Changes in Organization of *Crithidia fasciculata* Kinetoplast DNA Replication Proteins during the Cell Cycle

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**Abstract.** Kinetoplast DNA (kDNA), the mitochondrial DNA in kinetoplastids, is a network containing several thousand topologically interlocked minicircles. We investigated cell cycle–dependent changes in the localization of kDNA replication enzymes by combining immunofluorescence with either hydroxyurea synchronization or incorporation of fluorescein–dUTP into the endogenous gaps of newly replicated minicircles. We found that while both topoisomerase II and DNA polymerase β colocalize in two antipodal sites flanking the kDNA during replication, they behave differently at other times. Polymerase β is not detected by immunofluorescence either during cell division or G1, but is abruptly detected in the antipodal sites at the onset of kDNA replication. In contrast, topoisomerase II is localized to sites at the network edge at all cell cycle stages; usually it is found in two antipodal sites, but during cytokinesis each postscission daughter network is associated with only a single site. During the subsequent G1, topoisomerase accumulates in a second localization site, forming the characteristic antipodal pattern. These data suggest that these sites at the network periphery are permanent components of the mitochondrial architecture that function in kDNA replication.

Key words: cell cycle • kinetoplast DNA • DNA replication • DNA polymerase β • topoisomerase II

**Trypanosomatid** protozoa such as *Crithidia fasciculata* possess an unusual mitochondrial genome known as kinetoplast DNA (kDNA) that is comprised of ~5,000 minicircles (2.5 kb) and ~25 maxicircles (38 kb) topologically interlocked into a single network. An isolated kDNA network is a planar structure ~10 μm × 15 μm in size. In vivo, however, the network is condensed into a disk ~1 μm in diam and 0.4 μm in thickness within the mitochondrial matrix (see Shapiro and Englund, 1995 for discussion of network packaging in vivo). Like mitochondrial DNAs in other eukaryotes, maxicircles encode rRNAs and proteins required for mitochondrial energy transduction (Stuart, 1983). Maxicircle transcripts undergo RNA editing, the specific insertion or deletion of uridine residues at internal sites. The sequence specificity of RNA editing is dictated by guide RNAs that are encoded by minicircles (Benne, 1994; Simpson and Thiemann, 1995).

The complex network structure of kDNA dictates a novel replication mechanism (reviewed in Ray, 1987; Shlomai, 1994; Shapiro and Englund, 1995). In this process, covalently closed minicircles are individually released from the network and replicated as free minicircles via theta structure intermediates. Newly synthesized minicircles, which contain gaps, are then reattached to the network periphery by a topoisomerase. Thus replicating networks develop two zones, an outer zone of gapped, progeny minicircles and an inner zone of covalently closed minicircles that have not yet replicated. Once replication is complete, the network has twice the number of minicircles and all are gapped. After repair of the gaps in each minicircle, the network undergoes scission (presumably in another topoisomerase-mediated reaction) to form two daughter networks.

The intramitochondrial localization of kDNA replication enzymes has been crucial for our current understanding of network replication. The mitochondrial DNA primase is detected in regions adjacent to the two faces of the kDNA disk, suggesting that early stages of minicircle replication may occur in these locations (Li and Englund, 1997). In contrast, the mitochondrial topoisomerase II (topo II) and DNA polymerase β (pol β) localize to two sites flanking the network edge (Melendy et al., 1988; Fer-

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1. Abbreviations used in this paper: kDNA, kinetoplast DNA; pol β, DNA polymerase β; topo II, topoisomerase II; DAPI, 4,6-diamidino-2-phenylindole; TdT, terminal deoxynucleotidyl transferase; dUTP-F, fluorescein-5(6)-carboxamidomethyl-[5-(3-aminoallyl)-2'-deoxy-uridine-5'-triphosphate; RP-A, replication protein A.
A similar antipodal localization pattern is also observed for minicircle replication intermediates detected by fluorescence in situ hybridization (Ferguson et al., 1992). Furthermore, newly replicated minicircles are known to reattach to the network periphery at two antipodal positions (Simpson and Simpson, 1976; Pérez-Morga and Englund, 1993a). The similar localizations of top II, pol β, minicircle replication intermediates, and minicircle reattachment sites led to the hypothesis that these sites were involved in minicircle replication, particularly in the repair of highly gapped minicircles and the re-attachment of minicircle progeny to the network (Melendy et al., 1988; Ferguson et al., 1992; Li and Englund, 1997).

In most eukaryotes, the replication of mitochondrial DNA is independent of the nuclear cell cycle. Mitochondrial DNA in mammalian cells can replicate at any cell cycle stage (Bogenhagen and Clayton, 1977). In fact, some circles replicate multiple times during a cell cycle and others do not at all, with only the final copy number being accurately controlled (Shadel and Clayton, 1997). In contrast, kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. Each minicircle and maxicircle in kDNA replication is more stringently regulated. 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a 10 min wash in 2× SSC (0.3 M NaCl, 30 mM Na₂citrate). Slides were then washed three times for 5 min in PBS before proceeding with immunofluorescence.

**Results**

**Colocalization of Pol β and Topo II**

We found, as previously reported, that in asynchronously-growing cells pol β and topo II have similar patterns of localization to two antipodal sites flanking the kDNA disk (Melendy et al., 1988; Ferguson et al., 1992). To further characterize the antipodal localizations of topo II and pol β, we used double-label immunofluorescence to probe both enzymes. We observed that pol β and topo II colocalized to the same sites at the network edge, as shown in Fig. 1. In addition, we observed that in some cells with topo II concentrated in these antipodal sites, a small amount of the enzyme is also detected either between these sites, in the region of the kDNA network (Fig. 2d), or in a region adjacent to the flagellar face of the network (data not shown).

**Localization of DNA Polymerase β throughout the Cell Cycle**

Our next objective was to localize the pol β at different stages of the cell cycle using cells synchronized with hydroxyurea. We followed the growth of synchronized cultures through two cell division cycles, measuring the rate of incorporation of [3H]thymidine into total cellular DNA, the percentage of dividing cells, and the density of cells in

**Figure 1.** Colocalization of pol β and topo II. Cells from an asynchronous culture were fixed and processed for immunofluorescence to detect both pol β and topo II as described in Materials and Methods. (a) DAPI visualization of the dimly stained nucleus and brightly staining kDNA. (b) Detection of pol β with goat anti-rabbit secondary antibody conjugated to Texas red (Molecular Probes, Inc.). (c) Detection of topo II with goat anti-mouse secondary antibody conjugated to fluorescein (Boehringer Mannheim Corp.). Images were captured with a Photometrics slow-scan CCD camera (Tucson, AZ) using IPLab software. Bar, 3 μm.

**Figure 2.** Localization of pol β and topo II in hydroxyurea synchronized C. fasciculata. (a) Cells were treated with hydroxyurea and released into fresh medium at 0 min. Samples were analyzed as described in Materials and Methods for cell division (closed circles), cell density (open triangles), and for the rate of DNA synthesis (open squares). (b) Cells from another synchronized culture were processed for immunofluorescence to detect pol β and topo II and stained with DAPI. The percentage of cells undergoing division (closed circles), percentage of cells with topo II detected in two antipodal sites (open squares), and percentage of cells with pol β detected in two antipodal sites (closed triangles) are shown. At least 200 randomly chosen cells per time point were scored. Results were confirmed three times, and data shown are from a representative experiment. (c and d) Localization of pol β (c) and topo II (d) in cells synchronized with hydroxyurea. The pol β images are from the synchronization in a, and those of topo II are from the experiment in b. Top row, cells during interphase, 210 min after release from hydroxyurea arrest; bottom row, cells during the peak of cell division, 120 min after release from hydroxyurea arrest. Control experiments showed that the weak signal covering the entire cell results from non-specific binding of the secondary antibody (data not shown). Images were captured with a Zeiss 35 mm camera. Bar, 3 μm.
the culture (Fig. 2a). After release from hydroxyurea arrest, the rate of [3H]thymidine incorporation (Fig. 2a, open squares) peaks at 30 and 180 min, each peak indicating an S phase. This incorporation is due to synthesis of both nuclear DNA (75% of total) and kDNA (25% of total) (Englund, 1979) that replicate in approximate synchrony (Cosgrove and Skeen, 1970). Sudden increases in the percentage of cells with two nuclei (Fig. 2a, closed circles) at 120 and 270 min mark passage of the culture through mitosis and indicate the timing of the subsequent cell division. Cytokinesis occurs fairly quickly, and thus the populations of dividing cells also contain some mitotic cells and daughter cells already in G1. This characterization of the *C. fasciculata* cell cycle agrees with data previously reported (Pasion et al., 1994) and is reproducible between experiments.

Using a synchronized culture, we found that detection of pol β in the antipodal sites is cell cycle dependent. The number of cells with this antipodal localization pattern peaks at 30–60 and 240–270 min, during interphase (Fig. 2b, closed triangles; c, upper panels). The percentage of cells with this localization pattern decreases during the peak of cell division; however, when only 7% show this localization (Fig. 2b). Unexpectedly, when not localized in the antipodal sites, the pol β is not detectable by immunofluorescence (Fig. 2c, lower panels).

Our inability to detect pol β immunofluorescence in the population of dividing cells raised the possibility that the enzyme abundance may change during the cell cycle. To address this issue, we performed Western blots on whole cell lysates taken at 30 min intervals during synchronized growth of a *C. fasciculata* culture. We found that the enzyme is present at similar levels in all time points (Fig. 3a). Thus while the pol β protein is continuously present, it is detectable by immunofluorescence only at certain cell cycle stages. Possible explanations of the inability to immunolocalize this enzyme will be addressed in Discussion.

**Localization of Topoisomerase II throughout the Cell Cycle**

To determine whether this cyclic localization pattern was a feature of all enzymes detected in the antipodal sites, we also examined topo II localization at different cell cycle stages, using the same culture as was used for pol β. In this experiment synchronized cells were scored for the presence or absence of topo II in both antipodal sites. During interphase most cells have topo II in both antipodal sites (Figs. 2b, open squares; d, upper panels), but the percentage of cells with topo II concentrated in these sites decreases during cell division (120 and 300–330 min). Thus the periodicity of topo II localization to both antipodal sites is similar to that of the pol β. Unlike pol β, however, topo II is detectable by immunofluorescence in virtually all cells at all stages of the cell cycle. When topo II is not detected in the antipodal sites, it is present either in a diffuse pattern near the kDNA (Fig. 2d, lower panels) or in other localization patterns that will be discussed below.

These experiments indicate that both topo II and pol β are periodically detected in the two antipodal sites flanking the network edge. There are significant differences,

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**Figure 4.** In situ analysis of kDNA replication in fixed cells from an asynchronous culture. dUTP-F was incorporated into endogenous minicircle gaps by TdT as described in Materials and Methods. Top row, dUTP-F fluorescence of kDNA networks. Bottom row, DAPI fluorescence. (a–c) Pre-replication networks. (d–i) Replicating networks. (j–m) Post-replication networks. (n) Two fully repaired, post scission networks, both in a cell undergoing cytokinesis. The kDNA networks in c and i are positioned vertically. Images were captured with a Photometrics slow-scan CCD camera using IPLab software. Networks in cells treated with proteinase K to reposition the network for a horizontal view are indicated with an asterisk. Treatment with proteinase K was modified from a previous method (Ferguson et al., 1992). Before either immunofluorescence or dUTP-F labeling, fixed cells were treated at room temperature for 15 min with 1 μg/ml Proteinase K in 10 mM Tris-HCl, 1 mM EDTA, 150 mM NaCl, and 0.1% SDS. Slides were then given three 5-min washes in PBS containing 1 mM PMSF, and one 5-min wash in PBS. Without protease treatment, ~8% of cells contain tipped networks. Treatment with protease increases the frequency of horizontally positioned networks to ~25%. Bar, 1 μm.
however, in the timing of these changes in enzyme localization (Fig. 2 b, pol β, closed triangles; topo II, open squares). For example, at the 120-min time point 30% of cells have topo II localized in both antipodal sites, while <10% have pol β in this location. In addition, after cell division the accumulation of topo II in the two antipodal sites precedes that of pol β (Fig. 2 b, see 180-min time point). To further evaluate these differences between topo II and pol β it was necessary to determine the timing of the changes in localization with greater precision. We therefore developed a new method to determine the kDNA replication status in individual, fixed cells from an asynchronous culture. This method, coupled with immunolocalization of the two enzymes, allowed a correlation of enzyme localization with the stage of kDNA replication in individual cells.

A Method for Determination of kDNA Replication Stage in Asynchronous Cells

We determined the kDNA replication stage by fluorescence microscopy of individual cells in which fluorescein-labeled dUTP (dUTP-F) had been incorporated in situ into the endogenous gaps in newly replicated minicircles by terminal deoxynucleotidyl transferase (TdT). This analysis was facilitated by the well-characterized distribution of covalently-closed and gapped minicircles in pre-replication, replicating, and post-replication kDNA in both isolated networks (Englund, 1978; Pérez-Morga and Englund, 1993; Guilbride and Englund, 1998) and for networks in vivo (Ferguson et al., 1992).

Fig. 4 shows images of the kinetoplast region of dUTP-F-labeled cells from an asynchronous culture. In vivo, the kDNA network is condensed into a disk positioned perpendicularly to the flagellum, and therefore fluorescent images usually provide a view of the edge of the disk, as in Fig. 4, e, g, and h. In fixed cells the kDNA disk is occasionally re-oriented, or “tipped,” providing a view of the face of the disk. The frequency of tipped networks can be increased by a mild protease treatment (examples in Fig. 4, e, g, and h) (Ferguson et al., 1992). Since images of a tipped disk are more informative about the degree of replication in a partially replicated network, we chose most examples in Fig. 4 with the disk tipped.

Pre-replication networks do not label with dUTP-F because the minicircles are all covalently closed (Fig. 4, a–c). In contrast, networks undergoing replication have a peripheral ring of dUTP-F label and a central zone of unlabeled minicircles (Fig. 4, e–g). kDNA at an early stage of replication has a narrow peripheral ring (Fig. 4 e) that increases in thickness during the replication process (Fig. 4, e–g). Replicating networks (Fig. 4, d–i) are always flanked by terminal dUTP-F label, while post-replication networks (Fig. 4, c–e) have truncated peripheral rings (Fig. 4 c). Cells undergoing cytokinesis (Fig. 4, h–i) show immunofluorescence on the face of the disk. Observation of the whole cell (Fig. 4, i) is usually informative as these cells have both rims of the disk labeled with dUTP-F.
by two antipodally positioned, bright dUTP-F spots. These bright spots colocalize with pol β and topo II (see below), and their intense labeling by dUTP-F is probably due to the presence of highly gapped, free minicircle replication intermediates (Johnson, 1998). Fig. 4 d shows a network flanked by antipodal spots of dUTP-F, but without the peripheral ring of fluorescence. This image likely represents the earliest stage of kDNA replication before a significant number of gapped minicircles have been attached to the network periphery. Consistent with this possibility, the dUTP-F fluorescence in Fig. 4 d was weak and required a longer (58 s) exposure time than other images in Fig. 4, e–l (1.5–4 s). Some replicating networks, while flanked by bright dUTP-F spots, appear to lack an unlabeled central zone (Fig. 4 h). In these cases, in the final stages of kDNA replication, the central unlabeled zone is probably too small to have been observed with fluorescence microscopy. Since the network in Fig. 4 i is positioned vertically it is impossible to determine whether it is early or late in the replication process, but it is characterized as replicating because of the antipodally positioned bright dUTP-F spots. Postreplication kDNA networks label uniformly with dUTP-F because all of the minicircles are gapped, and these networks are not flanked by bright dUTP-F spots (Fig. 4, j–l). Some postreplication networks appear irregularly shaped, often like a “V,” and do not lie in a single plane of focus (Fig. 4 l). Networks with this irregular structure may be beginning the network scission process.

Repair of the remaining minicircle gaps occurs before network scission (Pérez-Morga and Englund, 1993b) and results in reduction and ultimately elimination of dUTP-F fluorescence (Fig. 4 m is probably a partially repaired network). Finally, after network scission, the cells contain two kDNA networks that do not incorporate dUTP-F as all the minicircles are covalently closed (Fig. 4 n).

**Localization of Pol β in Asynchronously Growing Cells**

We used immunofluorescence to determine the location of pol β in cells that had been first labeled with dUTP-F to determine the stage of kDNA replication (see Table I and Fig. 5). In cells which had not initiated kDNA replication, we detected no pol β immunofluorescence (Fig. 5, a–b). In contrast, 100% of the cells undergoing kDNA replication had pol β localized in the two antipodal sites (Fig. 5, c–e) that co-localize with the two bright spots of dUTP-F fluorescence. After replication is complete, different patterns of pol β localization were observed. In 54% of these cells, pol β was found in the two antipodal sites (Fig. 5 f). At this stage of kDNA replication, some weak pol β signal is also detected in the kDNA region (Fig. 5 f). The rest of the post-replication networks either contain no detectable pol β immunofluorescence (33%, Fig. 5 h) or the enzyme is in other locations (13% of post-replication cells, see Fig. 5 g).

![Figure 6. Analysis of topo II localization in asynchronous cells labeled with dUTP-F.](image)
Finally, in dividing cells which have two daughter kDNA networks (following gap repair and network scission), none showed detectable pol β immunofluorescence (Fig. 5 i).

**Localization of Topo II in Asynchronously Growing Cells**

At certain stages of kDNA replication the pattern of topo II localization in cells labeled with dUTP-F differed significantly from that of pol β (Table I; Fig. 6). Before the onset of kDNA replication all cells had topo II detectable in the kinetoplast region; in 22% topo II was found to be diffuse throughout the kDNA region (Fig. 6 a). In nearly half (43%) of pre-replication cells topo II was localized to both antipodal sites (Fig. 6 d), but surprisingly, 35% had topo II concentrated at only a single site at the network edge (Fig. 6, b and c).

As with pol β, all cells undergoing kDNA replication had topo II concentrated in both antipodal sites, which colocalized with the bright spots of dUTP-F fluorescence (Fig. 6, e–g). Additionally, some weaker topo II signal was detected in the region of the kDNA network. After network replication, topo II remains localized to both antipodal sites in 73% of cells (Fig. 6, h–j), but it has a more diffuse localization in the kDNA region in the other 27% (Fig. 6 k). Once the processes of network repair and scission are complete, 60% of cells still have topo II localized in two discrete sites, although now each daughter network is associated with only a single site of topo II localization (Fig. 6, l and m). The remaining 40% of these cells have topo II diffusely localized in the kDNA region (Fig. 6 n).

**Localization of DNA Primase throughout the Cell Cycle**

Given the cell cycle variation in localization pattern of pol β and topo II, we were interested to determine whether the localization of enzymes not detected in the antipodal sites also varied during the cell cycle. To this end, we performed immunolocalization of the DNA primase in conjunction with hydroxyurea synchronization. In asynchronous cells, primase is localized to regions adjacent to the faces of the kDNA disk (Li and Englund, 1997). Using synchronized cultures we found that the primase is detectable in virtually all cells (Fig. 7 a), and is similarly localized to the two network faces at all stages of the cell cycle (see Fig. 7 b for examples of cells in S phase; and c for examples of dividing cells).

**Discussion**

The specific location of replication enzymes in unique sites surrounding the kDNA network is one of the most striking features of the kDNA replication system. In this paper we demonstrate the colocalization of pol β and topo II, and use two independent methods to analyze the changes in location of three replication enzymes during the cell cycle. Surprisingly, we found that these enzymes act differently. The DNA primase, localized adjacent to the faces of the kDNA disk, was invariant during the cell cycle. In contrast, pol β and topo II colocalize to the antipodal sites during the period of kDNA replication, but behaved differently at other cell cycle stages.

![Figure 7](image-url)

**Figure 7.** DNA primase localization during the *C. fasciculata* cell cycle. Cells from a synchronized culture were processed for immunofluorescence and stained with DAPI. (a) Percentage of cells undergoing cell division (closed circles), and percentage of cells with DNA primase localized above and below the kDNA disk (open triangles). (b) Localization of DNA primase during S phase, 60 min after release from hydroxyurea arrest. (c) Localization of DNA primase from the peak of cell division, 120 min after release from hydroxyurea arrest. The same localization of primase was seen at other time points. *Left column,* DNA primase localization detected by goat anti–mouse antibody conjugated to fluorescein. *Right column,* cells stained with DAPI to visualize the brightly stained kDNA and dimly stained nuclear DNA. Images were captured as described in the legend of Fig. 1. Bar, 3 μm.

**Cell Cycle–dependent Changes in Pol β**

Although pol β is located in the antipodal sites throughout kDNA replication (Figs. 2 and 5; Table I), this localization pattern is not static. Despite constant levels of protein (indicated by Western blots, see Fig. 3), the enzyme ceases to be detectable by immunofluorescence shortly after network replication is complete. Pol β remains undetectable by immunofluorescence throughout G1, and then reappears in the antipodal sites abruptly at the onset of kDNA replication. This coincidence of renewed detectability of pol β and the initiation of kDNA replication is striking, and suggests these events are coordinately regulated. Furthermore, the appearance of pol β in the two antipodal sites precisely at the onset of kDNA replication suggests that this enzyme has an essential role in that process.
Given its lack of fidelity and low processivity (Torri et al., 1994), however, it is unlikely to be the major replicative enzyme, and we have recently detected and partially purified a second mitochondrial DNA polymerase which may play that role (Klingbeil, M., and P.T. Englund, unpublished observations). Since pol β enzymes efficiently fill gapped templates (Torri et al., 1994; Singhal et al., 1995), the *C. fasciculata* pol β is well positioned in the antipodal sites to function in the partial repair of the highly gapped minicircle progeny, a process known to occur just before reattachment to the network (Kitchin et al., 1985). Later in the Discussion section we will comment on the lack of detectability of pol β by immunofluorescence during part of the cell cycle.

**Topo II Localization throughout the Cell Cycle**

Topo II, like pol β, is always detected in the two antipodal sites during kDNA replication (Figs. 2 and 6, and Table I). Unlike pol β, however, topo II is detected by immunofluorescence at all cell cycle stages, and is antipodally localized in some pre-replication and postreplication cells. In addition, at these times before and after kDNA replication, topo II is frequently localized to a single site at the network edge in some cells. Presumably, after kDNA replication, a post-replication network associated with two antipodally-positioned sites divides into two daughter networks, each associated with a single site. Consistent with this hypothesis, we never observed a post-scission network flanked by two antipodal sites. Thus, during cell division, each daughter cell must inherit a single site and a daughter kDNA network, yielding a G1 cell with a single site associated with the pre-replication network. The second antipodal site must then form during G1.

**Significance and Function of the Antipodal Sites**

What is the nature of the two antipodal sites to which pol β and topo II colocalize? It is unlikely that the sites are comprised of stoichiometric complexes of these enzymes, since crude estimates of their relative abundance (based on recovery of activity during purification) indicate that pol β is ~50-fold more abundant than topo II.

Our detection of topo II in discrete site(s) at all cell cycle stages suggests that these sites are permanent structures in the mitochondrial architecture. The sites are consistently found in the same focal plane and antipodal position relative to the kDNA (Ferguson et al., 1992), indicating their fixed position within the mitochondrial matrix. Such organized sites for enzyme localization may ensure not only that newly replicated minicircles are attached at opposing edges of the network (Simpson and Simpson, 1976; Pérez-Morga and Englund, 1993a), but could also establish the spatial configuration of other processes in the maintenance of the kDNA network. For example, one could speculate a role for the sites in establishing the line of cleavage in the center of the network before network scission. That the primase is also localized in fixed positions throughout the cell cycle indicates that this enzyme may also associate with structural elements that are permanent components of the mitochondrial matrix. Clearly, to further understand the intramitochondrial organization of these proteins it will be necessary to identify other components of both the antipodal sites and the areas colocalizing with primase.

As fixed components within the mitochondrion, these localization sites may also associate with structures within the mitochondrial membrane, or even outside the mitochondrion. Experiments on *Saccharomyces cerevisiae* demonstrated that the mtDNA is anchored to the mitochondrial membrane, and such anchoring explains the non-random mtDNA inheritance patterns (Nunnari et al., 1997). In bacteria, accumulating evidence suggests multi-protein complexes bridging the cell membrane and the bacterial genome function in chromosome maintenance and inheritance (Firshiein and Kim, 1997; Mohl and Gober, 1997). In *T. brucei*, the kDNA network is physically attached (through the mitochondrial membrane) to the basal bodies (Robinson and Gull, 1991). Furthermore, kDNA segregation is dependent upon the microtubule-mediated basal body separation (Robinson and Gull, 1991). A basal body–kDNA network association is also found in *C. fasciculata* (Robinson, D.R., and P.T. Englund, unpublished observations). Unfortunately, antibodies specific to either basal bodies or the network attachment site in *C. fasciculata* are currently unavailable. However, experiments correlating the relative positions of the topo II–localization sites with sites of basal body–kDNA attachment could elucidate the possible functions of these localization sites in kDNA maintenance or segregation.

**The Pol β Conundrum**

We do not know why pol β is undetected by immunofluorescence at some cell cycle stages, since the steady state protein level does not change (Fig. 2). One possibility is that pol β is dynamically localized to the antipodal sites during the cell cycle. The enzyme could assemble at the antipodal sites during kDNA replication, subsequently disassemble, and disperse throughout the mitochondrial matrix. When not sequestered in the antipodal sites the enzyme may be too dilute for immunofluorescence detection. Our attempts to detect dispersed pol β signal with long exposures resulted only in amplification of background signal throughout the cell, and the result was not distinguishable from control experiments in which we used only secondary antibody. Another explanation is that during G1 and cytokinesis pol β epitopes may be obscured, or masked, by protein–protein interactions that make the protein unrecognizable to antibodies in situ. The “masked” pol β could be located either in the antipodal sites, or dispersed throughout the mitochondrion.

Candidates for proteins that may interact with pol β include DNA repair enzymes. Mammalian nuclear pol β has been well characterized for its role in base excision repair (Nealon et al., 1996; Sobol et al., 1996), and it is known to form complexes with other base excision repair enzymes such as AP endonuclease (Bennett et al., 1997) and DNA ligase (Prasad et al., 1996). The abundance of reactive oxygen species, which are known to damage DNA, in mitochondria may necessitate such a repair pathway in this organelle (Richter, 1995). In support of this possibility we have recently detected and partially purified an abundant AP endonuclease (a key enzyme in the base excision repair pathway) from *C. fasciculata* mitochondria (Sax-
owsky, T., and P.T. Englund, unpublished observations). Thus in C. fasciculata mitochondria, pol β may function in kDNA replication by filling in minicircle gaps, and then subsequently interact with base excision repair enzymes and fulfill a second role in DNA repair.

**Regulation of kDNA Replication**

This work has implications for the control of kDNA replication. Unlike mammalian mitochondrial DNA (see Introduction), kDNA replication has been known for many years to proceed during a discrete phase of the cell cycle, with timing close to that of the nuclear S phase (Cosgrove and Skeen, 1970; Simpson and Braly, 1970; Woodward and Gull, 1990). Little is known about either the molecular factors that trigger the initiation of kDNA replication or the synchronization of this event with nuclear DNA replication. Ray and co-workers have shown a cell cycle dependence in the periodic accumulation of mRNA levels of genes for certain proteins involved in kDNA and nuclear replication (Paston et al., 1994; Brown and Ray, 1997; Hines and Ray, 1997). Their finding obviously relates to kDNA replication control and the synchrony of replication of nuclear and kDNA. We suggest that the intramitochondrial reorganization of certain replication enzymes may also play a role in control of kDNA replication. It will be of great interest to identify the factors that trigger these enzymes to assemble at discrete sites around the kDNA, and to further understand the nature and function of these antipodal sites.

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