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Trends in Parasitology

Lexis and Grammar of Mitochondrial RNA Processing in Trypanosomes

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Abstract:	Parasitic protist <i>Trypanosoma brucei</i> sp. cause African human and animal trypanosomiasis. These hemoflagellates belong to the class Kinetoplastea, a taxon distinguished by the presence of a kinetoplast. This nucleoprotein body contains interlinked circular DNAs of two kinds. The maxicircles encode 9S and 12S rRNAs, two guide RNAs (gRNAs), and 18 proteins. The minicircles bear gRNA genes. Both maxicircle and minicircle genomes are transcribed from multiple promoters into 3' extended precursors which undergo 3'-5' exonucleolytic trimming. For translation, most pre-mRNAs further proceed through 3' adenylation, and often undergo U-insertion/deletion editing, and 3' A/U-tailing. The rRNAs and gRNAs are typically 3'

Highlights

- Mitochondrial RNA processing events in kinetoplastid protists include 5' modifications, 3'-5' degradation, internal sequence changes by U-insertion/deletion mRNA editing, and non-templated 3' extensions.
- The specificity of mRNA editing is dictated by guide RNAs while 5' modifications and 3' extensions are controlled by diverse PPR RNA binding factors.
- Antisense transcription plays a central role in delimiting 3'-5' trimming of primary transcripts.
- Macromolecular protein and ribonucleoprotein complexes and auxiliary factors involved in these processes have been identified and characterized to varying degrees. This review discusses recent developments and introduces a consensus nomenclature for mitochondrial RNA processing complexes and factors in *T. brucei*.

Outstanding Questions

- Recent understanding that mitochondrial mRNA, gRNA and rRNA genes are transcribed individually as 3' extended precursors places the onus on timing and mutual dependence of RNA synthesis and processing events. The nature of mitochondrial promoters and terminators, the composition of the transcription complex, the functionality of its components and the precise role of antisense transcripts remain to be elucidated.
- Many RESC subunits lack recognizable motifs and yet several bind RNA and most are essential for editing and cell viability. Understanding their functions would be a feast for structural biology.
- The mitochondrial ribosome likely selects fully-edited A/U-tailed mRNAs, but the mechanism of translation initiation at the 5' end is unclear.
- Mitochondrial proteomics remain a challenge due to extremely hydrophobic nature of mitochondrially-encoded proteins.
- The stage-specific patterns of mitochondrial transcription, RNA processing and translation are likely coordinated with nuclear gene expression by yet unknown mechanisms.

1 Lexis and Grammar of Mitochondrial RNA Processing in Trypanosomes

2

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46 polyadenylation; translation.

47

48 **Abstract**

49 Parasitic protist *Trypanosoma brucei* sp. cause African human and animal trypanosomiasis, a
50 spectrum of diseases affecting the population and economy in sub-Saharan Africa. These
51 dioxenous hemoflagellates belong to the class Kinetoplastea, a taxon distinguished by the
52 presence of a kinetoplast. This nucleoprotein body contains mitochondrial DNA of two kinds:
53 maxicircles and minicircles. The maxicircles (~25 per genome, each ~23 kb) encode 9S and
54 12S ribosomal RNAs, two guide RNAs (gRNAs), two ribosomal proteins and 16 subunits of
55 respiratory complexes. The minicircles (~5,000 per genome, ~1 kb each) bear gRNA and
56 gRNA-like small RNA genes. Relaxed maxicircles and minicircles are interlinked and packed
57 into a dense disc-shaped network by association with histone-like proteins. Both maxicircle
58 and minicircle genomes are transcribed by a T3/T7 phage-like RNA polymerase from
59 multiple promoters into 3' extended precursors which undergo 3'-5' exonucleolytic trimming.
60 To function in mitochondrial translation, most pre-mRNAs must further proceed through 3'
61 adenylation, and often undergo gRNA-directed uridine insertion/deletion editing, and 3' A/U-
62 tailing. The ribosomal and guide RNAs are typically 3' uridylated. Historically, the
63 fascinating phenomenon of RNA editing has attracted major research effort, and recent
64 developments provided insights into pre- and post-editing processing events and identified
65 key players in transforming primary transcripts into mature RNAs, and regulating their
66 function and turnover. Here, we introduce a unified nomenclature of mitochondrial RNA
67 processing complexes and factors in *T. brucei* and attempt to integrate known modalities of
68 kinetoplast RNA metabolism.

69 **Mitochondrial Gene Expression in Trypanosomes: A Trove of** 70 **Unconventional Biology**

71 Protist parasites of the genus *Trypanosoma* have occupied the research spotlight since 1895
72 when David Bruce identified *Trypanosoma brucei* as the causative agent of animal
73 trypanosomiasis (Nagana) [1], and later works linked these organisms to sleeping sickness in
74 humans [2]. Biomedical, economic and societal impact of parasite infections warranted in-
75 depth studies of the fascinating biology underlying *T. brucei* metabolism, development and
76 interactions with insect vector and mammalian hosts [3]. Among the most striking cellular
77 features of these parasites is the bipartite mitochondrial genome consisting of maxicircles and
78 minicircles, and aggregately referred to as the kinetoplast DNA (kDNA). In *T. brucei*,
79 maxicircles are catenated with minicircles into a single network and compacted by histone-
80 like basic proteins. The progress in elucidating kDNA maintenance and segregation has been
81 reviewed elsewhere [4-6]. Maxicircles, an equivalent of mitochondrial genomes (mtDNA) in
82 other organisms, encode 9S and 12S rRNAs, two ribosomal proteins [7], and 16 subunits of
83 respiratory complexes. Unlike most organellar genomes, kDNA lacks tRNA genes [8, 9], and
84 12 maxicircle genes are present as cryptogenes whose transcripts require RNA editing to
85 restore a protein-coding capacity [10]. The editing is mediated by hundreds of guide RNAs
86 (gRNAs) which are mostly encoded by minicircles, with only two gRNAs encoded by
87 maxicircles. The distinct, albeit interlinked, maxicircle and minicircle genomes are
88 transcribed independently, but the information converges at the post-transcriptional level
89 whereby minicircle-encoded gRNAs direct editing of maxicircle encoded pre-mRNAs. The
90 evolution of editing and whether this process confers a selective advantage to kinetoplastids
91 remain the subject of a debate [11], but the existence of alternatively edited mRNA sequences
92 and cognate gRNAs raises a possibility that editing-driven protein diversity may be
93 functionally relevant [12-15]. Historically, much attention has been focused on RNA editing

94 mechanism and composition of editing complexes [16-22] while more recently major
95 advances have been made in understanding transcription [23], primary RNA nucleolytic
96 processing [24-26], 5' [23] and 3' modifications [27-32], and ribosome biogenesis and
97 translation processes [7, 33-35]. The perceived complexity of mitochondrial gene expression
98 has been exacerbated by recent influx of new RNA processing factors and numerous names
99 often referring to the same entity. Bearing in mind that functions of only few proteins and
100 complexes are established beyond reasonable doubt, we nonetheless submit that the process
101 of discovering major players is close to completion. Here, we outline major stages in
102 kinetoplast RNA processing and build on previous attempts [36, 37] to introduce a unified
103 nomenclature for respective protein and ribonucleoprotein complexes, enzymes and factors in
104 *T. brucei*. Given that orthologs of nearly all *T. brucei* proteins listed in Table 1 are readily
105 distinguishable in related organisms, this nomenclature should be broadly applicable to other
106 *Kinetoplastea*.

107

108 **Nucleolytic Processing of Primary Transcripts**

109 In most organisms, primary polycistronic mtDNA transcripts are punctuated by tRNAs whose
110 excision by RNases P and Z defines functional RNA boundaries [38]. Although loss of
111 mtDNA-encoded tRNA genes renders such a mechanism inapplicable to *T. brucei*, it has
112 been held that mature mRNAs and rRNAs with uniformly monophosphorylated 5' and well-
113 demarcated 3' termini reflect maxicircle polycistronic precursor's partitioning by an unknown
114 endonuclease [39, 40]. To that end, a prominent transcription start site has been mapped
115 within the divergent region ~1,200 nt upstream of 12S rRNA [41] and transcription
116 proceeding through intergenic regions has been reported [42]. The uridylylated rRNA [43] and
117 adenylated mRNA [44] termini also typify distinct 3' end modification mechanisms for these

118 RNA classes. Conversely, short (30-60 nt) gRNAs maintain 5' triphosphates characteristic of
119 transcription-incorporated initiating nucleoside triphosphate and, similar to rRNAs, are 3'
120 uridylated [45]. However, the only explored candidate gRNA precursor processing
121 endonuclease KRPN1 (mRPN1) [46] is dispensable for axenically grown bloodstream stage
122 parasites [42]. Thus, it may be argued the essentiality of RNA editing [47], which requires
123 mature gRNAs [25, 48], renders KRPN1 an unlikely contributor to gRNA precursor
124 processing. The observations conducive to the endonucleolytic model have been re-examined
125 in light of mitochondrial 3' processome (MPsome) discovery [24]. These studies recognized
126 the MPsome-catalyzed 3'-5' exonucleolytic degradation as the major nucleolytic processing
127 pathway for mRNA and rRNA [29], and for gRNA [24] precursors. Composed of KRET1
128 TUTase [49], KDSS1 exonuclease [26, 50] and mitochondrial processome subunits (MPSS)
129 1-6 (Table 1), the purified MPsome displays 3'-5' RNA degradation, RNA hydrolysis-driven
130 double-stranded RNA unwinding, and 3' RNA uridylation activities. Although the
131 autonomous KDSS1 is inactive, incorporation into the MPsome converts this polypeptide into
132 a highly processive exonuclease capable of degrading structured RNAs to 5-6 nt fragments.
133 On the other hand, individual KRET1's robust UTP polymerization activity [51] is tamed
134 upon MPsome assembly to adding 1-15 Us, a pattern consistent with the U-tails observed in
135 steady-state RNA [15, 52]. Cumulatively, detection of *in vivo* uridylated precursors and
136 degradation intermediates [24, 31], stimulation of *in organello* KRET1-dependent RNA
137 decay by UTP [27], and MPsome's preference for U-tailed substrates suggest that uridylation
138 by KRET1 activates RNA degradation by KDSS1. It is unclear whether substrate tunneling
139 occurs within the same particle, but a coupling between RNA uridylation and degradation by
140 3'-5' RNase II/RNB-type exonuclease appears to be a highly conserved and phylogenetically
141 widespread mode of RNA decay [53, 54].

142 Exonucleolytic processing is often a case of regulated decay whereby mature 3'
143 termini are defined by a degradation blockade at a specific sequence or structure. In the *T.*
144 *brucei* mitochondrion, antisense transcripts cause MPsome pausing at 10-12 nt before double-
145 stranded region at which point MPsome-embedded KRET1 likely adds a U-tail causing
146 disengagement from the RNA [23, 24]. It follows that the precise transcription initiation site
147 on the antisense strand defines the position of mature 3' terminus of the sense transcript. The
148 antisense model of gRNA 3' end definition is consistent with bi-directional transcription from
149 converging promoters otherwise recognized as imperfect 18-nt inverted repeats [55] that
150 flank almost all gRNA genes in *T. brucei* minicircles [56, 57]. Identification of gRNA-sized
151 short antisense RNAs and accumulation of antisense precursors upon KRET1 and KDSS1
152 knockdowns [24] further indicates that sense and antisense precursors hybridize with their
153 complementary 5' regions. In the current model, the length of a double-stranded region,
154 which is a distance between respective transcription start sites, likely defines gRNA length
155 prior to uridylation [24]. However, most *T. brucei* minicircles encode 3-4 gRNA cassettes
156 [57] and primary RNAs may exceed a 1 kb linear length of a minicircle [24, 25]. Hence, an
157 extensive transcription of both strands may also generate much longer double-stranded RNAs
158 that are degraded by yet unknown mechanism. Conversely, *L. tarentolae* minicircle typically
159 contains a single gRNA gene and lacks recognizable inverted repeats [58]. Although both
160 strands are transcribed [59], the gRNA-flanking sequences are dissimilar from those of *T.*
161 *brucei*, which indicates a divergent nature of minicircle promoters among trypanosomatids.
162 The maxicircle promoters remain to be determined, but detection of antisense transcription
163 start sites near intergenic regions and presence of corresponding non-coding antisense
164 transcripts make a reasonable argument for a general mechanism of 3' end definition for
165 minicircle and maxicircle transcripts [23].

166

167 **Modification of the 5' End**

168 The 5' monophosphorylated termini of maxicircle-encoded rRNAs and mRNAs have long
169 been interpreted as indicative of endonucleolytic partitioning of polycistronic precursors. It
170 is, however, unfeasible to produce more than one monocistronic mRNA from a precursor by
171 3'-5' degradation. This logic dictates that: 1) each gene rests under control of a dedicated
172 promoter; 2) the 5' terminus is set by transcription initiation; 3) inorganic pyrophosphate
173 (PPi) is selectively removed from initiating nucleoside triphosphate in mRNAs and rRNAs,
174 but not in gRNAs; and 4) transcription may proceed across multiple genes and produce a 3'
175 extended precursor of which only the most 5' coding region is preserved after 3'-5' trimming.
176 Identification of the 5' pyrophosphate processome complex (PPsome) partially resolved the
177 question of differential phosphorylation status and linked 5' PPi removal to mRNA stability
178 [23]. A stable protein complex of MERS1 NUDIX [Nucleoside diphosphate linked to (X)]
179 hydrolase and MERS2 PPR RNA binding subunit, the MPsome selectively binds to
180 degenerate G-rich motifs found near mRNA 5' ends, but not in gRNAs. MERS1 hydrolase is
181 catalytically inactive as an individual protein while MERS2 confers both binding specificity
182 and affinity for RNA substrate. Remarkably, MERS1 downregulation or replacement with an
183 inactive version effectively eliminates most mRNAs but exerts negligible effects on gRNAs
184 and rRNAs. It appears that rRNA is stabilized by different factors, possibly those involved in
185 ribosome biogenesis [35]. Although PPsome-dependent mRNA protection against 3'-5'
186 degradation (see below) and the essential role of PPi removal are evident, the mechanistic
187 insights into these processes will likely come from understanding PPsome's interactions with
188 RNA editing substrate binding (RESC) and polyadenylation (KPAC) complexes discussed
189 below [32].

190

191 **Modifications of the 3' end**

192 Non-templated 3' nucleotide additions often wield profound influence on RNA processing,
193 function, trafficking and turnover [60]. In *T. brucei*, mitochondrial RNA 3' modifications are
194 classified into U-tailing by KRET1 TUTase (gRNAs and rRNAs), A-tailing by KPAP1
195 poly(A) polymerase (most mRNAs [31]), and A/U-extensions which require both enzymes
196 and a complex of Kinetoplast Polyadenylation Factors 1 and 2 (KPAF1/2, [30]). Lack of
197 pronounced RNA substrate specificity for KRET1 and KPAP1 raises the question of
198 accessory factors that enable modifications of distinct RNA classes, and the functionality of
199 these extensions. The presence of short U-tails in gRNAs and rRNAs, as well as non-
200 templated uridine residues sometimes found in mRNAs between the 3' UTR and the A-tail
201 [30], indicates that uridylation by the MPsome-embedded KRET1 is a default 3'
202 modification. It is plausible that U-tailing causes the MPsome to disengage from the
203 precursor when degradation pauses near double-stranded region formed by antisense RNA.
204 However, the U-tail itself does not exert an appreciable impact on mature gRNA or rRNA
205 stability [25] and its functionality beyond termination of processing remains debatable [61-
206 63]. Conversely, a short (15-30 nt) A-tail decorates most mRNAs and impacts stability
207 depending on transcript's editing status [28, 31, 64]. As demonstrated by KPAP1 poly(A)
208 polymerase loss-of-function studies [29, 31] and *in organello* decay assays [28], adenylation
209 mildly de-stabilizes pre-edited transcripts only to become essential for maintaining RNAs
210 that are edited beyond initial editing sites at the 3' end. A short A-tail also stabilizes never-
211 edited mRNAs (those that contain an encoded open reading frame and do not require editing).
212 The coupling between an mRNA's editing status and opposing effects of adenylation points
213 toward a surveillance system capable of both sensing the extent of internal U-
214 insertions/deletions and enabling 3' A-tail addition and function. In molecular terms,
215 sequence-specific activators and inhibitors would be expected to modulate mRNA

216 adenylation by KPAP1, and the resistance of such a modified molecules to decay. The
217 respective functions have been attributed to Kinetoplast Polyadenylation Factors (KPAF) 3
218 [29] and 4 [32], which belong to a family of 35 amino acid repeat-containing RNA binding
219 (PPR) proteins. Discovered in land plants [65], the helix-turn-helix PPR motif recognizes a
220 single nucleoside via side chains occupying cardinal positions 5 and 35 of the repeat (or the
221 last position in a longer structure). An array of adjacent PPR repeats often folds into a
222 superhelical domain capable of binding to specific RNA sequence and recruiting or blocking
223 various enzymes [66-68]. In this context, KPAF3 reportedly binds to G-rich pre-edited
224 mRNAs with sufficient affinity and coverage to stabilize these species following 3'-5'
225 trimming by the MPsome [29]. *In vitro* reconstitution experiments further demonstrate that
226 KPAF3 stimulates KPAP1 polyadenylation activity and this effect depends on the presence of
227 the G-rich site near the 3' end. Remarkably, KPAF3 binding is eliminated by the initiating
228 editing events leaving the stability of edited RNA reliant on A-tail added prior to editing [29].
229 Thus, KPAF3 functions as editing sensor and *bona fide* polyadenylation factor thereby
230 connecting the internal sequence changes and 3' modification [29-31].

231 The most apparent A-tail function would be protecting mRNA against degradation by
232 the MPsome. However, *in vitro* studies show that A-tailed RNAs can be degraded by the
233 purified MPsome, albeit less efficiently than uridylylated substrates [24]. The A-tailed
234 partially-edited pre-mRNAs are also somehow prevented from the post-editing addition of
235 the 200-300 nt-long A/U-tail. This modification marks fully-edited molecules [31] and
236 channels translationally-competent mRNA for translation [30, 33, 34]. Finally, the
237 mechanism of mRNA stabilization by the PPsome must reconcile binding of this complex to
238 the 5' end with blocking 3'-5' degradation [23]. To rationalize these observations, Mesitov et
239 al., envisaged a *trans*-acting factor that recognizes a nascent A-tail to enable an interaction
240 between PPsome occupying the 5' end and polyadenylation complex (KPAC) bound to the 3'

241 end [32]. It has been proposed that the resultant circularization increases mRNA resistance to
242 degradation and uridylation, and, therefore, to premature A/U-tailing and translational
243 activation of partially-edited transcripts [32]. Trypanosomal genomes apparently lack
244 mitochondrially-targeted canonical RRM motif-containing poly(A) binding proteins, but this
245 function is fulfilled by KPAF4. This PPR protein is almost entirely composed of seven
246 repeats of which five are predicted to bind sequential adenosine residues [69]. Co-purification
247 studies support KPAF4 interactions with KPAC components (KPAP1, KPAF1/2) and RESC-
248 mediated contacts with the PPsome. Accordingly, the A-tail has been identified as the
249 predominant *in vivo* binding site while *in vitro* KPAF4 selectively recognizes adenylated
250 substrates. Indeed, in the presence of KPAF4 adenylated RNA is more resistant to
251 degradation by the purified MPsome and uridylation by KRET1 TUTase.

252 Although direct demonstration of mRNA circularization is lacking, this event can be
253 imagined as a critical quality check point to ensure 5' end occupancy by the PPsome and
254 correct termination of 3'-5' trimming downstream from the KPAF3 binding site. In this
255 scenario, KPAF3 binding likely selects a correct 3' UTR among trimmed precursor isoforms
256 thereby distinguishing mRNA from rRNA and stimulating polyadenylation of the former by
257 KPAP1. KPAF4 binding to a nascent A-tail would enable interaction with 5' end-bound
258 PPsome, hence, mRNA circularization. Consequentially, only A-tailed mRNAs would
259 proceed though the editing cascade while the variants truncated beyond KPAF3 binding sites
260 become uridylated and degraded [29]. It follows that upon editing completion at the 5' end, a
261 signaling event must take place to release the circularization and enable access of KPAF1/2
262 factors and KRET1 TUTase in order to add long A/U-tail to the pre-existing short A-tail.
263 Although these postulates require further testing, it seems plausible that PPsome
264 displacement from the 5' end by final editing events may disrupt circularization and stimulate
265 A/U-tailing, a final processing step that renders mRNA competent for translation.

266 **U-insertion/deletion mRNA Editing**

267 *Editing Process*

268 In *T. brucei*, six of the 18 annotated mRNAs encode predicted polypeptides while the
269 remaining 12 transcripts undergo editing to acquire a protein-coding sequence. The extent of
270 editing varies from minor, typified by insertion of four Us into three closely-spaced sites
271 (COII mRNA, [10]), to moderate (e.g., *cyb* mRNA, 34 Us are inserted into a confined region
272 near 5' end [70]), to pan-editing during which hundreds of uridines are inserted or deleted
273 throughout the entire transcript (e.g. ND7 mRNA, [71]). The determinants of position-
274 specific U-insertions and deletions were discovered in the Simpson laboratory as short
275 patches of complementarity between edited mRNA and maxicircle DNA in *Leishmania*
276 *tarentolae* [45]. By allowing for G-U, in addition to canonical Watson–Crick base-pairing,
277 short (50-60 nt) RNAs transcribed from minicircles have been recognized as carriers of
278 genetic information and termed guide RNAs (gRNAs). *In vitro* experiments by the Stuart
279 laboratory directly demonstrated that gRNAs indeed constitute the necessary and sufficient
280 source of editing information [72-74]. Although the first gRNAs were discovered in the
281 maxicircle, further work in *Leishmania* established that most gRNAs are encoded in
282 minicircles [75]. In *T. brucei*, only two gRNAs have been identified in the maxicircle: a *cis*-
283 acting element embedded into 3' UTR of COII mRNA [76], and a *trans*-acting gRNA that
284 completes editing of the MURF2 mRNA. The secondary structure of gRNA-mRNA dictates
285 the editing site selection and the extent of U-insertions and deletions [72]. The initial gRNA-
286 mRNA interaction is accomplished via a short (10-12 nt) region of complementarity between
287 the gRNA's 5' anchor region and the pre-edited mRNA. The remaining guiding segment
288 forms an imperfect duplex with pre-mRNA resulting in looping out of single-stranded
289 uridines in mRNA (deletion sites) or purine nucleotides in gRNA (insertion sites). At either
290 site, the mRNA is cleaved at the first unpaired nucleotide adjacent to the 5' anchor duplex.

291 The resultant structures of deletion and insertion intermediates are distinct: single-stranded
292 uridines become exposed to a 3'-5' exonucleolytic attack in the former, while a single-
293 stranded gap is created between two helices in the latter. Upon trimming single-stranded
294 uridines from the 5' cleavage fragment in the deletion site or adding gap-specified number of
295 Us into the insertion site, the fragments are joined to restore mRNA continuity. Both types of
296 sequence changes extend the double-stranded anchor region. Pan-editing requires multiple
297 overlapping gRNAs and there is a method to it: sequence changes directed by the initiating
298 gRNA create a binding site for the next one to ensure an overall 3'-5' polarity along the
299 editing domain. However, editing may not always proceed strictly 3' to 5' as "mis-edited"
300 junctions are present at the leading edge of editing in the majority of steady-state mRNAs
301 [59, 77-79]. The role of junctions is not entirely understood, but they likely represent a
302 mixture of regions that undergo re-editing to canonical edited sequence, dead-end products,
303 or mRNAs with alternative non-canonical coding sequences [22].

304 A single editing domain may cover an entire mRNA [80], or an isolated region [71].
305 An individual gRNA can theoretically direct insertions and deletion at several closely spaced
306 sites (editing block), but as editing progresses within the block, the interaction between
307 gRNA and mRNA 5' cleavage fragment is supported by fewer base pairs. Stabilizing the 5'
308 cleavage fragment-mRNA tethering by additional base pairing stimulates cleavage and the
309 full editing cycle *in vitro* [81-83], but it is unclear how the problem of editing at distal sites
310 within one block or across sequential blocks is solved *in vivo*. An active displacement of a
311 gRNA with diminishing "3' tether" by RNA helicase is among possible solutions that would
312 enable binding of succeeding gRNA within a domain. Alternatively, the gRNA's U-tail may
313 help stabilize interaction with an mRNA [61, 84], but the conclusive evidence for this event
314 is yet to be obtained.

315

316 *Elemental Editing Reactions*

317 Editing reactions are catalyzed by enzymes embedded into the ~20S (~800 kDa) RNA
318 Editing Catalytic Complex (RECC), a remarkable example of a modular assembly that
319 enables broad functionality on distinct RNA substrates [85-89]. A common core particle
320 consists of U-insertion (KRET2 TUTase, KREPA1 zinc finger protein and KREL2 RNA
321 ligase) and U-deletion (KREX2 exonuclease, KREPA2 zinc finger protein and KREL1 RNA
322 ligase) sub-complexes and six structural and/or RNA binding proteins (KREPA3, KREPA4,
323 KREPA5, KREPA6, KREPB4, KREPB5). The U-insertion and U-deletion sub-complexes
324 likely function independently [48, 90] while most of the remaining components are essential
325 for assembly and/or integrity of the entire core particle [91-94]. The core particle is shared
326 among at least three RECC isoforms distinguished by association with endonuclease
327 modules. Each module is composed of an RNase III endonuclease and a partner protein(s)
328 and is primarily responsible for recognition and cleavage of insertion and deletion sites. The
329 U-deletion sites are recognized by the RECC isoform with KREN1+KREPB8+KREX1,
330 while U-insertion sites are recognized by the RECC isoforms with KREN2+KREPB7 or
331 KREN3+KREPB6, which display distinct and overlapping specificities [95-99]. The
332 canonical RNase III catalytic domain typically forms a functional homodimer with two active
333 sites that introduce four cuts into both strands of a double-stranded RNA [100]. In contrast,
334 editing endonucleases appear to cleave only mRNA. It seems plausible that RNA hydrolysis
335 is restricted to a single cut by heterodimer formation between KREN1, KREN2 or KREN3,
336 and catalytically-inactive degenerate RNase III domains in KREPB4 or KREPB5 [101]. A
337 contribution of RNase III partner proteins KREPB8, KREPB7 or KREPB6 to modulating
338 cleavage activity is also possible [102, 103]. Binding of KREN1, KREN2 and KREN3
339 modules to a common core containing U-deletion, U-insertion and ligase activities highlights
340 RECC's modular nature [98, 104, 105], but the nature of interactions responsible for mutually

341 exclusive contacts between the core and distinct modules remains unclear. Cross-linking
342 mass-spectrometry points to interactions involving RNase III domain dimerization between
343 editing endonucleases with partner proteins KREPB6, B7 or B8, and core proteins KREPB4
344 and KREPB5 [102, 103, 106].

345 Within the common core, the U-deletion and U-insertion cascades are spatially
346 separated by virtue of editing enzymes binding to zinc finger proteins, KREPA2 and
347 KREPA1, respectively [90, 107-109]. KREX1 and KREX2 proteins possess exonuclease-
348 endonuclease-phosphatase (EEP) catalytic domains and display single-stranded uridine-
349 specific 3'-5' exonuclease activity *in vitro* [108, 110]. However, their protein-protein
350 interactions are remarkably distinctive: the essential KREX1 belongs to the KREN1
351 endonuclease module, and is responsible for the main U-deletion activity; the dispensable
352 KREX2 probably represents a structural component of the U-deletion sub-complex [96, 106].
353 Fittingly, *L. tarentolae* KREX2 lacks a catalytic domain, but remains associated with the U-
354 deletion sub-complex [85]. In the U-insertion sub-complex, KRET2 TUTase binds to
355 KREPA1, which results in a mutual stabilization and stimulation of TUTase activity [48, 106,
356 111-113]. Selectivity of uridine incorporation is determined by KRET2's intrinsic specificity
357 for UTP [114] rather than the nature of the opposing nucleotide in the gRNA. To that end, the
358 +1U addition occurs equally efficient irrespective of the corresponding nucleotide in gRNA,
359 but the +2U addition occurs only if the +1U forms a base pair with either adenosine or
360 guanine. Consequentially, both purine bases in guiding positions direct U-insertions with
361 similar efficiency [81, 111] . RNA editing ligases 1 and 2 (KREL1 and KREL2) have been
362 identified as components of U-deletion and U-insertion sub-complexes, respectively [90, 106,
363 107]. Although spatial separation appears to suggest specialized roles, only KREL1, but not
364 KREL2, is essential for cell viability [47, 115, 116].

365

366 ***Editosome Definition***

367 From the early reports of RNA ligase-containing complexes sedimenting in glycerol density
368 gradients as particles with apparent 20S to 50S values [117, 118], the quest for an elusive
369 “editosome” evolved into a concept of an RNA editing holoenzyme. For the purposes of this
370 review, we shall equate the editosome and editing holoenzyme and define this entity as an
371 RNA-mediated assembly of the RNA editing catalytic (RECC), RNA editing substrate
372 binding (RESC) and RNA editing helicase REH2 (REH2C) complexes. It is a virtual
373 certainty that additional components are also involved [19, 21, 119]. This definition stems
374 from parallel lines of inquiry by the Stuart, Afasizhev and Lukeš laboratories that identified
375 an ~800 kDa protein complex (originally termed Mitochondrial RNA Binding Complex 1,
376 MRB1, and Guide RNA Binding Complex, GRBC), of which two components are essential
377 for gRNA stability [120-122]. Initially named GRBC1 and GRBC2, these homologous
378 polypeptides lack annotated motifs and similarity to any protein outside of Kinetoplastids
379 [63]. GRBC1 and GRBC2, also referred to as GAP2 and GAP1, respectively [122], form a
380 stable heterotetramer which binds gRNA *in vitro* and *in vivo* [52]. Extensive co-purification
381 and yeast two-hybrid screens further dissected MRB1 into two relatively stable protein
382 complexes: an ~20-component RNA Editing Substrate Binding Complex (RESC), which
383 includes RESC1 (GRBC1, GAP2) and RESC2 (GRBC2, GAP1), and three-subunit RNA
384 Editing Helicase 2 Complex (REH2C) (Table 1). It appears that both RESC and REH2C bind
385 editing substrates, intermediates and products, and engage in RNA-mediated interactions with
386 the catalytic RECC complex [52, 123-126]. All but five of the RESC subunits lack
387 discernible motifs or similarities to non-kinetoplastid proteins, although several exhibit *in*
388 *vitro* RNA binding activity [19, 21, 119]. Most subunits are essential for cell viability and
389 their knockdowns typically produce phenotypes consistent with an inhibited editing process.
390 Recently, a substantial progress has been made in deciphering roles of individual factors. The

391 RESC1/2 tetramer appears to be solely responsible for gRNA stabilization [121, 122]. Deep
392 sequencing studies showed that the RRM/RGG-containing RESC13 (RGG2) and proximal
393 protein RESC11A (MRB8180) contribute to editing processivity within an extended domain
394 [78, 127]. These two proteins strongly promote the formation of junctions, implying a critical
395 role of these regions in editing progression [78]. Conversely, the product of a duplicate gene
396 RESC12A (MRB8170) has been implicated in editing initiation and in constraining the
397 region of active editing [78, 128]. Biochemical attempts to refine RESC architecture indicate
398 a modular organization with potential protein clusters responsible for interaction with the
399 RECC and polyadenylation complexes [52, 129]. However, an unambiguous assignment of
400 specific polypeptides to functionally meaningful modules awaits elucidation of a high-
401 resolution structure of the RESC complex, which is likely to be heterogenous and dynamic.

402 RNA editing is an essential processing step for a subset of mitochondrial transcripts
403 and must be integrated into a general pathway of producing translation-competent mRNAs.
404 Accumulating evidence suggests that the RESC is responsible for coordinating pre- and post-
405 editing processing events via RNA-mediated contacts with 5' and 3' modification complexes,
406 and auxiliary factors. On the other hand, the catalytic RECC isoforms appear to act on RESC-
407 bound editing substrates in a transient manner. The RNA-mediated interaction between
408 RESC and PPsome has been deduced from co-purification of RESC1/2 (GRBC1/2) and
409 MERS1 hydrolase [23, 121, 130] whereas *in vivo* proximity biotinylation identified RESC19
410 (MERS3) as the most plausible adapter subunit [23]. An independent study predicted Z-
411 DNA-binding domains in RESC19 (termed RBP7910) and showed that *in vitro* this protein
412 preferentially binds RNAs containing poly(U) and poly(A/U)-rich sequences [131]. Likewise,
413 kinetoplast polyadenylation complex components have been consistently detected in various
414 RESC preparations and particularly in those with tagged RESC15-18 [29, 32, 52].
415 Furthermore, RESC13 (RGG2) and surrounding proteins likely mediate contacts between

416 RESC and RECC complexes [52]. Finally, RNA editing helicase 2 (KREH2, see below)
417 preferentially associates with RESC variants purified by tagging of either RESC1 or RESC2
418 [52, 121, 132-134]. Thus, the RESC complex functions not only in binding of RNA editing
419 substrates, intermediates and products, but also recruits mRNA modification complexes and
420 auxiliary factors via specific subunits.

421 The Kinetoplast RNA Editing Helicase 2 (KREH2) complex, termed REH2C, consists
422 of DEAH/RHA RNA helicase KREH2, and KREH2-associated factors 1 (KH2F1) and 2
423 (KH2F2). KH2F1 contains eight C2H2 zinc fingers while KH2F2 lacks any identifiable
424 motifs [135] (Table 1). Isolated REH2C exhibits ATP-dependent 3'-5' dsRNA unwinding
425 activity and co-sediments with a major peak of same activity in mitochondrial extracts [134].
426 Zinc finger protein KH2F1 emerged as an adaptor connecting KREH2 helicase with the
427 editosome while gRNA-mRNA hybridization has also been implicated in facilitating this
428 interaction [135]. KREH2 and KH2F1 knockdowns display consistent phenotypes of
429 increased editing pausing and reduced processivity of editing, which are indicative of REH2C
430 participation in continuous remodeling of gRNA-mRNA contacts and perhaps the entire
431 editosome [123, 135]. It must be emphasized that the reciprocal affinity purifications remain
432 the most salient evidence of the editosome being an RNA-based assembly of RECC, RESC
433 and REH2C protein complexes [120-122, 133, 134].

434

435 **Auxiliary RNA Processing Factors**

436 ***Putative Poly(A) Polymerase KPAP2***

437 A putative kinetoplast poly(A) polymerase KPAP2 has been identified by homology to the
438 human mitochondrial enzyme and apparently is not required for axenic *T. brucei* growth in
439 either bloodstream or procyclic life stages [136]. Although KPAP2 protein sequence is highly

440 similar to that of KPAP1, its enzymatic identity and *in vivo* substrates remain to be
441 established. Available proteomics data do not support KPAP2 association with KPAC [29,
442 31, 32].

443 ***REH1 RNA Helicase***

444 Editing reactions are expected to produce an mRNA-gRNA duplex wherein gRNA must be
445 eventually displaced to allow binding of a sequential gRNA, or before the edited transcript
446 can be translated. It stands to reason that active remodelers, such as DEAD/H-box RNA
447 helicases, would be involved and indeed two such proteins have been implicated in the
448 editing process. However, demonstrating their specific roles, RNA targets and mechanism of
449 action proved to be challenging. Knockdown of KREH1 (Hel61) helicase [137, 138] affected
450 editing mediated by two or more overlapping gRNAs [139], but the existing data do not allow
451 unambiguous placing of KREH1 into mRNA editing or processing complexes. The nature of
452 KREH1 substrates and the timing and purpose of KREH1-dependent RNA remodeling also
453 remain unclear. Although much remains to be understood about the helicases' involvement in
454 editing and other processes, the transcript-specific impacts of repressing KREH1 and KREH2
455 (discussed above) justify further efforts to dissect their molecular functions.

456

457 ***RNA Binding Factors***

458 Kinetoplast Mitochondrial RNA Binding Proteins 1 and 2 (KMRP1 and 2), originally called
459 gBP21 and gBR25, then MRP1 and MRP2, were identified independently in *T. brucei* by
460 UV-induced crosslinking with synthetic gRNA (gBP21, [140]), in *C. fasciculata* as poly(U)
461 binding proteins (gBP21 and gBP25, [141]), and in *L. tarentolae* via crosslinking to double-
462 stranded RNA resembling the U-deletion site (MRP1 and MRP2, [130]). Extensive
463 biochemical and structural studies concluded that KMRP1 and KMRP2 assemble into an

464 ~100 kDa heterotetramer, which binds both single- and double-stranded RNAs with high
465 affinity [130, 142, 143]. These RNA binding properties are manifested by an RNA annealing
466 activity, an attractive accessory function that may promote gRNA binding to cognate mRNA
467 targets [144, 145]. However, the transcript-specific impact of dual KMRP1/2 repression
468 suggests a contribution to stabilization of moderately edited and some never-edited mRNAs
469 rather than direct participation in the editing process [142, 146, 147]. In support of this
470 notion, RNAi experiments demonstrated that MRP1/2 depletion virtually eliminates the
471 edited form of the moderately-edited *cyb* mRNA, but exerts little impact on the pre-edited
472 transcript [146-148]. While much is known about the KMRP1/2 structure and *in vitro*
473 properties, the definitive function of this RNA binding complex remains to be established.
474 Much of the same narrative applies to KRBP16 (RBP16), which carries N-terminal cold
475 shock and C-terminal RG-rich domains [149]. RNAi studies revealed an overlap between
476 mRNA sets negatively affected by individual KMRP1/2 and KRBP16 knockdowns: edited
477 *cyb* mRNA, but not any other edited transcripts, was severely downregulated while never-
478 edited CO1 and ND4 transcripts also declined [146]. KRBP16 *in vitro* properties, such as
479 RNA binding affinity, RNA annealing activity and stimulation of editing activity [146, 148-
480 155], and the impact of RNAi knockdown on the initiation of *cyb* mRNA editing [148] are
481 consistent with participation in the editing process, although the mechanistic role remains to
482 be firmly established. Another enigmatic RNA binding protein, KRGG1 was serendipitously
483 discovered in a large (>50S) RNP of unknown nature [156], and subsequently demonstrated
484 to associate with the RESC complex [122, 157]. A different study identified a ribosome-
485 bound KRGG1 fraction, which would explain the observed sedimentation patterns, but found
486 no impact on RNA editing [52]. Another arginine-glycine-rich protein KRGG3, originally
487 identified by association with RESC1/2 proteins and termed MRB1820 [129], is essential for
488 parasite viability [158]. However, most KRGG3's interactions appear to be RNA-mediated

489 while the RNAi knockdown does not significantly impact major mitochondrial RNA classes.
490 A structural study identified an ABC-ATPase fold and potential RNA binding surface in a 72
491 kDa protein KRBP72, initially termed MRB1590 [159]. KRBP72 knockdown specifically
492 impacts editing of A6 mRNA [159]; however, an unequivocal functional placement of this
493 factor also requires further investigation. Finally, participation of KREAP1 in editing [160,
494 161] has been contravened by a report of general mitochondrial RNA upregulation upon its
495 knockdown and non-essentiality for parasite survival [162]. Mitochondrial RNA binding
496 proteins are abundant and notoriously promiscuous in their interactions and pleiotropic
497 effects on RNA steady-state levels [163], which makes an unequivocal definition of their
498 function a challenge worth meeting.

499 ***Ribonucleases***

500 Mitochondrial RNA processing most likely involves nucleolytic events beyond mRNA
501 cleavage by editing KREN1, 2 and 3 endonucleases, KREX1 exonuclease, and 3'-5'
502 degradation by the MPsome-embedded KDSS1. To that end, three distinct enzymes have
503 been identified and characterized to various degrees. The single-strand uridine specific
504 KRND1 3'-5' exonuclease [164] displays *in vitro* specificity for single-stranded uridines,
505 similar to that of KREX1 editing enzyme [108, 110], and yet possesses an RND rather than
506 EEP exonuclease domain. Given the diversity of U-tailed RNAs in the kinetoplastid
507 mitochondrion, it is tempting to speculate on KRND1 involvement in regulating the 3'
508 modification state, but its definitive function remains to be established. The same narrative
509 applies to KRPN1, an RNase III-like endonuclease with a characteristic double-stranded
510 RNA cleaving activity [42, 46] suggested to function in gRNA processing. Further studies are
511 required to reconcile KRPN1 activity and interactions with an exonucleolytic mechanism of
512 gRNA precursor processing by the MPsome. Finally, the discovery of PPR-repeat containing
513 proteinaceous RNase P (PROPR2 [165], re-named here as KRNP1) supported earlier reports

514 of RNase P activity-like which removes the 5' leader from a synthetic tRNA precursor in
515 mitochondrial lysate [166]. However, tRNAs are apparently imported into the mitochondrion
516 with 5' and 3' extensions removed [167, 168], which leaves the nature of KRNP1 *in vivo*
517 substrates open to future inquiry.

518

519 **Concluding Remarks**

520 This review compiles 74 processing enzymes, RNA binding proteins and factors with
521 unknown functionality that nonetheless are associated with RNA processing complexes.
522 Proteomics and interactions analyses allowed clustering most of these into 5-6
523 macromolecular assemblies albeit with various degree of confidence. Although the list is
524 almost certainly incomplete, with complexes and individual proteins, and their interactions
525 and functions being constantly re-examined, the overall flow of RNA processing in
526 trypanosome mitochondrion is taking shape and meaning. The key players responsible for
527 maturation of 5' and 3' termini have been defined and initial insights gained into the
528 molecular mechanism of internal sequence changes by editing. At this point, we suggest that
529 the RNA editing holoenzyme (editosome) represents an RNP that chiefly includes three
530 relatively stable protein complexes (RECC, RESC and REH2C) and RNA editing substrates,
531 intermediates and products. It is understood that the definition of protein complex is to large
532 extent a matter of purification techniques and we posit that future structural studies will shed
533 light on stoichiometry and functions of individual subunits and modules. Because of
534 sequence changes introduced by editing during mRNA processing, the overall picture is
535 emerging of the mRNA fate being dictated by diverse PPR RNA surveillance factors. These
536 proteins direct 5' pyrophosphate removal, transcript stabilization and pre-editing A-tailing,
537 monitor initiation and progression of editing, and signal its completion by stimulating the

538 A/U-tailing. Displacement of bound PPRs from pre-edited mRNA by the editing process
539 appears to be the principal quality control mechanism. It remains to be established whether
540 active RNP remodeling takes place or sequence changes alone suffice for this purpose. In any
541 event, the plurality of PPRs and their capacity to read linear sequences and modulate the
542 activity of RNA modification and degradation enzymes position this protein family as the
543 focal point of mitochondrial RNA processing.

544

545

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990

991 **Glossary**

992 **5' and 3' cleavage fragments:** mRNA fragments generated by gRNA-directed
993 endonucleolytic cleavage.

994 **Anchor:** 5' part of the gRNA that forms a continuous 10-15 nt duplex with pre-edited,
995 partially-edited or fully-edited mRNA; this region is responsible for initial gRNA-mRNA
996 interaction.

997 **Cryptogene:** a maxicircle gene with defective coding sequence; the defects are corrected by
998 U-insertion/deletion editing with concomitant restoration of protein reading frame.

999 **Editing block:** an mRNA segment covered by a single gRNA; often contains both U-
1000 insertion and U-deletion sites.

1001 **Editing domain:** an mRNA region covered by single or multiple overlapping gRNAs. In a
1002 multi-gRNA domain, sequence changes directed by the initiating gRNA create the binding
1003 site for the subsequent one. The hierarchical gRNA binding provides for the overall 3'-5'
1004 progression of editing events within a domain.

1005 **Editing site:** position of the gRNA-directed internal cleavage where uridines are either
1006 deleted from or inserted into the mRNA.

1007 **EEP domain:** endonuclease/exonuclease/phosphatase (EEP) domain in U-specific editing
1008 exonucleases.

1009 **Fully-edited mRNA:** a final editing product; contains a protein coding frame.

1010 **Guide RNA (gRNA):** a small non-coding RNA that specifies positions and extent of U-
1011 insertions and deletions by forming an imperfect duplex with pre-edited or partially-edited
1012 mRNA. Guide RNA is typically 30-60 nucleotides (nt) in length and possesses a 5'
1013 triphosphate and a 1-20 nt 3' U-tail.

1014 **Junction:** A region present in most partially-edited mRNAs at the 5' leading edge of editing;
1015 often displays mis-edited and non-canonically edited sequences. Junction-containing
1016 transcripts may represent intermediates that will be re-edited to canonical sequence, dead-end
1017 by-products, and mRNAs with a non-canonical protein coding sequence.

1018 **Kinetoplastids (class Kinetoplastea):** flagellated protists characterized by the presence of a
1019 kinetoplast. Phylogenetically positioned within the phylum Euglenozoa, this group includes
1020 the obligatory parasitic trypanosomatids (family *Trypanosomatidae*), free-living and parasitic
1021 bodonids, and more distantly related taxa.

1022 **Kinetoplast:** a densely packed nucleoprotein structure, disc-shaped and catenated in
1023 trypanosomatids, and dispersed to various degrees in most bodonids, that encloses
1024 mitochondrial DNA (kinetoplast DNA; kDNA). A non-dividing *T. brucei* cell contains a
1025 single mitochondrion with a single kinetoplast composed of catenated maxicircles (~23 kb
1026 each, few dozen copies) and minicircles (~1 kb each, ~5,000 units).

1027 **KPAC:** kinetoplast polyadenylation complex. A ribonucleoprotein complex of KPAP1
1028 poly(A) polymerase, and pentatricopeptide repeat-containing (PPR) RNA binding proteins
1029 designated Kinetoplast Polyadenylation Factors 1, 2, 3 and 4 (KPAF1, 2, 3 and 4).

1030 **Maxicircle:** an equivalent of a typical mitochondrial genome; includes a conserved ~15 kb
1031 region encoding 9S and 12S rRNAs, two guide (g)RNAs, and 18 protein genes. A variable
1032 region composed of repeated DNA sequences constitutes the rest of the molecule.

1033 **Minicircle:** the molecules forming the bulk of kinetoplast. Approximately 400 sequence
1034 classes present at various frequencies encode ~930 gRNAs required for the editing process
1035 and 370 gRNA-like molecules that likely participate in gRNA processing.

1036 **Moderately-edited mRNA:** a transcript with a few editing sites confined to a limited mRNA
1037 region.

1038 **MPsome:** mitochondrial 3' processome. A protein complex composed of Kinetoplast RNA
1039 editing TUTase 1 (KRET1), 3'-5' exonuclease KDSS1, and MPSS1-6 subunits lacking
1040 recognizable motifs.

1041 **MTRNAP:** mitochondrial RNA polymerase, a single-subunit T3/T7-like DNA-dependent
1042 RNA polymerase.

1043 **Never-edited mRNA:** a maxicircle transcript containing an encoded open reading frame
1044 which does not require editing.

1045 **Pan-edited mRNA:** a transcript that undergoes massive editing directed by multiple gRNAs.
1046 There can be two editing domains within a pan-edited mRNA.

1047 **Partially-edited mRNA:** an intermediate of the editing process. Partially-edited mRNAs
1048 often contain junctions whose sequences match neither pre-edited nor canonical fully-edited
1049 mRNAs.

1050 **PPsome:** 5' pyrophosphate processome. Protein complex containing MERS1 NUDIX
1051 pyrophosphohydrolase and MERS2 PPR RNA binding protein.

1052 **PPR:** Pentatricopeptide (35 amino acids) helix-turn-helix repeat. PPR arrays are present in
1053 many trypanosomal mitochondrial RNA binding proteins.

1054 **Pre-edited mRNA:** a 3' processed monocistronic cryptogene transcript that must undergo
1055 editing to acquire an open reading frame and/or translation initiation and termination signals.

1056 **RECC:** RNA Editing Catalytic Complex, formerly called ~20S editosome or RNA Editing
1057 Core Complex. A protein complex of 14 or more subunits, depending on the isoform;
1058 includes pre-mRNA cleavage, U-insertion, U-deletion and RNA ligation enzymes, and
1059 structural and RNA binding factors.

1060 **RESC:** RNA Editing Substrate Binding Complex, formerly called Mitochondrial RNA
1061 Binding Complex 1 (MRB1) and Guide RNA Binding Complex (GRBC). An ~20-subunit
1062 modular protein complex that likely exists in several isoforms; most components lack
1063 recognizable motifs. RESC binds RNA editing substrates, intermediates and products, and
1064 coordinates interactions of gRNA and mRNA with RECC, REH2C and other auxiliary
1065 factors during editing. RESC has also been implicated in coordination of pre-mRNA 5' and
1066 3' modification processes.

1067 **REH2C:** RNA Editing Helicase 2 Complex, a protein complex formed by an ATP-dependent
1068 DEAH/RHA RNA helicase KREH2, zinc finger protein KH2F1, and KH2F2 factor which
1069 lacks recognizable motifs.

1070 **RNA editing holoenzyme (editosome):** a ribonucleoprotein particle consisting of RECC,
1071 RESC and REH2C complexes, and several auxiliary factors.

1072 **RNA helicase:** a motor protein capable of harnessing the energy from NTP hydrolysis to
1073 unwind double stranded RNAs or to remodel ribonucleoprotein complexes.

1074 **RNase II:** Exoribonuclease II cleaves single-stranded RNA in 3' to 5'-direction yielding
1075 nucleoside 5' monophosphates.

1076 **RNase III:** Endoribonuclease III typically cleaves both strands in double-stranded RNA
1077 leaving 5' monophosphate and 3' hydroxyl groups. RNase III editing endonucleases cleave
1078 only the mRNA strand at an unpaired nucleotide adjacent to a gRNA-mRNA duplex.

1079 **TUTase:** terminal uridylyltransferase, UTP-specific nucleotidyl transferase which adds U-
1080 residues to the 3' end of RNA.

1081 **U-insertion/deletion mRNA editing:** a process by which U-residues are inserted into, or
1082 deleted from, a cryptogene transcript. Editing is directed by gRNAs and catalyzed by the
1083 RNA editing holoenzyme (editosome).

1084 **UTR:** untranslated region of mRNA.

1085

1086 **Table 1.** Proposed nomenclature of mitochondrial RNA processing complexes and factors.

1087 Gene identification numbers refer to *T. brucei* strain TREU927 predicted protein sequences

1088 (TriTrypDB, Release 45, September 5, 2019, <https://tritrypdb.org/tritrypdb/>).

Legacy	Assigned	Function	Motifs	TriTryp ID	References
Nucleolytic Processing: Mitochondrial 3' processome (MPsome)					
RET1	KRET1	KRET1	TUTase, 3' uridylation of primary and mature RNAs	TUTase, PAP associated	Tb927.7.3950 [24, 25, 48, 49, 169]
KDSS1	KDSS1	KDSS1	3'-5' exonuclease	RNB (Ribonuclease II)	Tb927.9.7210 [24, 26, 50]
MPSS1		MPSS1			Tb927.11.9150 [24]
MPSS2		MPSS2			Tb927.10.9000 [24]
MPSS3		MPSS3			Tb927.3.2770 [24]
MPSS4		MPSS4			Tb927.10.6170 [24]
MPSS5		MPSS5			Tb927.9.4810 [24]
MPSS6		MPSS6			Tb927.6.2190 [24]
Modification of the 5' end: Pyrophosphohydrolase complex (PPsome)					
MERS1		MERS1	PPi removal from 5' end	NUDIX hydrolase	Tb927.11.15640 [23, 121, 122]
MERS2		MERS2	Targets MERS1 to RNA	PPR	Tb11.02.5120 [23]
Modification of the 3' end: Kinetoplast polyadenylation complex (KPAC)					
KPAP1		KPAP1	Major poly(A) polymerase	NT/TUTase, PAP associated	Tb927.11.7960 [31]
KPAF1	PPR1	KPAF1	mRNA A/U-tailing	PPR	Tb927.2.3180 [30, 170, 171]
KPAF2		KPAF2	mRNA A/U-tailing	PPR	Tb927.11.14380 [30]
KPAF3		KPAF3	mRNA stabilization/A-tailing	PPR	Tb927.9.12770 [29]
KPAF4		KPAF4	Poly(A) binding protein	PPR	Tb927.10.10160 [32]
U-insertion/deletion mRNA Editing: RNA editing catalytic complex (RECC)					
REN1	KREPB1	KREN1	U-deletion endonuclease	RNase III, PUF, ZF-C2H2	Tb927.1.1690 [86, 172]
REN2	KREPB3	KREN2	U-insertion endonuclease	RNase III, PUF, ZF-C2H2	Tb927.10.5440 [86, 173]
REN3	KREPB2	KREN3	U-insertion endonuclease	RNase III, PUF, ZF-C2H2	Tb927.10.5320 [85, 86, 98]
REX1	KREX1	KREX1	3'-5' U-specific exonuclease	Exo/endo/phos (EEP)	Tb927.7.1070 [85, 86, 97]
REX2	KREX2	KREX2	3'-5' U-specific exonuclease	Exo/endo/phos (EEP)	Tb927.10.3570 [85, 86, 97]
RET2	KRET2	KRET2	U-insertion TUTase	TUTase, PAP associated	Tb927.7.1550 [48, 85, 86, 113]
REL1	KREL1	KREL1	RNA ligase (U-deletion)	RNA lig/RNL2	Tb927.9.4360 [47, 85, 86, 90]
REL2	KREL2	KREL2	RNA ligase (U-insertion)	RNA lig/RNL2	Tb927.1.3030 [85, 86, 90]
MP81	KREPA1	KREPA1		ZF-C2H2, OB fold	Tb927.2.2470 [85, 86, 174]
MP63	KREPA2	KREPA2		ZF-C2H2, OB fold	Tb927.10.8210 [85, 86, 175]
MP42	KREPA3	KREPA3		ZF-C2H2, OB fold	Tb927.8.620 [85, 86, 92, 176]
MP24	KREPA4	KREPA4		OB fold	Tb927.10.5110 [85, 86, 94]
MP19	KREPA5	KREPA5		OB fold	Tb927.8.680 [86]
MP18	KREPA6	KREPA6		OB fold	Tb927.10.5120 [85, 86, 91, 177]
MP46	KREPB4	KREPB4		RNase III, PUF, ZF-C2H2	Tb927.11.2990 [85, 86, 103]
MP44	KREPB5	KREPB5		RNase III, PUF, ZF-C2H2	Tb927.11.940 [85, 86, 176, 178]
MP49	KREPB6	KREPB6		RNase III, ZF-C2H2	Tb927.3.3990 [85, 86, 95, 102]
MP47	KREPB7	KREPB7		RNase III, ZF-C2H2	Tb927.9.5630 [95, 102, 104]
MP41	KREPB8	KREPB8		RNase III, ZF-C2H2	Tb927.8.5690 [95, 102, 104]
	KREPB9	KREPB9		RNase III, ZF-C2H2	Tb927.9.4440 [179, 180]
	KREPB10	KREPB10		RNase III, ZF-C2H2	Tb927.8.5700 [179, 180]
MEAT1		MEAT1	RECC-like associated TUTase	TUTase, PAP associated	Tb927.1.1330 [181]

U-insertion/deletion mRNA Editing: RNA editing substrate binding complex (RESC)						
GRBC1	GAP2	RESC1	gRNA binding/stabilization		Tb927.7.2570	[121, 122]
GRBC2	GAP1	RESC2	gRNA binding/stabilization		Tb927.2.3800	[121, 122]
GRBC3	MRB8620	RESC3			Tb927.11.16860	[52, 124, 129]
GRBC4	MRB5390	RESC4			Tb11.02.5390b	[52, 129, 182]
GRBC5	MRB11870	RESC5			Tb927.10.11870	[52, 129, 183]
GRBC6	MRB3010	RESC6			Tb927.5.3010	[52, 123, 129, 133, 184]
GRBC7	MRB0880	RESC7			Tb927.11.9140	[52, 129]
REMC1	MRB10130	RESC8	RNA binding	ARM/HEAT repeats	Tb927.10.10130	[52, 120, 125, 129]
REMC2	MRB1860	RESC9			Tb927.2.1860	[52, 129]
REMC3	MRB800	RESC10			Tb927.7.800	[52, 129]
REMC4	MRB8180 MRB4150	RESC11A RESC11B	RNA binding		Tb927.8.8180 Tb927.4.4150	[52, 78, 129]
REMC5	MRB4160	RESC12	RNA binding		Tb927.4.4160	[52, 128, 129, 185]
REMC5 A	MRB8170	RESC12A			Tb927.8.8170	[52, 78, 128, 129, 185]
TbRGG2	TbRGG2	RESC13	RNA binding	RGG, RRM	Tb927.10.10830	[52, 78, 127, 129, 182, 186-188]
	MRB7260	RESC14		PhyH	Tb927.9.7260	[126, 129]
PAMC1		RESC15			Tb927.1.1730	[52]
PAMC2		RESC16			Tb927.6.1200	[52]
PAMC3		RESC17			Tb927.10.1730	[52]
PAMC4		RESC18			Tb927.1.3010	[52]
MERS3	RBP7910	RESC19		Z-DNA binding	Tb927.10.7910	[23, 131]
U-insertion/deletion mRNA Editing: REH2 RNA Helicase Complex (REH2C)						
REH2		KREH2	RNA helicase, RNA binding	DEAH/RHA, HA2, DSRM, OB/NTP_binding	Tb927.4.1500	[120-123, 132-135]
H2F1	MRB1680	KH2F1		ZF-C2H2	Tb927.6.1680	[134, 135, 182]
H2F2		KH2F2			Tb927.6.2140	[132, 134, 135]
Auxiliary RNA Processing Factors						
KPAP2		KPAP2	Putative poly(A) polymerase	NT/TUTase, PAP associated	Tb927.10.160	[136]
REH1	mHEL61	KREH1	RNA helicase	DEAD/DEAH box helicase	Tb927.11.8870	[139, 189]
MRP1	gBP21	KMRP1	RNA binding		Tb927.11.1710	[130, 140-148]
MRP2	gBP25	KMRP2	RNA binding		Tb927.11.13280	[130, 142, 143, 146-148]
RGG1		KRGG1	RNA binding		Tb927.6.2230	[122, 156, 157]
RBP16		KRBP16	RNA binding	Cold-shock RNA binding	Tb927.11.7900	[146, 148-155]
	MRB1590	KRBP72	RNA binding	ABC-like ATPase domain	Tb927.3.1590	[159]
TbRGG3	MRB1820	KRGG3			Tb927.3.1820	[129, 158]
REAP-1		KREAP1	RNA binding		Tb927.10.9720	[160-162]
RND		KRND1	U-specific 3'-5' exonuclease	RND, ZF-C2H2	Tb927.9.12720	[164]
PRORP2		KRNP1	RNase P	PRORP, PPR	Tb927.11.3010	[165]
mRPN1		KRPN1	Endonuclease	RNase III	Tb927.11.8400	[42, 46]

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1091 **Figure 1.** A schematic diagram of mitochondrial RNA processing in *T. brucei*. The flow of
1092 processing reactions does not imply an experimentally established timing of these events. For
1093 example, the rRNA assembly into the ribosome or 5' pyrophosphate removal from mRNA
1094 may occur co-transcriptionally. Likewise, the mRNA may be edited prior to completion of 3'-
1095 5' trimming and 3' adenylation.

