Structure of discontinuities in kinetoplast DNA-associated minicircles during S phase in *Crithidia fasciculata*

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ABSTRACT

Kinetoplast DNA (kDNA) is a novel form of mitochondrial DNA consisting of thousands of interlocked minicircles and 20–30 maxicircles. The minicircles replicate free of the kDNA network but nicks and gaps in the newly synthesized strands remain at the time of reattachment to the kDNA network. We show here that the steady-state population of replicated, network-associated minicircles only becomes repaired to the point of having nicks with a 3′ OH and 5′ deoxyribonucleoside monophosphate during S phase. These nicks represent the origin/terminus of the strand and occur within the replication origins (oriA and oriB) located 180° apart on the minicircle. Minicircles containing a new L strand have a single nick within either oriA or oriB but not in both origins in the same molecule. The discontinuously synthesized H strand contains single nicks within both oriA and oriB in the same molecule implying that discontinuities between the H-strand Okazaki fragments become repaired except for the fragments initiated within the two origins. Nicks in L and H strands at the origins persist throughout S phase and only become ligated as a prelude to network division. The failure to ligate these nicks until just prior to network division is not due to inappropriate termini for ligation.

INTRODUCTION

*Crithidia fasciculata* is a protozoan parasite related to the pathogenic human parasites *Trypanosoma brucei*, *Trypanosoma cruzi* and *Leishmania*. These latter organisms cause devastating diseases such as African sleeping sickness, Chagas disease and leishmaniasis. These protozoan parasites diverged early in the course of eukaryotic evolution and have many unique features of their biology including antigenic variation, RNA editing, unusual genome organization, novel mechanisms of regulating gene expression and an unusual form of mitochondrial DNA termed kinetoplast DNA or kDNA (1,2). The kDNA is a large network of interlocked circular DNA molecules condensed into a disk-like structure within the mitochondrial matrix close to the flagellar basal bodies (3,4). The kDNA is physically connected to the basal bodies through a transmembrane filament system known as the tripartite attachment complex or TAC (5–7). The kDNA network consists of two types of circular DNAs, minicircles and maxicircles. The maxicircles are the equivalent of usual mitochondrial DNAs and encode ribosomal RNAs and genes for oxidative metabolism (8,9). The minicircles apparently do not code for proteins but rather encode small RNAs called guide RNAs that are involved in the editing of maxicircle transcripts by a process involving the insertion and/or deletion of U residues in the transcripts (10–12).

The kDNA in *C. fasciculata* consists of 5000 to 10000 minicircles and 20–30 maxicircles. The maxicircles replicate by a theta mechanism while remaining attached to the kDNA network (13). On the other hand, the minicircles are released from the network sequentially as covalently closed circular molecules and are replicated in the mitochondrial matrix in a space between the kDNA disk and the flagellum termed the flagellar attachment zone or KFZ (14,15). Following their replication the minicircles are rejoined to the kDNA disk at antipodal sites by a DNA topoisomerase (16,17). At any given time during S phase only a few percent of the minicircles are free and undergoing replication. The newly reattached minicircles become distributed around the perimeter of the kDNA network apparently as a result of rotation of the kDNA disk relative to the antipodal attachment sites (18–20). The annular ring of newly replicated minicircles grows inward as replication proceeds and the central zone of unrepli- cated covalently closed minicircles shrinks in size until all minicircles have been replicated. At that point the

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minicircles within the double size network all contain discontinuities in the newly synthesized strands.

_Crithidia_ minicircles have two origins of replication (oriA and oriB) located 180° apart and each origin contains three sequence elements (CSB-1, CSB-2 and CSB-3) that are conserved in kinetoplast protozoa (21). The longest of these conserved sequences (12 nt) is CSB-3, also known as the universal minicircle sequence or UMS. Replication of the free minicircles is unidirectional with one strand (the L strand) synthesized continuously and one strand (the H strand) synthesized discontinuously (22–26). Analysis of newly replicated free minicircles in an isolated kinetoplast system showed that the minicircle L strand initiates from either of two replication origins (A and B) but not from both in the same minicircle strand. The 5′ terminus of the L strand is within the sequence complementary to the UMS and is usually a nick or a gap with a few ribonucleotides (23–25, 27). The daughter molecule containing the new L strand has a single discontinuity at either oriA or oriB and a covalently closed circular H strand. In contrast, the nascent free minicircles containing newly synthesized H strands have a covalently closed circular L strand and have multiple discontinuities between Okazaki fragments in the H strand that all become repaired except for discontinuities at both oriA and oriB. Amazingly, the discontinuities in the minicircles newly synthesized _in vivo_ are not repaired until all minicircles have been replicated and reattached, then only closed as a prelude to division of the network to yield two daughter networks in which all minicircles are covalently closed (18, 19, 28). In light of the presence of multiple DNA polymerases and DNA ligases within kinetoplasts (29, 30) it is a mystery why the discontinuities in network-associated minicircles remain unrepaired until all minicircles have been replicated. To begin to address this question, we have undertaken an analysis of network-associated minicircles during kinetoplast S phase in synchronized cultures of _C. fasciculata_.

**MATERIALS AND METHODS**

**Cell growth and synchronization**

_Crithidia fasciculata_ was grown in a defined medium (31) at 28°C on a rotary shaker and synchronized by release from a hydroxyurea block as described previously (32). The cell number was determined using a Z1 particle counter (Beckman-Coulter, Miami, FL, USA) or pJH621 (pGEM-11Zf(+) containing DNA from plasmid pGEM-11Zf(+) (Promega, Madison, WI, USA) or pJH621 (pGEM-11Zf(+) containing a cloned XhoI-digested _C. fasciculata_ minicircle). Hybridization was performed as described (33).

**kDNA isolation**

kDNA was isolated from 40 ml of synchronized cells at each 30 min interval as described (34).

**Minicircle markers**

A mixture of linear, covalently closed circular and nicked circular minicircles was prepared by partial ligation of XhoI linear minicircles released from pJH621 DNA. Each form was gel purified on 0.7% agarose gels for use as markers on neutral and alkaline gels.

**Agarose gels and Southern blotting**

Neutral agarose gels contained 0.7% agarose in 0.5 × Tris-borate-EDTA (TBE) buffer and 0.25 μg/ml ethidium bromide and were electrophoresed with a running buffer containing 0.5 × TBE buffer and 0.25 μg/ml ethidium bromide. Gels were transferred to Hybond N membranes (Amersham) and probed with a minicircle DNA probe that had been 32P-labeled by random priming of an XhoI-cut minicircle DNA. Alkaline agarose gels containing 1% agarose were soaked in 30 mM NaOH, 1 mM EDTA for 1 h before running in 30 mM NaOH, 1 mM EDTA. Alkaline agarose gels were run and transferred to Hybond N membranes essentially as described (35). Duplicate samples were run on the same gel and after transfer the membrane was cut apart to allow separate hybridization to the L- and H-strand-specific probes and to a molecular weight marker probe (32P-end labeled DNA Plus ladder, Invitrogen).

**Strand-specific DNA probes**

pJH621 was digested with Sal I and transcribed with SP6 RNA polymerase and 32P-labeled ribonucleoside triphosphates to produce a probe for minicircle L strands. Similarly, a probe for H strands was produced by transcription of BamHI-digested pJH621 with T7 RNA polymerase. The labeled RNA transcripts were digested...
with pancreatic DNase and purified over G25 MicroSpin columns before use in hybridizations.

**RESULTS**

**kDNA replication in synchronized cultures of *C. fasciculata***

The present experiments investigate the possible basis for the delay in covalent repair of discontinuities at the replication origins of kDNA-associated minicircles until every minicircle in the network has been replicated. In order to examine the structure of kDNA network-associated minicircles during kinetoplast S phase we have synchronized *C. fasciculata* by release from a hydroxyurea block. Synchrony was shown by counting dividing cells (cells with two nuclei) (Figure 1A) and by monitoring both total DNA synthesis and kinetoplast minicircle DNA synthesis (Figure 1B) at 30 min intervals. The cells go through the first S phase during the first 90 min following release from the hydroxyurea block. Dividing cells are frequent from 150 to 210 min and the second S phase occurs from 210 to 300 min. Trinitiated thymidine pulse labeling shows that total DNA synthesis, which is largely nuclear, and kDNA minicircle DNA synthesis occur in synchrony in the culture.

**Figure 1.** DNA synthesis in a synchronous culture of *C. fasciculata*. *Crithidia fasciculata* was synchronized by release from a hydroxyurea block and sampled at 30 min intervals to determine (A) the total cell number (filled circles) and the percentage of dividing cells (filled squares) and (B) the rate of total DNA synthesis (filled circles) and the rate of kDNA synthesis (filled squares).

Fortunately, *C. fasciculata* minicircles have a nearly homogenous sequence (36), unlike minicircles in most other kinetoplastid species that can have hundreds of sequence classes. Thus with a unique restriction map available it is possible to map discontinuities in the kinetoplast-associated minicircles (23,37). Figure 2A shows a physical map of the *C. fasciculata* minicircle and sites of specific nicks observed previously at the replication origins of free minicircles replicated in an isolated kinetoplast system (22). The location of a static bend in the DNA, termed a bent helix, located approximately between nucleotides 1041 and 1240 is also shown. We have mapped the sites of nicks in each of the strands of the kinetoplast-associated minicircles by strand-specific probing of Southern blots of XhoI-digested kDNA networks run on alkaline denaturing gels. Cleavage of covalently closed minicircles at the unique XhoI site prior to denaturation releases unit-length strands (2.5 kb) of each type (H and L) but molecules with specific discontinuities at the replication origins will produce a 2.5 kb fragment of the circular strand and two or more fragments of defined size from the newly synthesized strand (Figure 2B).

We have analyzed the strands released upon denaturation of kDNA network-associated minicircles isolated throughout kinetoplast S phase. Figure 3 shows a Southern blot of a denaturing gel analysis of duplicate samples of XhoI-digested kDNA taken at 30 min intervals spanning the second S phase. One set was probed to detect L strands and one to detect H strands. The 2.5 kb L-strand fragments correspond to linear strands derived by XhoI cleavage of either covalently closed (unreplicated) minicircles or from replicated minicircles in which the L strand was the parental covalently closed circle. Two pairs of fragments (2.0 + 0.49 and 1.7 + 0.79; Table 1 and Figure 3) are consistent with the strands having been derived from nicked or gapped minicircles with a single discontinuity at oriA or oriB. This result indicates that L-strand replication initiates at either oriA or oriB but not from both in the same molecule. In addition, we observed a ladder of catenated minicircles and a band labeled denatured covalently closed (dcc) that represents denatured covalently closed minicircles. Both of these species represent a minor class of minicircles that are resistant to cleavage by XhoI and will be addressed later.

In the case of fragments derived from the H strand, two pairs of fragments (2.1 + 0.38 and 1.6 + 0.84) are consistent with the strands having resulted from cleavage of minicircles in which there is a single discontinuity at either oriA or oriB. In addition, a more abundant species corresponding to 1.25 kb half-length molecules must have resulted from replicated minicircles with a new H strand in which discontinuities are present at both oriA and oriB. Thus, in these molecules all Okazaki fragments have been covalently joined except for the fragments initiated at each of the origins. Two additional minor H-strand fragments of ~1.0 and 0.65 kb are also observed. These minor species could possibly be derived from minicircles containing discontinuities at oriB and at a previously unknown site at either 1.0 or 0.65 kb clockwise from the XhoI site. It is unclear what significance a discontinuity at either site...
might have. A smear of fragments from ~1.2 to 0.2 kb possibly reflects the presence of H-strand Okazaki fragments that have not yet been ligated.

**Repair of discontinuities in replicated minicircles**

The lack of covalent closure of discontinuities at the origin/terminus of strands in replicated minicircles until just prior to network division could be the consequence of inappropriate termini for covalent closure, among other things. To address this possibility we have treated kDNA samples with *Escherichia coli* DNA ligase prior to XhoI digestion and alkaline denaturation of network-associated minicircles containing nicks at the replication origins. Fragment sizes are based on mapping at the nucleotide level of nicks in free form II minicircles replicated in an isolated kinetoplast system (23). The observed values are estimates based on gel migration relative to DNA markers in Figure 3 and in earlier experiments (22).

**Table 1.** Predicted and observed XhoI fragments

<table>
<thead>
<tr>
<th></th>
<th>XhoI fragments</th>
<th>Observed</th>
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<tbody>
<tr>
<td><strong>L strand nicks</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>oriA</td>
<td>471 + 2044</td>
<td>490 + 2000</td>
</tr>
<tr>
<td>oriB</td>
<td>781 + 1734</td>
<td>790 + 1700</td>
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<tr>
<td>oriA and oriB</td>
<td>471 + 781 + 1263</td>
<td>n.o.</td>
</tr>
<tr>
<td><strong>H strand nicks</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>oriA</td>
<td>379 + 2136</td>
<td>380 + 2100</td>
</tr>
<tr>
<td>oriB</td>
<td>874 + 1641</td>
<td>840 + 1600</td>
</tr>
<tr>
<td>oriA and oriB</td>
<td>379 + 874 + 1250</td>
<td>380 + 840 + 1250</td>
</tr>
</tbody>
</table>

Minicircle H and L single-strand fragments produced upon XhoI digestion and alkaline denaturation of network-associated minicircles containing nicks at the replication origins. Fragment sizes are based on mapping at the nucleotide level of nicks in free form II minicircles replicated in an isolated kinetoplast system (23). The observed values are estimates based on gel migration relative to DNA markers in Figure 3 and in earlier experiments (22).
digestion, alkaline gel analysis and Southern blotting. Unlike T4 DNA ligase, the \textit{E. coli} enzyme cannot join a DNA 3' OH terminus to a strand containing 5' ribonucleotides (38,39). Termini of the latter type require the removal of the ribonucleotides(s) and gap filling before they can be ligated by the \textit{E. coli} ligase. The \textit{E. coli} ligase can only join a simple nick in DNA with a 3' OH and 5' deoxynucleoside monophosphate. The disappearance of the specific fragments and the increase of the denatured covalently closed circles upon ligase treatment (Figure 4) indicate that the specific discontinuities in both the L and H strands were repaired by \textit{E. coli} DNA ligase. Any RNA primers must have been removed and gaps filled soon after their reattachment to the kDNA network. We conclude that the lack of covalent closure of the origin-specific nicks \textit{in vivo} in the network-associated minicircles is not due to inappropriate termini for ligation.

**Nature of the catenated XhoI-resistant minicircles**

To address the possibility that the ladder of slower migrating minicircles seen on alkaline gels might result from head-to-tail catenated molecules rather than from chains of topologically interlocked unit-size minicircles we have analyzed a kDNA sample on a neutral gel after treatment with XhoI and topoisomerase as shown in Figure 5. The disappearance of the catenated molecules upon XhoI and topoisomerase treatment as shown in Figure 5 indicates that these molecules represent catenated, XhoI-resistant minicircles each of unit size. The basis for the resistance of this class of minicircles to cleavage by XhoI was investigated by PCR amplification and cloning of a DNA sequence spanning the region of the XhoI site in gel-isolated catenated molecules. DNA sequence analysis of four independent clones shows that these minicircles contained a T to C substitution in the XhoI site as well as two other single nucleotide substitutions, a single nucleotide deletion adjacent to the mutant XhoI site and a 10 nt insertion elsewhere (Figure 6).

**DISCUSSION**

Many of the details of the replication mechanism of kinetoplast minicircles are now known, yet much less is known about the regulation of network growth and division. A key feature of the regulation of network division is the lack of closure of discontinuities at the minicircle replication origins until the time of network division (18,19). The studies presented here imply that gaps between Okazaki fragments, except within the replication origins, are largely fully repaired soon after attachment of nascent minicircles to the kDNA network. The remaining discontinuities at the replication origins in the replicated minicircles were found to be simple nicks, based on their susceptibility to closure by \textit{E. coli} DNA ligase, that remain unrepaired throughout S phase.
RNA primers and gaps between fragments. Binding at the sites of the specific nicks could prevent binds to native minicircle origin fragments in vitro specifically to CSB-3 and to the complement of CSB-1. UMSBP is a good candidate for such a role since it binds possibly the UMS-binding protein UMSBP, may prevent by E. coli. Additionally, the remaining Okazaki fragments are ligated isolated kDNA networks with those in the L strands, can be repaired by treatment of inappropriate termini for ligation since these nicks, like the lack of covalent closure of the nicks in the H-strand 5 origin halfway around the circular DNA and likewise the synthesized H strand must also occur at the second site-specific initiation of the discontinuously replicated minicircles (23).

Replication of the H strand of the minicircle is discontinuous and initiates within the CSB-1 sequence of oriA and oriB. Although discontinuities between H-strand Okazaki fragments must be largely repaired shortly after reattachment of the nascent minicircles to the network, discontinuities remain in the newly synthesized H strands at the two origins. This result suggests that the 5' terminus of the first Okazaki fragment is somehow different from that of subsequent Okazaki fragments since the Okazaki fragments become covalently closed whereas the 5' terminus of the first H-strand fragment remains unligated. The abundant existence of minicircles with H-strand discontinuities at both oriA and oriB indicates that a second site-specific initiation of the discontinuously synthesized H strand must also occur at the second origin halfway around the circular DNA and likewise the 5' terminus is prevented from being ligated. The reason for the lack of covalent closure of the nicks in the H-strand discontinuities at the origins also cannot be due to inappropriate termini for ligation since these nicks, like those in the L strands, can be repaired by treatment of isolated kDNA networks with E. coli DNA ligase. Additionally, the remaining Okazaki fragments are ligated by E. coli DNA ligase as well, indicating that they lack RNA primers and gaps between fragments.

We suggest that it is more likely that a bound protein, possibly the UMS-binding protein UMSBP, may prevent closure of the discontinuities at the replication origins. UMSBP is a good candidate for such a role since it binds specifically to CSB-3 and to the complement of CSB-1 minicircle sequences in their single-stranded forms and also binds to native minicircle origin fragments in vitro (40). Binding at the sites of the specific nicks could prevent access to the nicks by DNA ligases. In addition, binding of UMSBP to an oriA fragment was found to require both CSB-1 and CSB-3 sequences suggesting a cooperative binding of UMSBP at the two sequences within each replication origin. It is particularly of interest to note that UMSBP binding to origin sequences involves the binding of monomers through the C-terminal region of the protein and that the binding is sensitive to the redox potential (41). Oxidation of UMSBP leads to dimerization and a concomitant inhibition of binding to DNA. Thus, if UMSBP is bound to the minicircle replication origins and thereby preventing closure of the nicks in the replication origins, oxidation of UMSBP might result in the release of UMSBP allowing the covalent closure of the nicks and permitting progression to network division. Networks with accessible nicks could serve to recruit a DNA ligase to the nicks for their repair. Such a model is consistent with the observation in C. fasciculata that DNA LIG kx is observed to be associated with the kDNA only at the end of S phase when cells have begun to undergo cell division.

Finally, the observation of XhoI-resistant catenated minicircles has implications for the organization of kDNA networks. Since the XhoI-resistant minicircles are quite rare, it is remarkable that many of these molecules are catenated to one another. This observation suggests that minicircles of this minor sequence class are not distributed randomly throughout the kDNA network but instead are clustered together within the network. Asymmetric distribution of newly replicated daughter minicircles to the antipodal attachment sites might possibly contribute to such a distribution.

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Conflict of interest statement. None declared.
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