ADF/cofilin-driven actin dynamics in early events of Leishmania cell division

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Summary

ADF/cofilin is an actin-dynamics-regulating protein that is required for several actin-based cellular processes such as cell motility and cytokinesis. A homologue of this protein has recently been identified in the protozoan parasite Leishmania, which has been shown to be essentially required in flagellum assembly and cell motility. However, the role of this protein in cytokinesis remains largely unknown. We show here that deletion of the gene encoding ADF/cofilin in these organisms results in several aberrations in the process of cell division. These aberrations include delay in basal body and kinetoplast separation, cleavage furrow progression and flagellar pocket division. In addition to these changes, the intracellular trafficking and actin dynamics are also adversely affected. All these abnormalities are, however, reversed by episomal complementation. Together, these results indicate that actin dynamics regulates early events in Leishmania cell division.

Key words: Leishmania, Actin dynamics, Kinetoplast, Basal bodies, Flagellar pocket, Cell division

Introduction

Actin is a highly conserved ubiquitous cytoskeletal protein that is essentially required in several important cellular processes such as cell division, cell motility, intracellular trafficking and endocytosis (Qualmann and Kessels, 2009; Kunda and Baum, 2009). It primarily exists in two forms: the monomeric globular form (G-actin) and the functional filamentous form (F-actin). The dynamics of actin-filament assembly and disassembly is regulated by a specific group of actin-binding proteins of which the actin-depolymerizing factor ADF/cofilin is an important component (Van Troys et al., 2008). ADF/cofilin is present in all eukaryotic organisms and has been implicated in several actin-based cellular activities, such as cell motility and cytokinesis (Ono, 2007; Pollard and Borisy, 2003). The main physiological function of these proteins is to depolymerize actin filaments from their pointed ends, and thereby promote filament dynamics (Carlert et al., 1997). In addition to depolymerization, these proteins also exhibit actin-filament-severing activity, which increases the number of filament positive ends and thereby enhances actin turnover (Van Troys et al., 2008). Higher eukaryotes express several isoforms of ADF/cofilin in a tissue-specific manner, which often have different cellular functions (Van Troys et al., 2008). These proteins have also been identified in a number of lower eukaryotic organisms, such as Acanthamoeba, Dictyostelium, Toxoplasma, Plasmodium and Leishmania (Blanchoin and Pollard, 1998; Aizawa et al., 1995; Allen et al., 1997; Schuler et al., 2005; Tammana et al., 2008).

Leishmania are an important group of flagellated, protozoan parasites that cause several diseases affecting millions of people worldwide, ranging from relatively mild cutaneous lesions to disfiguring mucocutaneous manifestations and fatal visceral disease (Desjeux, 2004). In these organisms, microtubules, rather than actin microfilaments, cover the inner surface of the plasma membrane enveloping the cell body. A small plasma-membrane invagination, called the flagellar pocket is the only site that is completely free of microtubule cytoskeleton, and therefore, it is the exclusive site where endocytosis and recycling of cell-surface molecules takes place (Gull, 1999).

Leishmania parasites express only one isoform of ADF/cofilin, which is essentially required in flagellar assembly and motility (Tammana et al., 2008). Detailed characterization of Leishmania ADF/cofilin (Cof) revealed that it readily binds both the monomeric and filamentous (Kapoor et al., 2008) forms of Leishmania actin and also depolymerizes filamentous actin into monomers (Tammana et al., 2008). Furthermore, it displays high nucleotide exchange but weak actin-filament-severing activities (Tammana et al., 2008). As the distributions of Cof and actin in Leishmania cells are largely polarized towards the flagellar pocket region where the basal body is located (Tammana et al., 2008), and as the cell division in trypanosomatids is initiated with basal body duplication followed by flagellum formation and flagellar pocket division (Ralston and Hill, 2008), we investigated the role of Cof-driven actin dynamics in Leishmania cell division. Results presented here indicate that actin dynamics has an important role in Leishmania cell division, especially in basal body separation and flagellar pocket division.

Results

Deletion of Leishmania COF gene results in impaired cell division

ADF/cofilins are actin-binding proteins that depolymerize F-actin into actin monomers and consequently enhance the actin dynamics (Ono, 2007). Our earlier studies have shown that deletion of COF gene in Leishmania promastigotes results in completely immotile, short and stumpy cells with highly reduced flagellar length and severely impaired flagellar beat (Tammana et al., 2008). In addition, we observed decreased growth of the mutant cells compared with wild-type cells in culture (Tammana et al., 2008). A detailed examination of these cultures revealed the presence of significantly higher numbers of dividing cells with two nuclei and two kinetoplasts (2N2K) or two nuclei and one kinetoplast (2N1K) in all the growth phases of COF−/− cells, compared with COF+/+ and
COF-/- cells complemented with GFP-Cof (COF-/-Comp) (Fig. 1). No multinucleated giant cells were, however, observed in these cultures at any given time. These results strongly suggest that, in addition to its essential role in the flagellar assembly, Cof is also involved in Leishmania cell division.

To test this possibility, we analyzed the cell cycle progression of log-phase cells that were synchronized at the G1-S boundary using hydroxyurea (HU). The progression of the cell cycle phases was dissected by flow cytometry at 2 hour intervals after releasing the HU block (Fig. 2, and supplementary material Fig. S1). About 90% of COF+/+, COF-/-Comp and COF-/- cells were arrested at the G1-S boundary by the overnight HU treatment (Fig. 2, and supplementary material Fig. S1). Upon releasing the HU block, the G2-M peak reached a maximum by 4 hours in COF+/+ (75.5±3.5%; n=3), COF-/-Comp (71.6±8.8%; n=3) and COF-/- cells (83.1±12.2%; n=3), in a synchronized manner. A decrease in the G2-M cell number (43.4±5.1%; n=3), with concomitant increase in the numbers of G1 phase cells (46.6±1.3%; n=3) occurred within 6 hours in synchronized COF+/+ culture. However, the subsequent progression of cell cycle from G2-M to G1 was significantly delayed in COF-/- cells, compared with COF+/+ cells (Fig. 2). In these cells, the G1 phase (43.8±13.1%; n=3) reappeared only after 10 hours, indicating a significant delay in progression of G2-M. However, GFP-COF complemented mutant cells showed a pattern that was similar to that observed with COF+/+ cells, except that an increased number of cells, compared with the wild type, were seen in the S phase of the cell cycle. This increase in S-phase cells could be due to overexpression of the GFP-Cof fusion protein in the mutant cells. The delay in the G2-M phase progression was reproducible in two randomly isolated COF-/- clones (see supplementary material Fig. S2).

Further analysis of the flow cytometry data revealed that COF+/+ cells returned to the G1 phase by 6 hours, but at this time point, COF-/- cells were largely arrested at G2-M. To ascertain whether the observed G2-M arrest was due to delay in mitosis, the COF-/- cells were labeled with DAPI 6 hours after HU block and then analyzed for nucleus and kinetoplast division (Fig. 3A). Interestingly, the percentage of 2N2K cells was significantly (P=0.009) higher (41.5±6.6%; n>600) in COF-/- cells, compared with COF+/+ cells (23±2.6%; n>600) and COF-/-Comp cells (27±5.2%; n>600), suggesting that the observed G2-M arrest was, in fact, post mitotic.

**Fig. 1.** Cell configurations according to the number of nuclei and kinetoplasts in Leishmania COF+/+ COF-/- and COF-/-Comp cells. Cells were counted in COF+/+ (black bar), COF-/- (gray bar) and COF-/-Comp (open bar) cell populations after DAPI staining during different phases of cell growth. 1N1K, cells with single nucleus and single kinetoplast; 2N1K, cells with two nuclei and one kinetoplast; 2N2K, cells with two nuclei and two kinetoplasts. Values are means ± s.d. of three independent experiments. In each experiment, at least 200 cells were analyzed. P-values were calculated using one-way analysis of variance and significance among the samples was analyzed by Tukey’s multiple comparison test.

**Fig. 2.** Cell cycle distribution of COF+/+, COF-/- and COF-/-Comp cells after removing the HU block. The log-phase cells (<10⁷/ml) were synchronized at the G1-S border with 200 µg hydroxyurea (HU) for 12 hours. DNA content was measured after staining with propidium iodide (PI) and the cell cycle phases were analyzed by flow cytometry at 2 hour intervals up to 12 hours. The G1, S and G2-M distributions in various cell cycle phases at different time intervals were calculated from the actual data using MOD-FIT software. Before the HU treatment, the characteristic profile of phase distribution observed in all the cell lines was: COF+/+: G0-G1, 53.1±0.9%; S, 25.0±1.27%; G2-M, 21.7±1.7%; COF-/-: G0-G1, 58.4±0.7%; S, 15.1±5.4%; G2-M, 26.5±6.0%; COF-/-Comp: G0-G1, 49.52±3.75%; S, 38.29±3.9%; G2-M, 12.28±6.45%. The values shown are means ± s.d. of three independent experiments.

**Cof-null cells show defects in basal body separation and cleavage furrow formation**

While observing the status of nuclei and kinetoplasts, it was noticed that the divided kinetoplasts in COF-/- cells were more closely positioned to each other than in COF+/+ and COF-/-Comp cells (Fig. 3B). As the kinetoplasts are physically linked with the basal bodies in trypanosomatids (Robinson et al., 2003), we also analyzed the distances between the basal bodies in the dividing cells. The
To further explore this problem, we labeled the furrow. Interestingly, most of the samples were treated with HU overnight and the samples were collected at 0 hours and 5 hours. Immunofluorescence images showing status of the nucleus and the kinetoplast. LdTUB, anti-tubulin (green); arrowheads indicate microtubule septa; arrows indicate closely spaced microtubules. DIC images merged with DAPI (cyan) show stage of the cell division. DIC images merged with DAPI (cyan) show stage of the cell division.

**Fig. 3. Analysis of COF−/− cells during cytokinetic arrest, compared with COF+/+ and COF−/−Comp cells.** (A) Different cell configurations in COF+/+ (+/+), COF−/− (−/−) and COF−/−Comp (−/−Comp) cell types were calculated based on DAPI staining. The values shown are means ± s.d. of three independent experiments. In each experiment, at least 200 cells were analyzed. P-values were calculated using one-way analysis of variance and significance among the samples was analyzed by Tukey’s multiple comparison test. (B) Immunofluorescence images showing the presence of higher numbers of microtubules in COF−/− cells, compared with COF+/+ and COF−/−Comp cells. All cells were treated with HU overnight and the samples were collected at 0 hours and 6 hours after removal of the HU block. The cells were then stained with anti-tubulin antibodies (LdTUB) for visualization of microtubules of cleavage furrow. Interestingly, most of the COF−/− cells were arrested during the furrow ingression stage of the cell division. DIC images merged with DAPI (cyan) show stage of the cell division. LdTUB, Leishmania tubulin (green); arrowheads indicate microtubule septa; arrows indicate closely positioned kinetoplasts; DIC, differential interference contrast. Scale bar: 5 μm. (C) Percentage of dividing cells with visible microtubule septum in COF+/+, COF−/− and COF−/−Comp cell populations, as calculated from immunofluorescence images after staining with anti-tubulin antibodies (LdTUB). The percentage of cells with visible microtubule septum at 6 hours is shown. The values are means ± s.d. of three independent experiments. At least 200 cells in each of three independent experiments were counted. P-values were calculated using one-way analysis of variance and significance among the samples was analyzed by Tukey’s multiple comparison test.

Average distance between the basal bodies, as measured by transmission electron microscopy, in dividing COF+/+ cells was 118.5±31.3 nm (n=10), compared with 184.0±41.5 nm (n=4) in COF−/− and 164.2±50.8 nm (n=4) in COF−/−Comp cells. These results indicated that the separation of the divided basal bodies and kinetoplasts was delayed in the dividing COF−/− cells. To further explore this problem, we labeled the COF+/+, COF−/− and COF−/−Comp cells with anti-tubulin antibodies and then analyzed for the formation of cleavage furrow microtubules at the divisional axis between the daughter cells. About 40% of COF−/− cells (39.4±6.5; n=600) showed a visible microtubule septum, compared with only ~20% of COF+/+(20.3±2.5%; n=600) and 23% of COF−/−Comp cells (23.0±4.0%; n=600) (P=0.0048) (Fig. 3C). However, none of the cells showed arrest in the abscission stage, suggesting that the cytokinetic arrest in COF+/+ cells was upstream of the furrow-ingression stage.

**Delayed furrow progression in COF−/− cells is caused by abnormalities in flagellar-pocket division and impaired vesicular movement**

The events upstream of furrow ingression include the formation of the new flagellum and flagellar-pocket division (Hammarton et al., 2007). As flagellum assembly was severely impaired in COF−/− cells (Tammana et al., 2008) and because flagellar biogenesis and flagellar-pocket organization have been shown to be coordinated with cell duplication in trypanosomes (Lacomble et al., 2009; Abasol et al., 2008), we envisaged that deletion of Cof could affect flagellar-pocket division. In trypanosomatids, flagellar-pocket division starts with an invagination of the flagellar-pocket membrane near the kinetoplast and proceeds towards the cell surface, eventually dividing the pocket into two. To examine flagellar-pocket division in COF−/− cells, we labeled the flagellar-pocket membrane with ConA-rhodamine and analyzed the labeled dividing cells using fluorescence microscopy (Fig. 4). About 80% of dividing COF−/− cells (81.6±3.5%; n=600) and COF−/−Comp cells (78.6±4.0%; n=600) showed two clearly separated flagellar pockets between the daughter cells. However, in case of COF−/− cells, only about 48% (48.0±2.6%; n=600) (P<0.0001) of dividing cells showed two separated flagellar pockets and the remaining 52% (52.3±2.6%; n=600) showed a single flagellar pocket between the daughter cells (Fig. 4A,B; also see supplementary material Fig. S3A), indicating delayed flagellar-pocket division in the mutant cells. Furthermore, in cells where the flagellar pockets were not divided, the two kinetoplasts also appeared closer to each other (Fig. 4A). In the case of COF−/−Comp cells, the flagellar pockets and the dividing kinetoplasts were clearly separated.

To further confirm these findings, we analyzed flagellar-pocket division by transmission electron microscopy (Fig. 5). Cross sections through the flagellar pocket region revealed a significantly higher number of COF−/− cells with two flagella within the same flagellar pocket (86%, n=43) compared with COF−/− cells (8%, n=50), confirming the delayed flagellar-pocket division in COF−/− cells. The flagellar pockets appeared enlarged, and a large number of membrane-bound vesicles were seen to accumulate close to the lumen of the flagellar pocket as well as in the cell body, suggesting disturbances in membrane trafficking in the COF−/− cells. To determine whether the flagellar pockets were still functional, we assessed the endocytic activity using the fluorophore N-(3-triethylammoniumpropyl)-4-(6-[4-(diethylamino)phenyl]-hexatrienyl) pyridinium dibromide (FM4-64), which has been widely used as a marker for assessing the flagellar pocket activity in trypanosomatids (Sahin et al., 2008; Mullin et al., 2001). In Leishmania, FM4-64 is readily internalized and targeted through endosomes to the final digestive compartment towards the posterior end (Waller and McConville, 2002). In COF+/+ and COF−/−Comp cells, endocytosis of the dye occurred with progressive labeling from flagellar pocket to the posterior end within 120 minutes (Fig. 6). Compared with COF−/− cells, the movement of FM4-64 in COF−/− cells was significantly slower, because the dye labeling was confined to the flagellar-pocket region for up to 60 minutes and it moved only as far as the kinetoplast and nuclear region by 120 minutes.
actin distribution was seen in the only at the later stages of cell division. However, no such polarized supplementary material Fig. S4) and its redistribution occurred anterior region of the flagellar pocket (Fig. 7, also see distribution in COF−/− cells. Actin, we analyzed the intracellular distribution of actin in dividing COF−/−, COF+/+ and COF−/− Comp cells. Closely positioned, divided kinetoplasts in the COF−/− cells were also clearly visible. N, nucleus; K, kinetoplast; arrows indicate flagellar pockets. Scale bars: 5 μm. (B) Statistical analysis of flagellar pocket status. COF−/−, COF+/+ and COF−/− Comp cells were counted after labeling their flagellar pockets with ConA-rhodamine. Dividing cells with two clearly separated flagellar pockets were counted under the category of divided pockets and dividing cells with a single flagellar pocket were categorized as undivided pockets. The values shown are means ± s.d. of three independent experiments. At least 200 cells in each of three independent experiments were counted. P-values were calculated using one-way analysis of variance and significance among the samples was analyzed by Tukey’s multiple comparison test.

**Fig. 4. Delayed flagellar-pocket division.** (A) Fluorescence images showing two divided flagellar pockets in the COF+/+ cells compared with single flagellar pockets in the COF−/− cell population. COF−/−, COF+/+ and COF−/− Comp cells were fixed with 4% paraformaldehyde in PBS and labeled with ConA-rhodamine (ConA-Rh, red) for flagellar-pocket staining. Nuclei and kinetoplasts are marked by DAPI staining (cyan). The presence of single flagellar pocket in COF−/− cells can be clearly observed compared with two separated flagellar pockets in COF+/+ and COF−/− Comp cells. Closely positioned, divided kinetoplasts in the COF−/− cells were also clearly visible. N, nucleus; K, kinetoplast; arrows indicate flagellar pockets. Scale bars: 5 μm. (B) Statistical analysis of flagellar pocket status. COF−/−, COF+/+ and COF−/− Comp cells were counted after labeling their flagellar pockets with ConA-rhodamine. Dividing cells with two clearly separated flagellar pockets were counted under the category of divided pockets and dividing cells with a single flagellar pocket were categorized as undivided pockets. The values shown are means ± s.d. of three independent experiments. At least 200 cells in each of three independent experiments were counted. P-values were calculated using one-way analysis of variance and significance among the samples was analyzed by Tukey’s multiple comparison test.

**Discussion**

The present study shows that deletion of the COF gene in *Leishmania* results in delayed basal body and kinetoplast separation, as well as in slower division-furrow ingression during cell division. It further reveals that the delayed furrow ingression in mutant cells is caused mainly by the delay in flagellar-pocket division, which is linked to reduced vesicular trafficking and impaired actin dynamics. These results indicate that Cof-driven actin dynamics is required to facilitate various early events underlying *Leishmania* cell division.

Various studies in *Saccharomyces, Drosophila, Dictyostelium, Caenorhabditis* and higher organisms have highlighted the important role of ADF/cofilin in cell survival and cytokinesis. Genetic studies in *Drosophila* show that ADF/cofilin is essential for cytokinesis and centrosome migration (Günsalus et al., 1995). Furthermore, deletion of the gene encoding cofilin-1 in *Dictyostelium discoideum* leads to cell death, indicating an important role of this protein in cell viability (Aizawa et al., 1995). Similarly, knockdown of the cofilin homologue UNC-60A in *Caenorhabditis elegans* causes embryonic lethality with cytokinesis and developmental defects (Ono et al., 2003). In higher organisms, ADF/cofilin is essential for depolymerization of the actin contractile ring that forms between two daughter cells during cell division (Abe et al., 1996). However, in the case of trypanosomatid parasites, cytokinesis is primarily a microtubule-mediated process, and despite the presence of actin and several actin-binding proteins (Sahasrabuddhe et al., 2004; Nayak et al., 2005; Kapoor et al., 2008; Tammana et al., 2008; Katta et al., 2009), their role in these parasites was largely unknown. Our recent studies (Tammana et al., 2008; Sahasrabuddhe et al., 2009) show that the actin-binding proteins Cof and CRN12 have essential roles in flagellar assembly and microtubule remodeling, respectively.

In trypanosomatids, the flagellum emerges from a specialized region called the flagellar pocket, a plasma-membrane invagination, which is also an exclusive site for endocytosis and exocytosis (Morgan et al., 2002a; Morgan et al., 2002b). Several studies have shown that the flagellar-pocket membrane is biochemically distinct from the flagellum or pellicular membrane and is central for trafficking of various GPI-anchored proteins (Schwartz et al., 2005).
These trafficking events involve recycling of several important proteins, such as clathrin, GTPase and Rab proteins (Garcia-Salcedo et al., 2004; Field et al., 2007), maintaining the flagellar-pocket membrane in a highly dynamic state. Since the flagellar-pocket membrane is devoid of microtubule cytoskeleton, it is possible that actin, as the flagellar-pocket cytoskeleton, regulates various dynamic activities of this organelle, such as endocytosis and membrane-furrow formation, during cell division. This is well supported by our observations that COF-null cells show absence of actin localization at the apical region of their flagellar pockets and highly reduced actin dynamics, as well as an increased number of flagellar pockets containing two flagella, compared with the COF+/+ and COF−/−Comp cells. Interestingly, despite the larger size of the flagellar pockets in the COF-null cells, the distances between the divided basal bodies are much smaller than in the wild-type cells, suggesting that the basal body separation is an active process, which requires the dynamic form of actin. Because, besides actin and Cof, myosin XXI is also present in the flagellar-pocket region (Katta et al., 2009), and because actin-based myosin motors essentially require the dynamic form of actin for their cellular activities (Zheng et al., 2009; Cramer, 2008; Semenova et al., 2008), we speculate that acto-myosin motor activity is involved in both basal body separation and vesicular trafficking in this organism.

In a closely related organism, Trypanosoma, it has been reported that ablation of flagellum formation results in short and immotile cells that fail to undergo cytokinesis, suggesting that the flagellum defines the position and direction of cleavage-furrow progression during cytokinesis (Kohl et al., 2003). Furthermore, ablation of flagellar biogenesis in these organisms has been shown to result in disturbances in flagellar-pocket organization (Absalon et al., 2008a). Flagellar elongation is essential for correct orientation and function of the flagellar pocket in these organisms (Absalon et al., 2008b). RNAi mutants of radial-spokes and central-pair proteins in Trypanosoma fail to undergo the final stage of cytokinesis, abscession, which leaves the daughter cells connected to each other at the posterior ends (Ralston et al., 2006; Branche et al., 2006), suggesting that the flagellar beating contributes the physical forces that are required to separate the daughter cells. Consistent with these observations, ablation of flagellar formation is accompanied by a reduction in cell growth in Leishmania cells (Cuvillier et al., 2000; Thiel et al., 2008; Tammana et al., 2008). However, it was hypothesized that because the Leishmania flagellum is mostly free from the cell body, a similar role to that of the Trypanosoma flagellum might not be expected (Kohl et al., 2003). This is supported by an earlier study which showed that the mutants of a dynein isoform (DHC2.2) in L. mexicana possess short flagella, but do not show any growth reduction (Adhiambo et al., 2005). Moreover, we did not observe any arrest of cell division at the abscession stage in Leishmania Cof mutants. Unlike in Trypanosoma, cytokinesis in these mutants proceeded more slowly than in the wild-type cells, but without any serious problems. Given these significant differences in cellular organization, a direct role of the flagellum in the Leishmania cell division can be ruled out at present.

It is intriguing to note that despite the completion of karyokinesis and formation of the microtubule septum, the daughter cells are still not separated. In trypanosomatids, the corset microtubules are
Role of actin in Leishmania cytokinesis

Fig. 7. Cell-cycle-dependant distribution of actin in cells labeled with antibodies against Leishmania actin. Immunofluorescence images show the enrichment of actin at the anterior tip of the flagellar pocket (arrows) during the initial biflagellate stage of the cell division in COF+/+ and COF−/− Comp cells. Redistribution of actin in these cells was observed at the later stages of cytokinesis. In case of COF−/− cells, most of the cellular actin is sequestered in the form of long cables in the cell body (arrowheads). Continuous extension of actin cables into the daughter cells can also be seen in dividing COF−/− cells. DIC images (gray) are merged with DAPI (cyan) images showing nucleus and kinetoplast. LdACT, Leishmania actin; N, nucleus; K, kinetoplast. Scale bar: 5 μm.

closely associated with the pellicular membrane (Gull, 1999). Therefore, the membrane ingression would need to follow an ‘interlocking-zipper’ mechanism along the cleavage-furrow microtubules from the anterior to the posterior end, up to the abscission stage to facilitate daughter-cell separation. Our results show that the cell division in Cof-null cells is arrested at the early stage of cytokinesis that includes flagellar-pocket division and cleavage-furrow formation. Although, no link between these two processes has so far been established, both these processes should require remodeling of the flagellar-pocket membrane. Apparently, this remodeling relies on the dynamic action of the actin network, and is delayed if the network is disrupted. During Leishmania cytokinesis, stages between the furrow formation and daughter-cell separation are so fast that the intermediate stages are rarely observed in culture. Interestingly, most of the Cof-null cells are arrested in the furrow-ingression stage and no dividing cells are encountered at the subsequent stages (i.e. after the furrow ingression and before the stage of abscission), indicating that once the furrow ingression is initiated, subsequent stages do not face any problem and the daughter cells separate normally. These results reveal that the actin dynamics is required only to trigger the initiation of furrow formation and is not involved in the downstream events during cytokinesis.

The process of membrane remodeling includes the addition of several specific proteins and lipids to the existing membrane, which in association with underlying cytoskeletal elements acquire different shapes and curvatures. The actin cytoskeleton has long been known to remodel the plasma membrane during various cellular processes, for example, formation of pseudopods or ruffles and cytokinetic furrows. During furrow formation, endocytic vesicles have been reported to serve as a pool of proteins and other membrane components in the mammalian cells (Barr and Gruneberg, 2007). This is consistent with our present observation that slower trafficking of endocytic vesicles, which is primarily caused by the impaired actin dynamics, is associated with delayed cleavage furrow formation in Cof-null cells. That actin in trypanosomatids is involved in intracellular vesicular trafficking and flagellar-pocket organization is further supported by an earlier study which showed that loss of actin prevents endocytosis and results in enlargement of the flagellar pocket in the bloodstream form of T. brucei (Garcia-Salcedo et al., 2004). However, it is important to mention here that the Leishmania F-actin by itself displays fast dynamics compared with conventional actins (Kapoor et al., 2008), and therefore some monomeric actin is always available to drive the actin dynamics, although at a reduced pace, even in the absence of any actin-dynamics-regulating protein. It

Fig. 8. Altered actin dynamics and cytoskeletal retention of actin in COF−/− cells. (A) Immunofluorescence images of cytoskeletons prepared by treatment of COF+/+, COF−/− and COF−/− Comp cells with 0.5% NP-40. The cytoskeletons were stained with anti-Leishmania actin antibodies (LdACT). COF−/− cells appear short and stumpy compared with long and slender COF+/+ cells (Tammama et al., 2008). Arrows indicate the presence of actin at the flagellar pocket region in COF+/+ and COF−/− Comp cells. Arrowheads represent the retention of actin cables in the COF−/− cells. LdACT, Leishmania actin (green). Scale bar: 5 μm. (B) Western blot analysis of cytoskeletal retention of actin in COF+/+, COF−/− and COF−/− Comp cells using antibodies against Leishmania actin. Anti-GRP78 (glucose-regulated protein of 78 kDa molecular mass) and anti-PFR antibodies were used as marker proteins for soluble and pellet fractions, respectively. +/+, COF+/+ cells; −/−, COF−/− cells; −/− Comp, COF−/− Comp cells; s, soluble fraction; p, pellet fraction.
might therefore be visualized that deletion of the Leishmania COF gene has only slowed down, but not abolished, the dynamics of filamentous actin, which has resulted only in the slowing down of the actin-based processes. This suitably accounts for the delayed flagellar-pocket division and impairment of the vesicular trafficking in COF−/− cells.

The cell cycle events of trypanosomatids broadly follow the general eukaryotic model, except that several checkpoints exist in these organisms (Woodward and Gull, 1990; Ploubidou et al., 1999). Defects in cytokinesis do not necessarily trigger mitotic checkpoints, as a result of which cells continue DNA replication and become multinucleated as a result of cytokinetic block checkpoints, as a result of which cells continue DNA replication and become multinucleated as a result of cytokinetic block (Ploubidou et al., 1999). However, we could not observe any multinucleated cells in the COF−/− cell population despite a delay in cytokinesis, suggesting an arrest of further nuclear division, as well as stimulation of the cell cycle block. Similar cytokinetic arrest has been observed in BILBO1 RNAi T. brucei cells, which results in apparent mitotic block (Bonhivers et al., 2008). The presence of significant numbers of undivided flagellar pockets in dividing COF−/− cells suggests that the flagellar-pocket division acts as a control point before daughter cell separation. Aberrations in flagellar-pocket separation trigger mitotic checkpoints, which results in a block of karyokinesis in the arrested cells. These results thus indicate that flagellar-pocket division has an important role in the Leishmania cell-division cycle. Finally, very little is known about the proteins that regulate the various steps involved in Leishmania cytokinesis (Hammarston et al., 2007). The results presented in this study show that ADF/cofilin-driven F-actin dynamics facilitates early events in Leishmania cell division.

Materials and Methods
Leishmania cultures and growth-curve analysis
Leishmania donovani COF−/− cells were maintained in high-glucose DMEM supplemented with 10% heat-inactivated fetal bovine serum (FBS) and 40 μg/ml gentamicin at 25°C. COF+/+ cells were grown in the same medium in the presence of 50 μg/ml G418 and 50 μg/ml hygromycin B and COF−/− cells (Tammana et al., 2008) were also grown in the same medium in the presence of 10 μg/ml tunicamycin. For growth analysis, COF−/+, COF+/− and COF−/− cells were grown in DMEM containing 10% fetal calf serum (FCS) without any antibiotics. Initially, the cells were seeded at a density of 1×105 cells/ml and the number of cells was counted at 24 hour time intervals with a Neubauer hemocytometer.

CoF-null mutants and COF−/− cells
Leishmania COF−/− cells were generated through sequential targeted replacement of the ADF/cofilin gene by selective marker genes conferring resistance to neomycin or hygromycin by site-specific homologous recombination, essentially as described earlier (Tammana et al., 2008). For the generation of COF−/− cells, we transfected the COF−/− cells with plasmid psh.5MCS containing the full-length COF gene tagged with GFP at its N-terminus (psh.GFP-COF) and the transfected cells were selected on DMEM agar plates containing 10 μg/ml tunicamycin, as described earlier (Tammana et al., 2008).

Cell cycle analysis
For flow cytometry analysis, cells were fixed as described (Dvorak, 1993) with slight modification. Briefly, about 106 cells from COF+/+, COF−/− and COF−/− cultures (asynchronous) were centrifuged separately at 3000 r.p.m. for 5 minutes, washed with cold PBS and resuspended in 50 μl phosphate-buffered saline (PBS) (pH 7.2). The cell suspension was mixed with 150 μl of fixative solution (1% Triton X-100, 40 mM citric acid, 20 mM sodium phosphate, 200 mM sucrose) and incubated at room temperature for 5 minutes. Finally, 350 μl of diluent buffer (125 mM MgCl2 in PBS) was added and the samples were stored at 4°C until further use. The fixed cells were treated with 50 μg RNase (5 mg/ml in 0.2 M sodium phosphate buffer, pH 7.0) for 3 hours at 37°C. Then, 50 μg/ml propidium iodide (5 mg/ml in 1.12% sodium citrate) was added and the tubes were incubated at 25°C for 1 hour. The samples were left overnight for equilibration at 4°C. The samples were analyzed on a FACS Calibur (Becton Dickinson), and the proportions of G1, S and G2-M populations were determined using ModFit software. Around 20,000 events were collected for each sample. For synchronization experiments, cells were maintained in exponential growth phase (≤1×105 cells/ml). About 1×106 cells/ml were centrifuged and transferred into fresh DMEM containing 200 μg/ml hydroxyurea and incubated at 25°C for 12 hours. The cells were washed twice with PBS and resuspended in fresh DMEM containing 10% FCS without hydroxyurea. Aliquots were taken at regular time intervals and the samples were processed for FACS analysis after addition of propidium iodide as described above.

Fluorescence microscopy
The nuclear and kinetoplast configurations of the cells were analyzed by staining the cells with 4′,6-diamidino-2-phenylindole (DAPI). Briefly, cells from COF−/+, COF−/− and COF−/− cells were centrifuged separately and washed with cold PBS twice and attached to poly-L-lysine (0.01% solution)-coated glass coverslips. The cells were fixed with 4% paraformaldehyde for 30 minutes at room temperature and washed thoroughly with PBS. The fixed cells were permeabilized with 0.5% Triton X-100 for 10 minutes at room temperature and washed twice with PBS. The coverslips were mounted using Fluorescein-FragEL mounting medium (Calbiochem) containing DAPI and analyzed on a Leica DM 5000B fluorescence microscope using ×63 1.4 NA (oil) Plan Apochromat lens. The cell configurations were categorized as 1N1K, 2N2K, 2N1K depending on the number of nuclei and kinetoplasts per cell and the percentage of each category was quantified.

For flagellar-pocket analysis, synchronized COF−/+, COF−/− and COF−/− cells were stained with ConA-rhodamine essentially as described earlier (Mores et al., 2008). Briefly, cells were treated with 50 μg/ml tunicamycin for 30 minutes at room temperature. The fixed cells were washed three times with cold PBS, resuspended in 1 ml PBS and then treated with 1:500 dilution of ConA-rhodamine (1 mg/ml in PBS) for 2 hours at room temperature. The cells were subsequently washed twice with PBS to remove excess ConA-rhodamine and then attached to poly-L-lysine (0.01% solution)-coated glass coverslips. The coverslips were mounted using Fluorescein-FragEL mounting medium (Calbiochem) containing DAPI and images were captured on a Zeiss LSM510 META confocal microscope using a ×63 1.4 NA (oil) Plan Apochromat lens. The specificity of anti-Leishmania actin antibodies has been established and reported by us earlier (Nayak et al., 2005; Sahasrabudde et al., 2004).

Transmission electron microscopy
Transmission electron microscopic analysis of COF−/+, COF−/− and COF−/− cells was performed essentially as described earlier (Tammana et al., 2008).

Intracellular vesicular trafficking
For monitoring vesicular trafficking activity, internalization of FM4-64 was performed essentially as described (Sahin et al., 2008). Briefly, 10×106 exponentially growing cells were incubated in DMEM containing 10% FCS and 2 μg/ml FM4-64 (Molecular Probes, 1:1000) and Cy3-conjugated goat anti-mouse secondary antibodies (Molecular Probes, 1:1000) were added and the mixture incubated at 4°C for 30 minutes. Thereafter the temperature was raised to 25°C. Small aliquots (50 μl) were harvested at different time intervals and mixed with 2% paraformaldehyde solution in PBS. To the fixed labeled cells was added Hoechst 33342 dye at 10 μg/ml concentration and the cells were immediately imaged under Zeiss LSM510 META confocal microscope using a ×63 1.4 NA (oil) Plan Apochromat lens and 3× digital zoom.

Cytoskeleton preparation
Leishmania promastigote cytoskeletons from COF−/+, COF−/− and COF−/− cells were prepared essentially as described earlier (Nayak et al., 2005) and analyzed by western blotting and immunofluorescence microscopy. Briefly, equal number of cells (about 106) of each sample were pelleted at 4°C and washed twice with chilled PBS. The washed cells were treated with 0.5% NP40 solution in PBS at 4°C for 5 minutes, and then centrifuged at 12,000 r.p.m. at 4°C. The soluble and pellet fractions were separated on polyacrylamide gels (10%) by electrophoresis and analyzed by western blotting using anti-Leishmania actin, anti-GRP78 and anti-PFR (mAb2E10) antibodies. mAb2E10 and GRP78 antibodies were a kind gift from Diane McMahon Pratt (Yale University, CT) and Emanuela Handman (Walter and Eliza Hall Institute of Medical Research, Melbourne, Australia), respectively. For immunofluorescence microscopy, the cytoskeletons were allowed to settle on poly-L-lysine-coated glass cover slips and processed for labeling as described above.

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Fig. S1

LdCOF<sup>+/+</sup> vs. LdCOF<sup>−/−</sup> vs. LdCOF<sup>−/−</sup> Comp

**ASYNC**

- **0 hr**
  - G1
  - S
  - G2/M

- **2 hr**
  - G1
  - S
  - G2/M

- **4 hr**
  - S
  - G2/M

- **6 hr**
  - S
  - G2/M

- **9 hr**
  - S
  - G2/M

- **12 hr**
  - S
  - G2/M

**COUNTS**

- **0 hr**
  - G1
  - S
  - G2/M

- **2 hr**
  - G1
  - S
  - G2/M

- **4 hr**
  - S
  - G2/M

- **6 hr**
  - S
  - G2/M

- **9 hr**
  - S
  - G2/M

- **12 hr**
  - S
  - G2/M

**FL2A (AFU)**
Fig. S3

A

![Bar graph showing the percentage of dividing cells in different genotypes](image)

- % Dividing cells
  - LdCOF^{+/+}
  - LdCOF^{-/-} 1
  - LdCOF^{-/-} 2

B

![Images showing cell division at 0 min and 60 min](image)

- 0 min
  - FP
  - K
  - N
- 60 min
  - LdCOF^{+/+}
  - LdCOF^{-/-} 1
  - LdCOF^{-/-} 2