Forward Genetic Analysis of the Apicomplexan Cell Division Cycle in Toxoplasma gondii

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Apicomplexa are obligate intracellular pathogens that have fine-tuned their proliferative strategies to match a large variety of host cells. A critical aspect of this adaptation is a flexible cell cycle that remains poorly understood at the mechanistic level. Here we describe a forward genetic dissection of the apicomplexan cell cycle using the Toxoplasma model. By high-throughput screening, we have isolated 165 temperature sensitive parasite growth mutants. Phenotypic analysis of these mutants suggests regulated progression through the parasite cell cycle with defined phases and checkpoints. These analyses also highlight the critical importance of the peculiar intranuclear spindle as the physical hub of cell cycle regulation. To link these phenotypes to parasite genes, we have developed a robust complementation system based on a genomic cosmid library. Using this approach, we have so far complemented 22 temperature sensitive mutants and identified 18 candidate loci, eight of which were independently confirmed using a set of sequenced and arrayed cosmids. For three of these loci we have identified the mutant allele. The genes identified include regulators of spindle formation, nuclear trafficking, and protein degradation. The genetic approach described here should be widely applicable to numerous essential aspects of parasite biology.

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Introduction

Apicomplexans are highly successful protozoan parasites infecting a tremendous variety of vertebrate and invertebrate animals. In humans they are responsible for several important diseases, including malaria, toxoplasmosis, and cryptosporidiosis. A key to their success is their adaptation to a unique intracellular niche, which allows them ready access to nutrients while sheltering them from the immune system. Asexual parasite replication is restricted to this intracellular part of the life cycle, and intermediate stages of intracellular replication in many species lack the machinery to infect new host cells. It is therefore critical for the parasite to time its cell division and the formation of invasive forms to coincide precisely with host cell clegs. To adapt to a variety of specific host cell niches, apicomplexa have developed several specialized cell division modes. These division modes are based on a ‘flexible’ cell and division cycle program that can actively coordinate DNA synthesis and chromosome segregation, while at the same time suspending nuclear division and/or cytokinesis until the last step of the replication cycle (see [1,2] for recent detailed reviews of apicomplexan cell division and cell cycle control). Thus, re-initiation(s) of DNA synthesis prior to the completion of cytokinesis occurs naturally in these parasites. The molecular mechanisms that count the rounds of DNA synthesis and provide the proper timing of parasite budding remain one of the compelling mysteries of these parasites. Database mining for factors commonly associated with eukaryotic cell cycle control has identified an extensive set of candidate regulatory proteins in apicomplexan parasites (e.g. cyclins, CDKs, MAPKs [3–6]). While these findings predict that cell cycle checkpoints exist in Apicomplexa, they do not provide information about where checkpoints function or how these controls operate to safeguard the diverse strategies utilized by these parasites.

Here we describe the genetic analysis of the apicomplexan cell division machinery in Toxoplasma gondii. While the simple binary division of Toxoplasma tachyzoites (also termed endodyogeny) offers an attractive model system, we expect these studies to apply broadly to the replication of other pathogens in this phylum, such as Plasmodium, Eimeria, and Cryptosporidium, where our knowledge of the parasite cell cycle is equally deficient. Using chemical mutagenesis and a high-throughput replica assay, we have isolated a large collection of temperature sensitive (ts) parasite mutants. Our phenotypic analyses map these mutants to specific steps of the parasite cell and division cycle. To identify the underlying genes we have developed a robust complementation model...
Author Summary

Parasites of the phylum Apicomplexa cause numerous important diseases, including malaria, toxoplasmosis, and cryptosporidiosis. The ability to modify the genome of these parasites by transfection has been the technological key to unlock the biology of parasitic diseases at a molecular level. In this study we further extend the experimental possibilities for the study of apicomplexans by adopting a classic forward genetic approach for Toxoplasma gondii. We have developed protocols and reagents to generate large numbers of mutant parasites, screens to hone in on a subset of mutants of particular interest, and tools to identify the mutated genes that are responsible for the phenotype. Using this new approach, we have genetically dissected the way the parasite divides and multiplies within its host cell. This effort has yielded a series of highly informative mutants along the progression of the apicomplexan cell cycle and more than 20 genes involved in orchestrating parasite cell division. Importantly, this approach should allow unbiased genetic analysis of any part of parasite biology for which a screen can be devised using the Toxoplasma model.

employing cosmid transformation. This approach allowed us to link mutant phenotypes to specific point mutations in parasite genes.

Results

Production and Isolation of a Large Collection of Conditional Growth Mutants

A pool of conditional growth mutants was established by chemical mutagenesis using N-nitroso-N-ethylurea (ENU), an agent used successfully in the past to generate point mutations in the T. gondii genome [7–10]. ENU was applied at a dose inducing 60–70% parasite killing (measured through plaque assay; results not shown). We estimate this dose to induce 10–100 mutations per genome based on the incidence of mutations in hypoxanthine-xanthine-guanosine phosphoribosyltransferase (HXGPRT [11,12]), which we measured by following the emergence of resistance to the HXGPRT activated prodru 6-thioxanthine (10-5 in our experiments, also see [7]). To enable high-throughput screening we developed a well plate replica assay (see Figure 1 for a schematic outline of the strategy). Following mutagenesis, parasites were immediately cloned into 384 (or 96) well plates seeded with HFF cells to avoid competition with wild type parasites were immediately cloned into 384 (or 96) well plates seeded with HFF cells to avoid competition with wild type parasites and allowed to expand at the permissive temperature, whereas the replicate was placed at the restrictive temperature. Two approaches were applied to detect temperature sensitivity: visual microscopic inspection of wells (mutagenizing parent strain RH/hxgprt–) and measurement of fluorescence (mutagenizing the autofluorescent reporter strain 2F-1-YFP2 [13]). Fluorescence was measured after 4 days (40°C) and 7 days (35°C). The values were normalized against the 2F-1-YFP2-parent line included in each plate, and corresponding wells were compared for differential growth at permissive and restrictive temperatures using an automated script (a fluorescence increase below 20% was scored as a growth phenotype). Parasites from wells exhibiting growth at 35°C but not at 40°C were expanded, and temperature sensitivity was re-confirmed (see Figure 1B for examples; mutants with an identifier that starts with a letter were obtained through the fluorescence screen, those that start with a number through the visual screen).

In total we have identified 165 ts mutants from ~60,000 clones produced (see Table S1). Approximately 5% of the primary clonal isolates selected in the screens were confirmed by secondary analysis to show conditional growth arrest at the restrictive temperature (165/2960). False positives arose equally using either detection method and were due to the liberal selection for potential growth mutants and also as the result of replica-pin transfer failures (~5%). The overall frequency of confirmed ts mutants generated by our combined screens is lower (0.26%) than the value obtained in an earlier pilot screen (1.1%, [14]), which was much smaller in scale (~3,600 total ENU clones screened) and employed less stringent criteria for validating conditional growth. The ts mutants produced here display a lower reversion to wild type growth than our earlier study with >85% of clones in the current screen having reversion frequencies that are <10-6 and many isolates revert at <10-7 (see Table S2). The ts mutants generated by this and our earlier study [14] display almost exclusively conditional-lethality. However, a few examples of mutants where viability is retained following temperature shift were observed; as noted in other eukaryotic models [15,16] these are mostly G1 mutants (see below).

Phenotypic Analysis of Cell Cycle Mutants

Mutants were analyzed for uniform population changes in DNA content measured by flow cytometry (FACS). Phenotypes were further characterized by immunofluorescence assays (IFAs) identifying distinctive cellular and nuclear morphologies that developed at the restrictive temperature (see Table S2 for detailed individual descriptions of all mutants examined). Specific antibody reagents used in IFA analyses were directed against centrin as a marker for the number and position of centrosomes [17–19], membrane occupation recognition nexus (MORN)1, a marker for spindle morphology and budding [19,20], TgPCNA1, an essential element of the nuclear DNA replication complex [21,22], and inner membrane complex (IMC)1 and 3, components of the membrane skeleton that served as a budding marker [23–25]. The majority of parasite mutants examined had phenotypes characteristic of growth arrest in a specific cell cycle phase (i.e. >75% of the parasite population examined by FACS or IFA show a similar phenotype, see Figures 2, 3, and 4 for representative examples). In comparison to similar efforts in Saccharomyces [26,27], fewer general growth mutants were produced by our screens (the overall yield of ts mutants is also lower). However, several non-cell cycle mutants were identified including e.g. F-P2, a mutant with normal intracellular development but a severe invasion and egress defect (M. J. Gubbels and B. Striepen, unpublished data).

Based on their shared terminal phenotypes, mutants were readily classified into groups. These groups express defects in mechanisms active across the full spectrum of events in parasite replication. As expected, cell cycle mutants were isolated that possess a dominant haploid DNA content (1N) at the restricted temperature. The Sytox Green-FACS histograms for mutant 88A5 (Figure 2B, 34°C versus 40°C) are representative of this large group of G1 phase mutants. In addition to a 1N DNA content, G1 mutants possessed a single
nucleus and had no or few internal daughter forms at the time of growth arrest (data not shown, see Table S2). Outside this core set of phenotypic features, G1 mutants in our collection express diverse secondary characteristics that include alterations in terminal cell size (e.g. mutant 73C1, Figure 3B), differences in the number of cell divisions before growth arrest, and variations in intermediate temperature sensitivity. The G1 class is the only group where non-lethal ts mutants were isolated in our screen. Mutants 63H4 and 31F1 stop within a single cell division in the G1 phase when shifted to 40°C and can be held at this temperature for 24 hours without significant loss of viability (116% or 92% plaques formed compared to controls maintained constantly at the permissive temperature. Note that most mutants show poor recovery in this assay e.g. 88A5 or 87A10 with 1 or 4%, respectively). Two unusual mutants classified in the G1 group display complete loss of nuclear proliferating cell nuclear antigen (PCNA)1 staining at the restrictive temperature (e.g. mutant 124H2, Figure 3A).

A second group of ts mutants in the collection arrest upon shift to 40°C with an intermediate DNA content (>1N but <2N by Sytox Green-FACS) consistent with S phase arrest. Five ts mutants representing this class (e.g. 150B8, Figure 2C) arrest with a 30% increase over the haploid DNA content at the restrictive temperature; this phenotype is very similar to RH3K− parasites blocked by thymidine treatment in early S phase [29]. Arrested cells had large, centrally located nuclei and the DAPI and nuclear PCNA1 staining patterns observed (Figure 3C) were consistent with the S phase assignment [22]. Two additional S phase mutants deserve mention here: mutant 150B10 possessed a mid-S phase DNA content at the time of growth arrest (Figure 2D), while mutant 104A4 (Figure 2E) showed a unique bimodal distribution of parasites into equal 1N and 1.8N subpopulations.

Internal budding is a unique feature of apicomplexan replication, and >30% of the initial ts mutants characterized by IFA and FACS display defects in cytokinesis, some of these show simultaneous defects in karyokinesis. Mutant 64D5 is representative of five mutants that have defects in chromosome segregation. A subpopulation with a sub-1N DNA content by FACS (Figure 2F) as well as microscopic evidence for zoid formation (anucleate daughter cell [29], Figure 3D) are key characteristics of these mis-segregation mutants. A second group of M-phase mutants shows defects in the early stages of budding. Multiple small IMC1 staining structures formed but failed to develop fully when the parasites were shifted to the restrictive temperature (Figure 4A). Other budding mutants developed defects that interfered with the resolution of the mature daughters from the mother cell. Apparently this did not prevent a second round of cell division from unfolding suggesting cell cycle counting mechanisms were still active in this late budding mutant (bud-within-bud mutant 7A11, Figure 4D). The early and late budding mutant examples shown here maintained normal numbers of nuclei per parasite (one or two), whereas in other mutants isolated by our screens a catastrophic breakdown of the coordination between cytokinesis and karyokinesis occurred at the restrictive temperature. The three examples of this type of mutant shown here were promiscuous for chromosome re-initiation, which led to the formation of multiple or very large nuclei (42D6 and PO-B3, Figure 4B and 4C, and V-A15 see below). In each mutant, the mitotic spindle apparatus appears disorganized based on MORN1 antibody staining (all three mutants had similar staining as shown for mutant V-A15 below). Interestingly, similar abnormal budding and massive DNA over-replication has been observed in parasites where the spindle has been disrupted pharmacologically [29], suggesting that spindle defects might be a key feature of the uncoupling phenotype expressed by these ts mutants.

A Toxoplasma Cosmid Library for Phenotypic Complementation

To identify the genes underlying the mutant phenotypes, we employed phenotypic complementation using a wild-type genomic DNA library. Several T. gondii libraries have been generated for this purpose and some success has been
reported [25,30–32]. However, in our initial experiments we found that the established cDNA library strategies were not sufficiently robust to provide complementation on a regular basis with the ts mutant pool produced in this study. This is likely due to limitations of cDNA based plasmid libraries with respect to pool redundancy and insert size (many cell cycle factors are encoded by low abundant mRNAs). To establish a complementation model that is independent of gene size and differential transcript levels, we generated a RH genomic DNA cosmid library.

*T. gondii* does not maintain stable episomes precluding a simple shuttle of cosmids between parasite and *E. coli* to isolate complementing sequences. We therefore adapted a strategy based on cosmid insertion and rescue of a sequence tag (see Figure 5 for a schematic outline). Our library was constructed in ToxoSuperCos, a double cos-site plasmid based on the commercial SuperCos1 construct (Stratagene). However, several features were engineered into ToxoSuper-Cos to facilitate rescue after insertion. The *T. gondii* pyrimethamine resistance marker DHFR-TSm2m3 [33] was included to select for stable cosmid integration into the parasite genome preserving the backbone. This is important as plasmid rescue requires a bacterial origin of replication and a drug resistance marker for selection in bacteria ([34], see Figure 5). We chose a kanamycin resistance gene, as most parasites used for screening already contain plasmid DNA harboring ampicillin resistance genes (due to previous genetic engineering to introduce reporters and markers). To further facilitate rescue, a polylinker was incorporated providing a broader choice of restriction sites for genomic excision of rescue tags.

The resulting library consists of $1.25 \times 10^6$ independent cosmid clones providing ~900-fold genomic coverage. Using pyrimethamine selection and parasite plaque assays, the transformation efficiency was determined to be 0.3% (which is comparable to plasmid based transfection taking the larger size of the cosmid construct into account). Using the protocol described in the materials and methods section, 12-fold coverage of the *T. gondii* genome should be achieved in each transfection experiment. To validate these calculations, we transfected the *T. gondii* RH-hxgprt-deletion mutant with the cosmid library and selected for complementation by treatment with mycophenolic acid [11] and pyrimethamine. Five out of five electroporations resulted in the isolation of viable and inheritably mycophenolic acid resistant parasites. Clonal lines were established from three independent transfections, and the wild type HXGPRT locus (absent from the mutant but present in the library) was detected by PCR and sequencing of the PCR product in ten out of fifteen clones (Figure S1). Further PCR-based analyses indicated that complementation occurs through heterologous insertion of an additional wild type copy rather than homologous gene replacement of the mutant locus (even in experiments where no selection for the pyrimethamine marker is applied).

**Figure 2.** Distribution of Parasite Genomic DNA Content at Permissive and Restricted Temperatures for Selected ts Mutants

Mutant clones were grown at 34°C and 40°C for various times (16–36 h, based on individual phenotype) prior to harvest and ethanol fixation. DNA was stained with SYTOX Green dye and measured using a FACS calibur (BD). The cytometer was set to mode fluorescence and calibrated to the 1N population of asynchronous RH wild type parasites; red dashed lines reference 1N and 1.8N fluorescence peaks in the asynchronous controls. DNA fluorescence was measured in FL-1 linear scale (x-axis) and 10,000 events were collected for each histogram. In comparison to asynchronous controls (A), a selection of ts mutants (grown at 34°C and 40°C) that cell cycle arrest with different genomic contents are presented: (B) mutant 88A5 arrests with a predominant 1N DNA content (G1 phase mutant), mutants 150B8, 150B10, and 104A4 show altered DNA contents that are intermediate with respect to normal haploid or diploid genomic contents (C, D, and E), mis-segregation mutant 64D5 shows significant chromosome loss ([F], <1N), while mitotic mutant 11C9 arrests with a predominant diploid DNA content (G).

**Complementation Analysis of the Mitotic Mutant V-A15**

Having established that the ToxoSuperCos library robustly complements the HXGPRT mutant, we then tested its ability to complement the mitotic mutant V-A15. This mutant, which arrests with a predominant diploid DNA content, was isolated using mycophenolic acid as selection. Using the protocol described above, we transfected the mutant with the ToxoSuperCos library and selected for complementation by treatment with mycophenolic acid and pyrimethamine. Five out of five electroporations resulted in the isolation of viable and inheritably mycophenolic acid resistant parasites. Clonal lines were established from three independent transfections, and the wild type HXGPRT locus (absent from the mutant but present in the library) was detected by PCR and sequencing of the PCR product in ten out of fifteen clones (Figure S1). Further PCR-based analyses indicated that complementation occurs through heterologous insertion of an additional wild type copy rather than homologous gene replacement of the mutant locus (even in experiments where no selection for the pyrimethamine marker is applied).
to genetically rescue ts mutants. Here we describe experiments with mutant V-A15 in detail as an example, but note that subsequently numerous additional ts mutants have been successfully complemented with this library (see below).

Mutant V-A15 shows tight temperature sensitivity, with a modest growth delay at 35°C, significant inhibition at 37°C and a severe defect at 40°C (Figure 1B; the reversion frequency of this mutant was measured by plaque assay to be $10^{-7}$). IFA and FACS analyses revealed that growth inhibition is due to severe mitotic defects. V-A15 parasites shifted to the restrictive temperature failed to complete mitosis which led to polyploid nuclei and/or chromosome loss (Figure 6B, 6D, and 6F; note that increase in nuclear size goes along with an increase in cell size). While internal daughter buds were readily observed in parasites grown at the permissive temperature by staining with IMC3 (~25% of all vacuoles, Figure 6A and 6C), no clear IMC3 structures were formed at the restrictive temperature. Because features of this phenotype are consistent with a defect in the mitotic spindle, we analyzed mutants for the cellular distribution of MORN1, a marker of the nuclear spindle compartment (centrocone [20]). In parasites grown at the permissive temperature, the centrocone was clearly detected in each nucleus, while under temperature restrictive conditions MORN1 nuclear staining appeared disorganized or was entirely absent from some of the nuclei.

To identify the underlying genetic locus, mutant V-A15 was transfected with the ToxoSuperCos library in five independent electroporations, inoculated into confluent HFF cultures and allowed to recover for 24 hrs at 35°C. The flasks were subsequently transferred to 40°C to select for growth restoration (phenotypic complementation), and pyrimethamine was added to select for stable cosmid backbone integration. Stable temperature and drug resistant parasites emerged in four out of five flasks. To identify the complementing cosmid sequence, we used plasmid rescue of a sequence tag (see Figure 5 for an outline of this strategy). Genomic DNA of complemented parasite lines was extracted

![Figure 3. Representatives Examples of G1, S, and Mis-Segregation Mutants](image-url)
and digested with SpeI, HindIII or BglII, self-ligated overnight, and then electroporated into E. coli. Kanamycin resistant colonies were recovered for all four complementations using at least two out of the three enzymes used. Plasmid DNA was isolated and the genomic inserts were end-sequenced using the T3 primer. Three out of the four sequence tags (each from independent complementation experiments) mapped to a 30,000 bp locus on chromosome IX and a region of 6.7 kb could be identified that was shared between all three rescued tags (see Figure 7A; the fourth complement mapped to Chr VIII 1225114 bp minus strand, and was not studied further). The overlapping region contains a single predicted gene model (80.m02355) encoding a putative NIMA-related kinase with close similarity to Plasmodium falciparum Nek1 [6,35].

Figure 4. Representative Examples of Mutants with Defects in Karyokinesis and Parasite Budding

Several mutants characterized in our screens display defects in daughter budding when shifted to the restrictive temperature. (A) Mutant 27D12 shows multiple small IMC1 staining bodies (often closely associated with the plasmalemma, arrows), indicating abortive budding at an early stage. When arrested at 40°C, these normal-sized and shaped parasites had single or duplicated nuclei (but never formed syncytial cells containing multiple nuclei). In time-matched 34°C controls, parasites were able to complete two to three divisions. (B, C, and D) Errors in karyokinesis and parasite budding were detected in several mutants, which led to an uncoupling of these processes in some cases. Mutants 42D6 and PO-B3 show multiple abnormal budding structures and show progressive nuclear reduplication leading to syncytial cells with many nuclei (cells with variable nuclei numbers are indicated in PO-B3 image). Mutant 7A11 forms more complete daughter buds that are associated with at least one round of nuclear division. Premature budding before completion of the previous cytokinesis is observed in many vacuoles, which causes retention of a mother cell and nuclear structure (arrow). Two independent example vacuoles are shown for mutants 7A11 and PO-B3. See Figure 3 for details on antibodies used.

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Locus Confirmation and Identification of a Point Mutation in TgNek1

Rescue of a sequence tag as described above readily identifies a candidate complementing locus, yet only a small portion of the complementing sequence is cloned into plasmid, which is often not sufficient to independently confirm the result by re-complementation. However, the ToxoSuperCos library (along with a second library constructed by Dan Howe and David Sibley, pSCBle, http://toxomap.wustl.edu/) has been end-sequenced in the course of the T. gondii genome project, and the cosmid tiling (covering essentially the entire genome) can be viewed through ToxoDB (http://www.toxodb.org/ancillary genome browser). To test if the locus identified is indeed sufficient to complement the mutation, V-A15 was transfected with cosmid ToxPJ50 (containing the TgNek1 locus, Figure 7A) or ToxP932 (an unrelated control cosmid; Chr VIIb 1,484,413bp-1,521,843bp). Stable transgenics were established by pyrimethamine selection at the permissive temperature. These parasites were scored for temperature sensitivity by plaque assay (parasite growth is indicated by host cell lysis resulting in clear plaques in the fibroblast monolayer) and fluorescence growth assay (measuring the fluorescence of the yellow fluorescent protein (YFP)-YFP transgene expressed by the parasites [13]). ToxPJ50 restores robust growth at 40°C, while the control transgenic is indistinguishable from the mutant (Figure 7B–7G, note that complementation with ToxPJ50 already confers a modest growth advantage at 35°C, panel F).

The complementation data indicate that the TgNek1 locus is the site of the mutation causing the V-A15 phenotype, or alternatively, that this locus can act as a suppressor. To distinguish these two possibilities, we amplified the locus by PCR from both wild-type parent and mutant V-A15, transfected the mutant with each allelic gene fragment and scored for complementation by plaque assay. PCR product from wild-type complemented in three out of three independent experiments, while no growth was observed with the V-A15 derived gene fragment (Figure 8A–8D), suggesting that the mutation is localized within this locus. Importantly, complementation with the wild type PCR fragment also restores the FACS DNA profile of parasites grown at 40°C to the typical wild type distribution (Figure 8F). The 7.2 kb PCR fragments were sequenced on both strands and compared to the genome sequence as well as each other. A single base pair change distinguishes the sequence of the mutant from RH wild-type. This change of a T to C lies within the Nek1 coding region and changes a cysteine to an arginine in a highly conserved portion of the predicted protein (Figure 8E shows the mutation along with a short alignment of Nek1 from T. gondii and P. falciparum). Taken together, these data indicate that a point mutation in TgNek1 is responsible for the severe mitotic defects observed in this mutant.

Cosmid Complementation of a Variety of T. gondii Cell Division Mutants

Encouraged by our success in complementation analysis of mutant tsV-A15, we have broadened our efforts to ultimately identify the genes affected in all of the 165 ts mutants isolated in this screen. Figure 9 summarizes phenotypic and genetic analyses for the first group of 41 mutants (7 additional mutants showed excessive reversion frequencies and were excluded from further analysis). While these studies are still ongoing, thus far, we have observed complementation (indicated by asterisks) for 22 out of 24 attempted mutants, and a candidate locus has been identified by marker rescue for 18 mutants (see Table 1). For 8 mutants the locus has been independently confirmed by successful re-complementation using the respective arrayed cosmid, and for three genes the
The mutant allele has been identified by sequencing as described above for V-A15 (underlined in Figure 9; this data is summarized in detail in Table 1).

The genes identified through our mutant analyses encode proteins with potential functions in a wide array of parasite cell cycle mechanisms. Eleven of the 14 *T. gondii* genes summarized in Table 1 have clear homologs in other apicomplexan genomes as identified by OrthoMCL. In several cases, they represent orthologs of known cell cycle factors from other eukaryotes. Orthologs of the NimA-related kinase that complements V-A15 have well described roles in centrosome biology and mitotic entry [35–38]. Other examples include the *Toxoplasma* protein encoded by 27.m00873, which appears to belong to the Sac3/GNAP family. The SAC3 gene product in yeast is a nuclear factor that is required for mitotic progression. Defects in this gene cause errors in yeast budding and mitotic delay [39], which compares with the failure of mutant 118G4 to properly progress into mitosis and budding. An AAA-ATPase (44.m0215) is present in the locus complementing S phase mutant 104A4. AAA-ATPases fold and unfold proteins and often act as gatekeepers of protein degradation [40]. Mutations in AAA-ATPases (most notably cdc48) have been shown to result in mitotic arrest [41]. A homolog of the nuclear actin ARP4a is present in the rescue locus of mitotic uncoupling mutant 20C2. Nuclear actins are involved in transcriptional control and DNA repair and are required for stable attachment of the kinetochore to the mitotic chromosome [42]. Mutations in ARP4a cause defects in the intranuclear spindle and lead to an arrest of the yeast cell cycle in the G2 and mitotic phases.

A second set of genes encodes products that harbor protein domains often found in regulatory proteins but which otherwise appear to be unique to Apicomplexa. Gene 583.m05476 features a TBC domain, an activator domain known to regulate rab-like GTPases and is a motif also found in the yeast spindle factor BUB2p [43]. Two mutants, 42D6 and PO-B3, were complemented by proteins harboring RCC1-domains (regulator of chromosome condensation, 25.m01896 and 72.m00409). RCC1 proteins control nuclear transport and mitotic progression through nucleotide exchange of ran-GTPases [44]. Interestingly, another unrelated *T. gondii* RCC1 protein was recently identified as a non-essential protein that when mutated attenuates the virulence of Type I strains in the mouse model [44].

Gene 20.m03766, which complements the G1 mutant 109C6, contains an RNA recognition motif (RRM) present in RNA splicing factors, among others. Key features of the mutant 109C6 phenotype share similarities to the defects observed in yeast splicing mutants [45], and epitope tagged TgRRM-protein exclusively localizes to the tachyzoite nucleus (not shown) consistent with a putative role in pre-mRNA splicing. Sequencing of the RRM-protein allele from mutant 109C6 reveals a single base change that alters a highly conserved tyrosine residue in the RRM motif (169-Y to N),
indicating that this mutation is likely responsible for the G1 arrest. In support of this hypothesis, we were unable to complement mutant 109C6 with a gene fragment carrying the mutant 20.m03766 allele.

**Discussion**

Mutant analysis and genetic complementation are powerful strategies to link specific genes to biological pathways [46–48]. Efforts to develop these methods in *Toxoplasma* have resulted in an episome-based protocol [49] and an insertion based approach [31]. In the latter, phage recombination [50] was used to mobilize cDNA fragments integrated into parasite transformants [30–32]. While these earlier methods achieved some success, the redundancy inherent to cDNA libraries limited the complementation success (B. Striepen and M. White, unpublished data). The genomic-DNA based approach introduced here benefits from the large inserts carried by cosmid vectors to deliver genes in their natural chromosome organization. As a consequence of these improvements, our success rate for genetic rescue of ts growth mutants has improved dramatically. Here we report on the complementation of a diverse selection of mutants (20 mutants, with a failure rate, 5%). The genes identified in these experiments represent a wide range of coding and genomic sizes. Consistent with a mostly regulatory role of their products, the level of transcript for these genes is generally modest based on microarray analysis of tachyzoite gene expression (see summary in Figure S2).

One of the mutants complemented in this study (11C9) was the subject of an earlier cDNA-based complementation experiment that yielded a suppressor (TgXPMC2) of the genetic defect in this mutant [32]. Multiple complementation experiments of 11C9 using the cosmid library have repeatedly identified a single *Toxoplasma* gene, 50.m03077 (Table 1), which was likely underrepresented in the previously used

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**Figure 7. Cosmids Complementing Mutant V-A15 Map to the Locus of the *T. gondii* NimA Kinase Nek1**

Mutant V-A15 was transfected with the ToxoSuperCos library and selected as described in Figure 5. (A) Sequence tags of complementing cosmids (from three independent transfections) were rescued and sequenced; they map to a locus on chromosome IX and show an overlapping region of 7,610 bp containing gene model 80.m002355 annotated as NimA related kinase. (B) Mutant V-A15 was transfected with cosmids ToxpJ50 (covering the identified locus on Chr. IX, [A]) or control cosmid Toxp930 (Chr. VIIb) and selected for pyrimethamine resistance at the permissive temperature. The resulting stable transgenic populations were tested for temperature sensitivity by plaque assay (B–E) and fluorescence assay (F and G). Note that ToxpJ50 complements while Toxp930 does not (as the mutant these transgenics show a slight growth defect even at the permissive temperature, resulting in smaller plaques [C] and slower increase in fluorescence [F]).

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A Point Mutation in T. gondii Nek1 Is Responsible for the Temperature Sensitive Cell Division Defect in Mutant V-A15

The locus of TgNek1 was amplified from wild type (A) and V-A15 (B) genomic DNA in three independent reactions. V-A15 was transfected using these PCR products and incubated at the restrictive temperature and evaluated by plaque assay. WT PCR product rescued in three out of three independent transfections (IC, a single representative example shown here), while no plaques were observed using V-A15 PCR product (D). Both PCR products were sequenced on both strands. (E) A single nucleotide change was observed changing a cysteine to an arginine codon in a conserved segment of the Nek1 coding sequence (a limited alignment with the P. falciparum Nek1 gene is shown). (F) The DNA profile of mutant V-A15 and its complement (derived from the transfection described in (C)) was evaluated at the 34°C and 40°C. Note that complementation restores the wild type profile.

cDNA libraries due to its large size and low expression level (<100 units average fluorescence intensity in tachyzoite microarrays for 50.m03077 versus ~15,000 units for GRA-1, which was the promoter source used to drive expression in the cDNA libraries [31,32]). Likewise, we have not re-isolated TgXPMC2 by cosmid complementation. Because gene expression from cosmids relies on native regulatory regions, we believe there is a higher likelihood that the genes identified through this approach will represent the defective gene rather than a suppressor. In the three ts mutants (109C6, VA-15, and FV-P6) where this question was examined so far sequencing and functional testing of the corresponding mutant alleles confirms this prediction. In summary, the cosmid system provides robust complementation for a broad range of genes. Furthermore, the identified loci can be readily biologically validated taking advantage of an extensive set of end-sequenced and tiled cosmids that provide essentially full genome coverage. The protocols and reagents developed in the course of this study should allow future forward genetic analysis of any essential aspect of parasite biology for which a mutant screen can be devised.

Tachyzoite growth rates differ dramatically among parasite strains and growth rate is a key virulence determinant in T. gondii [22,51]. Despite the obvious importance of growth control, how the parasite regulates growth and cell division remains largely unknown. A series of defined biochemical controls and checkpoints regulating progression through one cell cycle phase to the next have been established for a variety of eukaryotic models [52–54]. The catastrophic break down of cell cycle coordination observed in T. gondii in the course of certain drug treatments has lead to the hypothesis that there might be significantly fewer cell cycle controls in this microorganism [29,55]. By contrast, the phenotypic groups that have emerged from the collection of conditional growth mutants described in this study support the notion of specific mechanisms and checkpoints. For example, two ts mutants were isolated that reversibly arrest in the G1 phase when shifted to 40°C (mutant 63H4 and 31F1). The presence of such a natural G1 checkpoint is further supported by the observation that end-stage differentiated parasite forms (sporozoite and bradyzoite) show a uniform haploid DNA content [28,56], as do parasites that have been treated with the G1 phase inhibitor pyrrolidine dithiocarbamate [57,58]. Tachyzoites released from this drug block, grow synchronously through at least two division cycles, indicating that pyrrolidine dithiocarbamate is likely acting on the same G1 checkpoint affected in our mutants.

Another important checkpoint controls entry into S phase, and in Saccharomyces, this checkpoint (called START) also controls the initiation of spindle formation and budding [59]. We have previously argued that the tachyzoite cell cycle likely has a similar checkpoint based on the observation that dNTP depletion arrests tachyzoite growth at the G1/S boundary (1N DNA content, centrosomes largely duplicated but not yet separated [22,28]). We have isolated five ts mutants (e.g. mutant 150B8) that display a very similar phenotype at the restrictive temperature, suggesting that the G1/S transition is an important restriction point in the tachyzoite cell cycle as it is in yeast.

Mitosis in Apicomplexa has several unique features: the nucleus remains intact, the intranuclear spindle(s) reside in a peculiar elaboration of the nuclear envelope the so called centrocone, and daughter cells are scaffolded as internal buds which develop in close proximity and likely under the control of the extranuclear centrosomes [1,60]. There is significant evidence for the tight regulation of mitotic events in tachyzoites from three groups of mutants in our collection. These mutants arrest either in mitosis (11C9), display defects
in chromosome segregation (5 mutants), or loose the coordination of karyokinesis with cytokinesis at various stages in the replication timeline (12 mutants).

Mitotic mutant V-A15 becomes both aneuploid and polyploid at the restrictive temperature and fails to initiate internal budding. The gene affected in this mutant is an NIMA-related serine/threonine kinase (Nek). This kinase family was first identified as essential for division in *Aspergillus nidulans* [61] and its members have since been identified as cell cycle regulators throughout eukaryotes [62,63], including protozoa [35,36,64]. Neks have roles in microtubular dynamics in cilia, mitotic spindles and centrioles [63,65,66]. Consistent with the known roles for NIMA-related kinases, the mutation in V-A15 leads to defects in the spindle apparatus (reflected in the loss of MORN1 organization) and this causes chromosome mis-segregation. Apicomplexa encode a family of related NEK proteins, and in *P. falciparum* these genes have been shown to be expressed in a developmentally regulated fashion, making it likely that NEKs are critical to fine-tuning the cell cycle to different life-cycle stages and host cells.

At the non-permissive temperature mutants 42D6 and PO-B3 are promiscuous for nuclear reduplication leading to the formation of syncytial cells with multiple nuclei. Daughter budding is also abnormal in these mutants and uncoupled from the controls that ensure proper nuclear sorting into each daughter. Two distinct RCC1 domain proteins were found to rescue these mutants (25.m01896 and 72.m00409). In other eukaryotes, RCC1 domain proteins interact with Ran-GTPases to regulate spindle assembly as well as other mitotic progression controls through modulation of nuclear trafficking [67]. Like mutant V-A15, mutant PO-B3 (and also uncoupling mutant 42D6) looses MORN1 organization at the restrictive temperature pointing to a potential spindle defect (data not shown). Overall the phenotype of this mutant, as well as several other members of the uncoupling class produced here (e.g. 42D6, 20C2, and 7A11), are similar to the abnormal daughter budding and the induction of unregulated nuclear replication associated with the disruption of the parasite spindle by pharmacological microtubule ablation [29]. Collectively, these observations indicate that proper control over chromosome copy number and budding in Apicomplexa might critically rely on an intact intranuclear spindle or associated structures. In this context it is important to note that the unique centrocone structure that conducts the apicomplexan spindle into the nucleus appears to persist throughout the cell cycle at least in some Apicomplexa [20,68]. This model is further supported by preliminary electron microscopy studies of mitotic mutant 11C9, which upon temperature arrest retains an intact spindle and early daughter scaffolds ([32] and S. Halonen and M. White, unpublished data) and does not undergo nuclear reduplication as is seen in mutants 42D6 and PO-B3. Thus, these models predict that rather than the absence of cell cycle controls in the Apicomplexa, mitotic control mechanisms might require a strict physical context associated with the centrosome and/or the centrocone of the parasite nucleus. Breaking this physical context by drug treatment [29], overexpression of a centrocone structural component [20], or mutation (as in the uncoupling group of *ts* mutants) results in catastrophic loss of regulation. There is considerable precedence for spatial control of cell cycle checkpoint proteins through compartmental exclusion as well as physical tethering of factors to the centrosomes and spindle structure [69]. Strict compartmentalization of the cell cycle machinery could be a key to the cell cycle flexibility observed in these parasites. Further work is needed to validate this hypothesis. However, the large collection of mutants and genes identified in this screen provides an important pool of validated candidates for mechanistic dissection. Studies that link parasite cell cycle control to the adaptation to specific host cell niches and pathogenesis will be of particular interest.

**Figure 9. Summary of Phenotypic Classification of *ts* Cell Cycle Mutants**

A schematic outline of the tachyzoite cell cycle (60% G1, 30% S, and 10% M-phase) is shown at the top. Nuclei are shown in grey, microtubular structures in red (conoid, centrosome, and spindle; subpellicular microtubules were omitted for simplicity) and the inner membrane complex in purple. Mutants successfully complemented at this point in time are denoted by an asterisk.

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Materials and Methods

High-throughput isolation of temperature-sensitive mutants. Parasites were grown in human foreskin fibroblasts (HFF) as described [70]. All transgenic and mutant parasite lines are derivatives of the RH parasite line.

The non-fluorescent, visually screened mutants were generated from a RH-hxgprt-parasites parent line. Parasites grown overnight at 37°C in 150 mm plates are mutagenized with 400 l g/ml ethyl-nitrosourea (ENU) for 4 h at 37°C (70% killing) [9]. The plates were washed to remove the mutagen and harvested by needle-passage and purified by filtration. Purified parasites are diluted to 40 parasites/ml and 0.15 ml well plated directly into 96-well plates, which yields ~33 single plaques per plate (Poisson distribution.

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Table 1. Summary of Cosmid Complementation for a Selection of Distinct ts Mutants

| Mutant Phenotype | Clone Designation | Rescued Chromosome Locus* | Cosmid Clone Recomplementation Growth ++/ No Growth – | Candidate Gene(S) or Confirmed Gene | Apicomplexan Orthologs |
|------------------|-------------------|--------------------------|-------------------------------------------------|------------------------------------|-----------------------|
| G1 phase mutant  | 109C6             | Chr. VIIa                | TOX16F5 –                                      | 20.m03766h,c                        | PF13_0318             |
| Quick arrest     |                   | Nucleotide pos. 2,771,038 to 2,813,240 | PSBL5C6 + +                                  | RRM motif splicing factor? | cgd1_1070             |
|                  |                   |                          | PSBLF75 –                                      |                                    |                       |
|                  |                   |                          | PSBM7M1 + +                                    |                                    |                       |
| G1 mutant†       | 73C1              | Chr. XI                  | PSBL163                                       | 35.m00891                            | PFF 1345w             |
| Giant cell arrest|                   | Nucleotide pos. 6,071,712 & 6,071,100 | TOXO087                                       | Checkpoint shuttle regulator,(transportin) | cgd8_3440            |
|                  |                   |                          |                                               |                                    |                       |
| START mutant?    | FV-P6             | Chr. VI                  | TOXOA93 –                                     | 49.m00063h,c                        | PF08_0124             |
| G1/S phase boundary arrest | | Nucleotide pos. 3,182,879 & 3,223,879 | TOXOD43 –                                    | Transcription factor related to cactin | cgd5_1690             |
| S phase mutant | 60E5              | Chr. XI                  | PSBL314                                       | 583.m05476                           |                       |
| 1.3N arrest      |                   | Nucleotide pos. 2,542,112 | PSBL329                                       | TBC Domain-BUB2p-spindle factor     |                       |
|                  |                   |                          |                                               |                                    |                       |
| S phase mutant†  | 104A4             | Chr. VIII                | TOXP129                                       | 44.m02615                            | PF07_0047             |
| 1.8N pause       |                   | Nucleotide pos. 962,166 to 1,001,010 | TOXO764                                       | AAA-ATPase/cdc48 cell division regulator | cgd5_2010             |
|                  |                   |                          |                                               |                                    |                       |
| Budding mutant  | 118G4             | Chr. IV                  | TOXOW23                                       | 27.m00874                            | MAL8P1.92             |
| Early bud arrest |                   | Nucleotide pos. 1,533,747 to 1,564,960 | TOXO63 –                                     | Related to Sac3/GNAP               | cgd3_3570             |
| Budding mutant† | 154g11            | Chr. VIIa                | TOXO767                                       | 20.m03745                            | cgd1_2730             |
| Early bud arrest |                   | Nucleotide pos. 2,988,420 & 2,988,482 | TOXPC51                                       | RRM protein                          |                       |
|                  |                   |                          |                                               |                                    |                       |
| Budding mutant† | 7A11              | Chr. II                  | PSBLE12                                       | 41.m01273                            | PF13_0052             |
| Late bud arrest  |                   | Nucleotide pos. 40,305 & 43,810 | PSBMA57                                       | Cell division protein FtsJ          | cgd3_890              |
|                  |                   |                          |                                               | (ribosomal histidyl transferase)     |                       |
|                  |                   |                          |                                               |                                    |                       |
| Uncoupling mutant | 42D6              | Chr. 1b                  | TOXPG04 –                                     | 25.m01896                            |                       |
| Multiple nuclear division | | Nucleotide pos. 1,500,000 to 1,540,000 | TOXPD26 –                                    | RCC1 domain protein                 |                       |
|                  |                   |                          |                                               |                                    |                       |
| Uncoupling mutant | 50-B3             | Chr. VIIa                | TOXOG11 –                                     | 72.m00409                            | PF10975c              |
| Multiple nuclear division | | Nucleotide pos. 59,474 & 68,032 | TOXO961 –                                    | RCC1 domain protein                 |                       |
| Uncoupling mutant† | 20C2              | Chr. III                 | TOXOY49                                       | 52.m01560                            | PF14_0218             |
| Multiple nuclear divisions | | Nucleotide pos. 673,562 to 719,312 | TOXO430                                       | Actin-related protein ARF4a          | cgd8_2200             |
|                  |                   |                          |                                               |                                    |                       |
| Mitotic mutant   | VA15              | Chr. IX                  | TOXPJ50 –                                     | 80.m02355h,c                        | PF1370w               |
| Mis-segregation leading to zoids | | Nucleotide pos. 4,671,421 to 4,663,811 | TOXPJ92 –                                    | NimA related spindle assembly factor | cgd1_1490             |
| <1N arrest       |                   |                          |                                               |                                    |                       |
| Mitotic mutant† | 115C5             | Chr. X                  | PSBLJ36                                       | 44.m02780                            | MAL13PL294             |
| <1N arrest, chromosome loss | | Nucleotide pos. 2,563,769 & 2,547,691 | TOXOS31                                       | GTP-binding protein                 | cgd3_130              |
|                  |                   |                          |                                               |                                    |                       |
| Mitotic mutant   | 11C9              | Chr. XII                 | PSBL6H3 –                                     | 50.m03077                            |                       |
| 2N arrest        |                   | Nucleotide pos. 2,394,591 to 2,346,715 | TOXO457 –                                    | Unknown protein FtsK-related?       |                       |

aMarker rescue tags with opposite strand orientation bracket the complementation locus, while independent tags with the same strand orientation are listed in series.
bPCR fragments were used to further validate the identity of the complementing gene.
cThe gene from each mutant strain has been confirmed to be a ts-allele.
dRecomplementation to verify a single gene rescue is in progress.

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predicts 37.5% maximum). Following 10 days of growth, the master plates are scored for single colonies and then replica plated into two test plates (34°C and 40°C) using a 96-pin array that transfers 20 µl per pin (VP 403E Grooved Pin replicator, V&P Scientific) [13]. Growth/no-growth at 34°C and 40°C is scored 7 days later and putative ts+ mutants are passed into duplicate 24-well plates to reproducibly isolate putative ts+ clones (as described above). Following mutagenesis and release from the host cells, parasites were resuspended in medium without phenol red at a concentration of 250 parasites/ml and cloned directly into 384-well plates (transparent TC-coated plates; NUNC, RoskiSlde, Denmark) conditioned with 50% HFF cell supernatants using a plate filler (Genetics, New Milton, UK). Parasites were expanded for 7 days at 35°C. To suspend parasites, the plates were shaken 1 min at maximum speed on a Bellco orbital shaker (Vineland, NJ) placed inside a laminar flow hood. Parasites were grown for six more days until approximately 80% of the monolayer was lysed. Plates were shaken again and duplicated into two identical copies in black-with-optical-bottom TC treated 384-well plates (Falcon/BD, San Jose, CA) using a custom-made 384-pin tool with 5 µl slots (VP Scientific, San Diego, CA). Plates were seeded with a high multiplicity of infection (MOI) for parasites, washed with 4X saline for 3 times without phenol red. One plate was kept at 35°C and the second copy was incubated at 40°C. After 4 days YFP fluorescence was measured in each well using a Fluostar plate reader (BMG, Offenburg, Germany) as described [13]. Differential relative fluorescence units of wells in the 35°C versus the 40°C plate were calculated using a script in Excel (Microsoft, Redmond, WA). All identified ts+ clones were expanded into 24-well plates at 35°C and their growth arrest at 40°C was confirmed by microscopy in a 24-well plate format.

Phenotypic analysis of ts mutants. Parasite reversion frequencies were determined on plaques at 34°C versus 40°C using several parasite dilutions (10^7-10^4 parasites/T175cm2 flask).

DNA content and cellular and nuclear morphologies at the permissive and restrictive temperatures were used to characterize the phenotypes of individual ts+ clones. A culture of parasites in log phase (8–10 parasites per vacuoles) was used to seed duplicate T175cm2 flasks (5x10^4 parasites/flask) for SYTOxGreen flow cytometry and to inoculate 6 well plates with coverslips (1x100 parasites/well) for immunofluorescence assays. Following a 2 h incubation at 34°C, the cultures were washed 3x, allowed to grow at 34°C for 5 h and then one set of cultures was shifted to 40°C.

Co-staining of infected cultures with a monoclonal mouse antibody 45.15 for IMC1 (1:2000, a gift from Dr. Ward, University of Vermont) and a polyclonal rabbit antiserum for TgPCNA1 (1:5000) [71] was used to examine changes in cell morphologies and to determine the presence of intracellular daughter buds and to establish whether such buds were able to reactivate at the restrictive temperature (as indicated by the production of progeny parasites). Where indicated, further antibodies used are: monoclonal 20H5 specific for centrin I (1:2000; a generous gift from Dr. Salisbury (Mayo Clinic, Rochester, MN) [72]; rabbit anti-TgMorn1 (1:2000) [20]; rabbit anti-TgIMC3 (1:1000) [25]; rabbit anti-GFP (1:5000); Torrey Pines Biologs, CA); monoclonal antibody 19G10 anti-z-tubulin (1:10; a kind gift from Jaeck Garrigut, Univ. of Georgia, [73]); 4,6-diamidino-2-phenylindole (DAPI) staining was also performed to also gauge the qualitative changes in DNA content in parasite clones growing at 34°C versus 40°C. Briefly, parasite cultures grown on coverslips in well plates were fixed with 7% paraformaldehyde (pH 7.4) and then permeabilized with 0.25% Triton X-100. Coverslips were washed and incubated in 1X PBS pH 7.4 containing 5% FBS and 3% BSA (blocking solution) for at least 30 min. Primary antibodies diluted in blocking solution were incubated for 30–60 min, the coverslips were washed again with blocking buffer and then incubated for 30–60 min with secondary antibodies: Alexa Fluor 594 for mouse and Alexa Fluor 488 for rabbit diluted 1:2000 in blocking buffer. The secondary antibodies were removed and a DAPI solution (1 mg/ml stock) diluted 1:100 in blocking buffer was added for 5–10 min prior to washing and mounting in gel mounting solution. DNA content and epifluorescence microscopy (Eclipse TE300, Nikon Inc., Melville NY) and images collected with a digital camera (SPOTTM Dynamic Instruments Inc.). In addition, pictures were acquired on a DM RB inverted microscope (Leica) equipped with a PLAPLO 100x/1.4 lens and a Lambda 9000 3mW inverted microscope with a PRIOR stage and a NEOFULAR 100x/1.3 lens. The latter two microscopes are equipped with a Hamamatsu C4742-95 CCD camera, and both are controlled by Improvision software (Lexington, MA).

Flow cytometric determination of nuclear DNA content. Parasite nuclear DNA content was evaluated by flow cytometry using SYTOX Green (Invitrogen) staining of tachyzoites grown at the permissive and restrictive temperatures [2]. Briefly, parasites were harvested from the T175cm2 flasks by needle passage and filtration through 3 µm filters. Tachyzoites collected by centrifugation were resuspended in 300 µl of PBS and 700 µl of 0.5% paraformaldehyde (second wash). The fixed samples were stored at −20°C for at least 24 h prior to staining for flow cytometry. Fixed parasites were pelleted at 3000 x g and resuspended in 50 µM Tris pH 7.5 at a final concentration of 6 × 10^6 parasites/ml and stained with SYTOX Green cocktail (250 U; RNase A, RNase T1) and added the parasites incubated in the dark at room temperature for 30 min. Nuclear DNA content was measured based on fluorescence (FL-1) using a 488 nm argon laser flow cytometer. Fluorescence was collected in linear mode (10,000 events) and the results were quantified using CELLQuest V3.0 (Becton-Dickinson, CA) and the percentage of each DNA phase was calculated based on defined gates for each population.

Establishment of a genomic cosmid library. A cosmid library was constructed in ToxoSuperCos. This new cosmid backbone plasmid is composed of the double cos-sites from plasmid SuperCos (Stratagene, La Jolla, CA) combined with the kanamycin resistance gene and the origin of replication from pDONR201 (Invitrogen, Carlsbad, CA), the drug pyrithymethine resistant allele of T. gondii dihydrofolate reductase-thymidylate synthase (DHFR-TSm2m3) from pHDR-TSc3ABP [13]; pDONR-TgEF1α promotor from SuperCos with the super TgEF1α promoter; and bisulfate. To construct the cosmid library, genomic DNA was extracted from wild type RH strain. Parasites were lysed in 0.2% SDS in 10 mM Tris/100 mM NaCl pH 8.0 followed by digestion with 0.8 µg/ml proteinase K and 0.04 µg/ml RNase A for 4 hrs at 37°C. DNA was extracted once with phenol-chloroform-isomylalcohol (25:25:1) followed by chloroform extraction and precipitation using 0.2 volumes of 10 M Na2OAc and 2 volumes of 100% ethanol. DNA was resuspended using a glass rod and allowed to resuspend in two volumes of TE overnight at 7°C. 5–10 µl of this DNA preparation was digested with a serial dilution of SmaI. Reaction aliquots were analyzed by gel electrophoresis on field or gel electrophoreses on a BioRad CHEF system (15 hrs at 4°C using conditions optimized for separation from 5–200 kb at a gradient of 0.6 V/cm and an angle of 120°) using the New England Biolabs (Beverly, MA) Low Range PFGE marker as size standard. Conditions generating the highest amount of fragments between 40–55 kb (as well as one dilution step above and below) were chosen for preparative SmaI digestion. After verification by CHEF, the three digests were pooled and treated as described in the SuperCos manual (Stratagene). Briefly, digested genomic DNA was phenol extracted, precipitated, and reconstituted in TE. The purified plasmid DNA was used for electroporation of 8 x 10^10 parasites prior to inoculation into T175cm2 flasks. Parasites were allowed to recover for 24 hrs at 35°C prior to shifting to the 40°C and addition of 1 µM pyrimethamine to

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select for phenotypic complementation and cosmid backbone genomic integration, respectively. Individual cosmids from the tiled and sequenced plates were isolated from 5 ml bacterial cultures. Cells were lysed using 250 μl P1 buffer (Qiagen) supplemented with 70 μl 10 mg/ml lysozyme solution, followed by addition of Qiagen buffers P1 and P3 according to the manufacturer’s instructions. Following centrifugation, DNA was purified from the supernatant by addition of 0.7 volumes of isopropanol.

**Identification of complementing cosmids by plasmid rescue.** Genomic DNA was extracted from complemented parasite lines (polyclonal or clonal lines) and 2 μg of DNA was digested with one of the restriction enzymes in the linker region (20 U XhoI, HindIII, BglII or KpnI). Digested DNA was purified using the Quiagene Quick PCR purification kit (Qiagen) and an equivalent of 0.5 μg of DNA was self-ligated (overnight) in a volume of 20 μl at 16°C. After phenol extraction and ethanol precipitation, 15% of the total reaction (1.5 μl of 10 μl) was electroporated into 20 μl DH12S electromax E. coli (Invitrogen) and plated on LB-KAN (50 μg/ml, DNA can also be purified using a Qiagen spin column instead of phenol extraction). DNA minipreparations were isolated for multiple colonies and digested with BglII or NcoI. Inserts were sequenced using the T3 primer.

**Supporting Information**

**Figure S1.** Transfection of HXGPRT KO Mutant with the ToxoSuperCos Library Results in Complementation of the HXGPRT Locus

The RH HXGPRT knock out line was transfected with the ToxoSuperCos library and selected with mycophenolic acid for restoration of HXGPRT activity. Five out of five transfections resulted in stable drug resistant parasites. Clonal lines were established from three transfections and parasites were tested for the presence of diagnostic PCR products for the intact wild type gene (B), 0.48 kb, minor bands were sequenced and proved to be unrelated spurious amplification products, asterisks) and the KO locus (C), 0.7 kb. Note that restoration was confirmed in 10 out of 15 clones. Persistence of the KO locus suggests that this did not occur through homologous recombination. (A) Shows a simplified partial map of the HXGPRT locus indicating the positions of primers used. Found at doi:10.1371/journal.ppat.0040036.sg001 (2.5 MB TIF).

**Figure S2.** Expression Level and Size of Complemented Genes (A) Most of the genes identified as cell cycle regulators by complementation show modest expression judged by replicate (average of three independent samples) microarray analysis of Type I strain tachyzoites using the Affymetrix Toxoplasma GeneChip (see ToxoDB) (RNA normalized flourescent intensities are shown in red and ranked by expression level: 50, m0307, 32, m01150, 41, m01273, 44, m02615, 583, m0547, 25, m01896, 27, m00873, 35, m00891, 44, m0278, 80, m02355, 20, m0375, 72, m0409, 20, m03766, 49, m00633, see Table 1 for complementation details). The mRNA level for well-characterized genes in the groups/proteins class (in column, PCNA) abundance classes are shown for comparison in blue. (B) Complemented genes are of varying size (the size of the predicted protein product in number of amino acids is shown here) including numerous large genes. Note that these numbers are based on the current draft three annotation of the T. gondii genome (http://www.toxoDB.org/) and that the gene models have not been experimentally validated. The order of genes is the same as in (A).

Found at doi:10.1371/journal.ppat.0040036.sg002 (193 KB TIF).

**Table S1.** Summary of ENU Screens (Note that ts Mutants Obtained from the YFP Fluorescent Screen Are Indicated by –Y)

Found at doi:10.1371/journal.ppat.0040036.s001 (31 KB DOC).

**Table S2.** Summary of the Cell Cycle Analysis of Selected ts Mutants

Found at doi:10.1371/journal.ppat.0040036.s002 (147 KB DOC).

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**Competing interests.** The authors have declared that no competing interests exist.

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