Role of Centrins 2 and 3 in Organelle Segregation and Cytokinesis in *Trypanosoma brucei*

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Abstract

Centrins are calcium binding proteins involved in cell division in eukaryotes. Previously, we have shown that depletion of centrin1 in *Trypanosoma brucei* (*T. brucei*) displayed arrested organelle segregation resulting in loss of cytokinesis. In this study we analyzed the role of *T. brucei* centrin2 (*TbCen2*) and *T. brucei* 3 (*TbCen3*) in the early events of *T. brucei* procyclic cell cycle. Both the immunofluorescence assay and electron microscopy showed that *TbCen2* and 3-deficient cells were enlarged in size with duplicated basal bodies, multinuclei and new flagella that are detached along the length of the cell body. In both *TbCen2* and *TbCen3* depleted cells segregation of the organelles i.e. basal bodies, kinetoplast and nucleus was disrupted. Further analysis of the cells with defective organelle segregation identified three different sub configurations of organelle mis-segregations (Type 1–3). In addition, in majority of the *TbCen2* depleted cells and in nearly half of the *TbCen3* depleted cells, the kinetoplasts were enlarged and undivided. The abnormal segregations ultimately led to aborted cytokinesis and hence affected growth in these cells. Therefore, both centrin2 and 3 are involved in organelle segregation similar to centrin1 as was previously observed. In addition, we identified their role in kinetoplast division which may be also linked to overall mis-segregation.

Introduction

*Trypanosoma brucei*, a protozoan parasite of the order Kinetoplastida, is a causative agent of sleeping sickness in humans and Nagana in cattle in sub-Saharan Africa. The two replicating stages of the parasite, procyclic (in the vector gut) and the bloodstream form (in mammals) undergo a series of differentiation steps making the life cycle more complex. The parasite cell contains a single nucleus, one or two Golgi, one mitochondrion, one kinetoplast (containing mitochondrial DNA), one set of basal bodies (a mature basal body subtending the flagellum and a pro-basal body) and a flagellum. These must replicate and segregate in an orderly way in each cell cycle in order to form two similar daughter cells. The division and segregation of kinetoplast and nucleus depend on the prior duplication and segregation of basal bodies and flagellum complex [1,2,3,4].

Centrin is one out of several proteins involved in cell division. It is a calcium binding, centriole (in higher eukaryotes) and basal body (in lower unicellular eukaryotes and cells of tracheal epithelial, male gamete etc.) associated protein [5]. It is involved in duplication and segregation of these organelles [6,7,8,9,10,11]. Mouse centrin4 (MmCen4) is involved in the basal body assembly in the brain ependymal and choroidal ciliated cells [12]. Knockdown of human centrin2 by ribonuclease acid inhibition (RNAi) and a centrin gene deletion in *Tetranychusa thermophila* yielded defects in centrosome/basal body duplication and cell cycle progression [13,14], whereas disruption of *Chlamydomonas* centrin led to aberrant numbers of basal bodies that interfered with cytokinesis [7]. Centrins have also been found involved in other cellular processes such as maintenance of membrane integrity and cell morphology in yeast (yeast centrin, CDC31; [15]), homologous recombination and nucleotide excision repair in *Arabidopsis* (centrin2) and humans (HsCen2; [16,17]), nuclear mRNA export in yeast (CDC31; [18]), and genomic instability via increased chromosome loss in *C. reinhardtii* [19].

The recently completed genomes of protozoan kinetoplastid parasites, *T. brucei* and *Leishmania*, show that there are 5 centrin genes in these organisms. We have characterized centrin1 of *Leishmania donovani* (*L. donovani*) by ectopic expression of dominant negative gene mutation [20] and by gene knockout [8]. *L. donovani* centrin1 (*LdCen1*) was involved in the duplication of basal bodies only in amastigotes, an intracellular form and not in promastigotes, a form which is present in the sand fly vector [8]. On the contrary centrin1 in *T. brucei* (*TbCen1*; also named *TbCen4*) was found not to be involved in the basal body duplication but in the segregation of the basal bodies and other organelles [9,11]. However, *TbCen2* and *TbCen3* (also named *T. brucei* centrin1 by He et al., 2005) have been shown to be involved
in duplication of basal body [10]. In addition, TbCen2 was also shown to be involved in the duplication of Golgi [10]. In this report we have reexamined the functions of TbCen2 and TbCen3 in the basal body duplication. However, we did not analyze the role of TbCen2 in Golgi duplication. Similar to He et. al. 2005, our data suggests that TbCen2 and 3 have no role in nuclear division resulting in multinucleated enlarged cells. However contrary to the claim by He et al., 2005 that these two centrins have role in basal body duplication, upon re-examination, we observed that depletion of either TbCen2 or 3 had no effect on basal body duplication, but affecting the organelle segregation that may cause inhibition of cytokinesis as was observed with the depletion of TbCen1 [9,11].

**Results**

Both TbCen2 and 3 are essential for the growth of the parasite

In the present study we have characterized the functions of both TbCen2 and TbCen3 using RNAi methodology in *T. brucei* procyclics. Northern blot analysis of RNA obtained from the tetracycline induced cell cultures on day two revealed reduction of cognate mRNA levels of both TbCen2 and 3 (Figure 1A). Quantitation of the mRNA levels showed that there was ~78% reduction in the TbCen2 mRNA level and ~85% reduction in the TbCen3 mRNA level. There was no significant change in the mRNA levels of non-cognate centrins (Figure 1A). The effect of reduction of specific mRNA levels post induction on the growth of the cells in both cases was monitored by counting the cells in culture up to 5 days. RNAi induced TbCen3 depletion resulted in cell growth defect from day 2 (Figure 1B TbCen3 RNAi), whereas TbCen2 depletion showed cell growth defect only from day 3 (Figure 1B TbCen2 RNAi). The cell density in the induced cultures on day 3 was 69% for TbCen2 RNAi and 38% for TbCen3 RNAi compared to uninduced control cells. There was no substantial increase in the cell number in either case from day 4 onwards.

Depletion of centrins generates giant cells with multiple organelles

Under microscopic observation, both TbCen2 and 3 depleted cells on day 4 post induction were large, pleomorphic in shape, had new detached flagella not attached to the cell body, and were multinucleated, unlike the un-induced cells that were uniform in shape with one or two attached flagellum, and had single nuclei and kinetoplasts depending on the stage of the cell cycle (Figure 2A). In order to determine the cause of growth arrest, the RNAi-induced both TbCen2 and 3 cells along with uninduced cells as control were stained with propidium iodide (PI) and subjected to flow cytometry to analyze the relative DNA content (2C) at different time intervals in the cell culture (Figure 2B). The percent of cells having 2C, 4C and >4C were measured (Figure 2C). The analysis revealed that the RNAi induction for both TbCen2 and TbCen3 cells showed a gradual increase in the number of cells with >4C DNA content. Approximately 6% of TbCen2 and 8% of TbCen3 cells had >4C at day 3 compared with 3% of cells at day 0 in both the cultures (Figure 2C). The percentage of >4C cells increased up to 23% in the TbCen2 RNAi and 30% in the TbCen3 RNAi cells at day 5 (Figure 2C). The increase in the >4C cell population coincided with the simultaneous decrease in 2C cells (Figure 2B and C). In addition, to monitor the progression of replication of the other organelles, we counted the number of nuclei and kinetoplasts until day 5 in TbCen2 RNAi cells and day 4 in TbCen3 RNAi cells. A gradual increase in the number of cells with multinuclei and multikinetoplasts (>2K2N) was observed from day 4 onwards in the TbCen2 RNAi cells and day 3 onwards in the TbCen3 RNAi cells, coupled with a concomitant decrease in cells with one...
nucleus and one kinetoplast (1K1N; Figure 2D). We also observed cells with two kinetoplasts and one nucleus (2K1N), two kinetoplasts and two nuclei (2K2N), one kinetoplast and no nucleus (1K0N; zoid) and one kinetoplast and two nuclei (1K2N) as intermediate/minor populations before finally >70% of the TbCen3 RNAi cells and >25% of the TbCen2 cells on day 4 became >2K2N cells (Figure 2D).

Depletion of centrins did not affect duplication of the basal bodies

To analyze the organization of various organelles of both the control and TbCen2 and 3-depleted cells, they were stained with YL1/2 at day 3 for TbCen2 and day 2 for TbCen3 after RNAi induction to stain both pro and mature basal bodies, with anti-parafflagellar rod antibodies (L9C4) to stain the flagella and with DAPI to identify the nuclei and the kinetoplasts. The majority of the uninduced cells showed a single set of organelles of flagellum, basal bodies, kinetoplast (the mitochondrial genome) and nucleus (Figure 3A top panel). Both TbCen2 and 3-depleted cells displayed multiple nuclei and duplicated basal bodies and were also large and highly pleomorphic, as opposed to the control cells that were small and uniform in shape (Figure 3A middle and lower panels).

In either TbCen2 or 3 depleted cells, we noticed detached flagella in addition to one attached flagellum (Figure 3A middle and lower panels). We assume that the attached flagellum could be the initial flagellum of the cells before the induction of RNAi and the detached ones could be the newly formed after the RNAi induction as has been observed during RNAi induction for TbCen1 [9]. Careful examination of cells on either day 4 after TbCen2 RNAi or on day 3 of TbCen3 RNAi revealed that the very first new flagellum of more than 90% of such cells was detached type. The abnormal internal morphological characteristics of TbCen2 and 3-depleted cells were further confirmed by examining such cells by electron microscopy (EM). The features of multiple organelles observed in the centrin depleted cells from day 2–4 post induction were confirmed by the EM studies (Figure 3B b–h). Multi-basal bodies, multi-nuclei and abnormal kinetoplasts were clearly observed upon TbCen2 RNAi induction (Figure 3B b–d respectively) and upon TbCen3 RNAi induction (Figure 3B e–h respectively). Normal axoneme and parafflagellar rod were also noticed in the multiple flagella that appeared to be either attached or detached in a TbCen3 depleted cell (Figure 3B f).

The uninduced cells were mostly with a single nucleus, one kinetoplast and one basal body (Figure 3B a). To confirm the duplication of basal bodies does take place during depletion of both TbCen2 and 3, rat monoclonal antibodies against yeast tyrosinated-α-tubulin (YL1/2) stained basal bodies were analyzed by the immunofluorescence assay and quantitated. Most of the uninduced cells showed a single set of basal bodies. On the other hand, >85% of the TbCen2 or 3 depleted cells displayed duplicated basal bodies (Figure 3C) suggesting basal body duplication does proceed normally in these cells. Consistent with this observation, we also found 77% of TbCen2 depleted cells and 85% of TbCen3 depleted cells display more than one flagellum (Figure 3D).

Abnormal segregations of organelles in the T. brucei Centrin2 and 3 RNAi cells

The pattern of organelle segregation in the normal T. brucei procyclic cells has been described previously [3,4,21,22,23]. Briefly, during cell cycle, maturation and duplication of the basal bodies, division of kinetoplasts and the nucleus occurs successively followed by segregation. During segregation, one of the divided kinetoplasts migrates to the middle of the cell along with the linked basal bodies, followed by migration of one of the two divided nuclei posteriorly positioning between the segregated kinetoplasts allowing cell to initiate cytokinesis. In the uninduced cells, we observed clear segregation of the duplicated basal bodies along with the kinetoplasts followed by repositioning of organelles as described earlier prior to cell division (Figure 4A, panels 1 and 2). However, in the TbCen2 and TbCen3 depleted cells, we observed defect in the kinetoplast and basal body segregation in the two nucleated cells (Figure 4A panels 3–8). Careful observation among the TbCen2 and TbCen3 depleted cells resulted in the identification of three different configurations of organelle mis-segregation described here as “Types 1–3”. In ‘Type 1’ cells, in both TbCen2 and TbCen3 depleted cells, the duplicated basal bodies were observed with the kinetoplast, which was enlarged in its size without division, at the posterior end (Figure 4A panel 3 and panel 6) compared to the control cells (Figure 4A panel 2). In addition in the middle of the cell, 2 well divided nuclei were observed after mitosis in both the centrin depleted cells. These cells are designated K* NN cells, as per the positions of the nuclei and the kinetoplast from posterior to anterior direction (K* meaning kinetoplast enlarged in size). In ‘Type 2’ cells, duplication of the basal bodies and the kinetoplast were not affected. However, one group of the basal bodies and kinetoplast seemed to have migrated all the way to the anterior end followed by mitosis. As a result the divided nuclei remained in the middle of the cell (Figure 4A; panels 4 and 7). These cells are designated K NN K cells. In ‘Type 3’ cells, the duplicated basal bodies along with the kinetoplasts that were either enlarged and not divided in TbCen2 depleted cells (Figure 4A; panel 5) designated as N K* N or duplicated basal bodies along with divided kinetoplasts in TbCen3 depleted cells (Figure 4A; panel 8) designated as N K K N, migrated together to the middle of the cell allowing the posterior nuclei to shift to the far posterior end (Figure 4A; panels 5 and 8). The proportion of the three cell types varied among the TbCen2 and TbCen3 depleted cells. In TbCen2 depleted cells, on day 3 post induction the distribution of Type 1, Type 2 and Type 3 cells was 58%, 12% and 30% respectively (Figure 4B) with no significant change in the percentage of cell types at day 5 after RNAi induction. In TbCen3 depleted cells, on day 2 of induction, we did not observe Type 1 cells, but found 13% were of Type 2 and 87% were of Type 3 (Figure 4B). However, on day 4 of induction in TbCen3 cells, we did observe 27% of Type 1 cells with a slight decrease in Types 2 and 3 cells (Figure 4B). These observations demonstrate various distinct abnormalities in the organelle repositioning upon TbCen2 or TbCen3 depletion that may lead finally to the failure in cytokinesis.

Both TbCen2 and TbCen3 are also involved in the division of the kinetoplasts

Kinetoplast division is defined by the physical separation of the kinetoplasts into two after the kinetoplast S-phase (Figure 5A). In the uninduced control cells, we observed clear separation of the duplicated kinetoplast and subsequently their segregation as described (Figure 5A; panels 1–4). However, we observed defect in the kinetoplast division in the two nucleated cells in TbCen2 or 3 depleted cells (as in Figure 4A panels 3, 5, 6). Quantitation of such cells showed that at day 3, ~86% of the TbCen2 RNAi cells and at day 2, ~46% of the TbCen3 RNAi cells displayed kinetoplasts with defective division not observed in control cells (Figure 5B). However, the number of cell types with defective kinetoplast division decreased to 65% at day 5 in culture in TbCen2 RNAi cells, whereas the number remained nearly same till day 4 in TbCen3 depleted cells (Figure 5B). Further quantitation of cells with undivided kinetoplasts among the three
Figure 2. Analysis of cell structure and kinetoplasts and nuclei number in the RNAi cells. 

A: DAPI-stained images of the control and TbCen2 and 3-depleted cells, 4d after RNAi induction. DF, detached flagella; K, kinetoplast; N, nucleus. Scale bar, 5 μm. 

B & C: Fluorescence-activated cell sorting analysis for relative DNA content of TbCen2 and 3 depleted cells. 

B: Histogram plots of cell count by DNA content obtained directly from the flow cytometry analysis are shown. 

C: The percentage of cells that contain two times the DNA content (2C), four times (4C), and more than 4 times (>4C) were measured at each time point and are shown. Data represent the means ± SD of three independent experiments. 

D: Bar graph showing increase in the proportion of multinucleated and multikinetoplast cells with time after RNAi knockdown of TbCen2 and 3. Cells after different days of RNAi induction were stained with PI and examined by fluorescence microscope for tabulation of the cells with different number of kinetoplasts and nuclei. The time points at which the cells were analyzed after induction was up to day 5 for TbCen2 RNAi cells and day 4 for TbCen3 RNAi cells. Data represent the means ± SD of three independent experiments. For each TbCen2 and 3 RNAi studies, over 140 cells were manually counted and analyzed.

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Figure 3. Effect of ablation of centrins on cell shape and organelle number. **A**: Effects on the duplication and segregation of basal bodies, kinetoplasts and nuclei in TbCen2 and 3-depleted cells. The cells were stained with DAPI for the nuclei and kinetoplasts, YL1/2 for the basal bodies and L8C4 for the flagella. Note the RNAi induced cells are large and pleomorphic in shape with multiple organelles with more than one flagellum and the new flagella are of detached type (middle and lower panels) compared to the control cell with organelles in single number with one attached flagellum (top panel). Scale bar, 5 μm. **B**: Electron microscopy of centrin-depleted cells. (a) A typical control cell in this particular section shows single flagellum, pair of basal bodies, kinetoplast and nucleus. (b–d) TbCen2-depleted cells with (b) multi basal bodies and the kinetoplast with enlarged size compared to the control in ‘a’, (c) multi nuclei and (d) abnormal kinetoplast. (e–h) TbCen3-depleted cells with (e) multi basal bodies, (f) multi flagella. Inset is the cross-section image of a detached flagellum displaying the normal axoneme (with 9+2 microtubule structure) and the paraflagellar rod, (g) multi nuclei and (h) an abnormal kinetoplast. Scale bars, 500 nm (a, b, d–f and h) and 2 μm (c and g). **C & D**: Bar graphs showing the percent of *T. brucei* procyclic cells with duplicated basal bodies (C) and flagella (D). The cells were analyzed on day 3 after induction for
cell types from TbCen2 depleted cells showed that 100% of the Type 1 cells, which is the major cell type, and ~64% of the Type 3 cells had large kinetoplast without division (Table 1). On the other hand only 54% of the Type 3 cells, a major cell type in TbCen3 depleted cells, had undivided kinetoplast (Table 1). These results clearly indicate a defect in kinetoplast division after depletion of either of the two centrins. However, the overall effects were greater in TbCen2 depleted cells compared to TbCen3 depleted cells.

Discussion

In the procyclic stage of T. brucei mitosis occurs followed by one of the two nuclei being positioned between the two separated linked organelles, viz., basal bodies and kinetoplasts [4,9,21,24] as illustrated in Figure 6. In the current study upon TbCen2 RNAi induction, in the majority of the cells both duplication in the basal bodies and formation of new flagella occurred but failed in kinetoplast division (~86%). As a result, the duplicated basal bodies and the undivided kinetoplasts remained at the posterior end of majority (58%) of the cells (Type 1), despite normal division and separation of nuclei in these cells leading to defective

Figure 4. Images show direct effect of centrins’ depletion on kinetoplast biology and overall organelle segregation in T. brucei. A: Segregation patterns of the basal bodies along with the kinetoplasts during RNAi for TbCen2 and 3. The cells were stained with DAPI for nuclei and kinetoplasts and YL1/2 for basal bodies. Panels 1 and 2 display the typical segregation of basal bodies, kinetoplasts and nucleus in the control cells. Panels 3–5 and 6–8 display the 3 different segregation patterns of the organelles after RNAi of TbCen2 and 3 respectively, other than the typical pattern seen with the control cells (panel 2). Panel 1 is the ‘G’ stage of the cell cycle of the procyclic form with 1K1N, where both the basal bodies and kinetoplast are seen at the posterior region of the cell. Scale bar common for all the images, 5 μm. B: Histogram of cells with 2 basal body pairs displaying each of the 3 new segregation patterns of basal bodies and kinetoplasts in the RNAi cells. For each TbCen2 and 3 RNAi studies, over 100 biniucleated cells with abnormal organelle segregation were manually counted and analyzed. Data represent the means ± SD of three independent experiments. The time points at which the cells analyzed after induction was day 3 for TbCen2 RNAi cells and day 2 for TbCen3 RNAi cells. An, anterior end; B, basal body; K, kinetoplast; N, nucleus, Po, posterior end. Scale bars, 5 μm.

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segregation. Since there was no occurrence of zoids until day 3 after TbCen2 depletion and all the Type 1 cells had undivided kinetoplasts (K*N), it suggests that Type 1 cells were not the product of improperly placed furrow during cytokinesis, that usually leads to Type 1 (KNN) and zoids (K) cells as described by others [2,4,25,26]. In TbCen3 RNAi induced cells with no defect in basal body duplication, the majority (87%) of cells display Type 3 segregation defect and with overall only 46% of cells had undivided kinetoplasts. Both the basal bodies and kinetoplast were positioned in the middle of the cell (Type 3), with the two nuclei situated each at the ends of the cell which is different mis-segregation pattern from the majority of TbCen2 depleted cells (Type 1). Overall during TbCen2 depletion, only small percentage (~14%) of cells had divided kinetoplasts, on the contrary during TbCen3 depletion, majority of the cells (54%) showed divided kinetoplasts. Thus, our results not only highlight the differences in TbCen2 and TbCen3 functions in the organelle segregation but also identifying their function in kinetoplast biogenesis. It is possible that the differences in the nature of segregation among the organelles observed due to depletion of the two centrins (TbCen2 and 3) is through their mode of action, which could be influenced by other interacting components with the centrin, which are yet to be identified. In addition, centrin involvement in microtubule assembly could have an effect on the kinetoplast division/
segregation as was suggested in a recent study where segregation of minicircle and maxicircle DNA dependent on the dynamic remodeling of the cytoskeleton [22]. Therefore, the mechanism by which both centrin2 and 3 plays a role in kinetoplast biogenesis is not well understood at this time and needs further studying. Segregation defects, i.e. presence of Types 1–3 cells, seen in TbCen2 and 3 depleted cells have also been observed in cells expressing either altered polo-like kinase [27] or dynamin-like protein, necessary for mitochondrial fission in *T. brucei* [28].

Although the role of centrin in basal body or centriole biogenesis has been extensively demonstrated in various organisms [7,29,30,31,32,33] including our studies with *L. donovani* [8], we did not observe any effect on basal body duplication in the current study. The lack of effect by TbCen2 and TbCen3 depletion on basal body duplication in the present study is consistent with the effects of TbCen1 depletion (also named as centrin4 by Shi et al.) as indicated in our previous studies [9] and by Shi et. al., 2008 [11]. However, studies by He et al., 2005 have described, without any substantial experimental evidence, that depletion of Centrin2 and Centrin3 (Centrin1 by He et al., 2005) inhibit basal body duplication in *T. brucei* using stem-loop based RNAi system [10] as opposed to 2 T7 promoter based RNAi system that we used in the current study [9]. It is interesting to note that both we (in Selvapandiyan et al., 2007 [9]) and He group (in Shi et al., 2008 [11]) found that TbCen1/Centrin4 is not involved in the basal body duplication using same 2 T7 promoter based RNAi system. This suggests that differences in the RNAi systems used by He group in their two different studies could be the cause of the

| Type 1 configuration | Number of cells showing undivided kinetoplasts* | Type 3 configuration | Number of cells showing undivided kinetoplasts** |
|----------------------|-----------------------------------------------|----------------------|-----------------------------------------------|
| TbCen2RNAi | 85 (100%) | 22 | 14 (64%) |
| TbCen3RNAi | ---- | 95 | 51 (54%) |

DAPI stained cells were used in the analysis. The cells were analyzed on day 3 after induction for TbCen2 RNAi cells and day 2 for TbCen3 RNAi cells.

*Cells as observed in Figure 4A; panel 3.

**Cells as observed in Figure 4A; panel 5.

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Figure 6. Schematic diagram of the organelle segregation and the failure of segregation in the absence of TbCen2 or 3 in *T. brucei* procyclic form. Normal cell division and the three different types of abnormal segregation patterns of the organelles during centrin RNAi leading to the failure in cytokinesis and the generation of giant cells with multiple organelles are shown. Color coordination for organelles colored: red, basal body; green, kinetoplast; blue, nucleus.

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discrepancy in their observed functions. Furthermore, lack of involvement of TbCen1, TbCen2 and TbCen3 in basal body duplication in our studies supports a redundancy in gene function, which is a common hallmark of trypanosomatids.

In both TbCen2 and TbCen3 depleted cells besides lack of effect on basal body duplication, we also observed no effect on nuclear division indicating that nuclear division and cytokinesis are independent processes in the T. brucei procycls. An analogous situation has also been previously observed in T. brucei, when biogenesis and segregation of either basal body or flagella were affected leading to cytokinesis arrest even though mitosis was not affected [25,27,34,35]. Above mentioned observations are in conformity with the observations that cytokinesis was not affected when basal body and kinetoplast functions were normal even though mitosis was inhibited [11,21,36,37,38]. Taken together, it suggests that cytokinesis initiation depends on the combined function of duplication and segregation of the basal body and the kinetoplast and not on the mitosis.

The enlarged and pleomorphic shape observed in TbCen2 or 3 RNAi cells in the current study was also observed in TbCen1 depleted T. brucei procycls [9], LdCen1 knockout L. donovani promastigote cells [9] as well as in yeast with mutated centrin [39]. Similar effects were also seen with genes other than centrin e.g., depletion of T. brucei flagellar adhesion protein ( Fla) [35] and α-tubulin [40]. Whether the loss of shape in the centrin depleted cells is a direct effect or due to uncontrolled cell enlargement of size needs to be studied. TbCen2 and 3 depleted large cells also display multiple detached flagella as was observed with TbCen1 depleted cells [9]. It has been shown that Fla1 levels could control flagellar attachment [35]. However in TbCen1-3 depleted cells with detached new flagella, there was no effect on the level of Fla1 protein (Figure S1 and supplemental data in Selvapanidyan et al., 2007 [9] suggesting that the role of T. brucei centrins in flagellar attachment could be Fla1 independent. Similarly, FAZ is also involved in flagellum attachment and cytokinesis in T. brucei [2,35]. It has been suggested that during cell division the new FAZ, originates from the newly formed basal body in the posterior end and extends along the older FAZ to direct cytokinesis [41]. Hence, it is possible that the disruption of basal body biogenesis in the centrin depleted cells could simultaneously affect the origin of the new FAZ and in turn affect the biogenesis of the linked organelle kinetoplast. However, this hypothesis remains to be verified and is the subject of future studies. Taken together the data suggest the importance of all three centrins (TbCen1-3) in the normal shape, size and to the attachment of the flagella in T. brucei procycls. In T. brucei, flagellum position and polarity and the subpellicular cytoskeleton facilitate the morphogenesis of the flagellar pocket [42]. Depletion of cell cycle regulatory proteins, viz., dynamin-like protein [28] and polo-like kinase [27] in T. brucei results in enlarged flagellar pocket. On the contrary in our EM analysis depletion of centrins 2 and 3 had no effect on the overall structure of the flagellar pocket, again suggesting that the role of centrins in T. brucei cell biogenesis could be different from other cell regulatory proteins.

In conclusion re-examination of centrins 2 and 3 functions, in the current study, i.e. their noninvolvement in basal body duplication and nuclear division, their involvement in kinetoplast division and most importantly their role in segregation of organelles required to initiate cytokinesis is consistent with the functions of their counterpart centrins in the evolutionarily primitive organism T. brucei. Further studies are needed to elucidate the role of centrin association with the basal body to mediate kinetoplast biogenesis for cytokinesis and cell morphogenesis in the trypanosomatids.

Materials and Methods

In Vitro Culture of Parasites

T. brucei procycls form strain 29-13 that harbors integrated genes for T7 RNA polymerase and tetracycline repressor [43] was used. The parasites were grown and harvested as described previously [9,44].

Gene Cloning and Transfection of Parasites for RNAi

To amplify PCR fragments of TbCen2 and 3 genes for developing RNAi constructs, gene-specific forward and reverse primers were designed utilizing the putative centrin sequence from the T. brucei genome sequence databank [45]. Oligos were designed (Table S1) in which the ampiclon constitutes a portion from the 5’ untranslated region of the genes into approximately the middle of the open reading frame with HindIII and XhoI restriction sites added to the termini of the PCR fragments. The PCR amplified fragments were 347 bp for TbCen2 and 324 bp for TbCen3. These sequences are unique and share no significant sequence identity with the rest of the T. brucei genome sequences. The fragments were subcloned into the HindIII and XhoI sites of the pZJM vector [46]. Transfection of DNA in to the parasite, clonal selection of the parasite and the induction of RNAi were performed as described previously [9].

Isolation of RNA and Northern Blot Analysis

The cloned stable transfecteds, either uninduced or induced with tetracycline for 3 d, were analyzed for TbCen1 mRNA level using Northern blot analysis. The membranes were rehybridized with α-tubulin gene-specific probe as loading control. To confirm the specific inhibition of centrin transcripts, the TbCen2 membrane was also reprobed with TbCen1 and 3 gene specific probes and the TbCen3 membrane with TbCen1 and 3 gene specific probes. Probes were designed (Table S1) in such a way that the ampiclon constitute a portion from the middle of the ORF, not including the region selected for RNAi. The signal intensity was quantitated using a Phosphor Imager system (Molecular Dynamics, Amersham Pharmacia Biotech, Piscataway, NJ) as described previously [20]. As a loading control, the membranes were also reprobed with α-tubulin gene fragment and its mRNA levels were quantitated. After normalizing the centrin RNA intensity with the intensity of the α-tubulin mRNA control, the reduction in the amount of centrin mRNAs in the tetracycline induced cultures was compared with the uninduced cultures.

Flow Cytometry

T. brucei procycls inoculated at 1 x 10^5 cells/ml were allowed to grow and the cultures were harvested at 0, 3, and 5 d, fixed with paraformaldehyde, stained with propidium iodide (PI), and analyzed by flow cytometry according to the procedure described previously [47,48]. Cells were analyzed for populations with 2C, 4C, and >4C as a measure of relative DNA content. The stained cell samples were also examined with an Olympus phase-contrast and fluorescence microscope (Olympus, Melville, NY) to manually count the number of nuclei and kinetoplasts in individual cells.

Immunofluorescence Analysis

For the immunofluorescence experiments, cells were prepared and analyzed under fluorescence microscope following the procedure described previously [34]. Paraformaldehyde fixed mid-log T. brucei cells were stained with YL1/2 (rat mAb against yeast tyrosinated-α-tubulin from Chemicon, Temecula, CA; 1:400 dilution; [49]) for staining the basal body; L8C4 (antiparflagellar rod antibodies from Dr. Keith Gull, Oxford University; 1:4...
dilution; [50]) to stain the flagella. Appropriate secondary antibodies (anti-rat Alexa Flour 488 A-21200, anti-mouse Alexa Flour 568 A-21422, all from Molecular Probes [Invitrogen, Carlsbad, CA]) were used at 1:500 dilutions.

Electron Microscopy
Parasites harvested at appropriate time periods from culture were prepared and examined by electron microscopy as described previously [51]. Briefly, the parasite pellet was fixed overnight (4% formaldehyde and 1% glutaraldehyde in sodium phosphate buffer, pH 7.2), processed for transmission electron microscopy, sectioned and stained with uranyl acetate and lead citrate, and observed on a Philips CM110 Bio-twin electron microscope (Philips Electronic Instruments, Mahwah, NJ).

Statistical analysis
Statistical analysis of differences between means of groups was determined by two-sample t test assuming unequal variance. A p value<0.05 was considered as highly significant.

Supporting Information
Figure S1 Western blot showing the unaltered expression level of Flα1 protein using anti-Flα1 antibody in TbCen2 and TbCen3 depleted cells. In each well 20 μg of total extracted proteins were loaded. The cells were analyzed on day 3 after induction for TbCen2 RNAi cells and day 2 for TbCen3 RNAi cells. (TIF)

Table S1 Primer sequences used in the RNAi vector construction and probes for Northern blot analysis. All sequences are described in the 5’ to 3’ directions. Underlined regions are the restriction sites for the enzyme XhoI in the sequences 26F and 20F and HindIII in the sequences 27R and 21R. In sequence names ‘F’ stands for forward primer and ‘R’ stands for reverse primer. (DOCX)

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Author Contributions
Conceived and designed the experiments: AS HLN. Performed the experiments: AS PK JLS. Analyzed the data: AS HLN PK JLS CCW. Contributed reagents/materials/analysis tools: HLN JLS CCW. Wrote the paper: AS HLN.

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