-mass markers in kDa are also indicated on the right-hand side of the Western blots. PK, proteinase K; TX, Triton X-100.

with substantial efficiency, whereas a doubly modified form cannot. When this PEGylated TA region is modelled into the substrate-binding site of TRC40, our data are in good agreement with the proposed deep pocket [21–24] that could accommodate the addition of one centrally localized PEG-5000 to the substrate, but not two (see Figure 6A). Hence, we conclude that the doubly PEGylated form of Sec61 β OPG^{S77C.L79C} is incapable of membrane integration due to its inability to bind TRC40. When an alternative cysteine residue mutation was used to generate a doubly PEGylated variant, Sec61 β OPG^{S77C.V84C}, although the singly modified protein could bind TRC40, it was no longer membrane-integrated. Subsequent analysis of the single cysteine residue mutant Sec61 β OPG^{V84C} showed that PEGylation of Cys⁸⁴ specifically perturbs biosynthesis at a step occurring after TRC40 binding; for example, substrate release by TRC40 or the subsequent membrane integration step (Figure 6C). Our calculations of the free energy changes resulting from the insertion of a PEG polymer into a lipid bilayer favour the introduction of a constraint at the membrane insertion step. Hence, the free energy associated with the transfer of the unmodified Sec61 β OPG TMS into a lipid bilayer can be estimated as -11.5 kcal (1 kcal = 4.184 kJ)/mol (see Supplementary online data and Supplementary Table S2), assuming a non-Sec61 translocon-mediated pathway for insertion [7,40,41]. By the same criteria, the integration of Sec61 β OPG PEGylated at Cys⁷⁷ has an estimated energetic cost of -4 kcal/mol, and hence remains favourable (Figure 6B, S77C). In contrast, the energetic cost of integrating Sec61 β OPG PEGylated at Cys⁸⁴ is $+1$ kcal/mol (Figure 6B, V84C; see Supplementary

online data for detailed calculations). Thus our modelling supports the idea that the membrane integration efficiency of a PEGylated Sec61 β TA region is dependent on the precise location of the probe relative to the lipid bilayer (Figure 6B).

To further challenge this model, we created three further single cysteine residue mutants of Sec61 β OPG, Sec61 β OPG^{L74C}, Sec61 β OPG^{L87C} and Sec61 β OPG^{I10C} and analysed the membrane integration of their PEGylated species. As predicted, the modification of residue 74 towards the N-terminal side of the TA region did not prevent membrane integration, whereas the alteration of residue 87 towards the C-terminal end, and the addition of a novel cysteine at the extreme C-terminus, both did. Taken together, these observations suggest that thermodynamic constraints are a major factor in the membrane integration of TRC40-dependent TA proteins. Although we cannot exclude the possibility that the membrane insertion of Sec61 β requires a specialized integrase, we believe that the distinct behaviour of the differently PEGylated Sec61 β TA regions is more consistent with a simple partitioning model for membrane insertion. In this scenario, the recently identified membrane-bound receptor for TRC40 [42] would act to deliver TA proteins to the surface of the ER membrane, but subsequent integration would rely solely on the favourable thermodynamics of TA protein partitioning into the lipid bilayer (Figure 6C, cf. [7]).

In the case of Cytb5, the precise identity of any cellular components that mediate its membrane delivery and integration are uncertain [10,13,14,28,29]. Nevertheless, we see an equally flexible pathway for Cytb5 biogenesis that can tolerate both

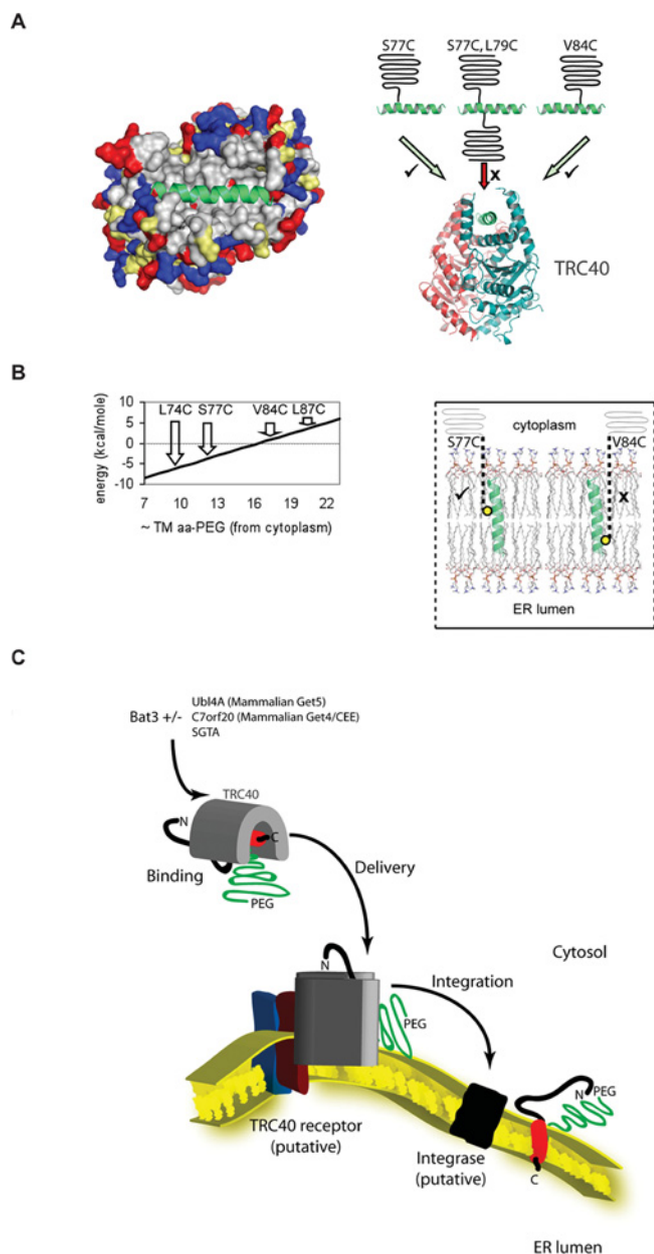


Figure 6 Membrane delivery and integration of PEG-modified Sec61 β

(A) Left-hand panel, human TRC40 was modelled using the *S. cerevisiae* Get3 dimer structure as a template (PDB code 2WOJ) and with the 23-amino-acid-residue α -helical TMS of Sec61 β (green) shown in the proposed binding groove. The surface of the TRC40 dimer is colour-coded by amino acid polarity: blue, positively charged; grey, non-polar; red, negatively charged; yellow, polar-uncharged. Right-hand panel, a schematic representation for the association of PEGylated Sec61 β with TRC40 showing binding of polypeptides PEGylated at residue 77 or 84, but not of a doubly modified Sec61 β ^{S77C,L79C} variant. (B) Left-hand panel, estimated energy changes (see Supplementary online data at <http://www.BiochemJ.org/bj/436/bj4360719add.htm>) for the incorporation of different singly PEG-modified TA-regions of Sec61 β OPG into a lipid bilayer. The relative ΔG values for PEGylation at residues 74, 77, 84 and 87 are indicated. Right-hand panel, representation of PEGylated Sec61 β TA regions within a model lipid bilayer using membrane co-ordinates taken from [47]. Experiments confirm the efficient membrane integration of a Sec61 β OPG PEGylated on Cys⁷⁴ and Cys⁷⁷, but not Cys⁸⁴, Cys⁸⁷ or Cys¹¹⁰. (C) Working model for the delivery of PEGylated Sec61 β to the ER membrane. A variety of upstream components including Bat3, SGTA and mammalian Get4 and Get5 [27,28,37] facilitate TRC40-mediated delivery to the mammalian ER membrane via its receptor [42]. Any requirement for a membrane integrase remains hypothetical (cf. [7]). Covalent modification of the TA region of Sec61 β could in principle perturb any one of the steps illustrated.

PEGylation and enforced dimerization. We conclude that any cytosolic factors that facilitate Cytb5 integration can tolerate these species without substantial perturbation (Figure 1). Once at the membrane, the rate at which Cytb5OPG can be N-glycosylated appears to be substantially higher than that of Sec61 β OPG (Supplementary Figure S2), and Cytb5OPG integration is also insensitive to trypsin treatment of the membranes (Supplementary Figure S5), further supporting the use of different pathways for integration into the ER [7,33,34]. Whether the flexibility of the post-translational pathways that underlie TA protein biogenesis is of physiological significance remains to be determined. However, our results are in good agreement with the finding that certain viral proteins are palmitoylated within their TMSs prior to membrane integration [43]. They are also consistent with the proposal that soluble SNAP-25 (25 kDa synaptosome-associated protein) is initially targeted to the ER membrane in a complex with the TA protein syntaxin [44], presumably via the binding of the complex to TRC40 prior to ER delivery [14]. It may even be the case that SGTA and TRC40 can bind to covalently modified TA proteins that are generated *in vivo* such as ubiquitinated species, consistent with reports of the co-translational ubiquitination of membrane proteins [45] and known links between the yeast homologue of TRC40, Get3 and the ubiquitin–proteasome system [46].

AUTHOR CONTRIBUTION

Pawel Leznicki conceived and carried out the experiments, analysed the data and contributed to the preparation of the manuscript. Jim Warwicker carried out the structural and thermodynamic modelling work and contributed to the preparation of the manuscript. Stephen High contributed to experimental design and the preparation of the manuscript for submission.

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SUPPLEMENTARY ONLINE DATA

A biochemical analysis of the constraints of tail-anchored protein biogenesis

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Calculations of the energetic cost for membrane integration of PEGylated Sec61 β OPG

Boundaries of the membrane-spanning region of Sec61 β OPG were determined using a prediction algorithm for changes in free energy, ΔG , upon Sec61 translocon-mediated protein integration into a lipid bilayer [1]. The transfer energy for the predicted 23-amino-acid segment (see Table S1) was calculated on the basis of the water/octanol partition scale [2] and a value of -11.5 kcal/mol was obtained.

$\log P$ values of atom fragments of the mPEG-5000 maleimide moiety were computed by best fit to a training set of 2473 compounds [3], and the sum obtained (-5.0 to -5.3) was in good agreement with the data available for the $\log P$ of PEG-400 derived from water/hexane partitioning (-4.824), assuming that PEG-400 is composed of, on average, eight or nine $\text{CH}_2\text{CH}_2\text{O}$

repeats, and taking into account any potential discrepancy between water/hexane and water/octanol systems. Making equivalent calculations for a 35 \AA ($1 \text{ \AA} = 0.1 \text{ nm}$) stretch of PEG, corresponding to the length of the predicted tail-anchor region, and converting the results to ΔG , the cost of transferring the PEG polymer into a lipid bilayer in a conformation that spans both leaflets was estimated as approximately $+3.6$ kcal/mol.

A rotameric entropy loss for an extended PEG chain was estimated on the basis of a comparison with studies of aliphatic side chain conformers in proteins [4] and the role of ligand rigidification in modulating binding affinities [5]. Hence, the entropy was calculated from the equation $S = -R\sum(P \ln P)$, where R is the universal gas constant, P the probability for each rotamer, and the sum is taken over all rotamers. For a torsional system with three stable rotamers, assuming a transition between equal probabilities for each, and just one (extended chain) rotamer,

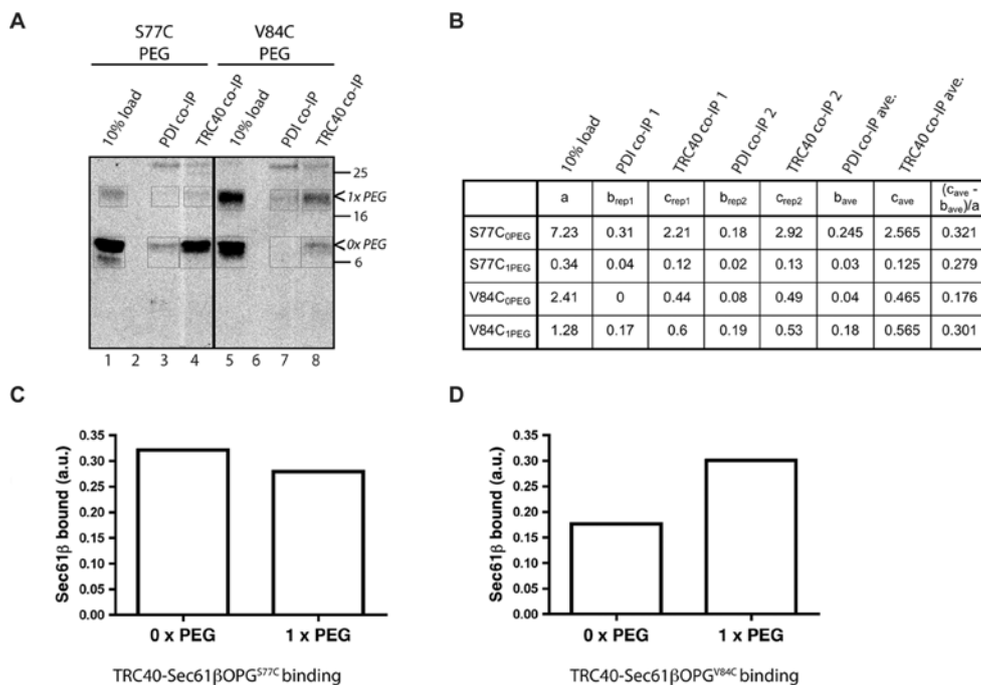


Figure S1 TRC40 binding to PEGylated Sec61 β OPG

(A) PEGylated Sec61 β OPG^{S77C} or Sec61 β OPG^{V84C} were incubated in the presence of rabbit reticulocyte lysate followed by co-immunoprecipitation with anti-TRC40 or control anti-PDI antibodies as indicated. Bound Sec61 β OPG variants were then eluted and detected by quantitative immunoblotting using a LiCor system (see the Experimental section of the main text) to estimate the association of the unmodified and PEG-modified forms with TRC40. Molecular-mass markers are shown in kDa on the right-hand side. (B) The actual values obtained from the LiCor system for each of the products analysed (see boxes in A) for two separate experiments are shown, together with the method for calculating Sec61 β OPG binding. Non-specific binding to Protein A–Sepharose coated with anti-PDI antibody was subtracted from the binding to anti-TRC40 antibody-coated Protein A–Sepharose, and the resulting values were normalized to the amount of each Sec61 β OPG species. These results are expressed graphically for Sec61 β OPG^{S77C} (C) and Sec61 β OPG^{V84C} (D).

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a free energy contribution of +0.6 kcal/mol at $T = 300$ K for each torsion was obtained. Over the 35 Å TMS length, this amounts to +17.0 kcal/mol for a fully extended chain.

A contribution of the mPEG-5000 maleimide linker region to the energetic cost of inserting PEGylated Sec61 β OPG into a lipid bilayer was estimated making an assumption, on the basis of the manufacturer's data, that in the experimental conditions used the maleimide-linker ring remains intact. Taking into account the nature of the maleimide-linker components and their bond rotamer status, the following parameters were calculated: length in the extended conformation, 10 Å; ΔG for octanol partitioning of -0.5 kcal/mol (calculated from $\log P$); rotamer entropy loss in extended conformation contributing +3.6 kcal/mol, i.e. a net value of +3.1 kcal/mol.

The estimated contribution of each component to the overall energetic cost for the membrane integration of the PEGylated forms of Sec61 β ^{L74C}, Sec61 β ^{S77C}, Sec61 β ^{V84C} and Sec61 β ^{L87C} is presented in Table S2. Although the ΔG values shown for the lipid partitioning of PEG attached to these distinct sites are estimates and absolute values may differ, our calculations clearly indicate that the integration of PEGylated Sec61 β is finely balanced, such that it changes from favourable to unfavourable as the site of PEG modification is moved deeper into the membrane from the cytosolic face. It should be noted that, in all calculations presented above, an extended conformation of the maleimide-PEG moiety, parallel to the TA of Sec61 β , was assumed. This conformation minimizes occlusion of the PEG chain from the solvent and provides a defined conformer for calculations. Although other conformations are theoretically possible, these would most likely require higher energy for their membrane integration, due to the increased number of the PEG repeats that would have to be accommodated by the lipid bilayer. In addition, it is assumed that the bulk of the PEG molecule does not translocate across the membrane, consistent with experimental data, so that PEG traversal of the membrane is always modelled from the cysteine/maleimide linker to the cytosolic side.

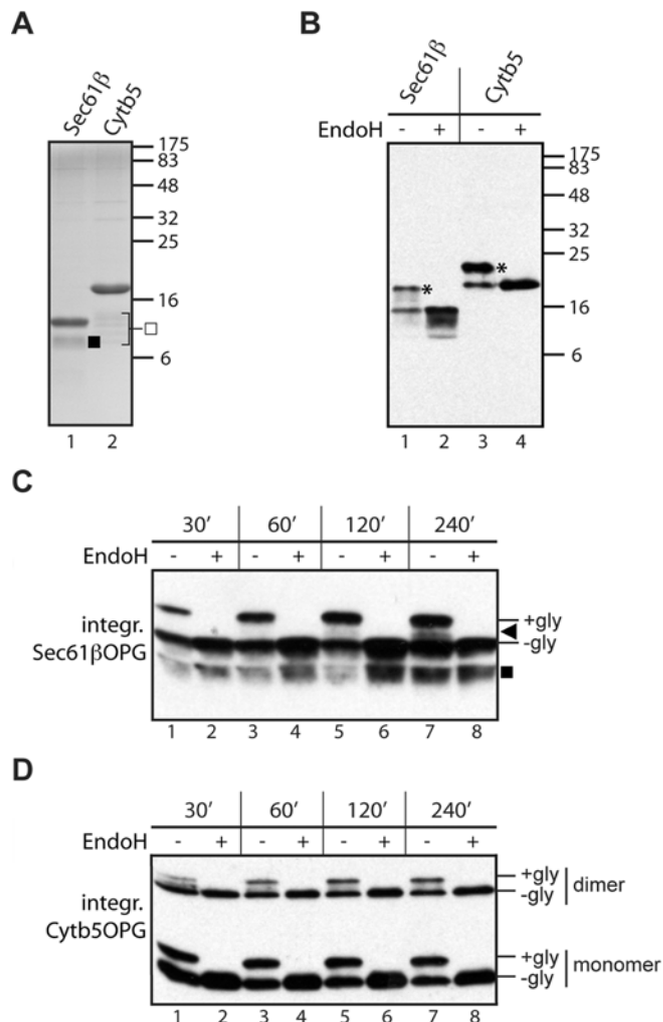


Figure S2 Characterization of recombinant Sec61 β OPG and Cytb5OPG

(A) Coomassie Blue-stained gel of purified recombinant Sec61 β OPG^{S77C} and Cytb5OPG^{S119C} both prepared using buffers supplemented with 0.75% (w/v) octyl- β -D-glucopyranoside (see also [6]). (B) Membrane integration reactions of recombinant Sec61 β OPG^{S77C} and Cytb5OPG^{S119C} carried out in the presence of reticulocyte lysate (see [6]), and using immunoblotting for detection and EndoH sensitivity of N-glycosylated products (*) to confirm membrane integration. (C and D) Time course for the membrane integration (integr.) of Sec61 β OPG^{S77C} (C) and Cytb5OPG^{S119C} (D) performed as described for (B). Truncated versions of Cytb5OPG (□); Sec61 β OPG (■) and N-glycosylated Sec61 β OPG (◄) are indicated. Molecular-mass markers in kDa are also indicated.

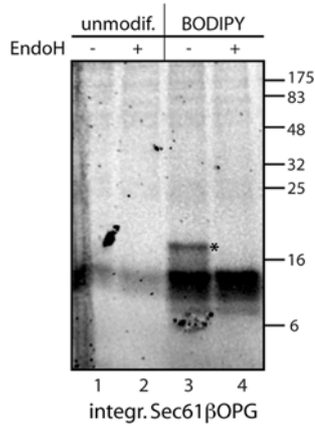


Figure S3 BODIPY® labelling of Sec61βOPG

Purified recombinant Sec61βOPG^{S77C} was modified with BODIPY® maleimide or not (unmodif.) as indicated, and membrane integration (integr.) was analysed as described in the text (see also Figures 1A and 1B of the main text). Fluorescently labelled products recovered with the membrane fraction were visualized directly using a FUJIFILM LAS-3000. N-glycosylated products are indicated (*). Molecular-mass markers in kDa are also indicated on the right-hand side.

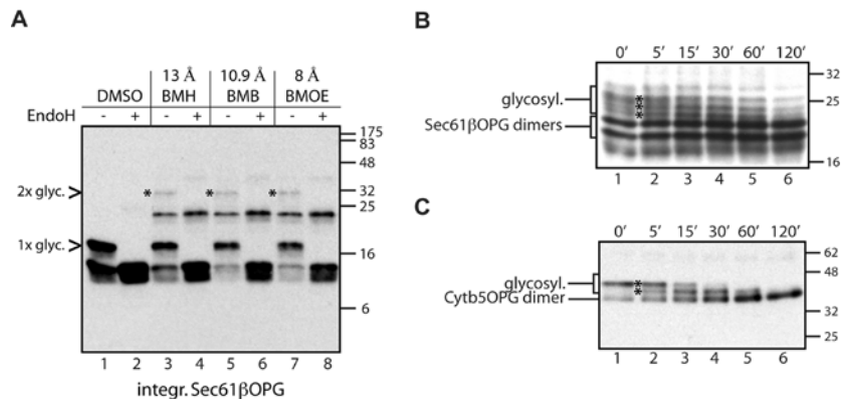


Figure S4 Both monomers of TA protein dimers are authentically integrated into the ER membrane

(A) Dimers of Sec61βOPG were generated by cross-linking the single cysteine mutant S77C (see Supplementary Table S1) with 1 mM of the indicated thiol-reactive reagents with a spacer length of 8–13 Å. The reaction mixtures were incubated for 10 min at 25°C, quenched with 15 mM 2-mercaptoethanol and the modified proteins were used in a membrane integration assay (see [6]). The membrane-associated fraction was isolated by centrifugation, treated with Endo H where indicated and products were analysed by immunoblotting. N-glycosylated monomers and dimers are indicated (1x glyco. and 2x glyco. respectively). BMB, 1,4-bis(maleimido)butane; BMOE, bis(maleimido)ethane. (B and C) BMH cross-linked single cysteine mutants of Sec61βOPG (B) and Cytb5OPG (C) were used in a membrane integration reaction [6], and the isolated membrane fraction subjected to partial digestion with Endo H (0.21 units/μl). At the indicated times, aliquots were mixed directly with Laemmli sample buffer, and products were subsequently analysed by immunoblotting. Migration of Sec61βOPG and Cytb5OPG dimers is indicated, and various N-glycosylated (glycosyl.) species of both proteins are shown (*). The doublet corresponding to the Sec61βOPG dimer results from a truncated form of the recombinant protein (see Figure S2), leading to a population of mixed dimers. These species are also responsible for the additional N-glycosylated protein species identified by EndoH treatment (B). Molecular-mass markers in kDa are indicated on the right-hand side.

Table S1 Sequence of TA proteins used in the present study

Sequences of TA proteins used in the present study are shown, with the predicted transmembrane segments in bold and the opsin-derived, C-terminal tag underlined. Mutations introduced into the amino acid sequences are indicated by asterisks (*). Amino acid numbering reflects that of the wild-type proteins; however, the recombinant versions lack an initiator methionine residue and have instead a GlySer dipeptide (*italics*) resulting from the thrombin-mediated cleavage of the N-terminal HisTrx tag.

Protein (species)	Sequence
Sec61 β OPG ^{S77C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMC*LLFIASVFMLHIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{V84C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMSLLFIASC*FMLHIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{L74C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVC*VMSLLFIASVFMLHIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{L87C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMSLLFIASVFMC*HIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{S77C.L79C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMC*LC*FIASVFMLHIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{S77C.V84C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMC*LLFIASC*FMLHIWGKY-TRSGPNFYVPFSNKTG
Sec61 β OPG ^{110C} (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPVPLVMSLLFIASVFMLHIWGKY-TRSGPNFYVPFSNKTGC*
Sec61 β OPG-TA ^{110C} eq. (human)	GSPGTPSGTNGSSGRSPSKAVAARAAGSTVRQRKNASS*GTRSAGRRTSAGTGGMWRFYTEDSPGLKVGPPGNFYVPFSNKTGC*
Cytb5OPG ^{S119C} (human)	GSAEQSDAEVKYITLLEIQKHNSKSTWLLHVKVYDLTKFLEEHPGGEVLRQAGGDATENFEDVGHSTDAREMSKTFIIGELHPDDRPKLNKPPETLIT-IDSSSSWWTNWVIPAIC*AVAVALMYRLYMAEDGNFYVPFSNKTG

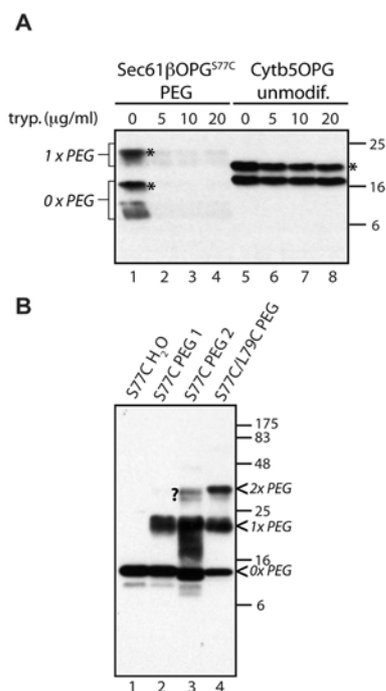


Figure S5 Characterization of PEGylated Sec61 β OPG

(A) PEGylated Sec61 β OPG^{S77C} or unmodified (unmodif) Cytb5OPG^{S119C} were incubated in the presence of reticulocyte lysate and sheep microsomes that had been pretreated with increasing amounts of trypsin (tryp.) as indicated. The membrane fraction was recovered and products were visualized via immunoblotting of the C-terminal opsin tag. N-glycosylated products are indicated (*). (B) Western blot showing purified Sec61 β OPG variants that were untreated (lane 1), treated with mPEG-5000 prequenched with glycine (lanes 2 and 4) or treated with mPEG-5000 without prequenching (lane 3). PEGylated forms are indicated together with a distinct unassigned product seen only in the absence of glycine prequenching (?). S77C, Sec61 β OPG^{S77C}; S77C/L79C, Sec61 β OPG^{S77C.L79C}. Molecular-mass markers in kDa are indicated on the right-hand side.

Table S2 Contribution of different factors to the energetic cost of membrane integration of the PEGylated variants of Sec61 β

See the text for more details. All values are in kcal/mol unless stated otherwise.

PEG link	Tail-anchor contribution	Maleimide linker	PEG distance (Å)	PEG contribution	Sum
L74C	-11.5	3.1	3.9	2.3	-6.1
S77C	-11.5	3.1	7.5	4.4	-4.0
V84C	-11.5	3.1	16	9.4	1.0
L87C	-11.5	3.1	19.6	11.5	3.1

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