Structures of the human mitochondrial ribosome recycling complexes reveal distinct mechanisms of recycling and antibiotic resistance

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Abstract

Ribosomes are recycled for a new round of translation initiation by dissociation of ribosomal subunits, messenger RNA and transfer RNA from their translational post-termination complex. Mitochondrial ribosome recycling factor (RRF<sub>mt</sub>) and a recycling-specific homolog of elongation factor G (EF-G2<sub>mt</sub>) are two proteins with mitochondria-specific additional sequences that catalyze the recycling step in human mitochondria. We have determined high-resolution cryo-EM structures of the human 55S mitochondrial ribosome (mitoribosome) in complex with RRF<sub>mt</sub>, and the mitoribosomal large 39S subunit in complex with both RRF<sub>mt</sub> and EF-G2<sub>mt</sub>. In addition, we have captured the structure of a short-lived intermediate state of the 55S•RRF<sub>mt</sub>•EF-G2<sub>mt</sub> complex. These structures clarify the role of a mitochondria-specific segment of RRF<sub>mt</sub> in mitoribosome recycling, identify the structural distinctions between the two isoforms of EF-G<sub>mt</sub> that confer their functional specificity, capture recycling-specific conformational changes in the L7/L12 stalk-base region, and suggest a distinct mechanistic sequence of events in mitoribosome recycling. Furthermore, biochemical and structural assessments of the sensitivity of EF-G2<sub>mt</sub> to the antibiotic fusidic acid reveals that the molecular mechanism of antibiotic resistance for EF-G2<sub>mt</sub> is markedly different from that exhibited by mitochondrial elongation factor EF-G1<sub>mt</sub>, suggesting that these two homologous mitochondrial proteins have evolved diversely to negate the effect of a bacterial antibiotics.
Introduction

The process of protein synthesis in all living cells is orchestrated by highly complex macromolecular assemblies called ribosomes, in coordination with mRNA, tRNAs and multiple translational factors. Mitochondrial ribosomes (mitoribosomes) and their associated translation machinery are distinct from those in the cytoplasm and display features reminiscent of prokaryotic translation (Pel and Grivell, 1994), in line with the assumption that mitochondria have evolved from endocytosis of an α-proteobacterium by an ancestral eukaryotic cell (Gray et al., 1999). However, cryo-electron microscopy (cryo-EM) structures have revealed that the mammalian mitoribosomes have diverged considerably from their bacterial counterparts and acquired several unique features (Amunts et al., 2015; Brown et al., 2014; Desai et al., 2017; Greber et al., 2015; Kaushal et al., 2014; Koripella et al., 2020; Koripella et al., 2019b; Sharma et al., 2003). A striking difference is the reversal in the protein to RNA ratio, as the bacterial ribosomes are high in ribosomal RNA (rRNA) whereas the mammalian mitoribosomes are high in protein. The increase in protein mass is the result of acquisition of multiple mito-specific ribosomal proteins (MRPs) and addition of extensions to many MRPs that are homologous to bacterial ribosomal proteins. Though the steps of mitochondrial translation closely resemble those for prokaryotic translation in the general sequence of events and the homologous accessory protein factors involved, they also show significant structural and functional differences (Christian and Spremulli, 2012; Sharma et al., 2013).

The complex process of protein synthesis is accomplished in four essential steps of initiation, elongation, termination and the ribosome recycling. Transitioning from translation termination to ribosome recycling has been best characterized in eubacteria. During translation termination, the nascent polypeptide chain attached to the peptidyl tRNA is released from the ribosome with the help of a class I release factor (RF) that interacts with the stop codon exposed at the ribosomal decoding site, or aminoacyl-tRNA binding site (A site) (Ito et al., 1996; Kisselev et al., 2003; Poole and Tate, 2000). Subsequently, the class I RF is dissociated from the ribosome with the help of a class II RF in a GTP hydrolysis-dependent manner (Freistroffer et al., 1997; Gao et al., 2007; Zavialov et al., 2002). At the end of the termination, the translated mRNA and the deacylated tRNA remain associated with the ribosome (Kaji et al., 2001; Karimi et al., 1999), a state referred to as the post-termination complex (PoTC). In order to initiate a new
round of protein synthesis, the ribosome must be split into its two subunits and its bound ligands must be removed. In eubacteria, the disassembly of the PoTC requires the concerted action of two protein factors, the ribosome recycling factor (RRF) and the elongation factor G (EF-G) (Ito et al., 2002; Janosi et al., 1998; Kaji et al., 2001; Rao and Varshney, 2001; Zavialov et al., 2005). RRF binds to the PoTC as the 70S ribosome adopts a ratcheted conformation (Barat et al., 2007; Gao et al., 2005), in which the small (30S) subunit of the ribosome rotates in an anticlockwise direction with respect to the large (50S) subunit (Frank and Agrawal, 2000). This is followed by the binding of EF-G in conjugation with guanosine 5’-triphosphate (GTP) to the RRF-bound PoTC and the dissociation of the 70S ribosome into its large and small subunits, a process that requires the hydrolysis of GTP on EF-G (Borg et al., 2016; Hirokawa et al., 2005; Peske et al., 2005; Zavialov et al., 2005). Though the involvement of a third factor, initiation factor 3 (IF3) in the recycling process is generally agreed upon, its precise function has been debated (Hirokawa et al., 2005; Iwakura et al., 2017; Karimi et al., 1999; Zavialov et al., 2005).

Unlike eubacteria, where a single form of EF-G participates in both the elongation and ribosome recycling steps (Borg et al., 2016; Savelsbergh et al., 2009), mammalian mitochondria utilize two isoforms of EF-G, EF-G1mt and EF-G2mt (Hammarsund et al., 2001; Tsuboi et al., 2009). While EF-G1mt specifically functions as a translocase during the polypeptide elongation step (Bhargava et al., 2004), EF-G2mt has been reported to act exclusively as a second recycling factor together with RRFmt (Tsuboi et al., 2009). Human RRFmt is about 25-30% identical to its eubacterial homologs but carries an additional 79 amino acids (aa) long extension at its N-terminus (Zhang and Spremulli, 1998). The recent high-resolution cryo-EM structures of RRFmt bound to the 55S mitoribosomes (Koripella et al., 2019b) and an in-vivo formed mitoribosomal complex (Aibara et al., 2020) revealed that the structurally conserved segment of the RRFmt is similar to its bacterial analog on (Agrawal et al., 2004; Barat et al., 2007; Dunkle et al., 2011; Fu et al., 2016; Gao et al., 2007; Gao et al., 2005; Weixlbaumer et al., 2007; Yokoyama et al., 2012; Zhou et al., 2020) and off (Kim et al., 2000; Nakano et al., 2003; Saikrishnan et al., 2005; Selmer et al., 1999; Toyoda et al., 2000; Yoshida et al., 2001) the 70S ribosome in terms of its overall size and domain composition. However, the unique mito-specific N-terminal extension (NTE) in RRFmt extends towards the GTPase-associated center and interacts with the functionally important 16S rRNA elements of the mitoribosomal 39S subunit, including the rRNA helices 89 (H89), H90 and H92 (Koripella et al., 2019b).
Valuable mechanistic inferences about the bacterial ribosome recycling process were made from the structures of the 70S•RRF (e.g., Agrawal et al., 2004) and dissociated 50S•RRF•EF-G (Gao et al., 2007; Gao et al., 2005) complexes. Capturing the simultaneous binding of both factors on the 70S ribosome is challenging, however, owing to the rapid rate of 70S ribosomes dissociation into subunits by the combined action of RRF and EF-G (Borg et al., 2016). To slow down this reaction, a heterologous system with \textit{T. thermophilus} RRF and \textit{E. coli} EF-G was used to capture both factors on the 70S ribosome by cryo-EM (Yokoyama et al., 2012). Subsequently, a time-resolved cryo-EM study was also able to capture various 70S•RRF•EF-G functional intermediates, albeit at low resolution (Fu et al., 2016). More recently, a bacterial ribosome recycling complex containing both RRF and EF-G was obtained by X-ray crystallography by stabilizing EF-G on the 70S ribosome through a fusion between EF-G and ribosomal protein bL9 (Zhou et al., 2020). All these structures conclude that binding of EF-G to the 70S•RRF complex induces rotation of RRF domain II towards the helix 44 (h44) region of the 30S subunit, destabilizing the crucial intersubunit bridges B2a and B3, and thereby facilitating the dissociation of the 70S ribosome into its two subunits.

With a molecular weight of 87 kD, human EF-G$_{2mt}$ is slightly larger than EF-G$_{1mt}$ (83 kD), as well as both the isoforms of bacterial EF-G (78 kD) and EF-G2 (73 kD). It should be noted that some bacterial species do carry two isoforms of EF-G, but the function for the second bacterial isoform remains undefined (Connell et al., 2007; Margus et al., 2007; Seshadri et al., 2009), except in case of a spirochaete (Suematsu et al., 2010). EF-G$_{2mt}$ has about 36 % aa sequence identity to EF-G$_{1mt}$ and about 30 % aa identity to both its bacterial homologues. Some mammalian mitochondrial translation steps are now better understood through determination of the cryo-EM structures of the initiation (Koripella et al., 2019a; Kummer et al., 2018; Yassin et al., 2011) and the elongation (Koripella et al., 2020; Kummer and Ban, 2020) complexes at high-resolution. Our previous study of the human mitoribosome recycling complex of the RRF$_{mt}$-bound 55S (Koripella et al., 2019b) provided useful insights into the mito-specific aspects of the recycling process, but a complete 55S mitoribosomal recycling complex comprising both RRF$_{mt}$ and EF-G$_{mt}$ remained elusive. To investigate the concerted action of RRF$_{mt}$ and EF-G$_{2mt}$ in splitting the 55S mitoribosome and to understand the detailed roles of mito-specific aa segments of RRF$_{mt}$ and EF-G$_{2mt}$ at the molecular level, we determined the key intermediate state structures of the 55S and 39S mitochondrial recycling complexes containing both RRF$_{mt}$ and EF-G$_{2mt}$. 


Results and discussion

Structure of the human mitoribosome recycling complex.
To investigate the molecular mechanism of ribosome recycling in mammalian mitochondria, we first prepared a model post-termination complex (PoTC) by incubating the human 55S mitoribosome with puromycin (Hirokawa et al., 2002; Koripella et al., 2019b). The model PoTC was briefly incubated with human RRFmt and human EF-G2mt-GMPPCP to obtain the mitoribosome recycling complex. (see Supplemental Materials and Methods). Single-particle cryo-EM analysis on this complex yielded three major classes that each represent a major functional state formed during human mitoribosome recycling, referred henceforth to as Class I, Class II and Class III (Fig. S1). Class I corresponds to the intact 55S mitoribosome that carries a strong density for RRFmt, and was refined to 3.5 Å (Fig. 1A,D, S2A). Class II, a relatively small class with only 28,929 particle images, corresponds to the 55S mitoribosome that carries both RRFmt and EF-G2mt, and was refined to 3.9 Å (Fig. S2B). In this class, the densities corresponding to the large (39S) mitoribosomal subunit, RRFmt, and EF-G2mt are well resolved, but the small (28S) mitoribosomal subunit appears to be loosely bound and present in multiple poses (Fig. 1B, S2). The most populated Class III corresponds to the dissociated 39S subunits that carry both RRFmt and EF-G2mt, and was refined to 3.15 Å (Fig. 1C,E, S2E). Class II likely represents an ensemble low-population intermediate states of mitoribosome recycling that occur between the states represented by Classes I and III. In addition to these three recycling complexes, we have also obtained a class of particles consisting of 55S mitoribosomes without either of the two factors where the 28S subunit was rotated by about 8° around its long axis such that its shoulder side moves closer to the 39S subunit while its platform side moves away from it (Fig. S3A). A similar orientation for the small subunit relative to the large subunit, termed as “subunit rolling” has been reported earlier for the 80S ribosomes (Budkevich et al., 2014) and the 55S mitoribosomes (Amunts et al., 2015; Koripella et al., 2019b).

RRFmt binding stabilizes the rotated state of the 28S subunit in the 55S mitoribosome.
Superimposition of our Class I complex with the RRFmt-unbound human (Amunts et al., 2015), bovine (Sharma et al., 2003) and porcine (Greber et al., 2015) 55S mitoribosomes showed that the small 28S subunit was rotated counter-clockwise by about 8.5° with respect to the large 39S subunit (Fig. S3B), similar to the “ratchet-like inter-subunit rotation” observed in the bacterial...
70S ribosome (Frank and Agrawal, 2000, 2001) and the 55S mitoribosomes complexed with translational factors (Koripella et al., 2020; Koripella et al., 2019b; Kummer and Ban, 2020; Kummer et al., 2018). In addition to the inter-subunit rotation, the head domain of the 28S subunit is rotated by about 4° towards the tRNA exit (E) site in a direction roughly orthogonal to the inter-subunit motion (Fig. S3B), similar to “head swiveling” in the bacterial 70S ribosomes (Ratje et al., 2010; Schuwirth et al., 2005). As expected, the structure of the Class I complex matches the previously published 3.9 Å resolution map of the analogous 55S•RRF\textsubscript{mt} complex (Koripella et al., 2019b).

The Class I map showed the characteristic “L” shaped RRF\textsubscript{mt} density and a density corresponding to a pe/E-state tRNA within the inter-mitoribosomal subunit space. The overall positioning and domain arrangement of RRF\textsubscript{mt} in the Class I map is similar to the bacterial RRF on (Agrawal et al., 2004; Barat et al., 2007; Dunkle et al., 2011; Fu et al., 2016; Gao et al., 2007; Gao et al., 2005; Weixelbaumer et al., 2007; Yokoyama et al., 2012; Zhou et al., 2020) and off (Kim et al., 2000; Nakano et al., 2003; Saikrishnan et al., 2005; Selmer et al., 1999; Toyoda et al., 2000; Yoshida et al., 2001) the 70S ribosomes, and also to the structures of RRF bound to the 70S chloroplast ribosome (Boerema et al., 2018; Sharma et al., 2007), and RRF\textsubscript{mt} bound to the human mitochondrial 55S in our previous study (Koripella et al., 2019b). As observed in bacteria, domain I is positioned close to the peptidyl transferase center (PTC) and extends towards the α-sarcin-ricin loop (SRL). A striking difference between the human RRF\textsubscript{mt} and its bacterial counterpart is the presence of a 79 aa long N-terminal extension (NTE) in RRF\textsubscript{mt}. We could model the last 14 aa residues of the NTE of RRF\textsubscript{mt} into an additional density contiguous with the α-helix1 from domain I. As discussed in our previous study (Koripella et al., 2019b), the NTE is strategically positioned in the intersubunit space between domain I and several functionally important 16S rRNA structural elements such as H89, H90, H92 (A-loop) and MRP L16 (Fig. S4) and interacts with several nucleotides (nts) and aa residues in its vicinity (Koripella et al., 2019b). Interestingly, unlike the α-helical nature inferred for the part of this segment of NTE (Koripella et al., 2019b), we find that its higher resolution density to be partially unstructured. A similar observation of a relatively unstructured NTE has been recently reported in an in-vivo state complex (Aibara et al., 2020). However, the mitoribosomal components interacting with RRF\textsubscript{mt} as described before (Koripella et al., 2019b) essentially remain unaltered.


We found a small density in a tight pocket surrounded by the outer bend of the junction between domains I and II of RRF<sub>mt</sub>, MRP uS12m and the small subunit’s 12S rRNA helix h44, and the large subunit’s 16S rRNA helices H69 and H71 (Fig. 2A). Except for our previous lower resolution map of the 55S•RRF<sub>mt</sub> complex (Koripella et al., 2019b), this additional density is not observed in any of the available 55S mitoribosomal structures, whether complexed with other translational factors (Koripella et al., 2020; Kummer and Ban, 2020; Kummer et al., 2018) or not (Amunts et al., 2015; Greber et al., 2015). Since our complex was reconstituted from purified components, this additional density should correspond to an RRF<sub>mt</sub>NTE segment that has been stabilized through interactions with multiple mitoribosomal components in its vicinity. Though bacterial and mitochondrial ribosomes exhibit significant differences in their overall shape, composition, and conformation, their internal rRNA core regions are largely conserved (Amunts et al., 2015; Greber et al., 2015; Kaushal et al., 2014; Sharma et al., 2003). Comparison of the RRF binding sites between the bacterial and mitochondrial ribosomes reveals that the H69 of 16S rRNA is slightly shorter in the mammalian mitoribosomes (Fig. 2B). This minor shortening of H69 is critical because it directly impacts the interaction of H69 with domain II of RRF<sub>mt</sub>. The small density most likely corresponding to N-terminus segment of the mito-specific NTE appears to compensate for the shortened H69 by mediating the interactions between RRF<sub>mt</sub> and H69 (Fig. 2B). Based on the available sidechain information from the small density, we generated a model that would account for the 10 aa residues (Ala2-Val11) at the N-terminus of NTE (Fig 2B). The present model is guided primarily by density for two consecutive large side chains of Phe8 and Arg9 within the first 11 aa (Fig. 2C). Since at least three more combination of two consecutive aa with large sidechains are present in the unmodelled segment of NTE, we refrain from analyzing the molecular interactions of this small segment of NTE with the rest of the complex.

It should be noted that the RRF<sub>mt</sub>-bound 55S mitoribosomes (present work and (Koripella et al., 2019b)) were never observed in the unrotated state, suggesting that the RRF<sub>mt</sub> binding locks the ribosome in a fully rotated state. This is in contrast to the bacterial 70S•RRF complexes that were found both in their rotated and unrotated conformational states (Agrawal et al., 2004; Barat et al., 2007; Dunkle et al., 2011; Fu et al., 2016; Gao et al., 2007; Gao et al., 2005; Weixlbaumer et al., 2007; Yokoyama et al., 2012; Zhou et al., 2020). The rotated conformation of the 55S mitoribosome seems to prime subunit dissociation by either destabilizing or completely breaking seven out of fifteen inter-subunit bridges in the unrotated 55S mitoribosome.
(Amunts et al., 2015; Sharma et al., 2003). The simultaneous interactions of N-terminus segment of the RRF_{mt} NTE with RRF_{mt}’s structurally conserved domain I, MRP uS12m, h44, H69 and H71 likely help prevent the back-rotation of the small 28S subunit. The rotated state of the 55S mitoribosome could serve as an ideal substrate for the subsequent binding of EF-G2_{mt} to complete subunit dissociation. In this context, it is important to note that the entire 79 aa long NTE is an integral part of the mature protein and is known to be essential for RRF_{mt} function during the mitoribosome recycling (Rorbach et al., 2008; Zhang and Spremulli, 1998).

**RRF_{mt} domain II motion helps split the 55S mitoribosome into its two subunits.**

Using fast-kinetics, it has been shown that the splitting/recycling of the 70S ribosome by the concerted action of RRF and EF-G happens in the sub-second time scale (Borg et al., 2016). Both RRF and EF-G have been captured on the 70S ribosome with time-resolved cryo-EM (Fu et al., 2016). It is more challenging without time-resolved techniques, but RRF and EF-G were also captured on the 70S ribosomes by using the factors from different species (Yokoyama et al., 2012) or by crosslinking the EF-G with one of the ribosomal proteins (Zhou et al., 2020). In the present work, collection of very large cryo-EM datasets (altogether 21,752 micrographs, (Fig. S1) enabled the isolation of a small subset of 55S particles (Class II) that contained both RRF_{mt} and EF-G2_{mt} (Fig. 1B). However, the 28S subunit density was found to be weak and present in multiple destabilized conformations relative to the 39S subunit in this complex.

Both Class II and Class III maps showed readily recognizable densities corresponding to RRF_{mt} and EF-G2_{mt}. The Class III complex had superior resolution, which enabled more accurate analysis of molecular-level interactions between the two factors and the mitoribosome, and their functional implications, while the Class II map was useful for interpreting large-scale conformational changes. In line with the bacterial 70S/50S•RRF•EFG complexes (Fu et al., 2016; Gao et al., 2007; Gao et al., 2005; Yokoyama et al., 2012; Zhou et al., 2020), the conformation of RRF_{mt} is substantially different between the Class I and Class III recycling complexes. The conformation of RRF_{mt} domain I remains unchanged among all three classes. In the Class II and III maps, domain II was rotated by about 45° towards the small subunit compared to its position in the Class I complex (Fig. 3A). This large conformational change is enabled by a highly flexible hinge regions between domain I and domain II in RRF_{mt}. Due to this rotation, the tip of domain II moved by about 40 Å towards the h44 of 12S rRNA. When the
maps of Class II and III were superimposed, this motion resulted in a major steric clash between the RRF\textsubscript{mt} domain II and the 28S subunit elements h44 and MRP uS12m (Fig. 3A). In the 55S mitoribosome, h44 is involved in the formation of two intersubunit bridges B2a and B3 by pairing with H69 and H71, respectively (Amunts et al., 2015; Sharma et al., 2003). By displacing h44, RRF\textsubscript{mt} disrupts these crucial intersubunit bridges, thereby splitting the 55S mitoribosome into its two subunits.

A well-defined 28S structure was not seen within the Class II 55S mitoribosomal complex, but several inter-subunit bridges do appear to be destabilized or broken, and the small subunit seems to be dissociating from the large subunit. To remove any possible large subunit contamination from the Class II map, extensive reference-based 3D classification was employed, but this class of 55S mitoribosomal particles with a well-resolved 39S subunit and a poorly resolved 28S subunit remained unchanged. This supports it being an ensemble of authentic, short-lived functional intermediates of mitoribosome recycling, where the small subunit is captured in multiple positions during its separation from the large subunit.

**EF-G2\textsubscript{mt} binding induces conformational changes in both RRF\textsubscript{mt} and the mitoribosome.**

The 55S•RRF\textsubscript{mt} complex (Class I) undergoes large conformational changes upon binding to EF-G2\textsubscript{mt} (Classes II and III). EF-G2\textsubscript{mt} would not be able to access its binding site on the mitoribosome without the significant movement seen in domain II of RRF\textsubscript{mt} in Class II and Class III complexes. This movement eliminates the direct spatial conflicts of EF-G2\textsubscript{mt} domains III, IV and V with the initial position of RRF\textsubscript{mt} domain II in the Class I complex (Fig. 3A,B; Fig. S5). This change could be induced either during or upon binding of EF-G2\textsubscript{mt}. The domain II of RRF\textsubscript{mt} is repositioned into a cavity created by domains III, IV and V of EF-G2\textsubscript{mt} (Fig. S6A) and an extensive network of interactions are formed between the two mitochondrial recycling factors. This is also in agreement with the location of RRF domain II reported in the bacterial 50S (Gao et al., 2007) and 70S complexes (Fu et al., 2016; Yokoyama et al., 2012). A majority of the interactions are formed between the hinge regions that connect the two domains of RRF\textsubscript{mt} and the loop regions from domain III of EF-G2\textsubscript{mt}. Several aa residues from the RRF\textsubscript{mt} hinge region (Pro183 -Thr186) interact with EF-G2\textsubscript{mt} domain III residues Glu495-Leu499 (Fig. S6B) via hydrogen bonds and hydrophobic interactions. Arg187 from the α-helix following the hinge region of RRF\textsubscript{mt} domain II has close hydrogen-bonding interactions with Tyr556 of EF-G2\textsubscript{mt}.
domain III (Fig. S6B). A second set of contacts formed between the two factors involves residues Ile109-Arg110 from the second hinge region of RRF\textsubscript{mt} and residues Ser527-Gln529 from the domain III of EF-G2\textsubscript{mt} (Fig. S6C).

Domain IV of EF-G2\textsubscript{mt} presses against domain II of RRF\textsubscript{mt} through multiple interactions. The surface residues of α-helix 1 (Asn629, Ser633 and Leu636) and α-helix 2 (Thr664, Met665, Ser667 and Ala668) from domain IV of EF-G2\textsubscript{mt} interact with residues in β-strand 3 (Ser134-Met138) and its adjoining loop region (Gln132 and Ile133) from domain II of RRF\textsubscript{mt} through a combination of electrostatic and hydrophobic interactions (Fig. S6D). Gln637 from the α-helix 1 of EF-G2\textsubscript{mt} also shares a hydrogen bond with Ser112 from the hinge region of RRF\textsubscript{mt} (Fig. S6D). Contacts are observed between the C-terminal α-helix of EF-G2\textsubscript{mt} domain IV and the α-helix 3 from the triple-helix bundle of RRF\textsubscript{mt} domain I. In bacteria, the analogous C-terminal α-helix of EF-G is often considered as part of domain V though the first atomic models of EF-G (AEvarsson et al., 1994; Czworkowski et al., 1994) grouped it with domain IV. While residues Ser776-Leu778 from EF-G2\textsubscript{mt} domain IV pair with residues Arg251 and Val255 from RRF\textsubscript{mt} domain I through hydrogen bonds (Fig. S6E), Arg775 from EF-G2\textsubscript{mt} domain IV strongly interacts with Glu259 from RRF\textsubscript{mt} domain I through a salt-bridge (Fig. S6E). Direct interactions of EF-G2\textsubscript{mt} domain III with RRF\textsubscript{mt} domain II at its hinge regions, known to confer interdomain flexibility to the bacterial factor (Nakano et al., 2003; Saikrishnan et al., 2005; Yoshida et al., 2001), likely help trigger the dissociation of 55S mitoribosomes into subunits by enabling the repositioning of RRF\textsubscript{mt} domain II. At the same time, the multiple interactions between domain IV of EF-G2\textsubscript{mt} and domain II of RRF\textsubscript{mt} appear to stabilize the RRF\textsubscript{mt} domain II in the altered position, which would push the 28S subunit away from the 39S subunit and prevent domain II from reverting back to its previous orientation, in order to maintain the 39S•RRF\textsubscript{mt}•EF-G2\textsubscript{mt} complex in a dissociated state.

In addition to aiding to the function of RRF\textsubscript{mt}, EF-G2\textsubscript{mt} plays a direct role in destabilizing the 55S mitoribosome. Superimposition of the maps of Class I and Class III complexes reveals a direct steric clash between the loop1 region of EF-G2\textsubscript{mt} domain IV and the 28S subunit component (rRNA h44) that participates in the formation of the inter-subunit bridge B2a (Fig. 3B). More importantly, the orientation of domain IV loop1 seems to be unique to EF-G2\textsubscript{mt} since the analogous region in EF-G1\textsubscript{mt} is positioned away from the intersubunit bridge B2a towards the decoding center (Koripella et al., 2020; Kummer and Ban, 2020).
EF-G2\textsubscript{mt} binding resulted in a prominent conformational change in the uL11m stalk-base region of the 39S subunit. The uL11m stalk-base region moved towards the CTD of uL12m, a component of the L10-L12 stalk, and assumed a unique conformation not reported previously (Amunts A, 2015; Greber BJ, 2015; Kummer E, 2018; Koripella RK, 2019; 2020). The 16S rRNA H43 of the uL11m stalk-base moved by 3 Å towards the CTD of uL12m, oriented parallel to the domain V of EF-G2\textsubscript{mt}, while the NTD of MRP uL11m moved about 5 Å away from the EF-G2\textsubscript{mt} domain V (Fig. 4A). This is in sharp contrast to the 55S•EF-G1\textsubscript{mt} translocation complexes (Amunts et al., 2015; Greber et al., 2015; Koripella et al., 2020; Koripella et al., 2019b; Kummer and Ban, 2020; Kummer et al., 2018), where the uL11m stalk-base region was observed to move 5 Å closer towards the domain V of EF-G1\textsubscript{mt} (Fig. 4B). The movement of uL11m away from the EF-G2\textsubscript{mt} domain V and towards the CTD of uL12m is essential for the binding of EF-G2\textsubscript{mt} in the present conformation, to avoid the steric clash between the NTD of uL11m and domain V of EF-G\textsubscript{mt} (Fig. 4A). It is also possible that the conformation of EF-G2\textsubscript{mt} observed in our 39S•EF-G2\textsubscript{mt} complex (Class III) was attained after the dissociation of 39S subunit from the 55S complex (Class II).

In addition to the unique conformation of the uL11m stalk-base region, the CTD of uL12m was also observed in a distinct conformation. In the 55S•EF-G1\textsubscript{mt} complexes (Koripella et al., 2020; Kummer and Ban, 2020), the CTD of uL12m is positioned close to EF-G1\textsubscript{mt} so that \( \alpha \)-helices 1 and 2 of uL12m CTD would have close interactions with the G’ subdomain of EF-G1\textsubscript{mt} (Fig. 4C), thereby providing stability to the otherwise flexible uL12m CTD. In contrast, the CTD of uL12m rotates by about 60° and shifts away by about 7 Å from the G’ subdomain of EF-G2\textsubscript{mt} in the 39S•EF-G2\textsubscript{mt} complex (Fig. 4D). As a result of this large rotational movement, interactions between the uL12m CTD \( \alpha \)-helix 2 and the G’ subdomain of EF-G2\textsubscript{mt} are lost, while the contacts between the \( \alpha \)-helix 1 and the G’ subdomain of EF-G2\textsubscript{mt} are maintained (Fig. 4D). Since protein uL12 is known to play a central role in the recruitment of translational factors to the bacterial ribosome (Datta et al., 2005; Diaconu et al., 2005; Helgstrand et al., 2007; Imai et al., 2020) the semi-stable conformation of uL12m observed in the 39S•EF-G2\textsubscript{mt} complex represents a late-stage conformation of uL12m prior to its detachment from the EF-G2\textsubscript{mt} as EF-G2\textsubscript{mt}’s participation in the 55S ribosome recycling process nears completion.

**Structural basis for use of EF-G2\textsubscript{mt} in mitoribosomal recycling instead of EF-G1\textsubscript{mt}**.
In most bacterial species, a single EF-G is involved in both translocation and ribosome recycling. Mammalian mitoribosomes have evolved to utilize the two isoforms EF-G1\textsubscript{mt} and EF-G2\textsubscript{mt} to cope with the significantly altered environment in mitochondria as compared to the bacterial cytoplasm (Hammarsund et al., 2001; Tsuboi et al., 2009) and perform two separate functions (Hammarsund et al., 2001; Tsuboi et al., 2009). EF-G1\textsubscript{mt} is used during translation elongation while EF-G2\textsubscript{mt} is used along with RRF\textsubscript{mt} for the recycling of the 55S mitoribosome. The overall position and domain arrangement of EF-G2\textsubscript{mt} is similar to that of EF-G1\textsubscript{mt} in the recently published human (Koripella et al., 2020) and porcine (Kummer and Ban, 2020) translocational complexes (Fig. S5). There are, however, some specific structural distinctions between the two factors that assign them specialized functional roles. The most striking of which is the presence of a C-terminal extension (CTE) in EF-G1\textsubscript{mt} domain IV (Fig. 5A) (Koripella et al., 2020).

Besides its CTE, the size of the conserved C-terminal α-helix of EF-G1\textsubscript{mt} domain IV is substantially longer (16 aa) (Fig. 5A) than the C-terminal α-helix (12 aa) of EF-G2\textsubscript{mt} domain IV (Fig. 5B).

The sturdier and longer C-terminal α-helix of EF-G1\textsubscript{mt} domain IV and its 11 aa CTE would not permit the coexistence of RRF\textsubscript{mt} on the mitoribosome due to a major steric clash between domain I of RRF\textsubscript{mt} and the C-terminal region of EF-G1\textsubscript{mt} (Fig. 5B). Even a reorientation of the C-terminal α-helix and its CTE away from the domain I of RRF\textsubscript{mt} would not resolve the problem as they would then clash with H89 of the 16S rRNA and the NTE of RRF\textsubscript{mt} that has been positioned in the inter subunit space between the domain I of RRF\textsubscript{mt} and 16S rRNA helix, H89 (Fig. 5B). Structural analysis of the bacterial EF-Gs from various species (Chen et al., 2010; Gao et al., 2009; Pulk and Cate, 2013) has revealed that the length of their C-terminal α-helices are about 12 aa long, suggesting that only EF-G2\textsubscript{mt} can function alongside RRF\textsubscript{mt} during subunit splitting. Moreover, the interaction between the C-terminal regions of EF-G2\textsubscript{mt} domain IV and RRF\textsubscript{mt} domain I is essential for stabilizing the bound RRF\textsubscript{mt}. This tight anchoring of RRF\textsubscript{mt} domain I to the mitoribosome would prevent RRF\textsubscript{mt} dissociation from the mitoribosome when its domain II undergoes substantial rotation to displace the h44 region of the 28S subunit. This agrees with the observation that deletion of the last few aa residues from the C-terminal region of bacterial RRF adversely effects its function during 70S ribosome recycling (Fujiwara et al., 2001; Fujiwara et al., 1999).
Now the question is why EF-G1\textsubscript{mt} mediates the translocation step during mitoribosomal elongation and not EF-G2\textsubscript{mt}? The most probable answer is that the CTE in the domain IV of EF-G1\textsubscript{mt} that limits its ability to participate in the mitochondrial ribosome recycling, is reported to be directly involved in the elongation step of mammalian mitochondrial protein synthesis (Koripella et al., 2020). By contacting the inner bend of the A-site tRNA, the CTE of EF-G1\textsubscript{mt} can help in translocating the CCA arm of the A-site tRNA into the P site, and by interacting with a 16S rRNA helix, H71, it prevents the P-site tRNA from reverting back into the A site (Koripella et al., 2020). This analysis is further supported by the fact that bacterial EF-Gs, which lack the CTE, are inactive on the 55S mitochondrial ribosomes while EF-G1\textsubscript{mt} is active on the 70S ribosomes (Eberly et al., 1985).

Specific sequence differences in a critical region (loop1) within the domain IV of EF-G2\textsubscript{mt} can also make it ineffective in driving translocation. During EF-G-catalyzed translocation in bacteria, the presence of two universally conserved glycine residues at the tip of domain IV (loop 1) region facilitate the insertion of domain IV into the decoding center (DC) (Gao et al., 2009). In the DC, the loop 1 of domain IV destabilizes the codon-anticodon interactions of the mRNA-tRNA duplex with the universally conserved 16S rRNA bases A1492 and A1493 in the 30S A site thereby aiding the A-site tRNA along with its associated codon to translocate into the P site (Zhou et al., 2014). The loop 1 region is conserved in EF-G1\textsubscript{mt} but is significantly altered in EF-G2\textsubscript{mt} (Fig. 5E). EF-G1\textsubscript{mt} retains both the glycine residues (Gly544 and Gly545) in its domain IV loop 1 region (Fig. 5C) whereas the second glycine is replaced by an aspartic acid in EF-G2\textsubscript{mt} (Fig. 5D) which alters the conformation of the tip of domain IV, rendering it structurally unfavorable for insertion into the groove between the P-site tRNA and the associated codon (Fig. 5D). Moreover, the Ala546 and Gly547 residues that follow the conserved glycines of loop 1 in EF-G1\textsubscript{mt} are substituted by the large polar sidechain residues lysine and arginine, respectively, in the corresponding region of EF-G2\textsubscript{mt} (Fig. 5E), thereby significantly altering the hydrophobicity of the loop 1 tip. Point mutations and deletions at the loop 1 tip region are known to have a pronounced effect on the function of EF-G during the elongation step of bacterial protein synthesis (Kolesnikov and Gudkov, 2003) but have a negligible effect on the activity of EF-G in bacterial ribosome disassembly (Zhang et al., 2015). Our study thus provides a structural rationale for the biochemical finding that mammalian mitoribosomes utilize two distinct EF-G-like factors during the translation elongation and recycling phases (Tsuboi et al., 2009).
Mammalian 55S mitoribosome recycling does not require GTP hydrolysis by EF-G2mt.

The overall G domain structure in the 39S•RRFmt•EF-G2mt complex (Class III map) is similar to the G domain in the 55S•EF-G1mt complex (Koripella et al., 2020; Kummer and Ban, 2020). The well-ordered density in our maps of highly conserved translational GTPase consensus motifs such as the P-loop, switch I and switch II regions, allowed complete modeling of these essential regions. A well-defined density corresponding to a bound GMPPCP molecule is also readily identifiable in the nucleotide binding pocket. GMPPCP is stabilized through interactions with universally conserved aa residues, such as Asp80 and Lys83 of P-loop, Thr122 of switch I and His145 of switch II (Fig. S7A). As in the 55S•EF-G1mt complex (Koripella et al., 2020), a crucial Mg^{2+} ion is positioned near the γ phosphate of GMPPCP and is coordinated by Thr84 and Thr122 from the P-loop and switch I regions, respectively (Fig. S7A). The catalytic His145 (His124 in EF-G1mt), known to play a central role in the hydrolysis of the bound nucleotide (Chen et al., 2013; Tourigny et al., 2013), is found oriented towards the γ phosphate of the bound GMPPCP (Fig. S7A), suggesting an active conformation of the factor prior to GTP hydrolysis.

The highly conserved α-sarcin-ricin stem-loop (SRL) region is known to be essential for the GTPase activity of all the translational G proteins (Chen et al., 2013; Gao et al., 2009; Tourigny et al., 2013). Base A3129 from the SRL was found to be contacting the Switch II His145 through hydrogen bonding interactions and thereby stabilizing His145 in its activated conformation poised to perform the hydrolysis reaction (Fig. S7B). While EF-G-dependent GTP hydrolysis is essential for an efficient splitting of the 70S ribosome into its subunits (Borg et al., 2016; Karimi et al., 1999; Zavialov et al., 2005), dissociation of the mammalian 55S mitoribosomes does not require EF-G2mt-dependent GTP hydrolysis, which is only needed for the release of EF-G2mt from the dissociated large subunit (Tsuboi et al., 2009). The above hypothesis is strongly supported by our observation that a significant proportion (82 %) of the EF-G2mt was found complexed to the 39S subunits as compared to the small proportion (18 %) that remained associated with the 55S mitoribosomes (see Fig. S1). Even though GTP hydrolysis by EF-G2mt is not necessary for 55S mitoribosome splitting, the presence of GTP or its non-hydrolysable analogues GDPNP/GMPPCP in the nucleotide binding pocket is essential, as the presence of GDP or the absence of any nucleotide does not split the 55S mitoribosome (Tsuboi et al., 2009).
Divergent mechanisms of fusidic acid (FA) resistance by EF-G$^{1\text{mt}}$ and EF-G$^{2\text{mt}}$.

Fusidic Acid (FA) is a fusidane class antibiotic that is used to treat bacterial skin infections along with chronic bone and joint infections. It is effective against several species of gram-positive bacteria and is clinically used to treat methicillin-resistant S. aureus (MRSA). FA prevents the release of EF-G•GDP from the 70S ribosome after GTP hydrolysis by preventing the switch II from attaining its GDP-bound conformation (Gao et al., 2009). Prior structural studies have demonstrated that FA binds in an interdomain pocket between the G domain, domain II and domain III of EF-G (Gao et al., 2009; Zhou et al., 2013). Stable binding of FA requires the switch I region to be disordered (Gao et al., 2009) because an ordered switch I would overlap with the binding site of FA. Biochemical studies have shown that a substantially higher concentration (10- to 100-fold) of FA is needed to inhibit the activity of EF-G$^{1\text{mt}}$ during mitochondrial elongation (Bhargava et al., 2004; Chung and Spremulli, 1990). Recent cryo-EM structures of EF-G$^{1\text{mt}}$ bound to the 55S mitoribosomes have presented the switch I in a well-defined conformation (Koripella et al., 2020; Kummer and Ban, 2020) (Fig. 6A), in contrast to bacterial 70S•EF-G complexes where the density for switch I has been consistently poorly resolved (Fig. 6B) (Chen et al., 2013; Gao et al., 2009; Tourigny et al., 2013; Zhou et al., 2013, 2014). It was proposed that the increased resistance observed for EF-G$^{1\text{mt}}$ towards FA resulted from a small insertion in the switch I region of EF-G$^{1\text{mt}}$ (Kummer and Ban, 2020). The two positively charged lysine residues (Lys80 and Lys82) in this insertion form salt bridges with the negatively charged phosphate backbone of the SRL from the 39S subunit (Fig. 6B), and hence were hypothesized to confer additional stability to the switch I of EF-G$^{1\text{mt}}$ (Kummer and Ban, 2020). Biochemical evidence showed EF-G$^{1\text{mt}}$ to be more resistant towards FA than its bacterial counterpart (Bhargava et al., 2004; Chung and Spremulli, 1990).

FA is known to inhibit both the translocation and the ribosome recycling steps in bacteria, though there is conflicting evidence on which step of translation is primarily targeted by FA (Borg et al., 2015; Borg et al., 2016; Savelsbergh et al., 2009). Comparison of the FA binding pocket between the bacterial EF-G, EF-G$^{1\text{mt}}$ and EF-G$^{2\text{mt}}$ revealed that key aa residues (Fig. S8A) reported to be necessary for the stable binding of FA (Gao et al., 2009; Zhou et al., 2013) are highly conserved (Fig. S8C), thereby suggesting a similar binding mechanism for FA for all the three EF-Gs. The three aa insertion in EF-G$^{1\text{mt}}$ that confers resistance to FA is not present in
EF-G2mt and the corresponding region in EF-G2mt does not contain any positively charged aa residues (Fig. 6E) that could strongly interact with the SRL. However, the cryo-EM map of EF-G2mt shows a well-resolved density for the switch I region (Fig. S8B) and enabled its modelling (Fig. 6C), indicating an alternative mechanism for the stabilization of switch I in EF-G2mt. Sequence alignment showed that the switch I region composition in EF-G2mt is significantly different as compared to EF-G1mt and the bacterial EF-G (Fig. 6E). Three new salt bridge interactions were identified within the switch I of EF-G2mt. Arg98 forms the first two salt bridges by pairing with Asp105 and Asp107 respectively, while Arg117 and Asp104 are involved in the formation of the third salt bridge (Fig. 6C). Furthermore, stronger interactions are observed between the switch I and domain II in EF-G2mt compared to EF-G1mt. Thr110 from switch I is placed in the close proximity of Asp419 of domain II with the possibility of hydrogen bond formation (Fig. 6C) while the corresponding interaction in EF-G1mt is between Asp404 and Val88, a much weaker interaction with no possibility of hydrogen bond formation. There is also potential for a tight T-stacking interaction between Phe417 and Tyr96 in EF-G2mt (Fig. 6C), while the corresponding residues in EF-G1mt being His402 and Arg72 offer no such interaction. Overall, through a combination of internal salt bridges and additional contacts with domain II, switch I gets highly stabilized in EF-G2mt. Since a stabilized switch I region occludes the binding site of FA (Fig. 6A,B), EF-G2mt is expected to exhibit strong resistance towards FA in the lines of EF-G1mt. To test this hypothesis, we measured the GTPase activity of EF-G2mt alongside EF-G1mt and E. coli EF-G in the presence of FA under multiple-turnover conditions (see Supplemental Materials and Methods). Our data shows that while 1µM FA has no effect on the GTPase activity in E. coli, significant inhibition is observed at higher concentrations of FA. In contrast, FA has almost no effect on the GTPase activity of either EF-G1mt or EF-G2mt even up to 10 mM FA. Our results are consistent with the earlier finding that EF-G1mt is highly resistant to FA compared to the bacterial EF-G (Bhargava et al., 2004; Chung and Spremulli, 1990), and also consistent with recent finding that showed EF-G2mt is not susceptible to inhibition by FA (Lee et al., 2020). Our results and analysis suggest FA resistance in EF-G2mt occurs by a structural mechanism fundamentally different from that in EF-G1mt.

In conclusion, structures of three distinct functional states formed during the process of human mitoribosome recycling are presented (Fig. 1). A previous biochemical finding that GTP hydrolysis is not required for the RRFmt•EF-G2mt-mediated splitting of the post-termination
mitoribosomal complex is corroborated. We also show that a mito-specific segment of the RRF_{mt}'s NTE compensates for the slightly reduced size of the H69 within the 16S rRNA of the mitoribosomal large subunit (Fig. 2). This is the first evidence showing a translational factor compensating for an rRNA segment lost during the evolution of the mitoribosomal translation machinery. Our structures reveal how RRF_{mt}'s domain II and EF-G2_{mt} domain IV directly help in disrupting the central inter-subunit bridge, B2a (Fig. 3), and suggest how the dynamics of the interactions among the uL11m, CTD of uL12m, and the G’domain of EF-G2_{mt} alters between elongation and recycling steps (Fig. 4). Structural analysis of domain IV of EF-G1_{mt} and EF-G2_{mt} explains their specific roles in two distinct steps of elongation and recycling, respectively (Fig. 5). Analysis of their GTPase domains complemented by GTPase assays reveal two distinct mechanisms of antibiotic fusidic acid resistance adopted by two homologous GTPases (Fig. 6). These observations help highlight the unique features of the main steps of human mitoribosomal recycling (Fig. 7). Future studies using time-resolved cryo-EM should help to resolve the short-lived intermediates that form during the transition from Class I to Class III states, which would help further characterize the functional roles of mito-specific segments of the two translational factors, RRF_{mt} and EF-G2_{mt}.

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Author Contributions: RKA conceived this study. PK purified human mitoribosomes, human RRF_{mt} and EF-G2_{mt}, and RKK prepared the human 55S P0TC•RRF_{mt}•EF-G2_{mt} complex. RKK and NKB collected cryo-EM data, and RKK and RKA performed image processing. RKK performed molecular modeling, using assistance from AD, EKA and NKB. AD, EKA and PK
performed the biochemical assays. RKK, AD, and RKA analyzed the data and wrote the manuscript. All authors read and approved the manuscript. Authors declare no competing financial interest in this work.

**Data Deposition:** The cryo-EM maps and atomic coordinates described in this study have been deposited in the Electron Microscopy and PDB Data Bank (wwPDB.org) under accession codes EMD-23096 and PDB ID: 7L08 for the RRF_{mt}-bound human 55S mitoribosome (Complex I), EMD-23114 for the RRF_{mt}•EF-G2_{mt}-bound human 55S mitoribosome (Complex II), and EMD-23121 and PDB ID: 7L20 for the RRF_{mt}•EF-G2_{mt}-bound human 39S mitoribosome (Complex III).

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Fig. 1. Cryo-EM structures of the human mitochondrial recycling complexes in three functional states. Segmented cryo-EM maps of the mitoribosomal (A) 55S•RRF\textsubscript{mt} complex (Class I), (B) 55S•RRF\textsubscript{mt}•EF-G2\textsubscript{mt}•GMPPCP complex (Class II), and (C) 39S•RRF\textsubscript{mt}•EF-G2\textsubscript{mt}•GMPPCP complex (Class III). In these three panels, the 28S subunit is shown in yellow, the 39S subunit in blue, E-tRNA in orchid, RRF\textsubscript{mt} in green and EF-G2\textsubscript{mt} in red. A lighter shade of yellow differentiates the 28S ribosomal proteins from the 12S rRNA, while a lighter shade of blue differentiates the 39S ribosomal proteins from the 16S rRNA. Landmarks of the 28S subunit: h, head; b, body. Landmarks of the 39S subunit: CP, central protuberance; Sb, stalk base; L1, MRP uL1m. (D) Cryo-EM density of RRF\textsubscript{mt} extracted from the Class I complex. (E) Cryo-EM density of RRF\textsubscript{mt} extracted from the Class III complex. (F) Cryo-EM density of EF-G2\textsubscript{mt} extracted from Class III complex. See Figs. S1 and S2, for overall and local resolutions, respectively, of densities corresponding to the mitoribosome, RRF\textsubscript{mt}, and EF-G2\textsubscript{mt} in these complexes. (G) Domain organization in RRF\textsubscript{mt} displaying domain I (light green), domain II (dark green), modelled region of NTE (pink) and unmodelled region of NTE (gray) (H) Domain organization in EF-G2\textsubscript{mt} showing G domain (orange), domain II (purple), domain III (cyan), domain IV (red) and domain V (blue).
Fig. 2. **Structures of two NTE segments of RRF\textsubscript{mt}**. (A) Density corresponding to the N-terminus segment of RRF\textsubscript{mt} NTE (pink) observed in the inter-subunit space sandwiched between the 28S SSU components, including MRP uS12m (cyan) and the 12S rRNA helix h44 (brown), the 39S LSU components, including 16S rRNA helices H69 (medium blue) and H71 (dark blue), and the outer bent of the junction between domains I and II of RRF\textsubscript{mt}. The dotted line (pink) depicts the connection between two structurally stabilized segments of the RRF\textsubscript{mt}’s NTE. (B) H69 superimposition from bacterial (gray) (Weixlbaumer et al., 2007) and human mitochondrial ribosomes (blue) reveals the shortening of H69 in the mitoribosome, also depicted in secondary structures of H69 on the lower left. Domain II of RRF\textsubscript{mt} interacts with the shorter H69 through its strategically positioned N-terminus of NTE (pink). (C) Sequence of NTE showing two modeled regions (pink). Thumbnails to the left depicts an overall orientation of the 55S mitoribosome, with semitransparent 28S (yellow) and 39S (blue) subunits, and overlaid positions of ligands. Landmarks on the thumbnail: h, head, and b, body of the 28S subunit, and CP, central protuberance; Sb, stalk base of the 39S subunit.
**Fig. 3. Direct involvement of RRF\textsubscript{mt} domain II and EF-G1\textsubscript{mt} domain IV in human mitoribosome recycling.**

(A) Comparison of the overall conformation of RRF\textsubscript{mt} between the Class I (gray) and Class III (green) complexes revealed that domain II rotates by about 45° towards the 28S subunit. Such a rotation would sterically clash with the 12S rRNA helix, h44 (brown) and MRP uS12m (cyan) and destabilize a crucial inter-subunit bridge, B2a (yellow). Inset shows the magnified view of the inter-subunit bridge B2a with RRF\textsubscript{mt} domain II residues V121-K127 (red) disrupting B2a by inserting between h44 residues (C1491 and A1492) and H69 residues (A2581 and A2582).

(B) Superimposition of the Class III complex with the Class I complex shows a direct overlap between the loop1 region (dark cyan) of EF-G2\textsubscript{mt} domain IV (red) and the intersubunit bridge B2a (yellow) formed between h44 (brown) and H69 (blue). Inset shows the magnified view of EF-G2\textsubscript{mt} domain IV residues L582-R585 (dark cyan) that will disrupt the bridge B2a by inserting between h44 residues (G1559 and U1560) and H69 residue (A2576) and pushing the 28S subunit (h44) away. Thumbnails to the left depict an overall orientation of the 55S mitoribosome, with semitransparent 28S (yellow) and 39S (blue) subunits, and overlaid position RRF\textsubscript{mt}. Landmarks on the thumbnails are same as in Fig. 2.
Fig. 4. EF-G2<sub>mt</sub> binding induces large-scale conformational changes in the uL11m stalk-base region and the CTD of uL12m. (A) In the 39S•RRF<sub>mt</sub>•EF-G2<sub>mt</sub> complex, the uL11m stalk-base region (blue) moves towards the CTD of uL12m and away from the domain V of EF-G2<sub>mt</sub>, as compared to its position in the EF-G2<sub>mt</sub>-unbound 55S mitoribosome (light gray) (Amunts et al., 2015). (B) In sharp contrast, in the presence of EF-G1<sub>mt</sub>, the uL11m stalk-base region was found to move towards the domain V of EF-G1<sub>mt</sub> in the translocation complex (dark gray) (Koripella et al., 2020). (C) In the 55S•EF-G1<sub>mt</sub> complex (Koripella et al., 2020), the CTD of uL12m (gray) is positioned in such a way that its α-helices 1 and 2 directly interact with the G' subdomain of EF-G1<sub>mt</sub> (yellow). (D) In our 39S•RRF<sub>mt</sub>•EF-G2<sub>mt</sub> complex, the CTD of uL12m (blue) is rotated by about 60° and shifted away by about 7 Å from the G' subdomain of EF-G2<sub>mt</sub>, resulting in the loss of contacts between its α-helix 2 and G' subdomain of EF-G2<sub>mt</sub> while a new set of interactions is formed between its α-helix 1 and G' subdomain of EF-G2<sub>mt</sub>. Thumbnails to the left depict an overall orientation of the 39S subunit (semitransparent blue), and overlaid positions of the ligands. Landmarks on the thumbnail: CP, central protuberance; Sb, stalk base; L1, MRP uL1m.
Fig. 5. **Structural basis for the exclusive roles of EF-G1\textsubscript{mt} in tRNA translocation and EF-G2\textsubscript{mt} in mitoribosome recycling.** (A) The presence of a substantially longer C-terminal α-helix (salmon) along with the presence of a unique CTE (blue), which is required for mitochondria tRNA translocation (Koripella et al., 2020), prevents the simultaneous binding of EF-G1\textsubscript{mt} and RRF\textsubscript{mt} (green) on the mitoribosome due to a major steric clash. Furthermore, any conformational repositioning the C-terminal α-helix of EF-G1\textsubscript{mt} will result in direct steric clash with the H89 (orange) of 16S rRNA and the NTE of RRF\textsubscript{mt} (pink). (B) Having a shorter C-terminal α-helix (red) and the absence of CTE allows the simultaneous binding of EF-G2\textsubscript{mt} and RRF\textsubscript{mt} (green) on the mitoribosome. (C) In EF-G1\textsubscript{mt} (Koripella et al., 2020), the presence of two universally conserved glycine residues (dark green) at the tip of domain IV loop1 region (salmon) facilitates its insertion between the mRNA-tRNA duplex at the decoding center (DC) during EF-G1\textsubscript{mt} -catalyzed translocation. (D) In EF-G2\textsubscript{mt}, the second glycine of domain IV loop 1 region (red) is substituted by an aspartic acid (dark green) altering its conformation and flexibility, and thereby making its insertion into the DC during translocation unfavorable. (E) The aa sequence

EF-G: DVEGKFIRQQQGQGQYCHVIPKVEPLPRG------SGFEFVNA 528
EF-G1\textsubscript{mt}: PFDPTHKQQGAGQYCKVIGLEPLPED--YTKLEFSDE 572
EF-G2\textsubscript{mt}: RATDQLRPLDKRHHTVEEARPIETSSVPVIEFEYAES 614
highlighted in green corresponds to loop 1 situated at the tip of domain IV, which is highly conserved between EF-G1_{mt} and the \textit{T. thermophilus} EF-G. The universally conserved glycine residues are shown in red while the aspartic acid substitution in EF-G2_{mt} is shown in blue.
Fig. 6. EF-G1\textsubscript{mt} and EF-G2\textsubscript{mt} follow diverse mechanisms to render resistance to the antibiotic fusidic acid (FA). (A) Stabilized Switch I region (salmon) (Koripella et al., 2020) in EF-G1\textsubscript{mt} blocks FA (dark blue) from accessing its binding site. Switch I stability is achieved by the presence of two unique lysine residues (K80 and K82) that strongly interact with the phosphates of SRL by forming salt bridges (light blue) (Kummer and Ban, 2020). (B) The absence of these lysine residues in the Switch I region (dark cyan) of bacterial EF-G disorders the Switch I (Gao et al., 2009) and hence makes it susceptible to FA binding and inhibition. (C) In EF-G2\textsubscript{mt}, the primary aa sequence of switch I (green) is highly altered enabling the formation of three salt bridges within its switch I, thereby stabilizing it. Furthermore, strong interactions are observed between the switch I and domain II (purple) in EF-G2\textsubscript{mt}. (D) EF-G1\textsubscript{mt} (salmon) and EF-G2\textsubscript{mt} (green) show strong resistance even at high concentrations of FA, while the bacterial EF-G (dark cyan) is susceptible to FA inhibition even at low concentrations. (E) The aa sequence alignment of the switch I region (highlighted in yellow) in three EF-Gs. The lysine residues

\begin{verbatim}
EF-G:  YYTGRHIKIGEVTHE---GAATMDMEQEGERGTTAAVTTCC 072
EF-G1mt: YYTGRIAKMHEVKGKDGVAGVMDSMELERQRGITIQSAAATYT 109
EF-G2mt: YYSGYTRSLGDVDD---GDTVIDPMAQGERGTTAAVTF 130
\end{verbatim}
shown in salmon are unique to mammalian EF-G1mt and confer stability to switch I by interacting with the SRL. Residues shown in green (except R117) are unique to EF-G2mt and confer stability to switch I by forming salt bridges within the switch I region.
**Fig. 7.** A sequence of main events in human mitochondrial ribosome recycling suggested by structures in this study. (A) The model post-termination mitoribosomal complex (PoTC), with mRNA and pe/E-state tRNA<sub>mt</sub> (Koripella et al., 2019). (B) Binding of RRF<sub>mt</sub> locks the mitoribosome in a partially destabilized state with rotated 28S subunit (Fig. 1A, Fig. S...). The mito-specific N-terminus of the RRF<sub>mt</sub>’s NTE occupies a unique site near the bridge B2a region (Fig. 2). (C) Subsequent binding of EF-G2<sub>mt</sub> further destabilizes the 55S complex with disruption of additional inter-subunit bridges, in a fast reaction leading to multiple short-lived intermediate states, with the 28S subunit present in multiple orientations relative to the 39S subunit (Fig. 1B). (D) 28S and 39S subunits are dissociated. While the 28S subunit still carries mRNA, the 39S subunit carries RRF<sub>mt</sub>, EF-G2<sub>mt</sub>, and surprisingly a tRNA<sub>mt</sub> in the 39S subunit’s E-site region (Fig. 1C). This is in sharp contrast to the tRNA dissociation mechanism in eubacterial ribosome recycling, where tRNA goes with the small 30S ribosomal subunit. (E) Depiction of final recycling products, steps of ligand release between panels D and E are yet to be characterized.