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Genome-wide Analysis Reveals Extensive Functional Interaction between DNA Replication Initiation and Transcription in the Genome of Trypanosoma brucei

Calvin Tiengwe, 1,5 Lucio Marcello, 1,5 Helen Farr, 3,5 Nicholas Dickens, 1 Steven Kelly, 4 Michal Swiderski, 1 Diane Vaughan, 2 Keith Gull, 3 J. David Barry, 1 Stephen D. Bell, 3,* and Richard McCulloch 1,*

1 The Wellcome Trust Centre for Molecular Parasitology
2 Cellular Analysis Facility
College of Medical, Veterinary and Life Sciences, Institute of Infection, Immunity and Inflammation, University of Glasgow, Sir Graeme Davies Building, 120 University Place, Glasgow G12 8TA, UK
3 Sir William Dunn School of Pathology, University of Oxford, South Parks Road, Oxford OX1 3RE, UK
4 Department of Plant Sciences, University of Oxford, South Parks Road, Oxford OX1 3RB, UK
5 These authors contributed equally to this work
*Correspondence: stephen.bell@path.ox.ac.uk (S.D.B.), richard.mcculloch@glasgow.ac.uk (R.M.)

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SUMMARY

Identification of replication initiation sites, termed origins, is a crucial step in understanding genome transmission in any organism. Transcription of the Trypanosoma brucei genome is highly unusual, with each chromosome comprising a few discrete transcription units. To understand how DNA replication occurs in the context of such organization, we have performed genome-wide mapping of the binding sites of the replication initiator ORC1/CDC6 and have identified replication origins, revealing that both localize to the boundaries of the transcription units. A remarkably small number of active origins is seen, whose spacing is greater than in any other eukaryote. We show that replication and transcription in T. brucei have a profound functional overlap, as reducing ORC1/CDC6 levels leads to genome-wide increases in mRNA levels arising from the boundaries of the transcription units. In addition, ORC1/CDC6 loss causes derepression of silent Variant Surface Glycoprotein genes, which are critical for host immune evasion.

INTRODUCTION

Faithful copying of the genetic material is central to life. However, the features that define origins, the loci at which DNA replication initiation occurs, are poorly understood for most eukaryotes (Masai et al., 2010; Méchali, 2010). Unlike in bacteria, where origins are DNA sequences that show both sequence and positional conservation in different species’ genomes (Robinson and Bell, 2005), only in Saccharomyces cerevisiae are similarly conserved sequences, termed the autonomously replicating sequence (ARS), found that define the sites of replication initiation (Wyrick et al., 2001). In other eukaryotes consensus sequences at mapped origins are lacking, and in particular in metazoa, it appears that epigenetic cues help define origins, which display an association with transcription the molecular basis of which is as yet unclear. Nevertheless, origins display mechanistic conservation. Eukaryotic chromosomes possess multiple origins, each bound by the six-subunit origin recognition complex (ORC; composed of Orcs 1–6), which recruits the replicative helicase (the MCM complex) via interactions with Cdc6 and Cdt1 (Duncker et al., 2009; Bochman and Schwacha, 2009). Initiation of DNA replication is regulated to occur once per cell division cycle through the actions of cyclin-dependent kinases (CDKs) and the Cdc7/DBF4 kinase complex (Difffley, 2010).

Nuclear DNA replication in African trypanosomes, such as Trypanosoma brucei, has been little characterized, and in common with all protists, origins of DNA replication have not been mapped genome wide. Intriguingly, trypanosomes appear to possess only a subset of the higher eukaryotic replication machinery, potentially having a single protein with homology to both Orc1 and Cdc6 (ORC1/CDC6) (Godoy et al., 2009; Tiengwe et al., 2012) and lacking detectable orthologs of Cdc7 and Dbf4 (data not shown). Trypanosomes also have an unusual genome organization, which may provide a particular perspective on replication. In T. brucei the majority of protein-coding genes are in 11 diploid megabase-sized chromosomes and are arranged in ~150 “directional gene clusters” (DGCs) that contain, on average, ~50 genes, but can encompass hundreds (Daniels et al., 2010). Transcription of each DGC is thought to initiate from a single RNA polymerase (pol) II promoter, yielding a primary multigene transcript from which individual mRNAs are generated by trans-splicing and polyadenylation. Transcription initiates bidirectionally between divergent DGCs (at so-called divergent strand switch regions: SSRs) and at some sites within some DGCs (Martinez-Calvillo et al., 2003; Koyle et al., 2010). These putative promoters display increased abundance of K10-acetylated histone H4 (H4K10Ac), K4-methylated H3, and variants of H2A and H2B (Respuela et al., 2008; Siegel et al., 2009; Wright et al., 2010; Thomas et al., 2009). Termination of transcription is also poorly understood (Martinez-Calvillo et al., 2004), but
the ends of DGCs (which are frequently, but not exclusively, convergent SSRs) display increased abundance of H3 and H4 variants (Siegel et al., 2009). A hypermodified base, J, also localizes to the boundaries of the DGCs (as well as telomeres, other repeats, and inactive Variant Surface Glycoprotein [VSG] expression units; see below) and influences transcription (Cliffe et al., 2010; Ekanayake et al., 2011). The T. brucei nuclear genome also has hundreds of VSG genes, switching between which is the basis of antigenic variation, a strategy for evasion of mammalian immunity. Transcriptionally silent VSGs are located proximal to telomeres in ~100 nuclear minichromosomes (Wickstead et al., 2004) and, more abundantly, in arrays in the subtelomeres of the megabase chromosomes (Marcello and Barry, 2007). To be expressed in the mammal, VSGs are recombined into 5–15 bloodstream VSG expression sites (BESs) at the telomeres of the megabase and intermediate chromosomes; only one BES is normally actively transcribed at a time (Horn and McCulloch, 2010). BES transcription is multigenic, with the VSG cotranscribed with approximately ten expression site-associated genes (ESAGs) (Hertz-Fowler et al., 2008). Another set of VSGs, termed the MVSGs, is expressed in the nonreplicating metacyclic trypanosome form, a transmission stage found in the tsetse fly vector. MVSGs are transcribed from ESs (MES) that lack ESAGs and are thus the only single-gene T. brucei transcription units (Ginger et al., 2002). All VSG transcription is mediated by RNA pol I.

Here, we present a global analysis of DNA replication initiation in T. brucei and show that DNA replication and transcription display remarkable levels of coordination, both positionally and functionally. We map ORC1/CDC6 binding sites and show that they localize to the boundaries of the transcribed DGCs in the chromosome cores. In contrast, we see high-density binding in the silent VSG array-containing subtelomeres. Mapping DNA replication initiation shows that early origins colocalize with a subset of the ORC1/CDC6 sites in the chromosome cores and display a spacing that is greater than in other eukaryotes. Finally, using RNA interference of ORC1/CDC6, we show that DNA replication functionally intersects with transcription because the absence of the initiator leads to increased levels of transcripts from the DGC boundaries and derepression of MVSG silencing.

RESULTS

Genomic Distribution of the Replication Initiator ORC1/CDC6
To address how DNA replication occurs in the unusual T. brucei genome, we first profiled the genomic binding sites for the candidate initiator protein ORC1/CDC6 in procyclic form (PCF; tsetse midgut stage) cells. A hemizygous ORC1/CDC6 T. brucei strain was generated in which one allele was deleted and the other modified to encode a C-terminal 12 Myc epitope-tagged ORC1/CDC6 variant (data not shown). Chromatin immunoprecipitation (ChIP)-chip was then performed using the myc tag, interrogating an ~385 K microarray covering the T. brucei TREU927 megabase chromosomes. Figures 1 and 2 show...
ORC1/CDC6-enriched sites in chromosomes 7 and 9, which are representative of all 11 chromosomes (Figure S1). As for ORC in other organisms (Eaton et al., 2010; MacAlpine et al., 2010), T. brucei ORC1/CDC6 ChIP enrichment (up to 4-fold) is low relative to that of other DNA binding proteins, such as modified histones (H4K10Ac is shown in Figures 1 and 2) or transcription factors (Siegel et al., 2010; Thomas et al., 2009). Applying a false discovery rate (FDR) of 0.2%, 953 ORC1/CDC6 binding sites were predicted, covering 640 Kbp (2.4%) of the megabase chromosomes. This revealed a striking, bimodal binding pattern, with dispersed ORC1/CDC6 binding in the interior of chromosomes (Figure 1) and denser binding at subtelomeres (Figure 2).

To analyze ORC1/CDC6 distribution globally, each megabase chromosome was demarcated into three categories (Figure 3A): “chromosome core,” the highly transcribed region containing housekeeping genes in DGCs as found in all chromosomes; “subtelomere-related,” a combination of nonfunctional and functional elements (e.g., ESAG [pseudo]genes, VSG-related genes, and RHS genes); and “VSG array,” which is also subtelomeric but has a more ordered arrangement, containing arrayed “cassettes” (Figure 2) composed of 70 bp repeats, VSG 5′-flanking sequence, VSG ORF (either functional or pseudogene), and VSG 3′ flank. The core (total size 22.3 Mb, 73% probe coverage) housed 38% (362) of ORC1/CDC6 binding sites (Figure 3A), with an average spacing of ~130 Kbp between adjacent members of 170 “unique” binding sites (those closer than 10 Kbp were grouped). In contrast 51% (484) of binding sites were in the VSG arrays (2.1 Mb, 61% probe coverage), at an average spacing of ~7 Kbp. Subtelomere-related sequence (1.3 Mb) is underrepresented by probes (29% coverage), and therefore, the 11% of binding sites (107) at ~12 Kbp spacing may be an underestimate.

Within the chromosome cores the ORC1/CDC6 binding sites clustered at the boundaries of the T. brucei multigenic transcription units. The promoters of these units have been inferred by H4K10Ac localization (Siegel et al., 2009), which identifies both divergent SSRs and intra-DGC transcription start sites. Thirty-eight percent of core ORC1/CDC6 binding sites (136 in total, mapped at FDR ≤0.2) mapped to divergent SSRs (Figure 3B), of which 62 (70%) out of the predicted 88 such loci showed evidence of at least 1 ORC1/CDC6 binding site. Fifty-four percent of ORC1/CDC6 sites (202 total, FDR ≤0.2) fell within DGCs, and virtually all localized to predicted transcription start sites: of the predicted 46 intra-DGC loci marked by H4K10Ac localization, 44 (96%) showed evidence of ORC1/CDC binding. Taken together, these data suggest an exceptionally strong association between ORC1/CDC binding and transcription initiation. Despite this close correlation in genome-wide localization (Figures 1, 3, and S1), a functional relationship between H4K10Ac modification and ORC1/CDC6 recruitment seems unlikely. We examined the localization of H4K10Ac by merging those of the predicted 367 binding sites that were closer than 10 Kbp, and discarding those that were <1 Kbp in size, yielding 158 unique binding loci. Of these, 108 were found within 15 Kbp of a predicted ORC1/CDC6 site, and 92 overlapped with at least 1 ORC1/CDC6 site. However, the two binding “footprints” do not overlap entirely. At 70% of ORC1/CDC6 sites within a DGC and proximal (within 10 Kbp) to H4K10Ac, the initiator binding site preceded the modified histone (examples can be seen at 0.35 and 1.25 Mbp in chromosome 7; Figure 1). In addition within divergent SSRs, where H4K10Ac localization is frequently seen as two distinct sites (e.g., 0.25, 0.71, and 1.0 Mbp in chromosome 7; Figure 1) (Siegel et al., 2009), ORC1/CDC6 binding was found centrally, between these sites. Thus, ORC1/CDC6 binding appears to localize close to H4K10Ac, but somewhat
upstream. It is also clear that the transcription initiation is not the sole determinant of ORC1/CDC6 binding: though only 7% of ORC1/CDC6 binding sites mapped to convergent SSRs, 24 of 64 such loci show evidence for ORC1/CDC6 binding (FDR < 0.2; 40%), and some clearly act as early replication origins (see below). It therefore seems likely that ORC1/CDC6 binds to all transcription boundaries, but less efficient localization to terminators is not always detected by ChIP-chip.

To investigate the high density of ORC1/CDC6 binding in the VSG arrays, individual sequence components (VSG 5’ flank, VSG ORF, and VSG 3’ flank), comprising 69% of the arrays, were demarcated, and the extent of binding was compared (Figures 2 and 3C; FDR ≤ 0.2). VSG 5’ flank and VSG ORF comprise the majority (40% and 50%, respectively) of VSG array sequence. ORC1/CDC6 sites were overrepresented in the 5’ flank (53% of sites) and underrepresented in the VSG ORF (15%), whereas VSG 3’ flank sequences bound the protein at a frequency commensurate with their contribution to the overall arrays. Despite these nonrandom distributions, we could not identify a consensus binding sequence (Figure S2), suggesting that ORC1/CDC6 binding in the VSG arrays, like in the core, is not a sequence-specific feature. Indeed, because the VSG...
arrays are not normally transcribed and lack known promoters and terminators, it is conceivable that ORC1/CDC6 localization in this region of the genome is functionally distinct from that in the core.

Identification of Early Replication Origins in the *T. brucei* Genome
To address whether or not the mapped ORC1/CDC6 binding sites correspond with replication origins, we employed marker frequency analysis (MFA). This technique compares the copy number of marker sequences in replicating and nonreplicating cells; origin proximal sequences will be overrepresented compared with origin-distal markers during replication. Early-mid S phase and G2-enriched populations of PCF *T. brucei* cells were prepared by cell sorting (data not shown). Following library preparation and Illumina sequencing, sequence reads were mapped to the *T. brucei* TREU927 genome; read abundance was then calculated in 2,500 bp windows as the ratio of replicating to nonreplicating DNA, revealing peaks of varying amplitude (Figure 4). We apply the following nomenclature to these

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**Figure 4. Mapping Replication Origins in the *T. brucei* Nuclear Genome**

Graphs show the distribution of replication origins in the megabase-sized chromosomes (numbered 1–11; numbers on the x-axis denote sizes in Mbp), determined by the extent of enrichment of DNA in S phase relative to G2. For each chromosome the top track displays CDSs (gray bars), and the relative position of predicted ORC1/CDC6 binding sites (color coding described in Figure 1) and centromeres (green ovals for chromosomes 1–8). The second track displays localization of H4K10Ac, derived from ChIP-seq data of Siegel et al. (2009), identifying positions of likely transcription start sites. The graphs in the lowest track display the ratio of the read depth between the S phase and G2 samples, where each dot represents 2,500 bp. The color of the dot refers to the mapping quality score at that position: black shows a score ≥50, whereas a heatmap of blue to red shows a score of 49–0. See also Figure S3.
in contrast, were found at intra-DGC sites where transcription both initiates and terminates. In all cases the slope of the peak was shallower when the direction of replication follows transcription and steeper when replication direction is counter to that of transcription. Taking the example at OBR:8:0.86 (Figure 4), replication proceeds 2.1-fold faster when traveling in the direction of transcription than counter to it. This indicates that T. brucei does not avoid head-on encounters between transcription and replication (Figure S3) and that such encounters impose a greater impediment to replication than when the two processes are co-directional (Merrikh et al., 2011). Fourth, there is a perfect correlation between the position of the mapped centromeres of T. brucei chromosomes (to date, described in chromosomes 1–8) (Obado et al., 2007) and the MFA peaks of highest amplitude. Thus, T. brucei replicates centromere-proximal loci early in S phase. Fifth, 39 of the 42 MFA peaks center on ORC1/CDC6 binding sites, supporting the proposed function of ORC1/CDC6 as an initiator protein. Nonetheless, it is clear from the two data sets that a large number of ORC1/CDC6 binding sites (~80% of the 170 sites grouped with 10 Kbp) do not correlate with active origins of replication (Figure 3). Although some of these sites may account for late-firing origins, there are many instances where binding sites lie within regions replicated from adjacent OBRs, suggesting potential redundancy.

Detailed examination of the positions of the OBRs reveals a striking relationship between the localization of origins and the transcribed domains. Most obviously, 19 of the peaks centered on divergent SSRs, correlating with sites of promoter activity. An additional three peaks were found at convergent SSRs, which were not always scored as significant ORC1/CDC6 binding sites (e.g., OBR:8:1.89) (Figure 4), consistent with our suggestion that the initiator binds transcription termination sites. Of the remaining 20 peaks, 2 were found in regions with a more broken distribution of genes between the 2 strands and 18 within DGCs, all at regions of H4K10Ac enrichment, indicative of transcription initiation and termination sites. Thus, all the observed origins coassociate with the boundaries of transcription.

**ORC1/CDC6 Influences Transcription**

In light of the association of T. brucei ORC1/CDC6 binding sites and origins with the boundaries of transcription units, and also the extremely high density at subtelomeric regions, we wished to address a possible role in transcriptional processes. We therefore examined mRNA abundance after ORC1/CDC6 RNAi. We first looked directly for a role in VSG expression by quantitative RT-PCR (qRT-PCR) (Figure 5). In T. brucei strain EATRO795 (Ginger et al., 2002), four MVSgs and promoters have been detailed and shown to be transcriptionally repressed in PCF cells. We therefore modified these cells to allow transcriptional alteration of RNAi and introduced an ORC1/CDC6 RNAi construct. As described previously by Godoy et al. (2009), inducing RNAi against ORC1/CDC6 led to only modest growth inhibition after 96 hr, despite loss of ~70% of ORC1/CDC mRNA (data not shown). At the same time point post-RNAi induction, significant increases (5- to 13-fold) in MVSg mRNA levels were seen (Figure 5), suggesting that loss of ORC1/CDC6 leads to derepression of the MESs. To ask if this
was true also for the BESs and in bloodstream form (BSF; mammal-infective) parasites, we analyzed RNAi in T. brucei strain Lister 427, where 14 distinct BVSGs in 15 BESs have been cataloged by Hertz-Fowler et al. (2008). In PCF cells of this strain, ORC1/CDC6 RNAi (70% mRNA loss) led to the same slow growth as in EATRO795 PCF cells (data not shown). However, derepression of the normally silent BESs was much less marked than seen for MESs (Figure S4A): of five BVSG genes examined, only three displayed significantly increased mRNA abundance, and only by 1.5- to 2.5-fold. In the BSF Lister 427 (90-13) cells used here, a single BES (427-2, expressing VSG221) is maximally transcribed, and the others are silent. In this life cycle stage, ORC1/CDC6 RNAi has a more profound effect on T. brucei survival, with cell death after 36–48 hr (Tiengwe et al., 2012). For this reason, BVSG mRNA abundance was measured 12 hr post-RNAi induction, at which point ~40% loss of ORC1/CDC6 mRNA was seen. No evidence was found for derepression of the silent BESs (Figure S4B).

To address whether the aforementioned response is due to locus-specific functions of ORC1/CDC6 or reflects a genome-wide influence on gene expression, we used RNA sequencing (RNAseq) to compare mRNA abundance in PCF Lister 427 cells 96 hr post-RNAi induction relative to uninduced cells. Read abundance was calculated in 250 bp windows across the genome and the extent of increased mRNA in the ORC1/CDC6 RNAi-induced sample relative to the uninduced sample examined as a function of position in the TREU927 genome. In many regions, transcripts were seen from loci where there was no detectable gene expression before ORC1/CDC6 RNAi; this is shown in Figure 6 for chromosome 10, which is representative of the whole genome (Figure S5). We also tested for loci that showed changes in mRNA abundance, from a baseline of detectable mRNA before RNAi, with strikingly similar findings (chromosome 10 is shown in Figure 6). These analyses revealed that ORC1/CDC6 RNAi had a marked, nonrandom effect on global gene expression, with increased mRNA abundance for genes positioned proximal to the boundaries of the transcription units. These changes matched closely the localization of ORC1/CDC6 protein, with increased mRNA from the vicinity of divergent SSRs (e.g., ~0.42 and 1.22 Mbp on chromosome 10), convergent SSRs (e.g., ~1.40 and 1.80 Mbp), and intra-DGC initiation/termination sites (e.g., ~0.80 and 2.35 Mbp) (Figure 6). The increased mRNA abundance was not greater at the regions of early origin firing (e.g., four origins are found in chromosome 10; OBR:10:0.80, OBR:10:1.25, OBR:10:1.92, and OBR:10:2.65; Figure 4), suggesting that it is not linked to replication.

The changes in transcript abundance did not normally result from elevated expression of genes within the DGCs (and proximal to the ends) but resulted mainly from increased abundance of RNA sequence reads upstream and downstream of the transcription start and stop points. However, in some cases, ORC1/CDC6 depletion did appear to undermine functionally significant gene expression controls (Table S1; genes showing significant changes in mRNA abundance; >1.4-fold), with increased transcript abundance of genes positioned at the end of both RNA pol I (e.g., PAG and GRESAG2) and pol II transcription units (e.g., ISG65, numerous PAG-like genes, and a putative iron/ascorbate reductase-encoding gene cluster). Expression of the ISG and PAG genes is known to be under transcriptional
However, sequence differences between the genic sequences, and not from the TREU927 (where the strain Lister 427 (which it was necessary to use for RNAi) and the VSG abundance after ORC1/CDC6 RNAi was also observed from control. Loss of ORC1/CDC6 may therefore result in readthrough is based on locus-specific termination or elongation PAG 192 Cell Reports control (Haenni et al., 2009; Leung et al., 2011), which at least for both transcription start and termination sites is elevated levels of transcripts at (clustered vertical black lines). The consequence of this lack of precision in (dotted horizontal lines). In addition, start site definition becomes less precise (red dotted lines), allowing a proportion of transcripts to elongate through them (dotted horizontal lines). In addition, start site definition becomes less precise (clustered vertical black lines). The consequence of this lack of precision in both transcription start and termination sites is elevated levels of transcripts at the boundaries of the DGCs.

control (Haenni et al., 2009; Leung et al., 2011), which at least for PAG is based on locus-specific termination or elongation control. Loss of ORC1/CDC6 may therefore result in readthrough transcription in all these cases. Increased RNA sequence read abundance after ORC1/CDC6 RNAi was also observed from the VSG arrays (Figures 6 and S5), which emanated from intergenic sequences, and not from the VSG CDS (data not shown). However, sequence differences between the VSGs of T. brucei strain Lister 427 (which it was necessary to use for RNAi) and TREU927 (where the VSG array genes have been positionally annotated) undermine attempts to map VSG RNAseq reads, and so we cannot discount the possibility that ORC1/CDC6 RNAi has global effects on VSG expression.

DISCUSSION

The data in this study reveal remarkably precise localization of the replication initiator ORC1/CDC6 to the boundaries of transcribed domains in the core of the diploid megabase chromosomes that constitute much of the T. brucei nuclear genome. In addition the data show that these ORC1/CDC6 binding sites act as replication origins, albeit with considerable redundancy, and that the coordination of transcription and replication is not simply positional. Following ORC1/CDC6 RNAi we show that transcript abundance is elevated upstream and downstream of the transcription units and occasionally affects gene expression within these units. ORC1/CDC6 therefore contributes to the global delineation of the transcription boundaries in T. brucei (Figure 7). Thus, analyzing this highly unusual genome has revealed an unprecedented level of interplay between transcription and DNA replication in a eukaryotic genome. It appears that both these crucial cellular reactions are delineated into discrete domains, most likely through the functional interaction and co-localization of the underlying machineries. We propose that this functional organization of the genome into modules is possible as a consequence of the near-exclusively posttranscriptional basis of gene regulation in kinetoplastid organisms and may have far-reaching consequences for genome maintenance. Moreover, our identification of the coassociation of highly active and early-firing replication origins with centromeres may contribute to the development of stably inherited extrachromosomal DNA molecules in T. brucei, thereby facilitating many forms of experimental analysis.

In fission and budding yeast, as well as in Drosophila melanogaster and Arabidopsis thaliana, origins have been identified through genome-wide mapping of the binding of ORC and MCM subunits by ChIP and by mapping sites where DNA synthesis begins by trapping newly formed replication forks or by isolating nascent leading strands (Gilbert, 2010). In mammals, ChIP of pre-RC components has not been reported, and therefore, origin identification has relied on initiation site mapping. Our work combines ORC1/CDC6 ChIP with MFA to define origins in T. brucei, adding information gleaned from this highly diverged protist to our knowledge of what constitutes an origin in eukaryotes, revealing both similarities and differences. In common with most other eukaryotes, we have not been able to identity consensus binding sequences for T. brucei ORC1/ CDC6 (data not shown), perhaps suggesting that epigenetic features underlie ORC1/CDC6 recruitment to origins (see below). Schizosaccharomyces pombe origins are found in AT-rich intergenic sequences (Dai et al., 2005; Hayashi et al., 2007), due to ORC DNA binding via an AT-hook domain on Orc4 (Chuang and Kelly, 1999). A preference for AT-rich sequences has also been reported for D. melanogaster ORC (MacAlpine et al., 2004), which additionally appears to localize predominantly at active promoters with open chromatin structure (MacAlpine et al., 2010), a feature of origins also observed in S. cerevisiae (Eaton et al., 2010). This appears compatible with T. brucei ORC1/CDC6, which shows preferential binding to genome regions that house the poorly defined promoters at the head of the multigene transcription units. These loci are marked by the deposition of variant and modified histones and display more mobile nucleosomes (Siegel et al., 2009). However, the association between origins and promoters is not absolute in eukaryotes. For instance in D. melanogaster a further study, mapping nascent strands, did not find enrichment at transcription start sites but instead at CpG-rich sequences frequently found in gene exons (Cayrou et al., 2011). CpG sequences may also be a feature of mammalian origins, which appear to be enriched at promoters, though are also found frequently within genes (Cadoret et al., 2008; Karnani et al., 2010; Cayrou et al., 2011; Sequeira-Mendes et al., 2009). Finally, A. thaliana ORC and CDC6 binding, and origin activity, have a preference for the 5’ body of genes (Costas et al., 2011). For T. brucei, ORC1/CDC6 displays localization to the ends of the transcription units, and these loci can act as origins, though with less efficiency than ORC1/CDC6 sites at the head of the transcription units. To our

Figure 7. Functional Interaction between DNA Replication and Transcription in T. brucei

A model of the effect of ORC1/CDC6 depletion on transcription and replication is shown. DGCs are shown as gray boxes, with origins of bidirectional replication as ovals. Transcription start sites are shown as vertical black lines, transcripts and their direction as horizontal black arrows, and termination sites as vertical red bars. Upon ORC1/CDC6 depletion, replication initiation efficiency reduces (small oval), and termination barriers become more porous (red dotted lines), allowing a proportion of transcripts to elongate through them (dotted horizontal lines). In addition, start site definition becomes less precise (clustered vertical black lines). The consequence of this lack of precision in both transcription start and termination sites is elevated levels of transcripts at the boundaries of the DGCs.
knowledge, origin activity at transcription terminator regions has not previously been seen.

Although our data reveal that replication and transcription in T. brucei functionally intersect, the basis for this is unclear, as is the mode of ORC1/CDC6 recruitment to origins. In A. thaliana, origins are associated with CG-poor sequences that are enriched for variants of histone H3 di- and trimethylated on lysine residue 4 (H3K4), as well as the H2A.Z variant (Costas et al., 2011). D. melanogaster ORC binding sites appear to be enriched in histone H3.3, and perhaps H2Av, variants (MacAlpine et al., 2010), whereas mono-methylation of H4K7 in mammals has been implicated in pre-RC assembly at origins (Tardat et al., 2010). In T. brucei, trimethylated H3K4 and H2A.Z each localize to transcription start sites (Siegel et al., 2009; Wright et al., 2010), perhaps suggesting some conservation in ORC localization with A. thaliana. However, the variant histones that have been mapped to date at transcription initiation and termination sites in T. brucei are distinct (Siegel et al., 2009), and our mapping and understanding of the function of all T. brucei histone modifications are undoubtedly incomplete (Figueiredo et al., 2009). We also cannot yet exclude other, nonchromatin modes of ORC recruitment to DNA (Donti et al., 2009). Thus, it is premature to say if epigenetic features that are common to the ends of the DGCs define ORC1/CDC6 binding and, indeed, are conserved with other eukaryotes. For the same reason further work will be needed to understand if the depletion of ORC1/CDC6 causes transcription changes indirectly, for instance due to changes in chromatin through loss of interaction, or if the T. brucei initiator plays a more direct role in transcriptional events (Sasaki and Gilbert, 2007).

A particularly striking feature of the replication profile of T. brucei lies in the relative paucity of replication origins. We estimate that fewer than 100 origins account for replication of the entire complement of megabase chromosomes of T. brucei. As discussed above, this is a far lower density of origins than seen in other eukaryotes. It is possible that the number of origins is constrained in part by the architecture of the megabase chromosomes, with their long DGCs. In this context it would seem intuitive that origins would localize to divergent SSRs, thus preventing clashes between transcription and replication machinery. Both bacteria and other eukaryotes have chromosome architectures that limit such conflicts. For example the rDNA loci in Escherichia coli are arranged in the same orientation as that of replication fork passage. Similarly, in yeasts and mammals specific replication fork barriers exist in the rDNA loci to prevent clashes between transcription and replication machineries (Labib and Hodgson, 2007). Interestingly, in trypanosomes we observe several examples where repulsion and RNA pol are in opposition, resulting in a marked slowing of the rate of replication fork progression (Figure S3). Thus, the low number of origins cannot solely be accounted for by architectural constraints. An alternative explanation lies in the possibility that trypanosomes branched early in the eukaryotic tree of life. Recently, Diffley has proposed that the increase in the number of replication origins required to replicate the increasingly large genomes that arose during eukaryotic evolution has resulted in selection for additional levels of control circuitry (Diffley, 2011). The apparent redundancy of control circuits in more complex eukaryotes reflects a fail-safe situation to ensure once per cell-cycle replication at any given origin. It is notable, therefore, that whereas trypanosomes possess cyclin and Cdk homologs, they may lack both the important Cdc7/Dbf4 and geminin/Cdt1 systems found in more recently branching eukaryas. Perhaps the limited control circuitry of trypanosomes has constrained the number of origins that can be employed to replicate their genome.

The small number of replication origins in T. brucei raises some further questions. The first of these relates to the excess of ORC1/CDC6 binding sites relative to origins. Even accounting for the fact that we have mapped to date only the origins that we estimate fire in the first half of S phase, it is likely that there are ~2-fold more ORC1/CDC6 sites than the 100 origins. Furthermore, it seems likely that some origins fire earlier, or more readily, than others, based on the variable sizes of the MFA peaks we map. It is well known in other eukaryotes that ORC binding sites are more abundant than origins and that a temporal order of origin firing exists (Masai et al., 2010; Douglass and Diffley, 2012). The basis for this remains unclear, but it is likely that early origins have a higher affinity for replication factors. In the context of the T. brucei genome, where it is likely that all promoters are equally active and initiate transcription constitutively, how differences emerge in the efficiency of ORC1/CDC6 binding or downstream replication events is unclear. The second question relates to replication of the VSG array-containing subtelomeres. Although we find exceptionally high density of ORC1/CDC6 binding in the VSG arrays, we see no evidence for origin activity. It is possible that, as in other eukaryotes such as Schizosaccharomyces pombe (Hayashi et al., 2007), these subtelomeric regions are late replicating and have not been captured by the MFAseq. Alternatively, the high-abundance ORC1/CDC6 sites may act as many dormant origins (Blow et al., 2011), perhaps because replication of these chromosome domains is distinct from the cores due to the lack of VSG array transcription. A final possibility is that ORC1/CDC6 provides a distinct, nonreplication function in this chromosome domain. Whether it acts as a silencer, as in yeast and mammals (Prasanth et al., 2010; Hickman and Rusche, 2010), is unknown, but this may be compatible with the increased expression on MVS genes after ORC1/CDC6 RNAi that we observe. Indeed, Orc1 in Plasmodium falciparum displays localization to telomeres (Mancio-Silva et al., 2008) reminiscent of what we describe here in T. brucei. Because both parasites populate their telomeres with genes central to antigenic variation, this perhaps suggests a common connection between immune evasion and the ORC machinery.

**EXPERIMENTAL PROCEDURES**

T. brucei PCF cells were of strain TREU927, Lister 427, or EATRO795 and were grown at 27°C in SDM-79 medium. BSF cells were of strain Lister 427 and were grown in HMI-9 medium at 37°C. To perform RNAi, T. brucei PCF strain 427 plew29-pLew13, and BSF strain pLew90-pLew13, developed by Wirtz et al. (1999), were used, constitutively coexpressing T7 RNA pol and Tet repressor. To generate a PCF derivative of EATRO795 in which RNAi could be performed, the individual pLew29 and pLew13 plasmids were introduced by transformation, and controllable expression was tested by monitoring expression of a reporter gene subsequently introduced under the control of...
the T7 RNA pol promoter (data not shown). To generate cells for RNAi, a fragment of ORC1/CDC6 was amplified by PCR (primers CTOL01 and CTOL02; sequences available on request) and cloned into the vector pZJM (Wang et al., 2000), where it is flanked by opposing T7 promoters and Tet operator sequences. The construct was digested with NotI, allowing integration into the DNA arrays after transformation. Transformant clones were selected with 10 μg/ml zeocin (PCF) and 2.5 μg/ml 1 phleomycin (BSF). To quantify levels of ORC1/CDC6 mRNA, primers CTOL07 and CTOL08 were used; GAP3 primers (CTOL27 and CTOL28) and tubulin primers (CTOL31 and CTOL32) were used as controls. To quantify levels of BSG mRNAs, primer pairs CTOL35/CTOL36, CTOL39/CTOL40, CTOL41/CTOL42, CTOL37/CTOL38, and CTOL33/CTOL345 were used to PCR amplify VSG13 (VSG427-13), VSG224 (VSG224/427-3), VSG800 (VSG800/427-18), VSG202 (VSG202/427-9), and VSG221 (VSG221/427-2), respectively. For MVSs, primers CTOL21/CTOL22, CTOL23/CTOL24, and CTOL25/CTOL26 were used to PCR amplify VSGs 1.2, 1.61, and 1.64, respectively. For all qRT-PCRs a master mix for 30 reactions was made in which each reaction had 12.5 μl of SYB Green PCR Master Mix (Applied Biosystems), 1.0 μl of each primer (300 nM stock), 9.5 μl of dH2O, and 1.0 μl cDNA (generated by SuperScript First-Strand Synthesis System for RT-PCR Invitrogen), according to manufacturer’s instructions. Reactions were run on an ABI Prism 7000 thermocycler and mRNA levels quantified from amplification according to the manufacturer’s instructions; conditions for all reactions were 50°C for 2 min, 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. Plasmids used for myc tagging of ORC1/CDC6, and the transformation conditions employed, are described elsewhere (Tiengwe et al., 2012).

ChiP and Microarray Hybridization

Microarray design was performed by Roche Diagnostics NimbleGen (USA) using a selected set of T. brucei TREU927 DNA sequence files obtained from the Pathogen Sequencing Group at the Wellcome Trust Sanger Institute (Cambridge), chosen to provide coverage of megabase chromosomes and some examples of telomeric sequences. The NimbleGen Tiling ChiP Service (design ID 19441, design name 090109_Tbru_LM_Chip) was used: 385,816 probes were tiled across all unique regions of the T. brucei genome at an average spacing of 61 bp. ChiP was performed from PCF T. brucei cells, essentially following Siegel et al. (2009). A total of 105 PF cells were used for each ChiP experiment in 40 ml of SDM/79 medium and were crosslinked by incubating them for 20 min at room temperature in 11% formaldehyde (in a buffer containing 50 mM HEPES [pH 7.55], 100 mM NaCl, 1.0 mM EDTA, and 0.5 mM EGTA). After crosslinking, glycine was added to the mix at a final concentration of 125 mM and centrifuged at 4,000 × g for 20 min at 4°C. The cells were then washed with 30 ml ice-cold PBS and centrifuged again at 4,000 × g for 20 min at 4°C. After the wash, the cells were resuspended in lysis buffer 1 (50 mM HEPES [pH 7.55], 140 mM NaCl, 1.0 mM EDTA, 10% glycerol, 0.25% Triton X-100, 0.5% NP-40, and protease inhibitors), vortexed thoroughly, and rocked on a platform rocker for 10 min. After rocking, the cells were centrifuged at 4,000 × g for 20 min at 4°C, and the supernatant was decanted. The cell pellet was next resuspended in lysis buffer 2 (50 mM Tris-HCl [pH 7.8], 200 mM NaCl, 1.0 mM EDTA, 1.0 mM EGTA, and protease inhibitor), and the vortexing and centrifugation were repeated. After decanting the supernatant from this step, the pellet was finally resuspended in 2 ml of lysis buffer 3 (10 mM Tris-HCl [pH 7.8], 100 mM NaCl, 1.0 mM EDTA, 0.1% Na-Deoxycholate, 0.5% N-lauroylsarcosine, and protease inhibitors). Next, the whole-cell lysates were sonicated for 60 cycles (30 s on/30 s off) using a Bioruptor (Diagenode), giving a DNA size range of 0.2–1.0 Kbp (as- determined by agarose gel). A total of 220 μl of elution buffer (50 mM Tris-HCl [pH 8.0], 1% SDS, 10 mM EDTA) was then added to the samples and incubated in a 65°C water bath for 30 min with vortexing every 2–5 min. The beads were then centrifuged for 1 min at 16,100 × g at room temperature. A total of 200 μl of the supernatant was removed after the spin step and transferred to a new microcentrifuge tube. This served as the eluate, from which DNA was reverse crosslinked from proteins. To do this, 3 vol of elution buffer was added to the “input sample” and mixed thoroughly. From here the “input sample” was treated like the eluted IP material. Both samples were reverse crosslinked by incubating at 65°C for ~9 h (not more than 15 h). After incubation, 8 μl of RNaseA (from a stock of 10 mg/ml-1) was added to each sample and mixed by inverting the tubes several times, and this was then incubated at 37°C for 2 h. Next, 4 μl of Proteinase K (from a stock of 20 μg.ml-1) was added to each sample and mixed by inverting the tube several times, and incubated at 55°C for 2 h. The DNA was then purified using a QIAAGen gel extraction kit and eluted with 60 μl elution buffer, following the manufacturer’s protocol. The purified DNA was then repaired using the Quick Blunting kit (NEB) according to manufacturer’s protocol. After the repair step, DNA was purified using the QiAgull PCR Purification Kit (QIAGEN), according to manufacturer’s instructions, and eluted with 41 μl of H2O. The purified DNA was then amplified using the Whole Genome Amplification kit (Sigma-Aldrich) and purified using the PCR purification kit from QiAgEN, both according to manufacturer’s instructions. A total of 5 μl of the eluted DNA was run on a 1.5% agarose gel, and the remaining material was quantified using a NanoDrop. For microarray analysis, ~5 μg of input and eluted DNA was used, and labeling, hybridization, and data acquisition were performed by NimbiGen. TbrORC1/CDC6-Myc IP DNA was labeled with Cy5 and input sample labeled with Cy3, and these were then cohybridized to the microarray. After cohybridization, data acquisition, and analysis by Nim- biGen, the scaled log2 ratio and peak data sets were aligned with the anno- tated genome sequence coordinates from T. brucei strain TREU927 using NimbiGen SignalMap software. For visualization the scaled log2 ratio data were manually adjusted to a minimum value of zero on the y axis of the plots, corresponding to an equal abundance of the Cy5-labeled sample and Cy3- labeled samples; positive log2 ratios correspond to enrichment of Cy5-labeled ChiP DNA relative to the Cy3-labeled input. To analyze the data, the scaled log2 ratio was normalized by searching for four or more probes whose signals were above specified cutoff values (percentage of a hypothetical maximum, which is the mean + 6 [SD]), ranging from 90% to 15%, using a sliding window of 500 bp. The ratio data were then randomized 20 times to evaluate the prob- ability of false positives, allowing the assignment of likely ORC1/CDC6 binding and sites, which were dependent on specified FDRs of ≤0.05 (5% of sites will be incorrect), 0.05–0.1 (5%–10% incorrect), and 0.1–0.2 (10%–20% incorrect).

MFA

PCF T. brucei TREU927 cells were used. To prepare cells for fluorescence-activated cell sorting (FACS), cells were grown in 200 ml of SDM-79 medium to a density of 5 × 106 cells.ml-1 and harvested by centrifugation at 500 × g for 20 min. The cell pellet was resuspended in 0.5 ml of PBS and fixed with 1% formaldehyde in PBS. After quenching the formaldehyde with glycine, cells were permeabilized with 0.01% Triton X-100 in PBS, then incubated for 1 hr at 37°C with 100 μg.ml-1 RNaseA, and stained with 10 μg.ml-1 propidium iodide. Cells were sorted by FACS using a Dako Cytomation Mo-Flo high-speed Sorter. A 488 nm laser at 100 mW was used, and emitted light collected through a 630/30 band-pass filter with a photomultiplier tube at 770V. Gating was set on side-scatter, then by the fluorescence measured on a linear scale. Sorted S phase and G2 cell fractions were centrifuged at 1,500 × g for 10 min. The cell pellet was resuspended in 300 μl of buffer ATL (QIAGEN) and incubated with 20 μl of Proteinase K solution (QIAGEN) at 65°C overnight to reverse crosslinking, after which DNA was extracted using a QIAamp DNA Micro Kit (QIAGEN) following the manufacturer’s instructions. The amount of DNA in each sample was determined using a Qubit 2.0 (Invitrogen), and 200 ng of each was sent for library preparation and Illumina sequencing. Paired-end 100 bp reads were obtained and mapped to the genome of T. brucei strain TREU927 (TrinityDB release 3.3) using Burrows-Wheeler Aligner (BWA) (Li and Durbin, 2010). The read depth for each sample was determined for each nucleotide position along each chromosome, by summing the reads that covered that nucleotide. The read-depth data for each position were then
binned into 2,500 bp sections, and the average value across each 2,500 bp bin was used to calculate the ratio of read-depth between the S and G2 samples, with the baseline for each chromosome normalized to one. For MFA by qPCR, primer pairs were designed to target amplicons of approximately 70 bp situated throughout chromosomes 4, 6, and 8 (primer sequences available on request). Reactions were prepared in quadruplicate using Power SYBR Green PCR Master Mix (Applied Biosystems). Each plate included the same control (an empirically determined region of chromosome 6 that showed no evidence for replication in the G2 sample) used to normalize the results and compare between plates. Reactions were run on an Eppendorf Realplex Mastercycler (in Oxford) and an Applied Biosystems 7500 Real-Time PCR system (in Glasgow), using the same two-step PCR-cycling conditions on each (10 min hold step at 95°C, followed by 40 cycles of 15 s at 95°C, 60 s at 60°C). The fluorescence intensity was recorded at the end of the annealing/extension phase in each cycle. For each amplicon the average Ct value was determined, and the relative amount of DNA present was calculated using the ΔΔCt method by Livak and Schmittgen (2001). Briefly, differences in the total amount of DNA present in each sample are determined by normalizing to a calibrator (the control region on chromosome 6), and the difference between the normalized values for S phase and G2 samples at the region of interest is used to determine the relative quantity of S phase DNA at that position.

**RNAseq Analysis of mRNA Abundance**

RNA was prepared from PCF Lister 427 cells 96 hr after tetracycline induction of RNAi against ORC1/CDC6 and from control cells that had not been treated with tetracycline. cDNA was then generated from polyA-selected RNA and Illumina Tru-Seq RNA libraries prepared, from which the samples were sequenced on an Illumina GAIIx (performed in the Sir Henry Wellcome Functional Genomics Facility, University of Glasgow). The Illumina reads from each sample were converted to Sanger-encoded quality scores and trimmed such that the median score for each read was >20. The reads were aligned to the genome using Bowtie (Langmead et al., 2009) with parameters that report the best 10 alignments for reads with multiple alignments but remove reads that map to more than 50 locations in the genome (-S -k10 -best -strata -m50). This allowed some sensitivity to families of closely related genes but did not allow them to be overrepresented in the further analysis. Gene expression values were calculated as FPKM (fragments per kilobase of exon per million reads mapped) with Cufflinks (Trapnell et al., 2010), using parameters that correct for read bias using the genomic background and reweight reads with multiple alignments (-N -u -b reference.fasta -G reference.gtf). Read depth per position in the genome was generated using the RNA-seq bam files using SAMTools (Li et al., 2009) (SAMTools mpileup-B) and awk. These were loaded into R (http://www.R-project.org/) and normalized using quantile normalization to make the read counts and distributions comparable. To detect those regions where mRNA was expressed and abundance changed, the ratio of read depth for the RNAi-induced to -uninduced samples was then calculated and log2 transformed. To detect those regions where there was no RNAseq reads in one sample, but where reads were detected in the other sample, the positions with zero in either of the samples were set to 0.00001; this highlighted peaks of expression. In each case, reads were binned into 250 bases along the chromosome, and the median depth for each bin was used as the center value. These data were then smoothed, using a running median with a window of 7 to remove some of the background noise, and visualized by SignalMap.

**Chromosome Annotation**

Megabase chromosomes were demarcated into discrete regions using a custom Perl script. Beginning from one chromosome-end, genes along the length of each megabase chromosome were analyzed sequentially, and upon arrival at the coordinates of the first VSG, a “VSG array” feature was generated. Identification of RHS, VSG related, and ESAGs (but not ESAG3 because this is also in VSG arrays) then generated a “subtelomere-proximal” feature. All hypothetical genes were omitted, and a “core” feature was generated that corresponds with the region encompassing all nonhypothetical genes, and gaps were then joined. Some short regions containing subtelomeric genes present at SSRs in the core of chromosomes were identified, corresponding with where chromosome fusions are believed to have occurred (El-Sayed et al., 2005), and these were annotated as subtelomere features, despite being in the chromosome cores. To demarcate the components of the VSG cassette, a distinct Perl script was used. Here, we first searched for 70 bp repeat features, and each repeat was annotated in the genome and manually checked. VSG ORF features were then looked for relative to the 70 bp repeats, and this annotation allowed the demarcation of the VSG 5’ and 3’ flanks.

**ACCESSION NUMBERS**

ORC1/CDC6 localization, MFAseq, and RNAseq after ORC1/CDC6 RNAi data are being archived at http://tritrypdb.org.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes five figures and one table and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2012.06.007.

**LICENSING INFORMATION**

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**REFERENCES**


