Functional characterization of cohesin subunit SCC1 in Trypanosoma brucei and dissection of mutant phenotypes in two life cycle stages

Eva Gluenz,1* Reuben Sharma,2† Mark Carrington2 and Keith Gull1
1Sir William Dunn School of Pathology, University of Oxford, South Parks Road, Oxford OX1 3RE, UK.
2Department of Biochemistry, University of Cambridge, 80 Tennis Court Road, Cambridge CB2 1GA, UK.

Summary
In yeast and metazoa, structural maintenance of chromosome (SMC) complexes play key roles in chromosome segregation, architecture and DNA repair. The main function of the cohesin complex is to hold replicated sister chromatids together until segregation at anaphase, which is dependent on proteolytic cleavage of the cohesin subunit SCC1. Analysis of trypanosomatid genomes showed that the core cohesin and condensin complexes are conserved, but SMC5/6 is absent. To investigate the functional conservation of cohesin in eukaryotes distantly related to yeast and metazoa, we characterized the Trypanosoma brucei SCC1 orthologue. TbSCC1 is expressed prior to DNA synthesis at late G1, remains in the nucleus throughout S- and G2-phases of the cell cycle and disappears at anaphase. Depletion of SCC1 by RNAi or expression of a non-cleavable SCC1 resulted in karyokinesis failure. Using the dominant negative phenotype of non-cleavable SCC1 we investigated checkpoint regulation of cytokinesis in response to mitosis failure at anaphase. In the absence of chromosome segregation, procyclic trypanosomes progressed through cytokinesis to produce one nucleated and one anucleate cell (zoid). In contrast, cytokinesis was incomplete in bloodstream forms, where cleavage was initiated but cells failed to progress to abscission. Kinetoplast duplication was uninterrupted resulting in cells with multiple kinetoplasts and flagella.

Introduction
The conserved structures of chromatin, chromosomes and the nucleus are defining features of eukaryotic cells. Trypanosomatids belong to a eukaryotic supergroup (Simpson and Roger, 2004; Adl et al., 2005) separate from fungi/animals or plants. Their chromatin is based on nucleosomes containing canonical core histones H2A, H2B, H3 and H4 (Hecker et al., 1994) although the linker histone H1 is atypical, lacking a globular N-terminal domain (Alsford and Horn, 2004). Higher-order chromatin structure remains uncharacterized but it is known that condensation to 30 nm fibres does not occur (Hecker et al., 1994) and there is no visible condensation of chromosomes during mitosis.

In Trypanosoma brucei, the chromosomes contain a single linear DNA molecule and there are 11 diploid pairs of ‘large’ chromosomes from 1 to 6 Mb. In addition there are several intermediate chromosomes, from 200 to 900 kb, and ~100 minichromosomes, from 50 to 150 kb (Ersfeld et al., 1999). The ploidy of these smaller chromosomes is uncertain. Other trypanosomatid species such as Trypanosoma cruzi (Gibson and Miles, 1986) or Leishmania (Ivens et al., 2005) do not contain the two categories of smaller chromosome and in T. brucei the smaller chromosomes are probably an adaptation to increase the number of telomeres which act as favoured substrates in the gene conversion-based system of antigenic variation (Robinson et al., 1999). The functional organization of the genome of T. brucei, and other Trypanosomatida, is unique (Berriman et al., 2005). On the large chromosomes, genes are arranged in polycistronic transcription units of tens of genes that are transcribed constitutively by RNA polymerase II. Monocistronic mRNAs are produced by post-transcriptional processing (Palenchar and Bellofatto, 2006). Putative discrete centromeric DNA sequences in T. brucei large chromosomes were recently identified by mapping topoisomerase II cleavage sites (Obado et al., 2007).
The behaviour of chromosomes during mitosis has been investigated in *T. brucei* where mitosis is closed without breakdown of the nuclear envelope (Ogbadoyi *et al.*, 2000). There is an intranuclear spindle (Ersfeld and Gull, 1997) and some spindle microtubules appear to terminate in trilaminar kinetochore-like structures (Ogbadoyi *et al.*, 2000). Segregation of chromosomes occurs in association with the spindle but large and mini-chromosomes segregate with different dynamics (Ersfeld and Gull, 1997; Gull *et al.*, 1998). The total number of chromosomes exceeds that of spindle microtubules and kinetochores so the segregation of smaller chromosomes is not straightforward. A current model proposes that the small chromosomes form a lateral association with anti-parallel microtubules in the central area of the pole-to-pole spindle and are transported along the spindle microtubules by motor proteins (Gull *et al.*, 1998). The 11 large chromosomes by contrast are thought to attach to microtubules via kinetochores and their segregation mechanism is more similar to that of yeast chromosomes.

In yeast and metazoa, sister chromatid cohesion from S-phase to anaphase is maintained by cohesin, a structural maintenance of chromosome (SMC) complex (Losada and Hirano, 2005; Nasmyth and Haering, 2005). SMC complexes are composed of a SMC heterodimer, a kleisin and specific non-SMC subunits. In the case of cohesin, the α-kleisin is SCC1 which together with SMC1 and SMC3 forms a tripartite ring structure necessary for sister chromatid cohesion (Gruber *et al.*, 2003). The fourth component of cohesin is the STAG domain protein SCC3. A second type of SMC complex, condensin, has a role in higher-order chromatid structure and contains SMC2 and SMC4, along with a β- or γ-kleisin and HEAT repeat proteins (Nasmyth and Haering, 2005).

The separation of sister chromatids prior to cell division is dependent on a chain of events starting with the activation of the anaphase-promoting complex (APC), an E3 ubiquitin ligase. One of the first proteins ubiquitinylated by the APC, and subsequently proteolysed, is separase which is a stoichiometric inhibitor of the proteolytic activity of separase (Ciosk *et al.*, 1998). One substrate of separase is the cohesin subunit SCC1 which is cut at one or both of two specific sites, an event both necessary and sufficient for the onset of anaphase A (Uhlmann *et al.*, 1999; Wirth *et al.*, 2006).

Yeast and metazoan represent a small fraction of eukaryotic diversity. Here, we provide evidence that orthologues of cohesin and condensin components are present in trypanosomatids and that basic mechanisms of sister chromatid cohesion are conserved and thus evolved early. A functional analysis of *T. brucei* SCC1 was performed and depletion of SCC1 decreased the growth rate and resulted in nuclear division defects without arresting the cell cycle. Expression of a separase-resistant mutant of SCC1 was used to produce a dominant negative phenotype with cells unable to separate sister chromatids and thus arrested in early anaphase. The cellular phenotype of the anaphase arrest was different in the two life cycle stages investigated. Procyclic-form cells completed cytokinesis in the absence of mitosis resulting in production of anucleate cells with one kinetoplast (zoids) whereas bloodstream forms initiated cytokinesis without progressing to abscission.

**Results**

**SMC complexes in trypanosomes – the mitotic cohesin complex is conserved**

To identify proteins with a function in chromosome organization in trypanosomatids, the genomes of *T. brucei* (Berriman *et al.*, 2005) and *Leishmania major* (Ivens *et al.*, 2005) were searched for components of SMC complexes. Iterative profile-based searches (Devaux *et al.*, 2007) identified candidate orthologues (Fig. 1A) and phylogenetic analysis clarified their relationship with the well-characterized SMC proteins of yeast and animals and plants. Four SMC homologues were found in both genomes and were most closely related to SMC1, SMC2, SMC3 and SMC4, components of cohesin and condensin (Fig. 1B). The non-SMC subunits of the core cohesin and condensin complexes were also present in both genomes (Fig. 1A). Some organisms have evolved meiosis-specific α-kleisin and STAG domain proteins, exemplified by the *Schizosaccharomyces pombe* proteins Rec8 and Rec11 (Ellermeier and Smith, 2005). No candidate genes for meiosis-specific cohesin proteins were identified in the kinetoplastid genomes. No components of other SMC complexes were detected. In particular no SMC5 or SMC6 orthologues and no homologues of any of the known subunits of the SMC5/6 complex (Nse1–6) were identified in trypanosomes.

**Expression profile of TbSCC1 in bloodstream-form trypanosomes**

In yeast, the expression of SCC1 is regulated during the cell cycle; it is first expressed around the beginning of S-phase and is degraded at anaphase. The expression of SCC1 during the *T. brucei* cell cycle was determined by immunofluorescence labelling of bloodstream forms using an antibody raised against recombinant trypanosome SCC1 (Sharma *et al.*, 2008). Trypanosomes have a single mitochondrion with an unusually structured genome, the kinetoplast, which contains a network of concatenated, tightly packed circular DNA molecules. Replication of the kinetoplast occurs in defined phases, in precise temporal co-ordination with the replication of the nuclear genome.

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Fig. 1. A. Orthologues of SMC1–4 and non-SMC subunits of the cohesin and condensin complex were identified in the genomes of Trypanosoma brucei and Leishmania major by (i) HMMER searches using seed alignments from Pfam and (ii) reciprocal best blast hits for the corresponding proteins from yeast. The accession numbers for GeneDB (http://www.genedb.org) and Pfam domains (http://pfam.sanger.ac.uk/) are listed.

B. Phylogenetic analysis was used to classify the trypanosome SMC proteins. Kinetoplastid genomes encode four SMC proteins that are orthologues of SMC1–4. A maximum parsimony phylogram was built using sequence alignments of the four T. brucei SMC sequences with SMC1–6 from Saccharomyces cerevisiae (Sc; NCBI Entrez Accession No. P32908, P38989, P47037, Q12267, Q08204, Q12749), Arabidopsis thaliana (At; NP_191027.2, NP_201047.1, NP_180285.2, BAB10693.1, BAB11444.1), Drosophila melanogaster (Dm; AAM5005.1, AAD52673.1, NP_523374.2, NP_728986.1, Q87386, AAF56254.1) and Homo sapiens (Hs; Q14683, O95347, NP_005436.1, Q9NIJ, Q8IY14, Q96SB8). Bootstrap values from 500 trees are indicated.
The number and relative positions of kinetoplasts (K) and nuclei (N) were used to stage cells within the cell cycle and 5-bromodeoxyuridine (BrdU) incorporation was used as an additional marker for S-phase. Nearly all trypanosomes in G1 (1K1N) showed no SCC1 labelling and no BrdU incorporation (Fig. 2A). Low levels of SCC1 were detected in some 1K1N cells in the absence of detectable BrdU incorporation (Fig. 2B); this may represent cells about to enter S-phase. The intensity of SCC1 staining increased with progression of DNA synthesis as judged by BrdU incorporation (Fig. 2C and D). Cells in G2 (2K1N) showed a strong SCC1 signal, some cells had also incorporated a small amount of BrdU (Fig. 2E), and these cells probably represent cells in the very last stages of S-phase when the labelling period started. The SCC1 expression pattern in bloodstream forms closely resembles that in procyclic forms (Sharma et al., 2008).

The antibodies were used to quantify the average number of SCC1 molecules per cell using Western blotting of cell lysates and serial dilutions of known quantities of recombinant TbSCC1 (Sharma et al., 2008 and data not shown). The estimate was that lysates from $1 \times 10^6$trypanosomes contain 2.5 ng of SCC1 protein. Taking into consideration the cell cycle-dependent expression pattern of SCC1 described above, and assuming that 40% of cells in an asynchronous culture of procyclic T. brucei are in nuclear S- or G2-phase (Woodward and Gull, 1990), these cells contain approximately one SCC1 protein per $1 \times 10^3$ bp of DNA.

**SCC1 is essential for normal karyokinesis**

In yeast, the absence of functional SCC1 leads to a failure in sister chromatid cohesion and a loss of viability (Uhlmann et al., 1999). To determine if SCC1 had an essential role in trypanosome mitosis, bloodstream forms were transfected with a construct for tetracycline-inducible RNAi based on p2T7-177 (Wickstead et al., 2002). By 6 h after induction of RNAi, SCC1 protein was reduced in the induced population, and remained below the detection limit between 12 and 48 h after induction (Fig. 3A and data not shown). Induction of SCC1 RNAi led to a slower population growth rate (Fig. 3B), and the calculated generation time increased from 5.6 h in uninduced cultures to ~11 h on the second and third day after induction of RNAi. Cells proliferated throughout the experiment and complete growth arrest was not observed.

Induction of SCC1 RNAi resulted in an accumulation of cells with abnormal K and N numbers and multiple...
flagella. There was a decrease in the fraction of 1K1N cells in the population (Fig. 3C). By 48 h post induction, 13% of cells had more than two kinetoplasts, and 4% of cells had more than two nuclei (Fig. 4). About a quarter of all cells in the population had nuclei with abnormal morphology evident by examination of 4,6-diamidino-2-phenylindole (DAPI)-stained cells. This included cells with normal combinations of K and N numbers. Typically, such aberrant nuclei appeared multi-lobed, with masses of DNA connected by thin DNA bridges (Fig. 3D).

The RNAi phenotype was transient and the growth rate and morphology of cells returned to normal in cultures induced for more than 3 days. This loss of phenotype correlated with a re-appearance of SCC1 protein in Western blots between 48 and 64 h after induction (data not shown) and is most likely due to outgrowth of cells refractory to RNAi.

Expression of a dominant negative SCC1

Proteolytic cleavage of SCC1 is sufficient to start anaphase in yeast whereas inability to cleave SCC1 results in a failure to separate sister chromatids and an anaphase arrest (Uhlmann et al., 1999). Separase is specific for a conserved sequence SxExxRx present once or twice, sites A and B, in all SCC1 orthologues, site A being present in all SCC1 homologues whereas site B is not universally conserved. In the trypanosomatids, T. brucei SCC1 contains sites A and B (Fig. 5A) whereas the L. major protein only contains site A. Three mutant versions of SCC1 were made with a R → D mutation in separase cleavage site A (mutA), or B (mutB) or both A and B (mutAB). The SCC1 transgenes were then expressed with C-terminal eYFP fusions from a tetracycline-regulated promoter from integrated constructs. Express-
SSION OF THE SCC1 TRANSGENES WAS USED TO TEST WHETHER THE MECHANISM OF SCC1 CLEAVAGE IS CONSERVED AND AS A TOOL TO STUDY THE CELLULAR PHENOTYPE OF AN ANAPHASE ARREST CAUSED BY INABILITY TO CLEAVE SCC1.

The effect of expression of the SCC1 transgenes on growth rate was determined. The wild-type SCC1-eYFP had no effect on growth (Fig. 5B). Expression of SCC1-mutA caused a slight reduction in population growth rate in both bloodstream forms and procyclic forms. Expression of SCC1-mutB had no effect on the growth rate. In contrast, expression of SCC1-mutAB caused a reduction in the growth rate of procyclic forms from 12 h post induction. By 50 h, the reduction in growth was 70% compared with wild type (Fig. 5B). The higher levels of SCC1-mutAB were probably due to accumulation of uncleaved protein. The simplest interpretation for the phenotype of SCC1 transgene expression is that mutation of both the potential separase cleavage sites in SCC1 has indeed rendered SCC1-mutAB resistant to cleavage and this in turn resulted in a slowing or cessation of population growth.

Western blot analysis of whole-cell lysates over a time-course after induction of the transgenes showed that expression of SCC1 peaked at 12–24 h, and that the steady-state levels of SCC1-mutAB were higher than wild type, SCC1-mutA or SCC1-mutB (Fig. 6). The higher levels of SCC1-mutAB were probably due to accumulation of uncleaved protein. The simplest interpretation for the phenotype of SCC1 transgene expression is that mutation of both the potential separase cleavage sites in SCC1 has indeed rendered SCC1-mutAB resistant to cleavage and this in turn resulted in a slowing or cessation of population growth.

On a Western blot, the anti-SCC1 antibody detects a doublet of ~75 kDa in T. brucei whole-cell extracts and a single band at ~75 kDa when tested against recombinant SCC1 (Fig. 6). The most likely explanation for this is that the lower band in the doublet represents unmodified TbSCC1, while the higher band is a phosphorylated or otherwise modified version. RNAi affects both bands in the doublet (Fig. 3) indicating that these proteins are products of the same gene rather than different proteins that share common epitopes. The idea that TbSCC1 is

**Fig. 4.** Detailed analysis of kinetoplast (K) and nucleus (N) numbers in bloodstream-form T. brucei after induction of SCC1 RNAi (A, 24 h post induction; B, 48 h post induction) and in cells expressing non-cleavable SCC1-mutAB (C, 24 h post induction; D, 48 h post induction). The number of K and N was counted in DAPI-stained cells and plotted. The proportion of cells in each category is represented by the area of the circle. At least 500 cells were scored at each time point.
Fig. 5. A. Identification of putative separase cleavage sites in TbSCC1 by homology with characterized sites from proteins in other organisms. The residues in the consensus motif are shaded grey and the two arginine residues mutated to aspartic acid in TbSCC1 are boxed.

B. The effect of ectopic expression of wild-type SCC1 and separase cleavage site mutants on the growth of *T. brucei* procyclic forms (left) and bloodstream forms (right). Gene expression was induced with tetracycline and the cell density in induced cultures (filled circles) and non-induced controls (open circles) was measured at regular intervals.
modified in vivo is further supported by the fact that all of the tagged versions of SCC1 examined also run as doublets in Western blots (Fig. 6 and data not shown). In yeast SCC1 is known to be phosphorylated at several residues and phosphorylation of serine residues near the separase cleavage sites are required for efficient cleavage of SCC1 by separase (Alexandru et al., 2001). There are serine residues in equivalent positions in TbSCC1.

The cellular phenotype in procyclic and bloodstream forms

Expression of SCC1 transgenes was induced and the nuclear and kinetoplast DNA visualized in individual cells over a time-course. Cells were categorized according to the number of kinetoplasts (K) and nuclei (N) to determine cell cycle stage (Woodward and Gull, 1990). In populations of procyclic trypanosomes expressing wild-type SCC1, SCC1-mutA or SCC1-mutB, ~80% of cells were 1K1N in G1 with 8–17% in S-phase. Zoids, which are anucleate cells with one kinetoplast, 1K0N, formed when cytokinesis occurs in the absence of mitosis (Robinson et al., 1995), were rare. In contrast, procyclic trypanosomes expressing SCC1-mutAB produced large numbers of zoids (20% at 12 h, 44% at 24 h and 36% at 48 h post induction) (Fig. 7A). In addition, 9% of cells expressing the SCC1-mutAB had abnormal combinations of K and N numbers and some cells had enlarged nuclei (Fig. 7B).

Bloodstream trypanosomes expressing wild-type SCC1, SCC1-mutA or SCC1-mutB were also similar to wild-type cells. Expression of SCC1-mutAB caused a sharp decrease in the proportion of cells with 1K1N, from 74% in uninduced populations to 30% at 24 h after induction. Cells with multiple kinetoplasts accumulated rapidly. At 12 h after induction the proportion of 2K1N cells had increased from 14% to 36%. By 48 h after induction, more than half of the cells in the population had abnormal combinations of K and N numbers (Figs 4C and D and 7C). The number of bloodstream-form zoids was small (9%).

Microscopic examination of DAPI-stained bloodstream forms expressing SCC1-mutAB showed multiple morphological abnormalities, with a single enlarged nucleus, multiple kinetoplasts and multiple flagella (Fig. 7D). The number of the kinetoplasts and flagella in individual cells indicated that new rounds of organelle duplication had occurred after a failed mitosis.

In both bloodstream-form and procyclic trypanosomes the result of SCC1-mutAB expression was a failure in nuclear division with no indication of progression beyond anaphase A. In procyclic forms cytokinesis still occurred and a zoid was produced; in bloodstream forms cell division was not successfully completed.

Examination of bloodstream-form trypanosomes expressing SCC1-mutAB by light and electron microscopy revealed that many had prominent cleavage furrows (Figs 8 and 9) and thus population growth arrest (Fig. 5B) occurred due to a failure to complete cell division as opposed to a failure to initiate cell division. At 24 h post induction, 38% of cells with more than one kinetoplast had one cleavage furrow and 11% had two or more (Fig. 8F). SCC1 knockdown by RNAi also resulted in cells with arrested cleavage furrows; 36% of cells with more than one kinetoplast had one cleavage furrow and 3% had two or more (Fig. 8F).

The posterior end of the stalled cleavage furrow was often in close apposition with the undivided multi-lobed nucleus (Fig. 8D and E; see also Fig. 3D). The proximity of the stalled furrow to the nucleus is illustrated at higher resolution in a thin section through a cell expressing SCC1-mutAB (Fig. 9). These observations suggest that, in bloodstream forms, the undivided nucleus may form a physical obstacle to further cleavage furrow ingestion.

After both expression of SCC1-mutAB and induction of SCC1 knockdown the cell nuclei are enlarged and irregularly shaped. Examination by transmission electron microscopy shows distinct abnormalities in nuclear ultrastructure. Electron-dense plaques are frequently observed (Fig. 9), often associated with spindle microtubules (Fig. 9B). Many nuclei are no longer of a regular

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Fig. 7.  A. Expression of the separase-resistant SCC1 (SCC1-mutAB) in procyclic forms causes a rapid accumulation of anucleate 1K0N cells (zoids).
B. The effect of SCC1-mutAB expression on cellular morphology and on the nucleus and kinetoplast in procyclic forms at 0, 12 and 24 h after induction; the arrows indicate one cell with three kinetoplasts and no nucleus and another cell with an enlarged nucleus.
C. Expression of SCC1-mutAB in bloodstream forms causes a rapid accumulation of cells with multiple kinetoplasts (category ‘others’) and yields a small number of zoids. Filled square: 1K1N; filled triangle: 2K1N; open square: 2K2N; open diamond: 1K0N (zoids); open triangle: 1K2N; filled circle: ‘others’.
D. Bloodstream forms at 12 and 24 h after induction; the arrows indicate the multiple kinetoplasts in a typical cell.
oval shape, but instead have multiple protrusions and invaginations of the nuclear envelope.

Taken together, these results provide evidence that the gene Tb927.7.6900 identified as the closest homologue of yeast SCC1 is indeed the functional orthologue. *T. brucei* is the most divergent eukaryote for which the mechanism of sister chromatid separation by SCC1 cleavage by separase has been demonstrated. The consequences of expression of a separase-resistant form of SCC1 were an arrest of mitosis prior to separation of sister chromatids. Mitotic arrest resulted in different cellular phenotypes in two life cycle stages; bloodstream forms initiated cytokinesis without progressing to abscission, while procyclic forms completed cytokinesis and produced zoids.

**Discussion**

The main findings of this study are: (i) the *T. brucei* genome contains homologues of all the components present in cohesin and condensin, two SMC complexes necessary for the correct segregation of chromosomes at mitosis, (ii) SCC1, the kleisin component of cohesin, is expressed from the beginning of S-phase until anaphase, (iii) knockdown of expression of SCC1 by RNAi impedes nuclear division, (iv) expression of a dominant negative

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**Fig. 8.** Bloodstream-form *T. brucei* expressing SCC1-mutAB and cells induced for SCC1 RNAi fail to complete cytokinesis.

A and B. (A) Morphology of a bloodstream-form trypanosome viewed by scanning electron microscopy before induction of SCC1-mutAB expression; (B) 12 h after induction. The arrow indicates a cleavage furrow. Scale bar represents 1 μm.

C–E. Phase-contrast images of cells with zero, one or two cleavage furrows, respectively, indicated by arrows. (C′)–(E′) show the DAPI-stained DNA in these cells. Images were taken 24 h after induction of SCC1-mutAB expression.

F. At 24 h post induction, cells with more than one k inetoplast were examined for the presence of cleavage furrows. The graph shows the proportion of cells with zero, one or two cleavage furrows. White bars: SCC1-mutAB-uninduced control; dark grey bars: SCC1-mutAB; light grey bars: SCC1 RNAi.

**Fig. 9.** Transmission electron microscopy shows the effect of expression of SCC1-mutAB on nuclear ultrastructure of bloodstream forms. The cell in (A) has a cleavage furrow (arrow) in close apposition to the undivided nucleus (N). Electron-dense plaques are visible in the nucleus in (A) and (B). The association of plaques with spindle microtubules can be seen in (B). (B′) shows the boxed area at a higher magnification. Scale bars represent 500 nm.
mutant of SCC1 that is resistant to cleavage by separase
blocks nuclear division, and (v) reduction of wild-type
SCC1 activity or expression of separase-resistant SCC1
resulted in a failure to complete cell division in blood-
stream forms but not in procyclic forms which divided to
produce one nucleated and one anucleate daughter cell.

Cohesin is the complex that physically holds the two
sister chromatids together from S-phase to anaphase.
Proteolytic cleavage of the SCC1 subunit by separase at
one of two specific sites is essential for sister chromatin
separation at anaphase. The results from this study
provide evidence that the gene Tb927.7.6900 identified as
the closest homologue to yeast SCC1 is the functional
orthologue; the expression pattern and nuclear phenotype
of both the knockdown and dominant negative are similar
equivalebent mutations in the SCC1 genes from yeast and
Drosophila (Uhlmann et al., 1999; Vass et al., 2003).
Knockdown of TbSCC1 by RNAi in bloodstream-form try-
panosomes resulted in a phenotype consistent with a
delayed progression through the cell cycle and aberrant
segregation of nuclear DNA. Expression of non-cleavable
SCC1-mutAB produced a more severe phenotype, char-
acterized in bloodstream-form trypanosomes by a growth
arrest at 12 h post induction and a decrease of cell
numbers 24 h post induction indicating cell death. During
mitosis, individual T. brucei chromosomes do not visibly
condense but the movement of DNA clusters to opposite
poles of the nucleus follows a defined pattern that can be
observed in DAPI-stained cells (Ogbadoyi et al., 2000).
The sharp increase in the proportion of cells with 2K1N
early after induction of SCC1-mutAB expression, and
accumulation of cells with > 2K1N indicate a block of
nuclear division early in mitosis. This phenotype is consis-
tent with a block in sister chromatid segregation resulting
in failure of anaphase A and the absence of anaphase B
events including spindle elongation and correct nuclear
positioning. Expression of separase-resistant SCC1 pre-
vented both procyclic and bloodstream-form trypano-
somes from partitioning nuclear DNA into two clusters of
equal size and no late mitotic, anaphase B, nuclei were
observed. The inducible expression of SCC1 with muta-
tions in both separase cleavage sites (SCC1-mutAB) thus
provided an experimental tool to investigate the phenotype
of cells with nuclei unable to separate sister chromatids
during mitosis. The conservation of function of SCC1, the
kleisin subunit of cohesin and the presence of homologues
for the other cohesin subunits in the genome provide
evidence for a conserved mechanism of sister chromatin
cohesion common to all eukaryotes. The mutational anal-
ysis of the separase cleavage sites showed that one func-
tional cleavage site is necessary and sufficient for normal
progression of T. brucei mitosis, a similar finding was
made in yeast (Uhlmann et al., 2000). Separase cleavage
of SCC1 as a trigger for anaphase was first discovered in
the budding yeast Saccharomyces cerevisiae (Uhlmann
et al., 1999; 2000) and subsequently in metazoans (Hauf
et al., 2001; Jager et al., 2001; Wirth et al., 2006). Our
data show that the mechanism whereby separase cleaves
SCC1 to release chromosome cohesion evolved early
in eukaryotic evolution, before the euglenozoan and
animal, fungal and plant lineages diverged, possibly
almost two billion years ago (Hedges et al., 2004).

Entry into mitosis and progression to metaphase is
dependent on the activity of the cyclin-dependent kinase
CDK1 in all eukaryotes studied to date. Progression from
metaphase to anaphase is dependent on activation of the
anaphase-promoting complex or cyclosome (APC/C). The
APC/C is the E3 ubiquitin ligase that indirectly activates
the protease separase by targeting its inhibitory subunit
securin for degradation (de Gramont and Cohen-Fix,
2005). Separase is then responsible for cleavage of
SCC1, permitting anaphase, and independently initiates
the release of the CDC14 phosphatase from the nucleolus
(Shou et al., 1999), which is necessary for mitotic exit and
probably reverses the CDK1 phosphorylation. This sim-
plified pathway was elucidated in yeast and, although
all steps have not been confirmed in trypanosomes,
orthologues of CDK1, cyclin and the APC/C have been
identified and homologues of separase (Tb927.1.3120)
(Mottram et al., 2003) and CDC14 (Tb11.01.4270) are
encoded in the trypanosomatid genomes. In T. brucei,
CYC6/CRK3 are the orthologues of cyclin B/CDK1. Depletion of CYC6 by RNAi results in a mitotic block; in
procyclic forms, cell division occurs in the absence of
nuclear division producing a zoid daughter cell whereas
in bloodstream forms there is an accumulation of cells
with a single enlarged nucleus and multiple kinetoplasts
(Hammarton et al., 2003; Tu and Wang, 2004). Similar
experiments with CRK3 produced a similar phenotype
(Kumar and Wang, 2005). The different phenotypes in
procyclic and bloodstream forms led to the suggestion
that regulation of cell cycle progression differs in the two
life cycle stages and that a mitosis to cytokinesis check-
point only operates in bloodstream forms (Ploubidou
et al., 1999; Hammarton et al., 2003; Tu and Wang,
2005). The depletion of APC/C activity through RNAi
knockdown of CDC27 or APC1 resulted in cell populations
enriched in G2/M cells; in procyclic forms cells arrested
with two kinetoplasts and a single enlarged nucleus con-
taining a short spindle whereas bloodstream-form cells
arrested with two kinetoplasts and a nucleus arrested in
late anaphase (Kumar and Wang, 2005).

In procyclic forms, the phenotype of cells expressing
separase-resistant SCC1 was similar to cells in which
CRK3 or CYC6 had been depleted with the production of
zoids resulting from cell division in the absence of nuclear
division. The accumulation of zoids in cultures expressing
separase-resistant SCC1 (40% after 24 h) was similar
both to that obtained using the microtubule assembly inhibitor rhizoxin (30% zoids after 8 h) (Robinson et al., 1995; Ploubidou et al., 1999) and to that obtained after RNAi depletion of CYC6 (~40% after 48 h) (Hammarton et al., 2003). In contrast, APC/C-depleted procyclic forms accumulated as 2K1N cells, the nucleus containing a short spindle consistent with an anaphase A arrest, but did not undergo cytokinesis.

Bloodstream-form cells expressing separase-resistant SCC1 had a different phenotype to CRK3- or CYC6-depleted cells. In both cases, cells arrested with an enlarged nucleus and multiple kinetoplasts but cells expressing separase-resistant SCC1 initiated cytokinesis whereas cells depleted of CRK3 and CYC6 did not. Depletion of APC/C activity caused a late anaphase arrest with an elongated spindle. The bloodstream-form phenotype observed in our study also differed markedly from the very specific precytokinesis cell cycle arrest observed when variant surface glycoprotein (VSG) transcripts were ablated by RNAi (Sheader et al., 2005). VSG RNAi caused a rapid accumulation of 2K2N cells with two external flagella and, in a minority of cells, additional internal flagella. No internal flagella were observed in SCC1 mutants. Clearly, not all bloodstream-form cytokinesis defects are the same and specific phenotype patterns are now emerging.

Previous RNAi studies manipulated CDK1 (CYC6/CRK3) and APC/C activity; both are regulators of the cell cycle and knockdown will have pleiotropic effects; for example, a reduction of CDK1 activity will not only affect phosphorylation of its substrates but also the substrates of downstream kinases and knockdown of the APC/C components will reduce separase activation which will prevent cleavage of SCC1 but also block the role of separase in the activation of CDC14 and mitotic exit. In contrast, SCC1 is a structural component with a single role in sister chromatid cohesion and is a substrate for cell cycle regulators. Manipulation of SCC1 was used to reveal the phenotype resulting from defects in sister chromatid cohesion. RNAi knockdown of SCC1 prevented a normal mitosis and expression of the separase-resistant SCC1 allowed an analysis of the phenotype of an early anaphase block. In both of the life cycle stages investigated cytokinesis was initiated in the presence of separase-resistant SCC1 or after SCC1 knockdown. Thus, the initiation of cytokinesis is not dependent on the completion of mitosis and there is no checkpoint capable of blocking initiation of cell division in response to incomplete mitosis in either life cycle stage. In T. brucei procyclic forms, complete cytokinesis occurred in the absence of nuclear division whereas bloodstream forms arrested with stalled cleavage furrows.

Why is cytokinesis in bloodstream-form cells incomplete after expression of separase-resistant SCC1? The trypanosome cytokinesis machinery and mechanisms of its activation are unknown (Hammarton et al., 2007) so it is possible in bloodstream forms that furrow ingression was blocked by a checkpoint mechanism in response to a signal indicating that mitosis had failed but this would be without precedent. Alternatively, the architecture of the bloodstream trypanosome may physically prevent cytokinesis in the SCC1 mutants. In procyclic trypanosomes, the position of the anterior nucleus remains fixed, through an as yet uncharacterized anchoring system, whereas the posterior nucleus ‘moves’ into the gap between the segregated basal bodies (Robinson et al., 1995). As the kinetoplast is physically linked to the basal body, prior to cell division the order of organelles from the anterior end is alternating kinetoplasts and nuclei – KNKN in bloodstream forms and KKKN in procyclic forms (Tyler et al., 2001). In cells expressing separase-resistant SCC1, the close apposition of the partial cleavage furrows with the nuclear membrane evident in the electron micrographs suggests that an undivided nucleus may form a physical barrier that prevents, or slows down, further cleavage furrow ingression. Thus, the different outcomes of separase-resistant SCC1 expression in the two life cycle stages investigated can be explained by the different geometry of organelle position in the two cell types.

The difference between the separase-resistant SCC1 dominant negative phenotype and the CYC6 and CRK3 knockdown phenotypes is informative. In procyclic forms, the phenotypes were similar whereas in bloodstream forms no initiation of cleavage was reported when CYC6 or CRK3 was depleted. One possible interpretation is that CYC6/CRK3 was required for initiation of cleavage in bloodstream forms, possibly through activation of the APC/C, but not in procyclic forms. However it would be worth confirming first the precise degree of CRK3 or CYC6 knockdown and whether there was any effect on APC/C activation. The absence of cell division in procyclic cells with depleted APC/C activity cannot be explained solely by a failure to licence SCC1 cleavage but suggests the APC/C activity may be required for initiation of cell division, possibly through its role in the activation of CDC14 as occurs in yeast. In bloodstream forms, depletion of APC/C led to a late anaphase arrest whereas expression of separase-resistant SCC1 resulted in an arrest in early anaphase with no elongation of the nucleus. The easiest explanation for these contrasting
observations is that the depletion of APC/C was not complete but was insufficient to activate the mitotic exit network (Sullivan and Morgan, 2007). In yeast mutation in mitotic exit network genes results in a late anaphase arrest (Jaspersen et al., 1998).

The experiments presented here provide no evidence for checkpoints that link the completion of mitosis to cytokinesis initiation in the two T. brucei life cycle stages investigated. Organelle duplication proceeds in the absence of a successful mitosis as several rounds of basal body and kinetoplast duplication occurred in bloodstream forms although not in procyclic forms. Re-initiation of nuclear S-phase was not directly determined but the nuclei were enlarged in procyclic and bloodstream forms after expression of separase-resistant SCC1 suggesting endoreduplication was occurring.

The results of the experiments above show that in the two T. brucei life cycle stages investigated, initiation of cytokinesis is independent of sister chromatid cohesion or cohesin release following SCC1 cleavage. The nuclear division defect caused by expressing non-cleavable TbSCC1 is reminiscent of the phenotypes of yeast and human cells that express non-cleavable SCC1. In S. cerevisiae, cytokinesis was delayed but not inhibited and progeny with abnormal DNA content were produced (Uhlmann et al., 1999). In human cells, cytokinesis was initiated but not completed and sister chromatid separation was shown not to be required for cyclin B destruction or mitotic exit (Hauf et al., 2001). Depletion of TbSCC1 by RNAi in bloodstream forms led to slower proliferation without cell cycle arrest and partial ingestion of cleavage furrows. In yeast and higher eukaryotes a spindle assembly checkpoint inhibits APC/C and prevents anaphase onset until tension generated by sister chromatid cohesion and bipolar attachment to the spindle microtubules is sensed. With the exception of Mad2p, no homologues of known spindle checkpoint proteins were found in the trypanosome genome (Berriman et al., 2005). Progression of the cell cycle in the absence of SCC1 could indicate that such a checkpoint is absent or, alternatively, that an arrest is only transient.

Experimental procedures

Bioinformatics

Kinetoplastid genome sequences were accessed via GeneDB (http://www.genedb.org/). Iterative profile-based searches (Devaux et al., 2007) were used to identify sequences in the kinetoplastid genomes. Seed alignments used to generate hidden Markov models (Eddy, 1998) were produced from sequences identified with simple BLAST searches and Pfam alignments (Bateman et al., 2004). For phylogenetic reconstruction, SMC protein sequences were aligned in CLUSTAL X, manually edited and a maximum parsimony tree was built with PAUP* 4.0b10 (Swofford, 1998).

Cells

Trypanosoma brucei brucei Lister 427-derived bloodstream-form cell line 90-13 (Wirtz et al., 1999) was cultured in HMI-9 (Hirumi and Hirumi, 1994) and procyclic cell line 29-13 (Wirtz et al., 1999) was grown in SDM-79 (Brun and Schonenberger, 1979).

Production of antibody and immunofluorescence microscopy

To produce anti-SCC1C, the 5′ end of the TbSCC1 coding sequence (encoding amino acids 390–584) was amplified with primers 5′-GACGGATCCCGCTCAAGGG-3′ and 5′-AAGCTTACACAGTGAGTTGCACCTC-3′ and cloned in the BamHI and HindIII sites of pQE-30 (Qiagen) to generate an in-frame fusion with a 6xHisptide tag. The protein was purified from Escherichia coli M15 PREP4 on Ni-NTA superflow resin (Qiagen) under denaturing conditions, precipitated with acetone and used to immunize rabbits. Immunofluorescence detection of SCC1 and BrdU was performed as previously described (Sharma et al., 2008).

For generation of the RNAi cell line, a 536 bp fragment of TbSCC1 was PCR-amplified from genomic DNA using primers 5′-TCTAGACCTTTTCTCCGGCTACG-3′ and 5′-TCTAGACTCATTTCCTCTCGGACGTCGACGTCGACGTCGAC-3′, digested with XbaI and cloned in the XbaI site of p2T7-177 (Wickstead et al., 2002). The resulting plasmid p2T7-177-SCC1 was linearized with NotI and used to transfect bloodstream-form 90-13 cells by electroporation. Transfected cells were selected with 2.5 μg ml⁻¹ Phleomycin and RNAi was induced using 1 μg ml⁻¹ doxycycline.

Cell growth was monitored using a CASY cell-counter and analyser system (Schärfe System GmbH). For DNA staining, cells were spread onto glass slides, fixed in 2–4% paraformaldehyde in phosphate-buffered saline (PBS) and embedded in Vectashield (Vector Laboratories) with DAPI. Images were captured on a Leitz DMRB fluorescence microscope (Leica Microsystems) using a CoolSnap FX Camera (Photometrics) and assembled in Adobe Photoshop.

Site-directed mutagenesis of SCC1 to produce SCC1-mutA, mutB, mutAB was carried out using standard procedures. All transgenes were fully sequenced after mutagenesis. Expression used either p2216 or p2280 (Kelly et al., 2007) in the procyclic-form cell line Lister 427 pLEW29 pLEW13 or the bloodstream-form cell line Lister 427 pLEW90 pLEW13 (Wirtz et al., 1999).

Electron microscopy

Bloodstream-form cells were fixed in 2.5% glutaraldehyde, 2% formaldehyde, 0.1% picric acid in 100 mM phosphate buffer pH 7.0 for 2 h. For thin-section electron microscopy, cells were then processed as described in Dawe et al. (2005). For scanning electron microscopy, fixed cells were left to adhere to glass coverslips and processed as described in Sharma et al. (2008).
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