Native mitochondrial RNA-binding complexes in kinetoplastid RNA editing differ in guide RNA composition

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ABSTRACT
Mitochondrial mRNAs in kinetoplastids require extensive U-insertion/deletion editing that progresses 3′-to-5′ in small blocks, each directed by a guide RNA (gRNA), and exhibits substrate and developmental stage-specificity by unsolved mechanisms. Here, we address compositionally related factors, collectively known as the mitochondrial RNA-binding complex 1 (MRB1) or gRNA-binding complex (GRBC), that contain gRNA, have a dynamic protein composition, and transiently associate with several mitochondrial factors including RNA editing core complexes (RECC) and ribosomes. MRB1 controls editing by still unknown mechanisms. We performed the first next-generation sequencing study of native subcomplexes of MRB1, immunoselected via either RNA helicase 2 (REH2), that binds RNA and associates with unwinding activity, or MRB3010, that affects an early editing step. The particles contain either REH2 or MRB3010 but share the core GAP1 and other proteins detected by RNA photo-crosslinking. Analyses of the first editing blocks indicate an enrichment of several initiating gRNAs in the MRB3010-purified complex. Our data also indicate fast evolution of mRNA 3′ ends and strain-specific alternative 3′ editing within 3′ UTR or C-terminal protein-coding sequence that could impact mitochondrial physiology. Moreover, we found robust specific copurification of edited and pre-edited mRNAs, suggesting that these particles may bind both mRNA and gRNA editing substrates. We propose that multiple subcomplexes of MRB1 with different RNA/protein composition serve as a scaffold for specific assembly of editing substrates and RECC, thereby forming the editing holoenzyme. The MRB3010-subcomplex may promote early editing through its preferential recruitment of initiating gRNAs.

Keywords: RNA editing; deep RNA sequencing; guide RNA; mitochondrial RNA-binding complex MRB1; Trypanosoma brucei

INTRODUCTION
Kinetoplastid protozoa include medically relevant species of Trypanosoma and Leishmania with life cycle stages that experience dramatic adaptations to host and changes in energy metabolism (Vickerman 1985). A unique mitochondrial process of RNA editing by uridylylate insertion and deletion is directed by guide RNAs (gRNAs) (Blum et al. 1990; Blum and Simpson 1990) and controlled in mRNA-specific and stage-specific manners by mechanisms that remain fundamentally unresolved (Stuart et al. 1997). The mitochondrial genome of Trypanosoma brucei (kinetoplast or kDNA) consists of several copies of a maxi-circle (~23 kb) and thousands of minicircles (~1 kb). Twelve out of 18 mRNAs encoded by the maxi-circle are edited. Minicircles encode most gRNAs, estimated at ~1200 on their “sense” strand (Simpson et al. 2000; Ochsenreiter et al. 2007). gRNAs are primary transcripts with three regions: a short 5′ anchor that anneals to pre-edited or edited mRNA next to the sequence to be edited, a guide sequence with complementarity to edited mRNA (editing block), and a ~10–15U 3′ tail added post-transcriptionally. An mRNA editing domain usually spans a set of juxtaposing and often overlapping editing blocks.

While the basics of the editing reaction catalyzed by RNA editing core complexes (RECCs) is well characterized, the regulatory aspects remain largely unknown (Madison-Anteuccci et al. 2002; Simpson et al. 2004; Carnes and Stuart 2007; Cruz-Reyes and Read 2013). Editing domains in mRNAs mature from 3′ to 5′ in small blocks, each directed
Differential gRNA partition in MRB subcomplexes

by a gRNA. Most mRNAs are extensively edited, while a few require limited editing or are never edited (Stuart et al. 1997). How a functional cognate substrate reaches the RECC is currently unknown. Accessory factors may facilitate selective binding of editing substrates, annealing, unwinding, and chaperone activities to stabilize precise base-pairing between the target mRNA and the gRNAs directing the U-insertions and U-deletions. Consistent with this idea, purified RECC was reported to be RNA-free and lacks the processivity and substrate specificity of editing in vivo (Rusche et al. 1997; Cruz-Reyes et al. 2001; Cifuentes-Rojas et al. 2005; Alatortsev et al. 2008; Carnes et al. 2011). Many non-RECC proteins impact editing (Aphasizhev and Aphasizheva 2011; Carnes et al. 2011; Cruz-Reyes and Read 2013; Hashimi et al. 2013). Some proteins preferentially affect a few mRNAs, while others affect a broader range. In a few cases, effects at initiation or during progression have been proposed, and RNA binding, annealing, and unwinding have been observed with recombinant protein or purified complexes (Ammerman et al. 2010; Hernandez et al. 2010; Kafkova et al. 2012). So, while much progress has been made, central long-standing questions remain perplexing, including the mechanisms of substrate recruitment, regulation of editing activity, and editing integration into mitochondrial RNA metabolism. The answer may involve a group of ribonucleoprotein particles of surprising complexity, known as the mitochondrial RNA-binding complex I (MRB1) (or gRNA binding complex, GRBC), which contain gRNA and a dynamic protein composition (Panigrahi et al. 2008; Weng et al. 2008; Hashimi et al. 2009; Hernandez et al. 2010; Ammerman et al. 2011). Subcomplexes of MRB1 (called here MRBs for simplicity) transiently associate via RNA linkers with several factors including RECCs, proteins that may affect processing and stability, and ribosomes. Our lab and others have proposed that these complexes may serve as “organizers” in the control of editing and its integration in mitochondria (Hernandez et al. 2010; Aphasizhev and Aphasizheva 2011; Cruz-Reyes and Read 2013; Hashimi et al. 2013). For example, the MRB3010 subunit (3010) affects an early editing step, whereas TbrGG2 impacts progression between blocks (Ammerman et al. 2010, 2011). We reported a helicase REH2 (H2) in an MRB that contains 3′-5′ unwinding activity. Copurification of REH2 with gRNA and unwinding activity is inhibited by mutation of its helicase or RNA binding domains (Hernandez et al. 2010). Importantly, RNAi-based repression of REH2 inhibits cell growth and RNA editing in trypanosomes (Hashimi et al. 2009; Hernandez et al. 2010; Madina et al. 2011).

We considered that MRBs of different protein composition could exhibit differences in their associated RNAs. We tested this hypothesis by performing the first next-generation sequencing (NGS) study of two native MRBs that contain either H2 or 3010. Analyses of the first editing blocks showed that the 3010-associated MRB is enriched in initiating gRNAs. We also found specific copurification of MRBs with mRNAs that undergo editing. Interestingly, we identified differences in gRNA expression, including alternative editing of mRNA 3′ ends, between the Lister strain 427 used here, EATRO 164 cells in a recent NGS study of total gRNA in procyclics (Koslowsky et al. 2013), and currently annotated sequences. This adds a level of complexity to the potential of alternative editing first observed by the Hajduk lab (Ochsnerreiter et al. 2008). Based on cumulative observations in our lab and others, we propose a model whereby subcomplexes of MRB1 with distinct RNA/protein composition serve as scaffolds that recruit editing substrates and route them into transiently associated RECCs. A dynamic higher-order MRB1 complex, formed by several subcomplexes with specialized roles, may provide the necessary context for concerted substrate selection or usage during editing initiation and progression.

RESULTS

Native subcomplexes of MRB1 containing either REH2 or MRB3010

MRB1 ribonucleoprotein complexes were previously isolated from procyclic trypanosomes expressing tagged copies of REH2 or MRB3010 (3010), and their protein composition examined in mass spectrometric studies (Hernandez et al. 2010; Ammerman et al. 2011), but the RNA composition of these complexes is not known. Furthermore, the large tag in the constructs or ectopic expression could induce unintended effects on the protein and RNA composition of MRBs. The current study characterized immunoprecipitated (IP) native MRBs from mitochondrial extracts using affinity-purified peptide polyclera against H2 or 3010 (Fig. 1A; Supplemental Fig. S1). Helicase H2 is often partially fragmented in extracts (Hernandez et al. 2010), whereas 3010 migrates slightly below IgG on SDS-PAGE. Interestingly, 3010 was not detected in Western analyses of native H2 IPs, whereas H2 is not detected or is barely visible in native 3010 IPs. We reproduced this observation multiple times using mitochondrial extract or whole-cell lysate in the IP s (e.g., Supplemental Fig. S2). Consistent with our analysis of native MRBs, H2 was not detected in a previous affinity-purification of tagged MRB3010 (Ammerman et al. 2011). So, loss of H2 in that study was not caused by the large tag in 3010 but instead is an intrinsic feature of native 3010-MRB.

A transient association between RECC and H2-purified MRB was previously observed (Hernandez et al. 2010). RECC subunits were almost undetectable with different antibodies in the current native H2 and 3010 purifications (Fig. 1B; data not shown). However, editing ligases can be radiolabeled by a sensitive auto-adenylation assay indicating substoichiometric levels of RECC (data not shown; Hernandez et al. 2010). As expected, both natively purified MRBs contain gRNA (Fig. 1C) and the proposed core GAP1 subunit (Fig. 1D), consistent with previous purifications of REH2 and MRB3010 (Hernandez et al. 2010; Ammerman et al. 2011).
Only endogenous gRNA has been reported to bind purified MRB1 complexes; however, several subunits have domains that suggest a role in RNA biology (Panigrahi et al. 2008; Weng et al. 2008). For example, our lab reported that H2 crosslinks with an mRNA editing site bearing a photo-reactive group (Hernandez et al. 2010). Importantly, immunopurified H2 and 3010 MRBs produced similar crosslinking patterns with this substrate (Fig. 1E), except for the crosslinks induced by H2 itself. The H2 crosslinks were confirmed by their mobility shift in a direct comparison of tagged vs. native H2 (Fig. 1F). This is the first time purified MRBs are directly compared by crosslinking, and the analysis suggests that H2 and 3010 MRBs share several RNA-binding proteins. Most of these crosslinks were not detected in the mock IP. Together, these data describe two native subcomplexes of MRB1 that stably copurify with gRNA, the core GAP1 subunit, and several common RNA-crosslinking proteins. However, these MRBs were readily distinguished by the presence of H2 or 3010. This result prompted us to investigate the RNA composition of our purified samples.

Native H2-MRB and 3010-MRB subcomplexes differ in gRNA composition

To address the possibility that MRBs of different protein composition may also differ in their gRNA complement, we compared the gRNA pools from H2, 3010, and mock IPs, and from procyclic (PF) parasites using next-generation sequencing. Barcoded Illumina libraries made in parallel with comparable amounts of gRNA scored in a capping assay (see Supplemental Fig. S3 and the Materials and Methods section) were multiplexed, and reads were aligned to edited mRNA sequences deposited at the KISS bioinformatics site (http://splicer.unibe.ch/kiss/) (Ochsenreiter et al. 2007). We filtered out fragmented RNA by scoring only gRNAs bearing a 3’ U-tail. Furthermore, because 3010 may impact editing at an early step (Ammerman et al. 2011), we focused on the first editing blocks of mRNAs (Fig. 2). In line with a recent characterization of the gRNA pool in PF cells (Koslowsky et al. 2013), we found that initiating gRNAs for several mRNAs occur in relatively low abundance. Because our libraries used limiting gRNA amounts extracted from the IPs, we only found initiating gRNAs for six mRNAs. The following analyses compare cumulative nucleotide frequency plots (NFPs) of edited mRNA coverage (Fig. 2) and individual frequency values and the ratio of major gRNAs in editing blocks 1 and 2 (B1 and B2, respectively) (Table 1). Some gRNAs predicted alternative editing sequences to those currently annotated (termed B1.alt or B2.alt) and are further discussed in Figure 3. Negligible gRNA levels in the mock IP library confirmed the specificity of our MRB libraries (Table 1).

**Cytochrome B (Cyb)**

mRNA Cyb has a small editing domain (~50 nt) with only two blocks (Fig. 2A). Notably, the coverage of the B1 and B2 blocks was similar in the parasites but different between MRBs. Relative to the B2 gRNA in each library, the initiating B1 gRNA appears enriched in 3010-MRB. A higher coverage of the B2 editing block in the H2-MRB libraries relative to the B1 block also suggests intrinsic differences between native MRBs. Table 1 shows the actual values and ratio of major gRNAs in these blocks (B1/B2), which is a simple parameter to evaluate relative B1 enrichment in MRBs independent of loading.

**NADH dehydrogenase subunits 7 and 8 (ND7 and ND8)**

mRNA ND7 has two separate editing domains with initiating gRNAs that were enriched in 3010-MRB (Fig. 2B,C). Interestingly, the 5’ domain has two possible initiating gRNAs, whose guide sequences have different 5’ ends but the same 3’ end: B1a and B1b (Fig. 2B). The relative values of B1b and B2 gRNAs suggested a preferential binding of B1b gRNA in 3010-MRB. In contrast, the B1a gRNA was too low in the MRB libraries and absent in the PF library (Fig. 2B; Table 1). As in mRNA Cyb, the H2-MRB library exhibited a higher coverage of the B2 gRNA for the 5’ domain. In the ND7 3’ domain (Fig. 2C), the B1 gRNA was dramatically increased in 3010-MRB. In contrast, the B2 gRNA in this domain was too low to quantify in the MRB libraries. In mRNA ND8 (Fig. 2D), we found an initiating gRNA but...
A relatively high value of the initiating gRNA in 3010-MRB when compared to upstream gRNAs also suggested a specific increased abundance in this complex. B1/B2 determinations assuming a value of ≤ 1 for nondetected gRNAs indicated B1 enrichment in 3010-MRB for ND8 and both ND7 domains, especially in the 3′ domain.

Ribosomal protein subunit 12 (RPS12), cytochrome oxidase 3 (CO3), and ATPase subunit 6 (A6)

Our libraries contain possible initiating gRNA for all three mRNAs RPS12, CO3, and A6 (Fig. 2E–G), whereas a B2 gRNA was missing for A6. In mRNA CO3, both B1 and B2 gRNAs were increased in 3010-MRB. However, the B1/B2 ratio for mRNA RPS12 suggested a preferential increase of B1 gRNA in 3010-MRB (Table 1). In A6 mRNA, we found an initiating gRNA in the MRB libraries but no B2 gRNA in any of the libraries. Nevertheless, the B1 gRNA was too low in our libraries to be examined quantitatively.

As a complementary approach to our above NGS studies, we performed Northern blots of B1 gRNAs for the ND7 3′ domain and A6 mRNA from direct IP samples of MRBs (Fig. 2H,I). Consistent with our findings in the libraries, the level of B1 gRNA in the ND7 3′ domain was much higher in the 3010 IP than in the H2 IP. The B1.alt gRNA in A6 mRNA was slightly higher in the 3010 than in the H2 IP, possibly reflecting a moderate but reproducible difference in capping signal in our unadjusted samples from direct IPs (see Supplemental Fig. S3 and the preparation of libraries in the Materials and Methods section).

Collectively, these data showed important differences between our MRB libraries in coverage NFPs of the first editing blocks. Relative comparisons of gRNAs (B1/B2) in the same library were consistent with an enhanced association of initiating gRNAs for mRNAs CYb, ND7 (both domains), ND8, and RPS12 in 3010-MRB. Furthermore, evidently not a B2 gRNA in the libraries. However, a relatively high value of the initiating gRNA in 3010-MRB when compared to upstream gRNAs also suggested a specific increased abundance in this complex. B1/B2 determinations assuming a value of ≤ 1 for nondetected gRNAs indicated B1 enrichment in 3010-MRB for ND8 and both ND7 domains, especially in the 3′ domain.

**FIGURE 2.** Analysis of edited mRNA coverage by gRNAs in Illumina libraries. (A–G) Steady-state gRNA from PF parasites and gRNA in immunopurified MRBs are annotated in blocks of edited sequence, each directed by a gRNA. Cumulative standard and G·U pairs between edited mRNA and gRNAs are scored in nucleotide frequency plots (NFPs Log2), including initiating gRNAs (i.e., block 1 or B1) and major upstream gRNAs (B2 or B3) in our libraries. Some gRNAs may guide alternative editing (e.g., B1.alt and B2.alt), and the ND7 5′ domain uses at least two similar initiating gRNAs (B1a and B1b). The entire editing domain in cytochrome B (CYb) and the 3′ terminus of other domains are plotted. The sequence of CYb (including all 34 U-insertions as lowercase t’s) and the ND7 5′ domain are shown. Equal protein loads were applied to the IPs, and the gRNA was gel-isolated and extracted. gRNA from IPs and total RNA were adjusted to apply comparable amounts in the libraries (e.g., Supplemental Fig. S3; see Materials and Methods). gRNA from the IPs was directly ligated to the adapters, whereas total gRNA from PF parasites was first treated with 5′ monophosphate specific Terminator exonuclease, which degrades rRNA but not 5′ triphosphate gRNA ends. (H,I) Northern blots of select initiating gRNAs from IPs and PF cells. The blot was hybridized with the A6 gRNA probe, stripped, and re-used with the ND7 gRNA probe. 15% UREA-PAGE was run as in Figure 1C.
different patterns in the coverage NFPs between procyclic and MRB libraries also indicated differential gRNA binding by MRBs. An increased selectivity of B1 gRNA by 3010-MRB implies that this complex may be more active in editing initiation than H2-MRB. Interestingly, the mRNAs ND7 (3′ domain) and ND8 exhibited the largest relative accumulation of initiating gRNA.

Alternative editing at mRNA 3′ ends and gRNA divergence in *T. brucei* strains

Some gRNAs in Figure 2 suggest alternative editing patterns compared to annotated edited mRNAs from studies in the early 1990s. All cases of alternative editing discussed here significantly extend the length and quality of the predicted duplex between guide sequence and edited mRNA. Importantly, the observed mismatches with annotated sequence were outside the gRNA anchor. Some guide sequences were well conserved between Lister 427 studied here and EATRO 164 cells in a recent study by Koslowsky et al. (Fig. 3), but others exhibited important strain-specific differences, consistent with a rapid evolution of minicircles. This section compares annotated and predicted alternative editing directed by B1 and B2 gRNAs in our libraries and examines the conservation of guide sequences between strains. Some gRNAs with very similar or identical guide sequences in both strains often exhibit highly diverged nonguiding termini.

**Cyb editing domain**

The B1 and B2 gRNAs in Lister 427 were identical to those reported by Koslowsky et al., including a 5′ terminal run of As (Fig. 3A). As indicated by that lab, an initiating gRNA, gCyb[560A] from an earlier study in EATRO 164 cells (Koslowsky et al. 1992) has the same guide sequence but a very different 5′ end. This gRNA was not present in their recent library from EATRO 164 cells or ours reported here.

**ND7 3′ editing domain**

The guide sequence of the B2.alt gRNA formed one mismatch (Fig. 3B[1]). However, simple alternative editing (one U-insertion and one U-deletion) would allow annealing with most of the gRNA (Fig. 3B[2]). Notably, this would also substitute two of the three C-terminal amino acids without changing the coding frame. The guiding sequence of B2.alt is nearly identical in Lister 427 and EATRO 164 cells (Fig. 3B[3]).

**ND7 5′ editing domain**

As mentioned above, this domain had two initiating gRNAs (B1a and B1b) (Fig. 3C[1]). The B1a gRNA was not found in the EATRO 164 library, but the B1b gRNA was the same in Lister 427 and EATRO 164 cells. An additional initiating gRNA, gND7(147–199), in EATRO 164 cells was not present in our libraries. These three gRNAs shared significant homology (Fig. 3C[2]) including a 7-nt identity at their 5′ termini. Based on this 5′ conservation, Koslowsky et al. suggests that gND7(147–199) could form an alternative anchor duplex (i.e., the same anchor by the B1a gRNA). However, the resulting alternative editing would introduce two A:C mismatches, decreasing the quality of the duplex (Koslowsky et al. 2013).

**CO3 editing domain**

The B1.alt gRNA in our libraries (Fig. 3D[1]) was not seen in the EATRO 164 library, and vice versa, we did not find the initiating gRNA in Koslowsky’s study. Alternative editing by the B1.alt gRNA would substantially modify the 3′ UTR sequence (Fig. 3D[2]). Coincidently, both the B1.alt gRNA

### Table 1. Relative amounts of B1 and B2 gRNAs

|        | PF | H2-IP | 3010-IP | Mock-IP |          | PF  | H2-IP | 3010-IP | Mock-IP | H2-IP | 3010-IP |
|--------|----|-------|---------|---------|-----------|----|-------|---------|---------|-------|---------|
| Cyb    | 8  | 1     | 8       |         |           |    | 10    | 5       | 3       | 1     | 0.2     |
| ND7 5′ dom | 1**| 1**   | 5**     |         |           |    | 494   | 180     | 82      | 8     | 0.006   |
| ND7 3′ dom | 60 | 179   | 2592    |         |           |    | 8     | 1       | nd      | nd    | 179     |
| ND8    | 3  | 9     | 147     |         |           |    | nd    | nd      | nd      | nd    | ≥29     |
| RPS12  | 3  | 2     | 4       |         |           |    | 3     | 4       | nd      | nd    | ≥26     |
| CO3    | 28 | 3     | 21      |         |           |    | 801   | 565     | 5208    | 22    | 0.005   |
| A6     | 0  | 1     | 1       |         |           |    | nd    | nd      | nd      | nd    | ≥1      |
| ND7 5′ dom | 0*| 1     | 1*      |         |           |    | 0     | 1       | nd      | nd    | ≥1      |

Frequency of a gRNA is defined as the value of the dominant 5′ end in a block. The 5′ end of guide sequences is generally more conserved than the 3′ end. (*) and (**) distinguish two analog initiating gRNAs (B1a and B1b, respectively) with a different 5′ end but the same 3′ end. A block may include one gRNA or multiple redundant gRNAs of equivalent guide sequence with conserved nucleotide polymorphisms. Redundant gRNAs may reflect genetic drift of a minicircle or homolog genes in different minicircles. (nd) Not detected. Calculations of B1/B2 assume “nd” to be ≤1.
FIGURE 3. (Legend on next page)
and both CYb gRNAs contain a 5′ A-run. The B2 gRNA, gCO3(935–977), was the same in both strains.

**ND8 editing domain**

The B1.alt gRNA formed a relatively short continuous duplex with annotated edited sequence (Fig. 3E[1]). However, alternative editing by a single U-insertion in the 3′ UTR extended the duplex dramatically (Fig. 3E[2]). This gRNA was similar to gND8(554–598) in EATRO 164 cells, except that the B1.alt gRNA in Lister 427 cells had one additional internal guiding “A” (see arrow, Fig. 3E[3]) that causes the alternative editing.

**RPS12 editing domain**

The guide sequence of the B1 gRNA (Fig. 3F[1]) was the same in Lister 427 and EATRO 164 cells. Alternative editing by the B2.alt gRNA would shorten the encoded C-terminus and substitute its last four amino acids (Fig. 3F[2]). The B2.alt gRNA was not seen in the EATRO 164 library, and we did not find the gRPS12(267–322) reported in the study by Koslowsky et al. (2013). However, these two gRNAs exhibit partial homology and have identical 5′ anchor sequences (Fig. 3F[3]). Interestingly, gRPS12(267–322) in EATRO 164 introduces an internal A:C mismatch that does not predict alternative edits (this “A” is noted by an arrow).

**A6 editing domain**

The B1.alt gRNA, gA6(774–822), formed two mismatches with annotated edited mRNA (Fig. 3G[1]) and is present in EATRO 164 cells. Alternative editing by the B2.alt gRNA would shorten the encoded C-terminus and substitute its last four amino acids (Fig. 3G[2]). As proposed by Koslowsky et al. (2013), allowed almost full annealing of the gRNA. This gRNA was almost identical in both strains (Fig. 3G[3]). Koslowsky et al. reported a second initiating gRNA gA6(770–822), a low abundance transcript not detected in our libraries.

**Native H2 and 3010 MRBs specifically copurify with pre-edited and edited mRNAs**

MRB1 complexes are only known to associate with gRNA (Weng et al. 2008; Hashimi et al. 2009; Hernandez et al. 2010). Because protein subunits of our immunopurified MRB samples including H2 photo-crosslinked with a synthetic mRNA (Fig. 1E, F), we considered that H2-MRB, but possibly also 3010-MRB, purifications could contain endogenous mRNAs. Mitochondrial mRNAs may associate with MRBs or with transiently bound factors including RECC or mitoribosomes (Weng et al. 2008; Hernandez et al. 2010; Aphasizheva et al. 2011). Initial primer extension assays of purified H2-MRB and 3010-MRB detected some mRNA ND7 but no other mRNAs (data not shown). Subsequent qPCR assays of purified MRBs and a mock IP revealed a robust enrichment of edited and pre-edited mRNAs (ND7, CYb, A6, and Murf2) relative to never-edited mRNA (COI), nuclear 18S rRNA, and cytosolic tubulin (Fig. 4A). Housekeeping transcripts, usually carried over during purification, served as a reference in our assays. Because editing progresses in a 3′-5′ direction and the amplicons (~50–100 bp) score 5′ edited (or pre-edited) targets, the corresponding downstream sequences are presumed to be correctly edited. Interestingly, both the relative abundance and ratios of edited and pre-edited targets differ substantially between MRBs. Also, while all tested edited and pre-edited mRNAs were significantly more enriched than COI, the trend appeared somewhat more consistent in edited transcripts. Relative to the mock IP, edited ND7 was enriched several thousandfold, while edited CYb was enriched only about a hundredfold. In contrast, COI (never-edited) and CYb and A6 mRNAs (edited and pre-edited) exhibit comparable steady-state levels (Fig. 4B). Although, edited ND7 was about one-thousandfold more abundant than other mRNAs at steady-state (i.e., it exhibits the lowest dCq), this difference was still several times smaller than the enrichment fold of edited ND7 in the

![Figure 4](image-url)
3010-MRB sample. Should the mRNAs described above be largely bound to mitoribosomes contaminating our samples, never-edited mRNAs would be enriched, and pre-edited mRNAs should not be present (Aphasizheva et al. 2011). Altogether, these data suggest a selective copurification of MRBs with mRNAs that undergo editing, relative to never-edited COI and housekeeping transcripts.

**DISCUSSION**

MRB1 complexes, isolated via epitope tags and specific antibodies, exhibit overlapping protein composition that has been discussed (Hernandez et al. 2010; Ammerman et al. 2011, 2012; Cruz-Reyes and Read 2013; Hashimi et al. 2013). However, the RNA component of these particles was unknown. Our current study of native MRBs with different protein composition indicated that these particles also differ in RNA composition. We distinguished two MRBs that contained either REH2 or MRB3010 subunits. Based on this work and our previous study (Hernandez et al. 2010), we propose that REH2 forms an MRB subcomplex that contains gRNA, unwinding activity, and several subunits that can be photo-crosslinked to RNA. REH2-associated unwinding may remodel RNA-protein interactions needed for efficient editing substrate association with REH2-MRB (Hernandez et al. 2010; V Kumar, B Madina, and J Cruz-Reyes, unpubl.). Whether or not this activity affects editing substrate binding to other MRBs needs to be examined. REH2-dependent unwinding could also control global intra-strand mRNA structure or gRNA exchange during editing progression, as it was proposed for TbRGG2 and REH1, respectively (Ammerman et al. 2010; Li et al. 2011). It was proposed that RNAi-repression of REH2 reduces gRNA stability (Hashimi et al. 2009). We also saw some gRNA reduction at Day 6 of REH2 repression (Hernandez et al. 2010 and Fig. 5D therein). However, this may be an indirect result of the late time point or a small but specific impact on metabolically stable gRNA. Revised analyses by us and by another lab indicate that robust REH2 depletion at Day 4 of RNAi does not significantly impact the gRNA steady-state level (Supplemental Fig. S4; R Aphasizhev, pers. comm.). MRB3010 may not bind gRNA but affects an unidentified early editing step (Ammerman et al. 2011). The REH2 and 3010 subcomplexes examined here contain the core GAP1 subunit. GAP1/2 homologs are known to bind and stabilize gRNA (Weng et al. 2008). Interestingly, native REH2-MRB and 3010-MRB may share multiple RNA crosslinking subunits, suggesting a common RNA-binding core. So, REH2-MRB, 3010-MRB, and other MRB subcomplexes may transiently associate in higher-order MRB1 complexes that assemble editing substrates and RECCs creating an editing holoenzyme (Fig. 5; Simpson et al. 2010; Cruz-Reyes and Read 2013; Hashimi et al. 2013), and dynamic contacts between the individual holoenzyme components may modulate the editing process, as well as productive interactions with mitoribosomes and other mitochondrial factors.

This study introduces NFPs (nucleotide frequency plots) of gRNAs, a simple but powerful tool that allows direct quantitative comparisons of gRNA content in purified complexes and total mitochondrial RNA. These NFP analyses provide (1) annotation of specific gRNAs at base resolution within editing domains, and (2) a measure of cumulative guide RNA potential at single and overlapping editing blocks in the parasite and purified complexes. This information is important because multiple redundant gRNA genes from polymorphic or entirely different minicircles often contribute to editing of the same block. Consistent with a role of the MRB3010 subunit in editing initiation, our NGS study of the first editing blocks in several mRNAs indicated a higher accumulation of initiating gRNAs in 3010-MRB than in H2-MRB. We note that similar amounts of gRNA (scored by capping of 5′ triphosphate ends) from IPs of REH2 and MRB3010 were used in the construction of Illumina libraries. This was largely confirmed given that counts go up in some gRNAs and down in others between the libraries. It is
possible that MRB3010 promotes specific recruitment or retention of accumulated initiating gRNAs in its subcomplex. A selective accumulation of initiating gRNA was especially clear for ND7 and ND8 but was also clear for other mRNAs. A partitioning of initiating gRNAs between MRBs, together with the observed low steady-state level of several initiating gRNAs in the parasite, both in our study and a recent report (Koslowsky et al. 2013), suggest that recruitment or usage of these particular gRNAs is controlled in vivo. Currently, we are working to determine the impact of initiating gRNA enrichtment on the target mRNAs. Interestingly, the study by Koslowsky did not find one or more gRNAs for half the mRNA substrates, implying that rare gRNAs may still editing progression.

Purified MRBs have only been reported to contain gRNA; however, we found between ten- and several thousand-fold specific enrichment of edited and pre-edited mRNAs in immunoselected MRBs relative to a mock IP. Housekeeping 18S rRNA and tubulin mRNA were not enriched, and the never-edited mRNA COI may be slightly increased in the H2-MRB sample. MRB subcomplexes may specifically bind edited and pre-edited mRNAs in addition to gRNA, because RECCs are stoichiometric in purifications of MRB1 (Fig. 1B; Fisk et al. 2008; Weng et al. 2008; Hernandez et al. 2010), and natively purified RECC was reported to be largely devoid of RNA (Rusche et al. 1997). In contrast to our purifications of native MRBs, mitoribosomes associate with never-edited, but not with pre-edited, mRNAs (Aphasizheva et al. 2011). So, assembled MRB1 complexes may store and route edited mRNAs from RECCs into ribosomes (Fig. 5).

REH2 and the GAP1/2 homologs are the only known RNA-crosslinking subunits in the context of purified MRBs, but other RNA-binding proteins and their cognate RNA targets in vivo need to be identified. The mRNA probe in our crosslinking assays may or may not reflect the cognate target’s specificity (Hernandez et al. 2010), and REH2 may potentially bind gRNA, mRNA, or both. Recombinant versions of TbRGG2 and the paralogs MRB8170 and MRB4160 were reported to preferentially crosslink in vitro with synthetic sequences resembling cognate mitochondrial mRNA (Fisk et al. 2008; Kafkova et al. 2012). Strong GAP1/2-dependent stability of gRNA in vivo and preferential crosslinking of a synthetic gRNA with GAPs in their purified MRB point to these transcripts as their natural target (Weng et al. 2008). The common ~100-kDa crosslink in native H2-MRB and 3010-MRB in our assays (Fig. 2A) may be induced by MRB8170, its paralog MRB4160, or both.

Several of the gRNAs, aligned to annotated mRNAs from early studies, predict alternative 3’ editing patterns within 3’ UTR or ORF regions that could impact mRNA stability, translation efficiency, or the encoded C-terminal amino acid sequence. Notably, our study in Lister 427 procyclic cells and recent analyses in a different strain (EATRO 164 cells) indicate important differences in minicircle content or expression, including strain-specific gRNAs, e.g., gRNAs detected in our study but not in Koslowsky’s. Continuing analysis of mRNA 3’ ends in our lab is consistent with our proposed alternative 3’ editing between strains (B Madina, V Kumar, and J Cruz-Reyes, unpubl.). The library of total gRNA examined by these authors achieved significantly higher depth than ours, as our preparations were limited by the gRNA amount extracted from purified MRBs. A high evolution rate of minicircles may create some variability in mitochondrial function among strains and thereby add adaptive potential in T. brucei. However, essential edits may be under strong selection pressure, as is illustrated by CYb gRNA genes in the same EATRO 164 cells used recently and in the early 1990s (Koslowsky et al. 1992, 2013) that have identical guide sequence but highly diverged nonguiding termini.

MATERIALS AND METHODS

Cell culture

T. brucei Lister strain 427 29-13 procyclic “PF” (tryps.rockefeller.edu) was grown axenically in log phase in SDM79 and harvested at a cell density of 1–3 × 10^7 cells/mL. A derived cell line expressing tetracycline-inducible TAP-REH2 was used as described (Hernandez et al. 2010).

Protein purification

Native MRB1 subcomplexes were immunoprecipitated from freshly prepared mitochondrial extracts (Harris and Hajduk 1992) using affinity-purified peptide antibodies raised in rabbit against REH2, MRB3010, and cytochrome oxidase 2 (mock) in T. brucei (Bethyl Laboratories, Inc), as we reported, with some changes (Hernandez et al. 2010). Briefly, specific antibodies of identical quality and concentration were conjugated to Protein A-Dynabeads (Life Technologies) that were pretreated with 5% BSA. Approximately 2 mg of mitochondrial extract was supplemented with 1× cOmplete Protease Inhibitor cocktail (Roche), SUPERase.In RNase inhibitor (Life Technologies), and precleared by passage over Protein A-Sepharose beads (GE Healthcare) before loading onto antibody-conjugated beads. Ectopically expressed TAP-REH2 was specifically immuno-purified using IgG-Dynabeads (Life Technologies) as reported (Hernandez et al. 2010). All washes were performed at 200 mM NaCl, 1 mM EDTA, 10 mM MgCl2, and 25 mM Tris, pH 8.

Western analyses and radioactive assays

Western blots were examined with rabbit affinity-purified anti-REH2 (Hernandez et al. 2010) and anti-MRB3010 polyclera, or rabbit anti-GAPI prebleed polyclera. Anti-MRB3010 antibodies were raised against the C-terminal peptide CPPLYQLYISRGSTPQA (Bethyl Laboratories, Inc), which is uniquely aligned to this protein in a Blast search of T. brucei. Monoclonal antibodies against REL1 ligase (a RECC subunit) were used as reported (Panigrahi et al. 2001). A 61-nt fragment of pre-edited A6 mRNA bearing a photo-reactive 4thio-U and ^32P at the first editing site was mixed with immunopurified MRBs and subjected to 365-nm UV irradiation on ice, as previously described (Sacharidou et al. 2006; Cruz-Reyes 2007). Radioactive capping of gRNA and adenylation of RECC...
ligases were performed as reported (Blum et al. 1990; Sabatini and Hajduk 1995). Northern blots of RNA extracted from IPs from whole-cell lysate with 5' labeled probes for gA6 B1.alt [1357]: CCACTGTAAMACTGATTTATTCATGGAG (Tm 57.9) and gND7 3' domain B1 [1358]: CGATATGAACTACGTGAGG ATTG (Tm 53.3) were performed in ULTRAhyb solution (Life Technologies) and 2× SSC washes at 40°C.

Illumina libraries of gRNA from purified MRBs and procyclic parasites

Identical immunoprecipitations were performed in parallel using REH2, 3010, and mock affinity-purified antibodies. The final washed beads were treated with proteinase K and SDS, and the associated RNA was treated with a DNA-free DNase kit and recovered by acid phenol-chloroform extraction and ethanol precipitation. RNA samples were treated with the ScriptCap capping system (CELLSCRIPT) and [α-32P]-GTP, and resolved on 15% UREA-PAGE to concentrate the gRNA in a narrow band. Size markers of gRNA mobility (~40–60 nt) were prepared with synthetic RNA fragments. Capped gRNA from IPs and total mitochondrial RNA were resolved in parallel with identical uncapped (unlabeled) samples (Supplemental Fig. S3). The labeled samples served as controls to adjust the amount of unlabeled gRNA used for library construction. We note that the total RNA sample was treated with Terminator 5’ Phosphate-Dependent Exonuclease (Epicentre) that degrades rRNA but not gRNA, as these transcripts carry a 5’ triphosphate (Blum et al. 1990). This step was not applied to the IPs because much nuclear rRNA and cytosolic mRNA were selected out in the sample IPs, relative to the mock IP, was calculated as follows:

Fold = 2^((-\Delta\Delta Cq) - 0.1) were averaged, and dilutions were adjusted to 100%.

Computational analyses. Read preprocessing

All base calls in the sequencing reads with a Phred score below 20 were converted into N’s and runs of trailing N’s were removed from the reads. Reads in which more than 10 N’s remained were discarded. Sequencing adapters were identified by requiring at least the first five bases of the adapter to be contained in the read and the sequence identity between the known adapter sequence and the part of the read containing the adapter to be at least 75%. Reads in which no adapter could be identified and reads with less than 10 bases after adapter removal were discarded. The bases immediately preceding the adapter were compared with the barcodes ATAC, ACCAA, CGAGA, and ATAGC, allowing at most one mismatch to distinguish reads originating from the procyclic, 3010, H2, and cytochrome oxidase 2 (mock) libraries, respectively, and the barcodes were removed. All reads with more than one mismatch to one of the four barcodes at the 3’ end were discarded.

Alignment to edited mRNAs

Reads from each library were separately aligned to the known edited mRNA sequences deposited at the KISS bioinformatics site (http://splicer.unibe.ch/kiss/). To this end, all reads with three or more consecutive N’s were discarded. Gapless semilocal alignments with a match score (including GU base-pairing) of 1 and a mismatch score of −2 were used to identify regions of local similarity between the reads and edited mRNA sequences. Reads with a local similarity score of at least 20 were parsed into a 3’ unmatched region, the matching region, a 3’ unmatched region, and a poly-U tail. These reads were post-filtered in order to only retain reads with at most two mismatches, a minimum length of the matching region of 25 bases, and a poly-U tail of at least three U’s. For RPS12, we also required that the length of the 5’ unmatched region was at most, 20 bases. For the remaining reads, we tallied the number of bases in matching regions covering each position in the edited mRNA.

Quantitative real-time PCR (qPCR)

RNA from the IPs and procyclic cells used in the preparation of Illumina libraries was used for cDNA synthesis, after treatment with a DNA-free DNase kit, in reactions with random hexamers in the iScript cDNA kit (Bio-Rad). Twenty microliters qPCR reactions were performed with reported primers specific for edited or pre-edited mRNAs, never-edited, and two reference transcripts, 18S rRNA and tubulin (Carnes and Stuart 2007), in a SsoFast EvaGreen Supermix (Bio-Rad). Diluted samples to score test transcripts (1:7) and reference transcripts (1:50) produced a single amplicon during linear amplification. All end-point amplicons described here were gel-isolated, cloned, and confirmed by sequencing. Fold-enrichment of mitochondrial and reference transcripts in the sample IPs, relative to the mock IP, was calculated as follows: Fold = 2^(-\Delta\Delta Cq), where \Delta\Delta Cq = Cq test IP – Cq mock IP. Steady-state mRNAs in mitochondrial extract relative to background tubulin and nuclear 18S rRNA (reference transcripts) were scored as follows: \Delta Cq = \Delta\Delta Cq – Ref Cq. Cq duplicates of each sample (STDEV ≤ 0.1) were averaged, and dilutions were adjusted to 100%.

DATA DEPOSITION

Deep sequencing data have been deposited in NCBI’s Sequence Read Archive under accession number SAMN02795843.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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