Mycolactone reveals the substrate-driven complexity of Sec61-dependent transmembrane protein biogenesis

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ABSTRACT
Mycolactone is the exotoxin virulence factor produced by Mycobacterium ulcerans, the pathogen responsible for Buruli ulcer. The skin lesions and immunosuppression that are characteristic of this disease result from the action of mycolactone, which targets the Sec61 complex and inhibits the co-translational translocation of secretory proteins into the endoplasmic reticulum. In this study, we investigate the effect of mycolactone on the Sec61-dependent biogenesis of different classes of transmembrane protein (TMP). Our data suggest that the effect of mycolactone on TMP biogenesis depends on how the nascent chain initially engages the Sec61 complex. For example, the translocation of TMP luminal domains driven by an N-terminal cleavable signal sequence is efficiently inhibited by mycolactone. In contrast, the effect of mycolactone on protein translocation that is driven solely by a non-cleavable signal anchor/transmembrane domain depends on which flanking region is translocated. For example, while translocation of the region N-terminal to a signal anchor/transmembrane domain is refractive to mycolactone, C-terminal translocation is efficiently inhibited. Our findings highlight the diversity of Sec61-dependent translocation and provide a molecular basis for understanding the effect of mycolactone on the biogenesis of different TMPs.

KEY WORDS: Endoplasmic reticulum, Membrane protein, Mycobacterium ulcerans, Mycolactone, Protein translocation, Sec61

INTRODUCTION
The exotoxin mycolactone is produced by Mycobacterium ulcerans and is the causative agent of Buruli ulcer, a disease characterised by necrotic skin ulcers and immunosuppression (George et al., 1999; Silva et al., 2009; Walsh et al., 2011). Mycolactone is linked to the underproduction of several key proteins involved in the inflammatory response (Pahlelev et al., 1999; Simmonds et al., 2009; Torrado et al., 2007) and the control of blood coagulation (Ogbechi et al., 2015) as a direct result of its effect on the Sec61 complex at the endoplasmic reticulum (ER) (Baron et al., 2016). Mycolactone does not interfere with SRP-dependent delivery of secretory proteins to the ER but rather prevents their co-translational translocation, most likely by stabilising the Sec61 complex in a closed conformation (McKenna et al., 2016) by interacting near the luminal plug of Sec61α (Baron et al., 2016).

A second major group of proteins that are initially targeted to the ER are the transmembrane proteins (TMPs) that, with the exception of tail-anchored proteins (Kutay et al., 1995), also depend on both SRP and the Sec61 translocon for entry into the ER (Cross et al., 2009b; High et al., 1993; Oliver et al., 1995). Following their initial delivery to the Sec61 translocon, TMP biogenesis involves, firstly, translocation of a hydrophilic region of the polypeptide into the ER lumen and, secondly, the stable integration of the polypeptide into the ER lipid bilayer. TMPs contain at least one transmembrane domain (TMD) that interacts with the Sec61 translocon, halting full translocation and mediating release of the TMP into the lipid phase via the lateral gate (Martoglio et al., 1995; Van den Berg et al., 2004). Consistent with its effect at the Sec61 translocon, mycolactone does not affect the membrane insertion of tail-anchored proteins but does affect the integration of the TMPs TNFa (also known as TNF) (Hall et al., 2014) and thrombomodulin (Ogbechi et al., 2015).

Single-pass TMPs can be sub-divided into three main classes based on whether or not they possess a cleavable N-terminal signal sequence, and their final topology in the ER membrane (see Table 1). Type I TMPs contain an N-terminal signal sequence as well as an internal TMD that anchors the polypeptide in the ER membrane with its N-terminus in the ER lumen and its C-terminus in the cytosol (‘N-lumenal–C-cytosolic’). Neither type II nor type III TMPs possess an N-terminal signal sequence and so depend on a single non-cleavable internal TMD (also known as a signal anchor) to both target them to the ER and anchor them in the membrane. While type II TMPs establish an N-cytosolic–C-lumenal topology, type III TMPs have an N-lumenal–C-cytosolic topology (Goder and Spiess, 2001).
The final topology of these different TMPs appears to be determined by their interactions with the Sec61 translocon during their biosynthesis. For example, type III TMPs favour a ‘headfirst’ mode of insertion into the Sec61 complex, where N-terminal translocation into the ER lumen occurs as soon as the TMD emerges from the ribosomal exit tunnel (Kida et al., 2000). Type II TMPs with sufficiently short (<20 residue) N-terminal domains may also enter the Sec61 translocon headfirst, with their N-terminus initially exposed to the ER lumen, before fully inverting to adopt their correct final (N-cytosolic–C-lumenal) topology (Devaraneni et al., 2011). Type II TMPs with longer (>20 residue) N-terminal domains, such as the mycolactone-sensitive TNFα (Hall et al., 2014), are most likely to engage with the translocon as a hairpin, where the N-terminus remains exposed to the cytosol and on-going translation provides the force necessary to translocate the C-terminal domain into the ER lumen (Kocik et al., 2012). Since type I TMPs possess an N-terminal signal sequence, it is assumed that the initial stages of their biogenesis are mechanistically similar to the full translocation of secretory proteins (Walter and Lingappa, 1986). However, whilst it has been demonstrated that the N-terminal signal sequence of type I TMPs initially inserts into the Sec61 translocon as a hairpin (Shaw et al., 1988), there is surprisingly little additional biochemical evidence to support this hypothesis. An obvious question is therefore: how does mycolactone affect the integration of these distinct classes of TMP?

In this study, we characterise the integration of type I, II and III TMPs and identify clear differences in how these processes are affected by synthetic mycolactone A/B (hereafter referred to as mycolactone). Our findings build upon our current understanding of the inhibitory effects of mycolactone on protein translocation at the Sec61 translocon and highlight the mechanistic diversity in the Sec61-mediated translocation of nascent polypeptides across, and insertion into, the ER membrane. Based on our findings, we propose a model where, in the presence of mycolactone, Sec61 is altered such that the headfirst insertion of polypeptides can still occur, but both hairpin insertion into and polypeptide inversion within the translocon are restricted.
RESULTS
TMDs of the type I TMPs CD3δ and GypA can partially rescue their membrane integration in the presence of mycolactone

We have previously demonstrated that co-translationally translocated secretory proteins are prevented from accessing the ER lumen due to the inhibitory effect of mycolactone at the Sec61 translocon (Hall et al., 2014; McKenna et al., 2016). Like secretory proteins, type I TMPs contain a hydrophobic cleavable signal sequence at their N-terminus (Table 1). In addition, type I TMPs contain a second hydrophobic domain (their TMD) that ultimately spans the ER membrane to generate an integral membrane protein with an N-luminal–C-cytosolic topology (High and Dobberstein, 1992). Using a reconstituted in vitro system, we tested the T-cell surface glycoprotein CD3 delta chain (CD3δ, encoded by CD3D; Fig. 1A) for its ability to integrate into ER-derived canine rough microsomes (RMs) in the presence and absence of mycolactone. As a type I TMP, successful insertion of CD3δ is indicated by modification of its two endogenous N-glycosylation sites contained within the region N-terminal to the TMD (Fig. 1A). In the presence of mycolactone, we observed a substantial reduction in the amount of glycosylated CD3δ, but note that a small but significant proportion of glycosylated substrate persists (Fig. 1B,C; Fig. S1A). This behaviour is distinct from the complete block on translocation of secretory proteins that we had tested previously (Hall et al., 2015; Voorhees and Hegde, 2016; see also Introduction).

Strikingly, we observed almost no loss in CD3δ glycosylation in the presence of mycolactone following this single amino acid substitution (Fig. 1C,G). We made similar observations regarding...
the mycolactone sensitivity of a second naturally occurring type I TMP, glycophorin A (GypA; Figs S1A-E). Together, these data demonstrate that type I TMPs are subject to mycolactone-dependent inhibition of the Sec61 translocon but, in contrast to the secretory proteins that we have studied previously, their subsequent TMD can influence this process.

**ER integration of the type I TMP CD3δ in the presence of mycolactone is driven by its TMD**

On the basis of their similarity to secretory proteins, it is generally assumed that the signal sequence of type I TMPs is sufficient to both co-translationally target the nascent chain to the Sec61 translocon and to enable the subsequent translocation of its luminal domain across the ER membrane. Here, the TMD simply acts as a ‘stop transfer’ sequence, halting further translocation of the type I TMP and enabling its release into the ER membrane (Walter and Lingappa, 1986) (Fig. 2Ci). The data presented in Fig. 1, however, suggest the TMD does not simply provide a stop-transfer sequence but indicate that in the presence of mycolactone the TMD can actively promote translocation of the type I TMP luminal domain. To further investigate the roles of the signal sequence and TMD, we generated a series of C-terminally truncated versions of the CD3δ-encoding mRNA that lack stop codons and so produce ribosome-trapped nascent chains that reflect different stages of biogenesis (Gilmore et al., 1991). In the absence of mycolactone, each of these truncations is capable of efficient membrane integration/translocation (Fig. 2A, top panel, lanes 1, 3, 5, 7, 9, 11), even when the TMD is predicted to be fully or partially obscured by the ribosomal exit tunnel (Cabrita et al., 2016) (Fig. 2A, bracketed area), or is missing completely (Fig. 2A, lane 11). These findings support a model in which the signal sequence of this type I TMP is normally sufficient for both ER targeting and translocation.

**Fig. 2. ER integration of the type I TMP CD3δ in the presence of mycolactone is driven by its TMD.** (A) Truncated mRNAs coding for CD3δ (top panel) and CD3δD111L (bottom panel) and lacking stop codons translated in the absence or presence of mycolactone (MYC) without puromycin-mediated release. The nascent chain length of each truncation is shown, as well as the number of residues synthesised C-terminal to the TMD to provide an estimate of its distance from the peptidyl-transferase centre (PTC) of the ribosome. Truncations where all or part of the TMD is likely obscured by the ribosomal exit tunnel (based on Cabrita et al., 2016) are indicated by the bracketed area. CD3δ158 is encompassed by a dashed bracket, since its TMD is likely on the border of having just fully emerged from the ribosomal exit tunnel. Arrowheads indicate maximal glycosylation resulting from the TMD-dependent rescue of integration in the presence of mycolactone. (B) Versions of (i) CD3δ and (ii) CD3δD111L lacking signal sequences (ΔSS) translated in the absence and presence of mycolactone, without or without subsequent EndoglycosidaseH (EndoH) treatment. (C) Predicted mechanism of type I TMP integration in the absence (i) and presence (ii) of mycolactone. Other symbols are as defined in Fig. 1 legend. ‘+g’, glycosylated; ‘0g’, non-glycosylated; ‘C’, C-terminus; FL, full length; ‘N’, N-terminus.
speculate that although the entire TMD of CD3δ, with the 173-residue protein (Fig. 2A, bottom panel, see VCAM1 in the presence of mycolactone (Fig. 3A; Fig. S1A), and 600 residues. Notably, we observed no membrane integration of protein 1 (VCAM1), which possesses an equivalent region of over 600 residues. Notably, we observed no membrane integration of mycolactone (Fig. 2A, lane 4). We assume that the low levels of glycosylation of shorter truncations of CD3δD111L in the presence of mycolactone reflect the spontaneous release of some nascent chains from the ribosome (Hentzen et al., 1972).

In the case of multi-pass TMPs, TMDs of insufficient hydrophobicity can enter the translocon and remain there until a second TMD arrives, at which point the two TMDs co-operatively open the lateral gate of the translocon and exit as a pair (Meindl-Beinker et al., 2006; Pitonzo et al., 2009). To investigate if the signal sequence and TMD of CD3δ might behave in a similar way, the signal sequence was removed from CD3δ and the CD3δD111L variant, and the integration efficiency examined. Both of these polypeptides lacking the signal sequence (DSS) are N-glycosylated (Fig. 2Bi,Bii), demonstrating that they retain the capacity to be targeted to the ER and inserted in the correct (N-lumenal–C-cytosolic) topology. Furthermore, the extent to which the integration of each construct was inhibited by mycolactone was qualitatively similar to their respective signal-sequence-containing versions (cf. Fig. 1B,G). We therefore conclude that the extent of mycolactone sensitivity of CD3δ integration is primarily determined by its TMD.

These findings lead us to propose that some type I TMPs can employ an alternative mechanism for ER translocation in the presence of mycolactone (Fig. 2Cii), thereby accounting for the portion of substrate that successfully integrates in vitro. In this instance, the signal sequence is sufficient to target the translating ribosome to the ER (McKenna et al., 2016) but is unable to mediate the translocation of the luminal domain. Instead, as translation continues, our data suggest that the polypeptide accumulates on the cytosolic side of the ER until its TMD can interact productively with the Sec61 translocon to retroselectively enable translocation of the luminal domain. While we assume that the signal sequence and TMD of such type I TMPs engage the same Sec61 translocon during this process (Gogala et al., 2014), we cannot rule out the possibility that the TMD engages a second translocon that is distinct from the one that has been unsuccessfully engaged by the signal sequence.

The large N-terminal domain of the type I TMP VCAM1 results in a complete block of its membrane integration with mycolactone

The regions separating the signal sequences and TMDs of both CD3δ and GypA are comparatively short (83 and 68 residues, respectively). For this reason, we chose to study the effect of mycolactone on the membrane integration of vascular cell adhesion protein 1 (VCAM1), which possesses an equivalent region of over 600 residues. Notably, we observed no membrane integration of VCAM1 in the presence of mycolactone (Fig. 3A; Fig. S1A), and hypothesised that its large luminal domain may explain the lack of a mycolactone-resistant pool. To this end, we generated a shorter version of VCAM1 with only 60 residues separating its signal sequence and TMD (see Fig. S2B and Fig. 3B, VCAM160).

Strikingly, we now observed a partial rescue of membrane integration of VCAM160 in the presence of mycolactone, as indicated by signal sequence cleavage and modification of an artificially engineered N-glycosylation site (Fig. 3C, C52N). Furthermore, this TMD-dependent effect is enhanced when the hydrophobicity of the VCAM160 TMD is increased by altering a single amino acid residue (Fig. 3D). In contrast, this more hydrophobic TMD has no effect on the mycolactone-sensitivity of full-length VCAM1 (Fig. S2A). Upon extending the region between the signal sequence and TMD of VCAM160 by an additional 50 residues (VCAM1110), the partial rescue of protein integration in the presence of mycolactone is lost, even in combination with the more hydrophobic TMD (Fig. S2B,C). These findings highlight the importance of both TMD hydrophobicity and luminal domain size in conferring sensitivity to mycolactone.

Our characterisation of VCAM1 as a mycolactone-sensitive substrate allows us to draw comparisons with the mechanisms of CAM741, which is a well-defined small molecule that inhibits Sec61-dependent translocation in a highly substrate-specific manner (Besemer et al., 2005). Previous studies have demonstrated that VCAM1 integration is selectively and efficiently blocked by CAM741 on the basis of its signal sequence composition (Harant et al., 2006). In contrast to mycolactone, CAM741 efficiently inhibits the translocation of both VCAM1 and VCAM160, but not of a third – the CAM741-resistant substrate CecOPG2, an opsin-tagged construct derived from Hyalophora cecropia cecropin-A (Johnson et al., 2013) (Fig. 3E). Furthermore, increasing the net hydrophobicity of the VCAM160 TMD does not reduce the effectiveness of CAM741 inhibition (Fig. 3F). Therefore, our studies of VCAM1 highlight key differences in the inhibitory mechanisms of the polyketide mycolactone and the cyclopeptolide CAM741 (see Discussion).

Mycolactone does not interfere with type III TMP integration

We next sought to study the effect of mycolactone on a naturally occurring membrane protein that has the same final topology as a type I TMP but that lacks an N-terminal signal sequence. Glycophorin C (GypC) is a naturally occurring type III single-pass transmembrane protein (type III TMP; High and Tanner, 1987) and so depends on an internal TMD to both target the nascent polypeptide to the ER and anchor it in an N-lumenal–C-cytosolic topology within the ER membrane (see Table 1). Strikingly, the integration of GypC into RMs is unaffected by mycolactone (Fig. 4A,B; Fig. S1A). Furthermore, versions of GypC possessing point mutations that reduce net TMD hydrophobicity remain resistant to mycolactone (Fig. 4B,C,D), suggesting that hydrophobicity per se is not sufficient to explain this observation. Similar findings were also made using a second naturally occurring type III TMP, synaptotagmin 1 (Syt1) (Kida et al., 2000; Perin et al., 1991) (Fig. S3A).

To establish if mycolactone affects GypC at a stage before it has completed membrane integration, we again used truncated mRNAs to generate ribosome-trapped nascent chains (Fig. 4E). Furthermore, in order to specifically focus on the translocation status of nascent ribosome-associated nascent chains (Fig. 4E). Furthermore, in order to specifically focus on the translocation status of nascent ribosome-associated nascent chains (Fig. 4E). Furthermore, in order to specifically focus on the translocation status of nascent ribosome-associated nascent chains (Fig. 4E).
Fig. 3. The large N-terminal domain of the type I TMP VCAM1 results in a complete block of its membrane integration by mycolactone. (A) VCAM1 translated in the absence or presence of mycolactone (MYC) and treated with EndoglycosidaseH (EndoH). (B) VCAM1 and VCAM1<sub>60</sub> constructs (wild type and S707L/S707L* mutants) used in this study. (C) VCAM1<sub>60</sub> and a version containing an artificial N-glycosylation site (C52N) translated in the absence or presence of mycolactone, without or without EndoH. (D) VCAM1<sub>60</sub> and a variant with a more hydrophobic TMD (VCAM1<sub>60</sub>-S707L*) translated in the absence or presence of mycolactone, without or without subsequent EndoH treatment. Estimated TMD hydrophobicities (kcal/mol) are indicated in D. Graph shows the reduction in the amount of +g VCAM1<sub>60</sub> and VCAM1<sub>60</sub>-S707L* in the presence of mycolactone, relative to control samples, as described in the legend to Fig. 1. Graph is also shown in D (graph). The statistical test performed was one-way ANOVA. Error bars show mean±s.d. VCAM1<sub>60</sub>, n=3; VCAM1<sub>60</sub>-S707L*, n=3). P-values are as defined in Fig. 1 legend. (E) Translation of VCAM1, VCAM1<sub>60</sub> and the secretory protein cecropin, possessing a C-terminal opsin tag (CecOPG2), performed with increasing concentrations of CAM741 or an equivalent volume of DMSO (−). (F) VCAM1<sub>60</sub> and VCAM1<sub>60</sub>-S707L* translated in the absence or presence of 250 nM CAM741. Other symbols are as defined in Fig. 1 legend.
for GypC110, for which the TMD is predicted to be partially obscured by the ribosomal exit tunnel (Cabrita et al., 2016), as well as GypC122 peptRNA, for which the TMD is likely to be fully exposed (Fig. 4E,F). In the presence of mycolactone, however, glycosylation of GypC110 peptRNA is lost whereas that of GypC122 peptRNA is maintained (Fig. 4F). GypC is therefore sensitive to mycolactone at some point during its biogenesis but achieves mycolactone-resistance when the TMD is sufficiently beyond the ribosomal exit tunnel.

To investigate the effect of mycolactone on the interacting partners of these GypC integration intermediates, we introduced a single cysteine residue at either the N-terminal [*(52)] or C-terminal [*(84)] side of its TMD (Fig. 4E). This allowed us to generate cysteine–cysteine crosslinks upon addition of bis-maleimidohexane (BMH). In the absence of mycolactone, we observed crosslinking from the N-terminal cysteine probe to Sec61α with the shortest GypC truncation constructs (GypC110 and GypC115) (Fig. 4G, top left panel, see arrowheads; Fig. S3B). These adducts are N-glycosylated (Fig. S3C) and therefore represent N-terminally translocated nascent GypC polypeptides. Furthermore, these adducts are lost upon disruption of the ribosome–nascent-chain complex following treatment with puromycin (Fig. S3D), thus demonstrating that they reflect the environment of bona fide trapped integration intermediates. Extension of the nascent chain by just seven residues (to GypC122)
results in a concomitant loss of adducts with the N-terminal cysteine probe and an increase in crosslinking from the C-terminal cysteine probe to both Sec61\(\alpha\) and Sec61\(\beta\) (Fig. 4G, top two panels; Fig. S3B). When the same analysis is performed in the presence of mycolactone, the Sec61\(\alpha\) adducts with GypC\(_{110}\) and GypC\(_{115}\) are lost, while those of the two longer intermediates are maintained (Fig. 4G, bottom panels; Fig. S3D), consistent with the acquisition of mycolactone resistance at these longer chain lengths (Fig. 4F).

Taken together, these data show that integration of type III TMPs at the ER is highly resistant to mycolactone, even when these substrates possess TMDs of relatively modest hydrophobicity. Additionally, our data suggest that only a portion of the GypC TMD needs to be exposed in order for N-terminal translocation to occur but that these truncated polypeptides are transiently sensitive to mycolactone. Acquisition of mycolactone resistance occurs when the C-terminus of GypC is extended by just a few residues, at which point the TMD is most likely to be fully exposed and not protected by the ribosome.

**Mycolactone efficiently blocks integration of type II TMPs**

While N-terminal signal sequence-driven translocation of polypeptides through the Sec61 translocon is efficiently blocked by mycolactone, its effects on the TMD-driven translocation of the substrates we have so far investigated appears to be more variable. One possible explanation for this observation is that TMDs tend to be longer and more hydrophobic than N-terminal signal sequences (von Heijne, 1985, 1986), and this may enable them to overcome the effects
of mycolactone more successfully. Alternatively, the differences we observed may be determined by whether the region that is being translocated is N- or C-terminal to the hydrophobic ER-targeting sequence that engages the Sec61 complex. For example, while an N-terminal signal sequence translocates the region to its C-terminus, the TMDs of type III TMPs (Kida et al., 2000), and type I TMPs when analysed in the presence of mycolactone (this study), translocate regions that are to their N-terminals.

We therefore studied the MHC class II-associated invariant chain (Ii; Fig. 5A), a type II TMP with a single TMD that enables the translocation of its C-terminus, resulting in a final N-cytosolic–C-lumenal topology. Despite having a long and relatively hydrophobic TMD (compared to CD3δ and GypC; cf. Figs 1F, 4C and 5B), correctly integrated Ii is barely detectable in the presence of mycolactone (Figs 5C,D; Fig. S1A). Furthermore, integration of Ii remains highly sensitive to mycolactone when the hydrophobicity of the TMD is increased to a ΔG value similar to that for GypC (cf. Figs 4C and 5B, and Fig. 5E). Likewise, mycolactone efficiently blocks the integration of Ii with a truncated C-terminus (IC125, Fig. 5F), thus ruling out the possibility that its sensitivity is due to the larger size of its lumenal domain.

Despite the strong inhibitory effect of mycolactone on Ii integration, we observed surprisingly little change in the crosslinking profile of ribosome-trapped nascent chains of various lengths when mycolactone is present (Fig. 5G,H). What we do find is that the full membrane integration of Ii following puromycin-mediated release from the ribosome is prevented by mycolactone. Instead, the nascent chain is retained in proximity to the translocon, as indicated by continued cross-linking to Sec61α in the presence of mycolactone (Fig. 5I). We therefore conclude that mycolactone prevents the type II TMP Ii from assuming its correct N-cytosolic–C-lumenal topology, and instead causes the nascent chain to be retained at a pre-integration step (see Discussion).

**Mycolactone sensitivity is dependent upon which TMD-flanking region is translocated**

The observations in this study with type II and type III TMPs suggest that mycolactone sensitivity may be dependent on whether the region that is translocated through the Sec61 complex lies N- or C-terminal to the TMD. These comparisons have so far relied on substrates that vary in several factors known to contribute to final protein topology, including TMD hydrophobicity, relative position of the TMD, and the number and location of charged residues flanking the TMD. To address these issues, we generated a chimeric protein that contains Ii downstream of a pre-prolactin signal sequence (PPL-Ii; Fig. 6Ai) with the intention of promoting the unnatural N-terminal translocation of this artificial protein across the ER membrane. We therefore expected to observe either cleavage of the N-terminal signal sequence (Fig. 6Aii) or N-glycosylation of the C-terminal region (Fig. 6Aiii), depending on whether the region that is translocated is N- or C-terminal to the TMD. By introducing an artificial N-glycosylation site between the signal sequence and TMD, we exclude a third possible scenario in which the chimeric protein translocates the domain N-terminal to the TMD but signal sequence cleavage does not occur (Fig. S4A,B). In the absence of mycolactone, we observed both signal-cleaved and N-glycosylated...
PPL-Ii (Fig. 6B, lane 1), indicating that this artificial protein assumes a mixed transmembrane topology. In the presence of mycolactone, however, almost no glycosylation is detected, yet signal sequence cleavage is still observed (Fig. 6B, lane 3, and 6C). Increasing the hydrophobicity of the TMD of PPL-Ii (see Fig. 5B) results in the complete insensitivity of signal sequence cleavage to mycolactone, whereas N-glycosylation remains almost completely blocked (Fig. 6B, lane 6, and 6C). Since the PPL signal sequence is incapable of overcoming the inhibitory effect of mycolactone (McKenna et al., 2016) (Fig. 1D), and we observed no contribution of the signal sequence to the mycolactone resistance of TMD-containing proteins (Fig. 2B), both final topologies of PPL-Ii in the presence of mycolactone must result from membrane integration enabled by the TMD. We therefore conclude that even when membrane integration in two different topologies can be driven by the same TMD, translocation of its C-terminal region is preferentially inhibited by mycolactone.

**Mycolactone traps headfirst-inserting type II TMPs in an N-lumenal–C-cytosolic topology**

Previous studies suggest that type II TMPs can engage the Sec61 translocon in one of two ways, depending largely on the size of their N-terminal domain. Those with an N-terminal domain longer than ∼20 residues, like our model type II TMP, Ii, are generally believed to insert as a hairpin, while those with an N-terminal domain shorter than ∼20 residues are proposed to insert headfirst, like a type III TMP, before fully inverting within the translocon (Devaraneni et al., 2011; Kocik et al., 2012; Table 1). To look more closely at the effect of mycolactone on these two alternative modes of insertion of type II TMPs, we used the asialoglycoprotein receptor subunit H1 (Fig. 7A, ASGPR H1), which possesses a 40 residue N-terminal domain, in combination with a truncated form that possesses an N-terminal region of just four amino acids (Fig. 7A) (ASGPR H1Δ; Wahlberg and Spiess, 1997). Previous studies suggest that ASGPR H1 employs a hairpin mechanism for membrane insertion whilst the truncation of its N-terminal domain promotes the insertion of ASGPR H1Δ via a headfirst mechanism (Kocik et al., 2012; Wahlberg and Spiess, 1997). N-glycosylation of both ASGPR H1 and ASGPR H1Δ is strongly inhibited by mycolactone (Fig. 7B,C,D, see ‘+g’; Fig. S1A), supporting our conclusion that mycolactone efficiently prevents type II TMPs from achieving their correct N-cytosolic–C-lumenal topology. Notably, the TMD of the type II TMP ASGPR H1 is more hydrophobic than that of Ii (ΔG of −2.828 kcal/mol versus −0.756 kcal/mol), reinforcing that hydrophobicity alone is not sufficient to explain the effects we see of mycolactone on TMP biogenesis. Interestingly, the amount of non-glycosylated ASGPR H1Δ increases in the presence of mycolactone, even after the membranes are treated with alkaline sodium carbonate to remove non-integrated polypeptides (Fig. 7C, cf. lanes 1 and 3, and 7D, see ‘0g’). We therefore conclude that the non-glycosylated ASGPR H1Δ species most likely represents successfully integrated substrate that has failed to invert and hence remains in a N-lumenal–C-cytosolic topology [cf. Fig. 7Eii, Eiii; see also Wahlberg and Spiess, 1997]. In contrast, the amount of non-glycosylated full-length ASGPR H1 does not increase in the presence of mycolactone under similar conditions (Fig. 7B, cf. lanes 1 and 3, and 7D, see ‘0g’), suggesting that in this case, the nascent polypeptide has simply failed to integrate (Fig. 7Ei). These results therefore suggest that type II TMPs that employ a headfirst-inversion mode of insertion into the ER may be prevented from inverting within the Sec61 translocon by mycolactone and, instead, become integrated into the ER membrane in an N-lumenal–C-cytosolic topology (Fig. 7Eii, Eiii).

**DISCUSSION**

In this study, we explore the inhibitory mechanism of mycolactone at the Sec61 translocon by investigating the integration of three distinct classes of single-pass transmembrane proteins (TMPs). We conclude that mycolactone binding restricts the Sec61 complex, leading to disruption of TMP biogenesis. However, distinct classes of TMP respond differently to mycolactone, most likely reflecting the precise nature of their initial engagement at the translocon. Thus, whilst the headfirst insertion of polypeptides is unaffected, their hairpin insertion into and inversion within the translocon are inhibited.

**What makes a protein mycolactone-sensitive?**

Secretory and membrane proteins are key therapeutic targets, and it is important to understand the molecular basis for the selectivity of inhibitors that affect their biogenesis (Kalies and Römisch, 2015). For the majority of small molecules known to modulate ER translocation, they appear to do so by acting directly at the Sec61 complex (Kalies and Römisch, 2015). Compounds such as eeyarestatin and apratoxin inhibit the Sec61-dependent translocation of a broad range of substrates (Cross et al., 2009a; Paatero et al., 2016). In contrast, and despite their common Sec61 target, HUN-7293-derived compounds such as CAM741 and the cotransin family display clear substrate specificity (Besemer et al., 2005; Garrison et al., 2005; Maifeld et al., 2011). Exactly why certain compounds have different substrate-specific effects on ER translocation is unclear, and it is in this context that we are studying the effects of mycolactone, which we have previously showed to have a broad ranging effect on the co-translational translocation of secretory proteins across the ER membrane (Hall et al., 2014; McKenna et al., 2016).

We now present several lines of evidence that support a model in which the inhibition of protein translocation by mycolactone is strongly influenced by how the nascent polypeptide initially engages the Sec61 translocon. Firstly, we have already shown that secretory proteins are efficiently blocked by mycolactone (Hall et al., 2014; McKenna et al., 2016). We now establish that removal of the TMDs of type I TMPs that otherwise retain some degree of resistance to mycolactone generates artificial secretory proteins that are strongly inhibited by mycolactone as a consequence of their now wholly N-terminal signal sequence-dependent ER translocation. Secondly, we show that type I TMPs can integrate into the ER via one of two mechanisms: in the absence of mycolactone, the N-terminal signal sequence is sufficient to mediate translocation of the region C-terminal to it, consistent with the generally assumed mechanism of integration for type I TMPs (Walter and Lingappa, 1986); whilst in the presence of mycolactone, translocation driven by the N-terminal signal sequence is efficiently blocked and, instead, translocation of sufficiently small lumenal domains can be mediated by the TMD. Here, we see that removal of the N-terminal signal sequence from the type I TMP CD3δ has no effect on the extent to which its membrane insertion is inhibited by mycolactone. Thirdly, although both type II and type III TMPs contain a single TMD, the former translocate their C-terminal region and are strongly inhibited by mycolactone, while the latter translocate their N-terminal region and are unaffected by mycolactone treatment. Fourthly, by generating a chimeric protein whose TMD can translocate either its N- or C-terminus, we show that C-terminal translocation is inhibited by mycolactone to a much greater extent.
Lastly, we use a truncated version of ASGPR H1 as a model to study the so-called headfirst-inversion mechanism of some type II TMPs (Devaraneni et al., 2011; Kocik et al., 2012; Wahlberg and Spiess, 1997). Our data suggest that mycolactone does not prevent the headfirst insertion of this model substrate into the Sec61 complex but inhibits its subsequent inversion, most likely fixing it in a "type II" topology.
III-like (N-lumenal–C-cytosolic) topology. Interestingly, the N-terminal signal sequences of secretory proteins and type I TMPs have been shown to engage the Sec61 translocon in a hairpin conformation (Mothes et al., 1994; Shaw et al., 1988; Voorhees and Hegde, 2016). Taken together, our findings therefore suggest that polypeptides that initially engage the Sec61 translocon as a hairpin are effectively prevented from correctly integrating/translocating into the ER. In contrast, headfirst insertion of polypeptides into the Sec61 translocon can still occur, but those substrates that require a subsequent inversion step within the translocon are prevented from doing so.

Our conclusion that the type II TMP Li is trapped at a ‘pre-integration’ stage by mycolactone is supported by a previous study of membrane protein biogenesis using the cotransin CT8 (MacKinnon et al., 2014). Li closely resembles TNFα, the model substrate used in that previous study (Fig. 5G). In both cases, similarly positioned cysteine residues form adducts with components of the Sec61 translocon that are lost following full integration into the ER membrane but that are maintained in the presence of mycolactone or CT8. Moreover, mutagenesis of Sec61α and competitive binding assays have shown that CT8 and mycolactone most likely have overlapping binding sites (cf. Baron et al., 2016 and MacKinnon et al., 2014). Whilst these similarities suggest the inhibitory mechanisms of these two small molecules are related, there is notably little effect of CT8 on the biogenesis of other mycolactone-sensitive substrates, such as pre-prolactin (Maifeld et al., 2011; McKenna et al., 2016), highlighting clear differences between them. Furthermore, our investigation of the type I TMP VCAM1 distinguishes the mechanism of inhibition of mycolactone from that of the cotransin-like compound CAM741. While CAM741 blocks integration in a signal sequence-specific manner that prevents productive interactions of a subsequent TMD with the Sec61 translocon, mycolactone indiscriminately interferes with signal sequence-mediated protein translocation, but its effect can be partially overcome by sufficiently hydrophobic TMDs.

To date, the focus for identifying mycolactone-sensitive substrates in a cellular context has been on mediators of the immune response and blood anticoagulation. Both secreted mediators, such as cytokines and chemokines, as well as TMPs, such as receptors, are under-produced in cells treated with mycolactone, while cytoplasmic substrates appear to be consistently unaffected (Coutanceau et al., 2007; Hall and Simmonds, 2014; Simmonds et al., 2009). Our current study suggests that type I TMPs with N-terminal domains of more than ~100 residues are highly sensitive to mycolactone treatment in vitro. Notably, most of the type I TMPs that are mycolactone-sensitive at a cellular level have N-terminal domains of a similar or larger size (cf. Hall and Simmonds, 2014), including thrombomodulin, consistent with the previously suggested mycolactone-dependent inhibition of its ER translocation (Ogbechi et al., 2015). Our limited in vitro analysis of type II TMPs suggests that their membrane integration is highly sensitive to mycolactone, and this is consistent with previous observations using TNFα (Hall et al., 2014). In contrast to our observations with type I and type II TMPs, we find that type III TMP integration appears to be completely resistant to mycolactone. Strikingly, the only type III TMP to be studied at a cellular level, linker for activation of T-cells family member 1 (LAT), has been previously shown to also be unaffected by mycolactone treatment (Boulkroun et al., 2010). Estimates suggest that human type III TMPs represent only ~2% of all single-spanning membrane proteins (Uniprot), and hence the majority of single-spanning TMPs are likely to display some degree of sensitivity to mycolactone. In line with this, a recent study has reported that the majority of Sec61-dependent substrates that could be identified are sensitive to mycolactone, with single-pass TMPs being particularly enriched (Baron et al., 2016). In short, our findings provide a potential molecular basis for the complex changes in membrane protein expression that underlie the pathogenesis of Buruli ulcer.

What can mycolactone tell us about Sec61-dependent translocation?

The substrate-specific inhibition of Sec61-dependent translocation by mycolactone that we describe in this study supports the idea that the TMD of a nascent polypeptide plays an important role in opening the Sec61 lateral gate. A recent study using native membranes concludes that full opening of the Sec61 translocon appears to occur upon the docking of even non-translating ribosomes (Pfeffer et al., 2015). In contrast, our findings align more closely with a model in which ribosome docking primes the Sec61 translocon by partially opening its lateral gate. In this model, full opening occurs only when a sufficiently hydrophobic signal sequence/TMD of the nascent polypeptide chain positionally replaces one of the Sec61α TMDs located at the lateral gate, thereby exposing the hydrophobic region to the lipid phase of the ER membrane (Voorhees and Hegde, 2016). An interesting implication of our study is that opening of the lateral gate may occur more readily when a substrate engages the translocon in a particular orientation (namely headfirst), regardless of its hydrophobicity. Recent structural studies of the engaged Sec61 translocon have greatly advanced our knowledge of ER translocation (Gogala et al., 2014; Voorhees and Hegde, 2016) but have so far only investigated a narrow range of Sec61-dependent substrates. Our findings demonstrate the substrate-driven complexity of Sec61-dependent protein translocation and therefore highlight the importance of considering a broad range of substrates in future structural studies aiming to characterise this process.

MATERIALS AND METHODS

Synthetic mycolactone A/B was a gift from Yoshito Kishi, Harvard University, Cambridge, MA (Song et al., 2002). CAM741 was generously provided by Boehringer. Unless otherwise stated, all standard laboratory reagents were obtained from Merck or Sigma.

DNA constructs

Unless otherwise stated, all cDNAs matched human sequences. GypC, GypA and Li (Mus musculus; Oliver et al., 1995), PPL (Bos taurus; McKenna et al., 2016) and CecOPG2 (Hyalophora cecropia; Johnson et al., 2012) have all been described previously. VCAM1 was a gift from Hanna Harant (Ingenetix). CD3δ was a gift from Cornelia Wilson (Canterbury Christ Church University, Kent, UK). ASGPR H1 was a gift from Martin Spiess (University of Basel, Switzerland) and Syt1 (Rattus norvegicus) a gift from Alan Morgan (University of Liverpool, Liverpool, UK). The PPL-Li chimera was generated by cloning Li downstream of PPL using an engineered EcoRI site. Point mutations were generated using primers from Eurogentec. cDNAs were generated by PCR and transcribed with T7 polymerase (Promega).

Antibodies

Rabbit antiserum against GypC has been described previously (Elliott et al., 1997). Rabbit antiserum against canine Sec61α was from Richard Zimmermann (University of Saarland, Homburg, Germany). Rabbit antiserum against Sec61β and Li were from Bernhard Doberstein (University of Heidelberg, Heidelberg, Germany).

In vitro translation and translocation assays

In vitro translations used rabbit reticulocyte lysate (Promega) to synthesise radiolabelled proteins in the presence of canine rough microsomes as...
Previously described (McKenna et al., 2016), except that reaction volumes were 20 µl and were incubated at 30°C for 15 min. For crosslinking assays, translation reactions were carried out at 30°C for 10 min. All samples were then treated with 0.5 mM puromycin and incubated at 30°C for 5 min unless stated otherwise in the figure legends. For CAM741 inhibition assays, CAM741 (in DMSO) was added to give the appropriate concentration at the beginning of the reaction. The control used an equivalent volume of DMSO.

Membrane recovery and visualisation
Membranes were recovered as described previously (McKenna et al., 2016). Unless stated otherwise in the figure legends, the membrane pellet was immediately resuspended in 30 µl SDS sample buffer [100 mM Tris-HCl, pH 6.8, 100 mM DTT, 4% (w/v) SDS, 20% (v/v) glycerol, 1% (v/v) L-methionine, 10 mM EDTA, Dromophenol blue]. Where indicated, samples were also treated with Endoglycosidase H (EndoH; New England Biolabs) as described by the supplier. The resulting samples were analysed and processed as described previously (McKenna et al., 2016). Data were quantified using AIDA software (Raytek), and statistical analyses (one-way ANOVA or two-way ANOVA) were performed using GraphPad (Prism). The exact sample size (n) for each experimental group is provided in the appropriate figure legends. In each case, n was defined by the number of times the substrate was tested in the same experimental system and so represents technical replicates.

Crosslinking and carbonate extraction
After recovery, the entire pellet was resuspended in 20 µl low-salt buffer [100 mM sucrose, 100 mM KOAc, 5 mM Mg(OAc)2, 50 mM Hepes-KOH pH 7.9, 1 mM DTT]. Crosslinking [using BMH (1 mM final)] and carbonate extraction were performed as described previously (McKenna et al., 2016). Samples were then either analysed directly by SDS-PAGE or were first immunoprecipitated under denaturing conditions (see below).

Denaturing immunoprecipitation
Following carbonate extraction and recovery of the membrane fraction, pellets were resuspended in 20 µl of 1% (w/v) SDS and incubated for 10 min at 70°C. Ten volumes of Triton immunoprecipitation (IP) buffer [10 mM Tris-HCl pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 5 mM PMSF, 1 mM methionine] with the appropriate antiseraum (1:200) were added. Samples were incubated for 15 h at 4°C with constant agitation. Protein-A-Sepharose beads (Genscript) were added to 10% (v/v), and samples were incubated at 4°C for a further 2 h. Protein-A-Sepharose beads were then recovered by spinning at 13,000 g for 1 min and washed with Triton IP buffer before being heated at 70°C for 10 min in SDS sample buffer.

Native immunoprecipitation of PPL-ii
Instead of recovering RMs by ultracentrifugation, PPL-ii translations (see Fig. 6B) were subjected to an anti-Ii immunoprecipitation. Following translation, nine volumes of Triton IP buffer with anti-Ii antiseraum (1:200) were added. Samples were incubated for 15 h at 4°C with constant agitation. Protein-A-Sepharose beads (Genscript) were added to 10% (v/v), and samples were incubated at 4°C for a further 2 h. Protein-A-Sepharose beads were then recovered by spinning at 13,000 g for 1 min and washed with Triton IP buffer before being heated at 70°C for 10 min in SDS sample buffer.

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Competing interests
The authors declare no competing or financial interests.

Author contributions
All authors conceived and designed the experiments. M.M. performed the experiments. All authors contributed to the analysis of the data and the preparation of the manuscript.

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Supplementary information
Supplementary information available online at http://jcs.biologists.org/lookup/doi/10.1242/jcs.2016.bisupplemental

References
Hall, B. S., Hill, K., McKenna, M., Ogbechi, J., High, S., Willis, A. E. and Simmonds, R. E. (2014). The pathogenic mechanism of the Mycobacterium


Figure S1

(A) Graph showing the amount of N-glycosylated substrate in the presence of mycolactone for each of the wild-type transmembrane proteins used in this study, relative to samples treated with a vehicle control, which were taken as 100%. For all samples, n > 3. *Quantification of N-glycosylated ASGPR was performed following carbonate extraction of the membrane fraction of the translation. All other samples were quantified without carbonate extraction. Phosphorimage of in vitro translated GypA (B), GypA_{ΔTMD} (C), and GypA_{T106I} (D) in the absence and presence of mycolactone. (E) Graph showing the proportion of glycosylated material of GypA and related constructs in the presence of mycolactone, relative to control samples. Error bars show mean±s.d (for GypA n = 9, for GypA_{ΔTMD} n = 3 and for GypA_{T106I} n = 5).
Figure S2

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**Supplementary Figure 2.**

(A) VCAM1 and a version containing a more hydrophobic TMD (VCAM1 \(_{S707L}\)) were translated in the absence or presence of mycolactone. (B) Diagram showing the amino acid sequences of VCAM1\(_{60}\) and VCAM1\(_{110}\). Signal sequences are shown in blue, transmembrane domains in pink. Artificially introduced N-linked glycosylation sites and S707L/S707L\(^*\) point mutations are also indicated. (C) VCAM1\(_{110}\) and a version containing a more hydrophobic TMD (VCAM1\(_{110, S707L^*}\)) were translated in the absence and presence of mycolactone.
Figure S3

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Supplementary Figure 3.

(A) The naturally occurring TIII TMP, synaptotagmin 1 (Syt1), was translated in the absence or presence of mycolactone. (B) GypC truncations containing a cysteine residue at either position 52 (*52) or 84 (*84) were translated in the absence or presence of mycolactone without puromycin-mediated release. Samples were then treated with the cross-linking reagent BMH or a vehicle control. Following carbonate extraction, samples were subjected to denaturing immunoprecipitation (IP) using antisera raised against either glycophorin C (GypC), Sec61α (α) or Sec61β (β). (C) GypC_{110} *(52) and GypC_{122} *(84) lacking stop codons were translated in the absence or presence of mycolactone. Samples were treated with BMH or a vehicle control following membrane recovery. Samples were then treated with EndoH where indicated. (D) GypC_{110} *(52) and GypC_{122} *(84) lacking stop codons were translated in the absence or presence of mycolactone. Prior to membrane recovery, samples were treated with puromycin where indicated. Samples were then treated with BMH or a vehicle control.
Supplementary Figure 4.

(A) Translation of chimeric proteins containing pre-prolactin (PPL) signal sequence upstream of either the first 125 or 98 residues of li. Chimeric proteins are either unmodified (WT) or contain a point mutation that introduces an artificial N-glycosylation site between the PPL signal sequence and the li TMD (Q13S*) to report entry into the ER lumen. Following translation, samples were treated with puromycin and/or EndoH as indicated. (B) Diagram of PPL-li125 and PPL-li98 constructs used in this study.