Capture and delivery of tail-anchored proteins to the endoplasmic reticulum

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Tail-anchored (TA) proteins fulfill diverse cellular functions within different organellar membranes. Their characteristic C-terminal transmembrane segment renders TA proteins inherently prone to aggregation and necessitates their posttranslational targeting. The guided entry of TA proteins (GET in yeast)/transmembrane recognition complex (TRC in humans) pathway represents a major route for TA proteins to the endoplasmic reticulum (ER). Here, we review important new insights into the capture of nascent TA proteins at the ribosome by the GET pathway pretargeting complex and the mechanism of their delivery into the ER membrane by the GET receptor insertase. Interestingly, several alternative routes by which TA proteins can be targeted to the ER have emerged, raising intriguing questions about how selectivity is achieved during TA protein capture. Furthermore, mistargeting of TA proteins is a fundamental cellular problem, and we discuss the recently discovered quality control machineries in the ER and outer mitochondrial membrane for displacing mislocalized TA proteins.

Introduction

The protein components of biological membranes expand their functionality beyond physical barriers by acting as gateways, allowing intercompartment communication as well as facilitating transport and other membrane-associated processes. Membrane proteins collectively constitute ∼30% of the proteome of most organisms (Krogh et al., 2001), and their biogenesis represents a major challenge for cells. Their hydrophobic transmembrane domains (TMDs), essential for integration into the lipid bilayer and functionality, render such proteins inherently prone to aggregation in the aqueous cytosolic environment. Dedicated targeting strategies for chaperoning such proteins to their target membranes are therefore necessary. Proteins destined for the ER that carry short signal sequences at their N-terminal end and/or internal TMDs are typically recognized cotranslationally by the signal recognition particle (SRP). This arrests translation and induces relocation of the ribosome nascent chain complex (RNC) to the ER-bound Sec61 translocon, where the newly synthesized protein is channeled directly into the ER lumen and/or membrane (reviewed in Akopian et al., 2013; Rapoport et al., 2017). In both yeast and mammals, a macromolecular ER membrane protein complex (EMC) cooperates with the translocon by assisting the cotranslational folding and biogenesis of polytopic membrane proteins in the ER as well as itself acting as a membrane insertase, mediating the correct topological insertion of the first TMD of specific G-coupled receptors (Bai et al., 2020; Chitwood et al., 2018; Shurtleff et al., 2018). Furthermore, an SRP-independent ER targeting pathway (SND) has recently been revealed in yeast, where proteins containing TMDs in their central regions are captured by Snd1 and directed toward a Sec61 translocon associated with Snd2 and Snd3 (Aviram et al., 2016). This pathway appears to have a broad client spectrum and has been suggested to functionally compensate when other ER targeting pathways are impaired.

Tail-anchored (TA) proteins represent a specific class of membrane proteins characterized by a single TMD close to their C-terminus (reviewed in Kutay et al., 1993). The TA protein family contains >50 members in yeast (Beilharz et al., 2003), >500 in plants (Kriechbaum et al., 2009), and >300 in humans (Kalbfleisch et al., 2007), which populate different membranes (ER, Golgi, and inner nuclear, outer mitochondrial, and peroxisomal membranes). These proteins fulfill diverse membrane-associated functions ranging from regulating intracellular vesicular trafficking (SNARE proteins) to apoptosis, autophagy, lipid biosynthesis, and protein degradation. The topology of TA proteins dictates their posttranslational targeting, as translation termination occurs concurrent with emergence of the TMD from the polypeptide exit tunnel of the ribosome. A major route to the ER for TA proteins is the evolutionarily conserved guided entry of TA proteins (GET) pathway in yeast and the homologous transmembrane recognition complex (TRC) pathway in mammals (Figs. 1 and 2; Borgese et al., 2019; Schuldiner et al., 2008; Stefañovic and Hegde, 2007). The established view of the GET/TRC pathway (reviewed in Borgese et al., 2019; Chio et al., 2018) represents a major route for TA proteins to the endoplasmic reticulum (ER). Here, we review important new insights into the capture of nascent TA proteins at the ribosome by the GET pathway pretargeting complex and the mechanism of their delivery into the ER membrane by the GET receptor insertase. Interestingly, several alternative routes by which TA proteins can be targeted to the ER have emerged, raising intriguing questions about how selectivity is achieved during TA protein capture. Furthermore, mistargeting of TA proteins is a fundamental cellular problem, and we discuss the recently discovered quality control machineries in the ER and outer mitochondrial membrane for displacing mislocalized TA proteins.
2017) involves the posttranslational capture of TA proteins by a pretargeting complex composed of the homodimeric, cytosolic chaperone Sgt2 (yeast)/SGTA (mammals) and the Get4-Get5 heterodimer (yeast)/TRC35-UBL4A-BAG6 complex (mammals; Jonikas et al., 2009; Mariappan et al., 2011; Wang et al., 2010). Interaction of the pretargeting complex with an ATP-bound form of the ATPase Get3 (yeast)/TRC40 (mammals) results in transfer of the TA protein from Sgt2/SGTA to Get3/TRC40 such that the TMD is shielded within a hydrophobic pocket of Get3/TRC40 (Bozkurt et al., 2009; Mateja et al., 2009; Stefanovic and Hegde, 2007; Suloway et al., 2009). ATP hydrolysis triggered by interaction of Get3/TRC40 with the TA protein, coupled with conformational changes in Get3/TRC40 induced by interaction with Get4-Get5/TRC35-UBL4A-BAG6, drives dissociation of TA protein-bound Get3/TRC40 from the pretargeting complex, allowing delivery to the ER-bound GET receptor composed of Get1 and Get2 (yeast)/tryptophan-rich basis protein (WRB) and calcium-modulating cyclophilin ligand (CAML; mammals; Mariappan et al., 2011; Mateja et al., 2015; Stefer et al., 2011). Receptor binding triggers ADP release and conformational rearrangement of Get3/TRC40, allowing TA protein insertion into the membrane and recycling of Get3/TRC40 (reviewed in Mateja and Keenan, 2018). Although much less is currently known about the GET pathway in plants, homologues of the GET pathway components have been identified or are predicted (Xing et al., 2017; Srivastava et al., 2017; Asseck et al., 2021), and two TA SNARE proteins have been shown to be affected by lack of Arabidopsis thaliana (At)GET3 (Xing et al., 2017). A growing body of evidence indicates functional redundancy between different pathways for targeting TA proteins to the ER (Figs. 1 and 2;

Figure 1. **TA protein targeting to the ER in yeast.** Nascent TA proteins emerging from the ribosome can be captured by alternative ER-targeting machineries. A major route to the ER is via the GET pathway, involving ribosome-associated capture by Sgt2, followed by Get4/Get5-mediated handover to the Get3 ATPase and insertion into the ER membrane by a heterotetrameric GET receptor complex composed of Get1 and Get2.
Casson et al., 2017). Although some TA proteins of the secretory pathway fulfill essential cellular functions, deletion of GET pathway components is not lethal in yeast or plants, and for many TA proteins, lack of GET pathway components reduces but does not abolish ER targeting. Consistent with this notion, both the EMC insertase (Guna et al., 2018) and the SND pathway (Aviram et al., 2016) have been shown to support the ER targeting of TA proteins.

Despite a wealth of knowledge on some aspects of TA protein targeting to the ER by the GET/TRC pathway, other features and mechanistic details remain enigmatic. How TA proteins can be captured posttranslationally but also reliably avoid aggregation during handover from the ribosome to the pretargeting complex or other chaperones is an inherent conundrum of their biogenesis. Knowledge on the architecture of the GET/TRC receptor complex and mechanistic understanding of how the TA protein, delivered to the receptor by Get3/TRC40, is inserted into the ER membrane, have been limited by the technically challenging nature of structural analyses of membrane-bound complexes. Furthermore, how the fidelity of TA protein targeting to different membranes is ensured is poorly understood, and little is currently known about how the actions of TA protein targeting and quality control are coordinated. In this review, we describe recent advances that address these key aspects of the GET/TRC pathway and TA protein biogenesis.

**Capture of nascent TA proteins**

In contrast to cotranslational protein targeting, where capture of client proteins and their delivery to the ER-bound receptor complex is performed by the SRP complex, posttranslational targeting of TA proteins to the ER by the GET/TRC pathway is a more stepwise process, involving a dynamically assembling.

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**Figure 2. TA protein targeting to the ER in mammals.** Mammalian TA proteins are predominantly targeted to the ER by the TRC pathway. After capture by SGTA, together with the BAG6 complex (BAG6, UBL4A, and TRC35), the TA protein is passed to the TRC40 chaperone for delivery to the ER-bound receptor complex formed by WRB and CAML. BAG6 has dual functions bridging ER targeting and ubiquitination of TA proteins and can be antagonized by SGTA. Ubiquitinated TA proteins can be deubiquitinated by ER-associated UPS20/UPS33.
pretargeting complex that mediates initial capture but then hands over substrates to another chaperone for delivery to the membrane-bound receptor. The existence of a modular pretargeting complex appears to be a feature of the GET/TRC pathway conserved throughout eukaryotes, but compositional and structural differences between species are apparent. In yeast, Get4-Get5 form an obligate heterodimer, whereas the homologous TRC35 and UBL4A interact via an additional component BAG6 (Chang et al., 2010; Chartron et al., 2010; Mariappan et al., 2010; Mock et al., 2015). Biochemical evidence shows that a Get4/TRC35 homologue exists in plants, and homologues of Get5/UBL4A and Sgt2/SGTA are predicted from in silico analyses (Srivastava et al., 2017; Xing et al., 2017). A BAG6 homologue has also been identified in plants (https://www.arabidopsis.org/servlets/TairObject?id=35038&type=locus); however, it remains unknown if this protein associates with components of the GET pathway and/or contributes to TA protein targeting. These differences likely reflect subtle variations in the mechanism of TA protein capture between species and/or the greater need for regulation and surveillance in multicellular organisms.

**Ribosome binding of pretargeting complex components**

Due to the position of the TMD at the C-terminus of TA proteins, the GET/TRC pathway must capture clients posttranslationally. However, posttranslational capture has the inherent caveat of protein aggregation in the narrow window between emergence of the TMD from the ribosome exit tunnel and protein capture by a chaperoning factor. The first hint how this issue may be overcome came with the intriguing discovery of Get5 in a high-throughput screen for ribosome-associated proteins in yeast (Fleischer et al., 2006). This observation raised the possibility of a physical connection between the upstream components of the GET pathway and the translation machinery, despite the posttranslational nature of the final capture event. Detection of the Get4-Get5 heterodimer associated with polysomes importantly confirmed recruitment of these proteins to actively translating polysomes (Zhang et al., 2016), further supporting that the GET pathway pretargeting complex might be poised on the ribosomes ready to shield nascent TA proteins directly as they emerge from the exit tunnel. In vitro reconstitution confirmed a high-affinity interaction between Get4-Get5 and ribosomes, and protein−protein−protein−RNA cross-linking analyses pinpointed the polypeptide exit tunnel of the ribosome as the Get4-Get5 binding site (Fig. 1; Zhang et al., 2021). Get4-Get5 bridge interactions between Sgt2 and Get3 to facilitate handover of the TA protein to the downstream chaperone, and therefore functional significance of Get4-Get5 ribosome binding must be coupled to TA protein capture by Sgt2. Indeed, it was recently shown that Get4-Get5 act as a binding platform for recruitment of Sgt2 to ribosomes and that the presence of Get4-Get5 on ribosomes enhances TA protein capture by Sgt2 (Zhang et al., 2021). Crystal structures of GET pathway subcomplexes demonstrate that the N-terminal domains of the Sgt2 homodimer interact with the central UBL domain of Get5, while the N-terminal region of Get5 mediates interaction with C-terminal region of Get4 (Chang et al., 2010; Chartron et al., 2012a; Simon et al., 2013). As Get5 appears to simultaneously interact with Get4, Sgt2, and ribosomes, it is tempting to speculate that ribosome binding of the pretargeting complex may be mediated by the C-terminal region of Get5. However, due to the multimeric nature of the pretargeting complex and evidence that the Get5 C-terminal region also mediates homodimerization (Chartron et al., 2012b), structural analyses of Get4-Get5-Sgt2–bound ribosomes will be necessary to resolve in detail the architecture of GET pathway pretargeting complex–bound ribosomes.

The mammalian pretargeting complex components UBL4A, TRC35, BAG6, and SGTA also associate with RNC complexes (Fig. 2; Leznicki and High, 2020; Mariappan et al., 2010), implying that the mechanism of ribosome-associated capture of TA proteins is also employed in mammalian cells to circumvent protein aggregation upon the TMD encountering the aqueous cytosol during targeting. However, the mammalian-specific pretargeting complex component BAG6, rather than the Get5-homologous UBL4A, appears to act as the key ribosome adaptor, and SGTA can also interact with RNC complexes independently of TRC35-UBL4A (Leznicki and High, 2020; Mariappan et al., 2010).

Notably, Get3/TRC40, which receive the TA protein from the pretargeting complex, are not ribosome associated (Mariappan et al., 2010; Zhang et al., 2016). This implies that following Get4-Get5–facilitated capture of the TA protein by ribosome-associated Sgt2, the pretargeting complex should dissociate from the ribosome to encounter Get3. It is possible that a conformational change in the pretargeting complex, induced by TA protein binding, triggers release from the ribosome.

**Recognition of ribosomes synthesizing TA proteins**

The discovery of ribosome-associated populations of the GET pretargeting complex and the BAG6 complex and SGTA in yeast and mammals, respectively, raises the question of how these complexes, which are significantly less abundant than cytosolic ribosomes, are able to identify ribosomes synthesizing TA proteins. Analogous to SRP, which probes the translating ribosome population, preferentially binding ribosomes translating proteins with a signal sequence (Berndt et al., 2009; Holtkamp et al., 2012; Ogg and Walter, 1995; Voorhees and Hegde, 2015), yeast Get4-Get5 show increased association with RNCs containing a TA/TMD in the exit tunnel (Zhang et al., 2021). Mammalian SGTA is likewise selectively recruited to ribosomes synthesizing membrane proteins (Leznicki and High, 2020). This implies that a similar substrate-scanning mechanism, involving transient association events followed by high-affinity docking only onto appropriate ribosomes, is employed in both the co- and posttargeting pathways. The underlying mechanism of how the GET/TRC pretargeting components sense the presence of the TA/TMD in the exit tunnel still remains to be elucidated. Intriguingly, the newly identified ribosome-binding site of the GET pathway pretargeting complex overlaps with that of SRP, implying mutually exclusive ribosome occupancy. Consistent with this, the presence of Get4-Get5 on RNCs with a TMD in the exit tunnel reduces the amount of SRP bound, and strong binding of SRP to an exposed TMD leads to displacement of Get4-Get5 (Zhang et al., 2021). This suggests a compelling model
where the GET/TRC pretargeting complex, SRP, and potentially ribosome-bound Snd1 (Fleischer et al., 2006) constantly screen the translating pool of ribosomes and outcompete each other for ribosome binding upon encountering one synthesizing an optimal substrate.

Additional players in initial TA protein capture
It is well established both in vitro and in vivo that TA protein targeting by the GET/TRC pathways involves a chaperone exchange in which the TMD, initially shielded by a hydrophobic patch within the C-terminal region of Sgt2, is handed over to Get3, where it is protected within a dedicated hydrophobic groove formed in the ATP-bound state. While Sgt2 has long been considered the upstream component of the GET pathway, it has recently emerged that the abundant, Hsp70-like chaperone Ssa1 can also act as a highly efficient nascent TA protein chaperone and that its effectiveness in TA protein trapping is enhanced by the J domain–containing cochaperone protein Ydj1 (Fig. 1; Cho et al., 2021; Cho and Shan, 2018). Ssa1 and numerous other chaperones physically interact with Sgt2 via its tetra-tricopeptide repeat domain (Cho and Shan, 2018; Krysztowskiska et al., 2017). As this domain of Sgt2 is important for ER targeting of TA proteins in vivo, this supports a potential role of other chaperones alongside Sgt2. Transfer of TA protein cargoes from Ssa1 to Sgt2 is energetically favorable and stimulated by the J domain proteins Ydj1 and Sis1. Interestingly, Ydj1 and Sis1 appear to function redundantly in targeting of a reporter TA protein to the ER (Cho et al., 2021), perhaps suggesting that the chaperone cascade protecting TA proteins during their biogenesis can be more extensive than initially anticipated. However, it still remains to be determined how much of a contribution these proteins make to endogenous TA protein targeting within the native cellular environment. Interestingly, human cells lacking SGTA or BAG6 are viable, and TA protein targeting to the ER can still be accomplished in the absence of these factors (Culver and Mariappan, 2021). It is possible, therefore, for capture by the pretargeting complex to be bypassed, and perhaps in this context, other chaperones, analogous to those characterized in yeast, contribute to the initial protection of nascent TA proteins before their association with TRC40. Along this line, TA proteins with less hydrophobic TMDs, which use the EMC rather than the TRC pathway, have been shown to be chaperoned through the cytosol by calmodulin (Fig. 2; Guna et al., 2018). The existence of alternative, partially redundant, targeting pathways, supported by the nonlethality of yeast, plant, and human GET pretargeting factor knockouts (Stefanovic and Hegde, 2007; Schuldiner et al., 2008; Jonikas et al., 2009; Srivastava et al., 2017; Xing et al., 2017; Culver and Mariappan, 2021), likely helps ensure high-fidelity and robust targeting of TA proteins in vivo.

Selectivity of TA protein capture
It has become increasingly clear that nascent TA proteins emerge into a crowded environment where encounters with different machineries can direct them toward different fates, such as targeting to the ER via different routes, targeting to different organelles, or potentially, degradation. These observations suggest a finely tuned process of nascent TA protein capture to direct different proteins to the appropriate destinations and highlight the question of how TA proteins are selectively captured. While all TA proteins share the common characteristic of a single TMD close to the C-terminus that can serve as a targeting signal, the length and hydrophobicity of this TMD, as well as the net charge of the downstream sequence (also termed C-terminal element [CTE]), vary significantly, and these features are important determinants of the ultimate destination of the protein (Bellharz et al., 2003; Borgese et al., 2019). TA proteins of the outer mitochondrial membrane (OMM) and peroxisomes are typified by short, less hydrophobic TMDs, and positively charged CTEs are characteristic features of peroxisomal TA proteins (Horie et al., 2002). In contrast, ER and Golgi TA proteins generally have relatively long, hydrophobic TMDs, and the charge of their CTEs varies considerably (Rao et al., 2016; Borgese et al., 2019). The physicochemical properties of ER TMDs favor capture by Sgt2/SGTA and binding by Get3/TRC40, whereas TA proteins with less hydrophobic TMDs are poor substrates (Coy-Vergara et al., 2019; Guna et al., 2018). This implies that an important layer of capture selectivity is already encoded within the proteins themselves. Strict categorization of different organelar TA proteins based on physicochemical properties is not possible, however, as differently localized TA proteins have partially overlapping properties, indicating that this criterion alone is insufficient to ensure correct targeting.

TA proteins not only need to be directed to different target membranes, but they also need to be recognized as bona fide membrane proteins. The hydrophobic sequences that are integral features of membrane proteins must be distinguished from exposed hydrophobic patches of misfolded, nonmembrane proteins that serve as signals for recruitment of the protein quality control machinery. Interestingly, the BAG6 component of the mammalian pretargeting complex sits at the nexus between the alternative fates of targeting and degradation; a minimal C-terminal BAG domain in BAG6 scaffolds interactions between TRC35 and UBL4A and is sufficient for TA protein targeting to the ER (Mock et al., 2015), while the N-terminal UBL domain of the protein promotes recruitment of the ubiquitination machinery to mediate quality control of aberrant proteins (Fig. 2; Rodrigo-Brenni et al., 2014). In this context, BAG6 has been implicated in rerouting SGTA-bound TA proteins that are not efficiently relayed to TRC40 toward the degradation pathway (Shao et al., 2017). SGTA has emerged as another central player in determining the fate of TA proteins, as it is able to antagonize BAG6-facilitated protein ubiquitination. Interestingly, SGTA not only reduces the likelihood of protein ubiquitination by shielding the hydrophobic TMD but also promotes active deubiquitination of BAG6 complex–associated proteins (Leznicki and High, 2012). In this way, SGTA could contribute to the recovery of nascent TA proteins aberrantly marked for degradation by BAG6–associated ubiquitin ligases. Notably, the interplay between the BAG complex and SGTA in determining protein fate extends beyond TA proteins to other membrane proteins targeted cotranslationally (Leznicki and High, 2020). Intriguingly, it was recently shown that ubiquitination of TA proteins can occur independently of...
BAG6, and that ubiquitinated TA proteins can still be handled by TRC40 and directed to the ER, where they are fully deubiquitinated by USP20/USP33 before or after membrane insertion (Fig. 2; Culver and Mariappan, 2021). It remains unclear mechanistically how exactly ubiquitinated TA proteins evade proteasome-mediated degradation in the cytosol, but it is possible that either the nature of the ubiquitination and/or rapid capture by TRC40 enables them to efficiently reach their destination and be deubiquitinated. It will be interesting to determine if this ubiquitination-deubiquitination cycle simply represents a futile mislabeling and recovery process or whether it fulfills a specific function during TA protein biogenesis.

Delivery of TA proteins into the ER
Both the GET/TRC receptor and the EMC complex have emerged as gateways into the ER for TA proteins (Guna et al., 2018; Schuldiner et al., 2008; Vilardi et al., 2011; Yamamoto and Sakisaka, 2012). TA proteins destined for ER insertion via the GET/TRC receptor converge on the homodimeric cytosolic chaperone Get3/TRC40, which escorts them to the ER-bound GET/TRC insertase (McDowell et al., 2020; Wang et al., 2014). Docking of ADP and TA protein–bound Get3/TRC40 onto the GET receptor allows transfer of the TA protein to the receptor, which subsequently mediates their insertion into the membrane (Wang et al., 2014). Then, upon ADP release, Get3/TRC40 is recycled for another round of ATP binding and TA protein targeting (reviewed in detail in Chio et al., 2017).

Evolutionary conservation of the GET receptor complex
The Get1 component of the GET receptor is a member of the Oxa1 superfamily of insertase proteins, which includes bacterial VldC and eukaryotic EMC3 (Anghel et al., 2017; McDowell et al., 2021), and it shares its three-TMD topology with other members of this family. Get1 sequence conservation among different phyla readily facilitated the identification of homologues in mammals (WRB) and plants (AtGet1; Srivastava et al., 2017; Xing et al., 2017). In contrast, Get2 homologues are more divergent, and sequence conservation is limited to the functionally essential N-terminal Get3-binding sequence and, to a lesser extent, the three TMDs (Borgese, 2020). CAML has nevertheless been identified as a functional and structural homologue of Get2 in mammals (Yamamoto and Sakisaka, 2012), and demonstrated to complement phenotypes associated with loss of Get1/Get2 when coexpressed with WRB in budding yeast (Vilardi et al., 2014). The existence of Get2 orthologues or functional homologues in other phyla were uncertain for a long time. The recent identification of the archaeaplasmid Get2 homologue AtGet2 (Asseck et al., 2021) and in silico prediction of Get2 homologues in mollusks and arthropods (Borgese, 2020), however, now strongly support that not only Get1/WRB, but the GET receptor complex as a whole is conserved among eukaryotes.

Architecture and stoichiometry of the GET receptor
The interaction between Get1/WRB and Get2/CAML is mediated by their TMDs, which are also necessary for the insertase function of the complex (Vilardi et al., 2014; Wang et al., 2014). Moreover, mammalian WRB and CAML, and likely other homologues as well, are thought to exist as an obligate complex, mutually stabilizing each other. Indeed, the expression levels of WRB or CAML decrease in the absence of the other subunit, likely because of destabilization of the remaining subunit (Colombo et al., 2016; Rivera-Monroy et al., 2016). However, the effects of the individual components seem to be asymmetric on each other. Namely, WRB can insert into the ER membrane correctly in the absence of CAML and is later degraded as an orphan subunit, whereas WRB is required for CAML to assume a correct topology in the first place (Carvalho et al., 2019; Inglis et al., 2020). More specifically, the second TMD of CAML remains exposed to the ER lumen in the absence of WRB, where it acts as a degron, triggering protein degradation. Interestingly, its topology can be corrected posttranslationally, and its degradation prevented when WRB is available in the membrane (Inglis et al., 2020). It remains to be seen whether a similar inter-subunit interplay also occurs in other species; however, results from A. thaliana imply that correct assembly of the GET receptor may be differently controlled in plants, as ectopically expressed AtGet2 remains stable in the absence of AtGet1 (Asseck et al., 2021).

Although formation of a Get1/Get2 heterodimer is recognized as a prerequisite for a minimal functional receptor complex, a higher-order stoichiometry of the insertion-competent GET receptor complex has long been actively discussed. Due to the symmetric, homodimeric nature of Get3 in the TA protein-loaded complex, two analogous binding sites for both Get1 and Get2 are offered (Stefer et al., 2011). Despite a partial overlap of the Get1 and Get2 interaction sites on Get3, simultaneous binding of Get3 by Get1 and Get2 has been observed in crystal structures of Get3, together with the cytosolic domains of Get1 and Get2 (Stefer et al., 2011). This gave rise to the notion of a heterotetrameric structure of the GET receptor composed of two Get1 and two Get2 subunits, which could bind a single Get3 dimer with high affinity (Mariappan et al., 2011; Stefer et al., 2011). However, results obtained with in vitro reconstituted proteoliposomes demonstrated that a single dimer of Get1/Get2 can be sufficient for insertion of TA substrates into the membrane (Zalisko et al., 2017). Nevertheless, the heterotetrameric arrangement is supported by recent high-resolution cryo-EM structures of Get3/TRC40 homodimers docked onto the yeast and mammalian GET receptors, which reveal the formation of a heterotetrameric receptor complex upon Get3/TRC40 binding (Figs. 1, 2, and 3; McDowell et al., 2020). These new structures further consolidate the previously proposed model (Mariappan et al., 2011; Stefer et al., 2011) that Get3/TRC40 is initially captured by the extended cytosolic domains of Get2/CAML before contacting Get1/WRB (Figs. 1, 2, and 3). This arrangement, coupled with the finding that Get1-Get2 can simultaneously bind two Get3 molecules (McDowell et al., 2020), opens the possibility for a relay system where translocation of a first Get3/TRC40-ATA protein complex to Get1 immediately allows capture of a second substrate complex by Get2, potentially increasing the efficiency of receptor complex loading and minimizing the risk of mistargeting. Interestingly, the GET/TRC receptor seems to be stabilized at the heterotetramer interface by not only protein–protein but also protein–lipid interactions, indicating that the lipid environment of the ER membrane may also influence the
olegic state of the receptor. Importantly, complementation experiments in yeast demonstrate that disrupting lipid binding and thus the formation of the heterotetramer leads to in vivo loss of function of the receptor manifesting in TA protein mislocalization (McDowell et al., 2020). Therefore, although a Get1/Get2 dimer appears to be sufficient for the insertase function of the receptor in vitro, it is highly likely that a tetrameric complex is required to ensure efficient and accurate TA protein targeting within the cellular environment.

**Mechanistic view of TA protein insertion into the lipid bilayer**

Recent structural advances provide exciting mechanistic insights into the details of TA insertion, both by the GET/TRC receptor (McDowell et al., 2020) and the EMC complex (Bai et al., 2020; Miller-Vedam et al., 2020; O’Donnell et al., 2020; Pleiner et al., 2020). In the case of the GET/TRC receptor, within the membrane, the TMDs of WRB, together with TMD3 of CAML are arranged such that a hydrophilic groove, sealed at the luminal face but accessible from the cytosol, is assembled (Fig. 3). This likely serves as a substrate entry point with the charged, extreme C-terminus of the TA protein drawn in by interactions with the numerous hydrophilic residues of the receptor channel, thus bringing the TMD in close proximity to the destabilized bilayer, allowing insertion. This corroborates previous results obtained by cross-linking nascent TA substrates to the receptor in vitro, which pinpointed the region around the hydrophilic groove as the entry point for TA proteins into the membrane (Wang et al., 2014). Interestingly, assembly of the GET receptor as a heterotetrameric complex means that tandem hydrophilic grooves generated by each Get1/WRB-Get2/CAML pair represent two alternative routes into the membrane. The asymmetric binding of the TA protein within the Get3 dimer (Mateja et al., 2015), means that depending on the orientation of the docking, insertion via a particular channel will be favored. Dynamic modeling of interactions between the receptor and the TRC40 dimer in different conformations/nucleotide-bound states (Mateja et al., 2015; Stefer et al., 2011) suggest that transition of Get3/TRC40 to the open conformation leads to rearrangement of complex such that the C-terminus of the released TA protein is oriented toward the hydrophilic groove (McDowell et al., 2020). Complementary structural analyses of the EMC complexes (Bai et al., 2020; Miller-Vedam et al., 2020; O’Donnell et al., 2020; Pleiner et al., 2020) reveal analogous hydrophilic groove features, indicating that a common insertion mechanism is used by evolutionarily distant membrane receptors to accomplish insertion of a diverse set of membrane proteins (Bai and Li, 2021; McDowell et al., 2021).

For the GET/TRC receptor, it is not yet clear how exactly the TA protein transits from this hydrophilic groove to become fully immersed in the membrane, but an amphipathic helix of Get1/WRB that lies close to the membrane has been suggested to cause membrane distortions that could facilitate TA protein integration.

Another key feature of the GET receptor revealed by the new structures is a short helix, α9, present in the cytosolic domains of both yeast Get2 and human CAML, which binds the TA-binding domain of Get3/TRC40 and serves a gating function for the hydrophilic groove (Fig. 3; McDowell et al., 2020). Contact between the Get2/CAML helix and the Get3/TRC40 TA-binding domain induce conformational rearrangements crucial to the release and insertion of TA substrates in vivo, further underlining the role the GET receptor plays in stimulating substrate release from Get3. It remains to be determined whether, mechanistically, this helix facilitates TA protein insertion by preventing backsliding of the TMD out of the hydrophilic groove or actively driving the TA protein into the channel.

**Quality control and rescue of TA protein targeting**

The multiple possible destinations for TA proteins, together with the broad spectrum of physical properties of secretory pathway TA proteins, the existence of alternative ER targeting and insertion strategies, and the complexity of ensuring optimal capture, renders targeting of TA proteins to the ER inherently prone to errors. Defects in this process can manifest as either redirection of ER TA proteins to other membranes or the spurious ER insertion of non-ER TA proteins. Given the importance of TA protein functions and the toxic effects of cytosolic protein aggregates, either of these scenarios can have seriously detrimental effects on cells, necessitating robust surveillance, recovery, and degradation pathways (Jiang, 2021).

**Removal of TA proteins misdirected to the OMM**

In contrast to ER TA proteins, some TA proteins of the OMM have been proposed to be inserted directly, potentially due to the specific lipid composition of the OMM (Figueiredo Costa et al., 2018). The hydrophobicity of their TMDs are lower than that of secretory pathway TA proteins (Borgese et al., 2001; Beilharz...
et al., 2003), which, considering that high TMD hydrophobicity is necessary for recognition by the GET pathway (Guna et al., 2018), helps explain how they avoid capture and subsequent delivery to the ER by Get3. In contrast, peroxisomal TA proteins can either use the Pex19-Pex3 machinery to directly reach peroxisomes (reviewed in Mayerhofer, 2016) or be first targeted to the ER by the GET pathway (Schuldiner et al., 2008) before reaching peroxisomes (Fig. 4).

It has been observed that ER-inserted TA proteins can mislocalize to mitochondria when the functionality of the GET pathway is impaired (Schuldiner et al., 2008). A salient example is yeast Pex15, which is a peroxisome-destined TA protein first inserted into the ER but that mislocalizes to mitochondria not only when the C-terminal 30 amino acids or Pex19 are lacking, but also in the absence of Get3 (Schuldiner et al., 2008; Okreglak and Walter, 2014; Li et al., 2019). Furthermore, a basal level of mistargeting of secretory pathway TA proteins, such as yeast Gos1, to the OMM is observed even in the presence of a functional GET/TRC pathway, implying that some level of mistargeting is unavoidable (Chen et al., 2014). This therefore raises the question of how mistargeted TA proteins are recognized and cleared while the appropriately localized proteins are retained.

A mechanism by which TA proteins incorrectly inserted into the OMM can be extracted has recently been described; the highly evolutionarily conserved mitochondrial and peroxisomal AAA-ATPase Msp1 (ATAD1 in mammals) has been shown to be essential for the removal of mitochondrially mislocalized Pex15 and is thought to act as a general dislocase of TA proteins mislocalized to the mitochondria (Fig. 4; Chen et al., 2014; Okreglak and Walter, 2014). This explains why double-mutant yeast strains lacking both GET pathway components and Msp1 mislocalize TA proteins noticeably to mitochondria (Li et al., 2019). Accordingly, GET pathway components and Msp1 show a synthetic negative genetic effect (Chen et al., 2014; Okreglak and Walter, 2014), emphasizing the importance of high-fidelity protein targeting and quality control. Msp1 forms hexamers in the OMM, and in vitro, its ATPase activity is sufficient to drive the removal of TA proteins from proteoliposomes (Wohlever et al., 2017), demonstrating that Msp1 alone can both recognize its substrates and drive their subsequent extraction from the membrane. Recent visualizations of Msp1-substrate complexes provide insights into the mechanism of membrane extraction; the TA protein substrate enters via a single, hydrophobic site and is then anchored within the hydrophobic pore by a network of aromatic amino acids (Wang et al., 2020). ATP hydrolysis, the driving force for membrane extraction, is coordinated with subunit positioning within the complex via specific elements at the subunit interfaces (Wang et al., 2020). Interestingly, based on in vitro experiments with MBP-tagged Pex3, it has been suggested that the unfoldase activity of Msp1 may be regulated by Pex3 on peroxisomes (Castanzo et al., 2020), but it remains to be determined whether a similar regulatory mechanism exists in the OMM as well.

The stringency of recognition of mitochondrially mislocalized TA proteins relies on at least a twofold recognition mechanism by Msp1. First, basic residues typical of the luminal tails of peroxisomal TAs are recognized when exposed in the mitochondrial intermembrane space (Li et al., 2019). Second, exposed hydrophobic amino acids close to the membrane, present in various secretory pathway and peroxisomal TA proteins, also act as a recognition signal for Msp1 (Li et al., 2019). Furthermore, there is evidence indicating that orphan TA proteins lacking binding partners within the OMM are more readily recognized and displaced by Msp1 (Weir et al., 2017;
This indicates that besides the biophysical properties of Msp1 substrates, their ability to form functional protein–protein interactions with other membrane proteins is an important discriminating factor that determines whether a TA protein is retained in or removed from the OMM. Mechanistically, lack of a binding partner may render mislocalized TA proteins less stably anchored within the membrane, or, when separated from their normal interaction partners, mistargeted proteins may be more likely to expose hydrophobic patches to the cytosol, both of which would increase the chance of expulsion by Msp1.

It is possible that after removal from the OMM by Msp1, secretory pathway TA proteins returned to the cytosol could be retargeted if they are captured by Get3/TRC40 or other chaperones capable of directing them to the ER-bound insertase machineries. However, a cellular machinery also needs to exist that degrades excess TA proteins in the cytosol or at the ER to ensure protein homeostasis. Indeed, the E3 ubiquitin ligase Doa10, an important player in ER-associated degradation, has recently emerged as a quality control factor responsible for sensing and ubiquitination of spurious and excess TA proteins ejected from the OMM. It has been suggested that Doa10-mediated ubiquitination may take place either in the cytosol (Dederer et al., 2019) or after retargeting to the ER, where their extraction and degradation is facilitated by the AAA-ATPase Cdc48 (Matsumoto et al., 2019; Fig. 4).

**Mistargeting of mitochondrial TA proteins to the ER**

It is not only the case that nonmitochondrial TA protein are misdirected to the OMM; also, reciprocal events can occur as OMM TMD-containing proteins are observed in the ER, especially when mitochondrial targeting signals are masked, when the mitochondrial import machineries are overloaded, or upon mitochondrial dysfunction (Hansen et al., 2018; Xiao et al., 2021; Vitali et al., 2018). More specifically, the GET pathway itself has been shown to contribute to the ER mislocalization of OMM TA proteins (Vitali et al., 2018; Xiao et al., 2021). While the high efficiency of mitochondrial targeting appears to keep such mistargeting to a minimum, perturbation of the equilibrium between OMM targeting and ER mistargeting by the GET pathway can lead to aberrant accumulation of clients in the wrong membrane. Similar to nonmitochondrial TA proteins ejected from the OMM by Msp1, a parallel ATP-dependent mechanism for displacing TA proteins incorrectly introduced into the ER has recently been discovered (Fig. 4; McKenna et al., 2020; Qin et al., 2020). Structural and biochemical analyses of the ER-bound P5A-type ATPase Spf1 (yeast)/ATP13A1 (human)/CATP-8 (*Caenorhabditis elegans*) have identified it as a major quality control factor in the ER. Spf1 contains a substrate-binding pocket, laterally accessible from the membrane, that has been proposed to flip ER-associated proteins, promoting their release back into the cytosol or their topological rearrangement. Precisely how specificity for mislocalized/missinerted proteins is achieved remains unclear, but it is suggested that the lower hydrophobicity of non-ER TMDs may favor their dislocation. Furthermore, in cells lacking Spf1, the ergosterol content of the ER is altered to more closely resemble the OMM, implying that membrane composition may contribute to TA protein distribution and that Spf1 could also influence TA protein mislocalization by regulating the lipid composition of membranes (Krumpe et al., 2012). In yet another analogy to Msp1-mediated rejection of mislocalized TA proteins from the OMM, it is likely that mitochondrial proteins extracted from the ER may then be successfully retargeted to their desired destination. Indeed, the recently described ER-surface-mediated protein targeting pathway sets a precedent for such a route (Hansen et al., 2018).

**Concluding remarks**

The initial momentum of the TA protein-targeting field following the discovery of the GET/TRC pathway has not abated. Recent years have seen not only a deepening mechanistic understanding of the intricacies of this pathway but also growing knowledge on its interplay with other targeting pathways and the safety nets present to ensure the fidelity of TA protein targeting. Building on the early structural analyses of individual proteins and protein domain complexes, recent advances in cryo-EM have now enabled visualization of larger complexes, such as the targeting factor-bound membrane insertase, allowing the route of TA proteins from the cytosol into the ER membrane to be anticipated. Alongside high-throughput microscopy-based screens, further analyses of TA protein targeting in cells have uncovered alternative routes that TA protein can take to reach the ER and have highlighted functional redundancies. The existence of different pathways for TA protein targeting and insertion into the ER, on the one hand, suggest a robust system with backup strategies in place in case a client protein escapes its normal targeting route, but on the other hand, emphasize the complexity of the capture process and selection of the optimal targeting pathway. In turn, these additional layers of complexity underscore the need for quality control pathways. The discovery of active removal of mislocalized TA proteins not only provides a mechanism for such quality control but perhaps also suggests a dynamic aspect to TA protein targeting wherein TA proteins within membranes can be removed and then either retargeted to their correct destination or reinserted if appropriate. Altogether, a picture emerges of an array of capture, insertion, and ejection machineries that are finely balanced in terms of substrate preferences to optimize TA protein localization within the cell.

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