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Identification of Obscure yet Conserved Actin-Associated Proteins in *Giardia lamblia*

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Consistent with its proposed status as an early branching eukaryote, *Giardia* has the most divergent actin of any eukaryote and lacks core actin regulators. Although conserved actin-binding proteins are missing from *Giardia*, its actin is utilized similarly to that of other eukaryotes and functions in core cellular processes such as cellular organization, endocytosis, and cytokinesis. We set out to identify actin-binding proteins in *Giardia* using affinity purification coupled with mass spectroscopy (multidimensional protein identification technology [MudPIT]) and have identified >80 putative actin-binding proteins. Several of these have homology to conserved proteins known to complex with actin for functions in the nucleus and flagella. We validated localization and interaction for seven of these proteins, including 14-3-3, a known cytoskeletal regulator with a controversial relationship to actin. Our results indicate that although *Giardia* lacks canonical actin-binding proteins, there is a conserved set of actin-interacting proteins that are evolutionarily indispensable and perhaps represent some of the earliest functions of the actin cytoskeleton.

n addition to being a major parasite infecting more than 280 million people each year, Giardia lamblia (synonymous with Giardia intestinalis and Giardia duodenalis) belongs to one of the earliest diverging groups of eukaryotes (1-4). Therefore, investigation of Giardia biology has the potential to uncover evolutionarily deep cellular mechanisms. However, the placement of Giardia near the root of the eukaryotic tree, in addition to the placement of the root itself, is contentious (5). Nevertheless, Giardia is the most divergent eukaryote that can be manipulated in the laboratory with molecular and genetic tools (4, 6-10). In addition, many pathways in Giardia have fewer components than in the well-studied model eukaryotes (4). Thus, the combination of Giardia's highly divergent and minimalistic genome provides a unique perspective through which cellular processes may be examined. This perspective may potentially define both the minimal requirements for function and the portions of cellular mechanisms constrained throughout evolution.

A major point of divergence between Giardia and other eukaryotes is the cytoskeleton (11). Giardia lacks the canonical actin-binding proteins, once thought common to all extant eukaryotes, which perform critical functions in other eukaryotes (12). Their absence may indicate a split from the last eukaryotic common ancestor before the canonical set of actin-binding proteins was established. Alternatively, Giardia may have evolved a novel set of actin-interacting proteins that allowed for the gradual loss of the canonical set of actin-binding proteins (11, 13). Our previous work has shown that despite the lack of canonical actinbinding proteins, Giardia actin (giActin) is required for conserved cellular functions, including membrane trafficking, cytokinesis, polarity, and control of cellular morphology (13). The Giardia cytoskeleton is also quite elaborate, suggesting the presence of cytoskeletal regulators (Fig. 1). That giActin performs similar functions to actin in other eukaryotes suggests these processes were already associated with actin at the time Giardia split from the other eukaryotes (13). We have also shown that giRac, the sole Rho family GTPase in Giardia, regulates actin despite the absence of all actin-binding proteins known to link G-protein signaling to

the actin cytoskeleton (Arp2/3, formin, wave, myosin, and cofilin) (13). Therefore, *Giardia* must contain a novel set of actin-interacting proteins comprised of ancient yet undiscovered and/or *Giardia*-specific actin regulators. We sought here to identify actinbinding proteins using actin affinity chromatography coupled with multidimensional protein identification technology (Mud-PIT) (14). The discovery of *Giardia*-specific actin-binding proteins with essential functions would open an avenue to potential therapeutic targets, while the discovery of conserved proteins would highlight an ancient relationship between actin and the identified protein, worthy of further exploration.

MATERIALS AND METHODS

Strain and culture conditions. *Giardia lamblia*, strain WBC6 was cultured as described previously (15). Morpholino knockdown experiments and quantitative Western blotting were performed as described previously (9, 13). Large volume high-yield cultures required a method to increase the surface area. We filled standard wide-mouth media bottles with cut-to-length "jumbo drinking straws" and autoclaved them before filling with media (see Fig. S1 in the supplemental material). *Giardia* cell counts increased by ~30% in straw-filled 500-ml bottles versus those without. After 3 days of growth, we did not observe unattached cells at the bottom of straw-filled culture vessels that are typical of overgrown cultures, while bottles without straws had a layer of cell sediment. After 72 h of growth 1-liter cultures regularly reach 2.5×10^6 cells/ml, exceeding maximum trophozoite concentrations obtained with Farthing's roller bottles, without needing specialized equipment (16).

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FIG 1 *Giardia* cytoskeletal organization. (A) Maximum projection of a Z-stack. Actin is green, tubulin is red, and DNA is blue. (B) Diagram of *Giardia* trophozoite with all of the prominent cytoskeletal structures labeled.

Constructs. The TS-Actin vector was constructed by modifying pGFPapac (17). A BamHI site was first introduced between BsrGI and NotI of enhanced green fluorescent protein (eGFP) using an oligonucleotide adapter; all primer sequences can be found in Table S1 in the supplemental material. The glutamate dehydrogenase (GDH) promoter was exchanged for the actin promoter by excising GDH with HindIII and NcoI, the actin promoter was subsequently ligated into the same position. Next. eGFP was excised with NcoI and BamHI, allowing for the TwinStrep tag to be ligated into the same position. Finally, the vector was digested with BamHI and NotI so that the actin gene could be ligated into the vector. All PCR amplification steps were performed with iProof high-fidelity polymerase (Bio-Rad), and the resulting vectors were verified by sequencing. The putative actin-interacting proteins were PCR amplified from genomic DNA and inserted into the pKS 3HA.NEO vector (10) using the restriction sites indicated in Table S1 in the supplemental material. All resulting constructs except for TS-Actin, GL50803_6744, and GL50803_13273 were linearized and integrated into the genome by homologous recombination to generate endogenously tagged proteins.

Actin affinity chromatography. One-liter straw-packed and sterilized bottles were filled with medium and inoculated with two 13-ml confluent cultures containing wild-type (WT) or TwinStrep-giActin cell lines. After 3 days the cultures were incubated in an ice water bath for 1 h to detach cells. The media and unattached cells were transferred to centrifuge bottles and pelleted at 750 \times g for 15 min. The resulting cell pellet was washed in 10 ml of cold HEPES-buffered saline, transferred to 15-ml conical tubes, and pelleted again. The cell pellet was resuspended in an equal volume (~2 ml) of lysis buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 7.5% glycerol, 0.25 mM CaCl₂, 0.25 mM ATP, 0.05 mM dithiothreitol [DTT], 0.5 mM phenylmethylsulfonyl fluoride [PMSF], 2× Halt protease inhibitors [Pierce]). The pellet was stored overnight at -80°C and, after thawing, the cells were sonicated, and the extract cleared at $10,000 \times g$ for 10 min. The lysate was added to 200 μ l of Streptactin-Sepharose beads (IBA) previously equilibrated with lysis buffer. Binding was performed for 1.5 h with end-over-end mixing at 4°C. The beads were washed once in batch (100 mM Tris, pH 8.0, 150 mM NaCl, 7.5% glycerol, 0.25 mM $\mathrm{CaCl}_2, 0.25$ mM ATP, 0.5 mM DTT) and then moved into a chromatography column (Bio-Rad) and washed four additional times with one column bed volume of wash buffer. Protein was eluted with 6 half-column bed volumes with elution buffer (100 mM Tris, pH 8.0, 150 mM NaCl, 7.5% glycerol, 0.25 mM CaCl₂, 0.25 mM ATP, 0.5 mM DTT, 2 mM D-biotin).

Actin pelleting assay. TwinStrep-Actin was purified as described above and then dialyzed for 2 h in G buffer (5 mM Tris, pH 8.0, 0.2 mM ATP, 0.2 mM CaCl₂, 0.5 mM dithiothreitol). After a buffer exchange, the actin was dialyzed overnight. The dialyzed actin was cleared by centrifugation at 100,000 × g for 30 min to remove aggregates. A 1/10 volume of 10× KMEI80 (800 mM KCl, 10 mM MgCl₂, 10 mM EGTA, 100 mM imidazole [pH 7.0]) was added to the cleared actin, followed by incubation for 30 min at room temperature. The KMEI80-actin mixture was then centrifuged at 100,000 × g for 30 min.

Mass spectroscopy. Mass spectrometry was performed by the Vincent J. Coates Proteomics/Mass Spectrometry Laboratory at UC Berkeley. The protein solution was adjusted to 8 M urea, subjected to carboxyamidomethylation of cysteines, and digested with trypsin. The sample was then desalted using a c18 spec tip (Varian). A nano-LC column was packed in a 100-µm-inner-diameter glass capillary with an emitter tip. The column consisted of 10 cm of Polaris c18 5-µm packing material (Varian), followed by 4 cm of Partisphere 5 SCX (Whatman). The column was loaded by using a pressure bomb and washed extensively with buffer A (see below). The column was then directly coupled to an electrospray ionization source mounted on a Thermo-Fisher LTQ XL linear ion trap mass spectrometer. An Agilent 1200 high-pressure liquid chromatograph equipped with a split line so as to deliver a flow rate of 300 nl/min was used for chromatography. Peptides were eluted using an eight-step MudPIT procedure (14). Buffer A was 5% acetonitrile-0.02% heptafluorobutyric acid (HBFA); buffer B was 80% acetonitrile-0.02% HBFA. Buffer C was 250 mM ammonium acetate-5% acetonitrile-0.02% HBFA; buffer D was the same as buffer C, but with 500 mM ammonium acetate. The programs SEQUEST and DTASelect were used to identify peptides and proteins from the Giardia database (18, 19).

Immunoprecipitation and Western blotting. Immunoprecipitation began with a single confluent 13-ml tube per cell line. After detachment, cells were pelleted at 900 \times g and washed once in HBS. The cells were resuspended in 300 µl of lysis buffer (50 mM Tris 7.5, 150 mM NaCl, 7.5% glycerol, 0.25 mM CaCl₂, 0.25 mM ATP, 0.5 mM DTT, 0.5 mM PMSF,

0.1% Triton X-100, 2× Halt protease inhibitors [Pierce]) and sonicated. The lysate was cleared by centrifugation at 10,000 × g for 10 min at 4°C and then added to 30 μ l of anti-HA beads (Sigma). After 1.5 h of binding, the beads were washed four times with wash buffer (25 mM Tris 7.5, 150 mM NaCl, 0.25 mM CaCl₂, 0.25 mM ATP, 5% glycerol, 0.05% Tween 20) and then boiled in 50 μ l of sample buffer. Western blotting was performed as described previously (13). Rabbit anti-giActin polyclonal (13) and anti-HA mouse monoclonal HA7 antibody (Sigma-Aldrich) were both used at 1:3,000. Fluorescent secondary antibodies (Li-Cor) were used at 1:15,000, horseradish peroxidase-linked anti-rabbit antibodies (Bio-Rad) were used at 1:7,000.

Microscopy. Fixations were performed as described previously (13). Anti-HA mouse monoclonal HA7 antibody (Sigma-Aldrich) was used at 1:125, and anti-mouse and anti-rabbit secondary antibodies were used at 1:200 (Molecular Probes). Images were acquired on a DeltaVision Elite microscope using a 100×1.4 NA objective and a CoolSnap HQ2 camera. Deconvolution was performed with SoftWorx (API, Issaquah, WA). Maximal projections were made with ImageJ (20), and figures were assembled using the Adobe Creative Suite (Mountain View, CA).

RESULTS

We set out to identify giActin interactors via an affinity chromatography approach utilizing the TwinStrep Tag (21–23). Actin is notoriously sensitive to chimeric fusions, because epitope or fluorescent protein fusions may cause steric interference or otherwise affect filament formation and dynamics (24). Thus, we devised a strategy to test whether our TwinStrep-Actin fusion (TS-Actin) was functional in vivo. Previous work demonstrated that actin can be effectively depleted with translation-blocking morpholinos (13). These antisense morpholinos bind to the start of the transcript and block translation initiation machinery from recognizing the start codon (9, 13). Therefore, by fusing Twin-Strep to the N terminus of giActin, we generated a morpholinoinsensitive version of giActin. In this case, morpholino treatment is expected to block translation of endogenous actin, whereas it should have no effect on the transgenic version. We also sought to maintain actin levels near endogenous levels by driving expression of our TS-Actin fusion with the native actin promoter.

Quantitative Western blotting with an anti-giActin polyclonal antibody (13) indicated that although we used the native promoter, there was roughly a 4-fold increase in total actin levels compared to nontransgenic controls; ca. 75% of this was TS-Actin (Fig. 2A). The higher levels of transgenic actin are presumably due to the copy number of our episomally maintained construct exceeding the number of endogenous actin genes. Morpholino treatment of the TS-Actin-expressing cell line behaved as predicted; the N-terminal epitope tag protected TS-Actin from being depleted by anti-actin morpholinos, while the endogenous actin was depleted to $\sim 20\%$ of control levels (Fig. 2A). Further, we examined the morpholino-treated cells for morphological defects associated with actin depletion such as abnormal cell shape, outof-position flagella, and multiple or out-of-position nuclei (13). In the control-treated cell line we observed a slight increase in the number of abnormal cells: 6.6% for TS-Actin (n = 600) versus 1.9% for wild-type (n = 400), indicating that the increased actin levels and/or the epitope tag mildly interfered with normal actin function (Fig. 2B). The transgenic line was, however, resistant to morpholino depletion since the proportion of abnormal cells remained at 6.8% (n = 600) after morpholino treatment. In contrast, 35.2% of the WT cells (n = 500) treated with the anti-actin morpholinos had abnormal morphology. Therefore, we conclude



FIG 2 TS-Actin is functional *in vivo*. (A) Multiplex Western blot (actin, green; tubulin, red) showing that TS-Actin is morpholino resistant, while endogenous actin is significantly reduced. (B) Reducing endogenous actin results in cellular disorganization, morpholino-resistant TS-Actin can substitute for endogenous actin.

that TS-Actin can partially rescue endogenous actin depletion, indicating that TS-Actin is functional *in vivo*.

A particular challenge of producing large-scale Giardia cultures, sufficient for biochemical analysis, is the need to provide surface area for adherent growth. Giardia is an extracellular parasite that colonizes the host intestine by attaching via its "suction cup" organelle, the ventral disc (25, 26). Likewise, in the laboratory Giardia trophozoites grow attached to the sides of the culture tubes. Cultures cease to proliferate after the culture tubes are confluent with cells. Free-floating cells are often observed to have an aberrant morphology, indicating the importance of surface attachment, possibly because Giardia divides by an adhesion-dependent mechanism (27, 28). Custom "inside-out" roller bottles have been used by others to grow high-yield Giardia cultures, but these are not commercially available (16). We developed a lowcost high-yield method of growing Giardia by inserting common polypropylene drinking straws into wide mouth bottles (see Fig. S1 in the supplemental material and see Materials and Methods). Using our high-surface-area culture system, 1-liter cultures of WT and the TS-Actin transgenic cell lines produced ~2-ml cell pellets. Extracts from these cell pellets were affinity purified in parallel. The elutions from a pilot experiment were concentrated before sodium dodecyl sulfate (SDS) analysis so that \sim 50% of the eluted protein could be analyzed by SDS-PAGE. Many unique bands are apparent in the TS-Actin sample (Fig. 3A). The purification was repeated for mass spectroscopy analysis; Fig. 3B represents 5% of the elutions that were analyzed by mass spectroscopy. Table 1 lists 57 proteins that were unique to the TS-Actin sample and had a minimum of five detected peptides. The complete list, including low-abundance hits and proteins also identified in our mock control, is given in Table S2 in the supplemental material. Bioinfor-



FIG 3 Isolation of TS-Actin and interacting proteins. (A) Elutions from streptactin columns for both WT and TS-Actin purifications were concentrated and then analyzed by SDS-PAGE. Note that several bands are unique to the TS-Actin cell line. Actin is marked with an asterisk. (B) Five percent of the TS-Actin purification used in the MudPIT analysis was loaded onto a 4 to 16% gradient gel and stained with SYPRO Ruby.

matics analysis was utilized to place these hits into six categories (Table 1; see Table S2 in the supplemental material).

We identified several hits that support the quality and relevance of this data set. For example, we identified all eight subunits of the TCP-1 chaperonin complex, which has an important role in folding actin (29). In addition, two proteins, p28 dynein light chain (p28 DLC) and centrin, were found in the giActin interactome, which we had previously hypothesized to be conserved actin-interacting proteins (13). Genetic and biochemical analysis of flagellar components has demonstrated that actin has an important role in flagella, where it functions in the inner dynein arm complexes (30-32). Within the inner dynein arms, p28 DLC and centrin, have been demonstrated to directly interact with actin (32). In Giardia, actin is readily detectable within all eight flagella, and both p28 DLC and centrin are conserved (13). In terms of peptides per molecular weight, p28 DLC was the most abundant interactor identified in our analysis. In addition to these examples, homologs of several other proteins that have been reported to complex with actin in other eukaryotes were identified and are indicated in Table 1.

The genome of Spironucleus salmonicida, another diplomonad and close relative of Giardia, was recently released (33). As part of our analysis, we compared our list of putative actin interactors to the S. salmonicida genome (Table 1) (33). Although most of the identified proteins are present in S. salmonicida, several appear to be specific to Giardia. We also searched the S. salmonicida genome for the presence of canonical actin-binding proteins. Intriguingly, we found that S. salmonicida contains several actin-binding proteins not found in Giardia; these include formin, cofilin, and coronin (see Table S3 in the supplemental material). S. salmonicida, however, lacks many canonical actin-binding proteins, including the Arp2/3 complex, nucleation-promoting factors, dynactin, capping protein, and myosin. Nevertheless, the subset of canonical actin-binding proteins in S. salmonicida suggests the loss of such proteins from Giardia. Without additional genomes, we can only speculate whether the diplomonads ever had the full complement of actin-binding proteins; however, the absence of myosin in

Giardia, *S. salmonicida*, and *Trichomonas vaginalis* (a nondiplimonad excavate) remains consistent with the idea that a subset of excavates may have split from the other eukaryotes before the full complement of actin-binding proteins was established (4, 34). This possibility could help explain how *Giardia* could have lost proteins that are essential in the model eukaryotes.

Next, we sought to validate a subset of the conserved interactions through reciprocal immunoprecipitations. We selected nine representative proteins, at least one from each of the conserved categories; these are indicated by an asterisk in Table 1. In each case, we tagged the identified protein with a C-terminal triple HA tag. We were able to verify complex formation with giActin for p28 DLC, centrin, HSP70, ARP7, TIP49, ERK2, and 14-3-3 (Fig. 4A). Attempts to validate dynamin (Fig. 4A) and myeloid leukemia factor (MLF; data not shown) were unsuccessful. Both dynamin and MLF have been shown to interact with actin and alter filament organization in other eukaryotes (35, 36). Although these hits may be false positives, it is also possible that the C-terminal tag disrupted interaction or that the lower concentration of cell extracts in our immunoprecipitation experiments versus large-scale affinity chromatography failed to maintain integrity of the complex.

To better understand the relationship between these conserved interactors and actin, we colocalized actin and the tagged interactors (Fig. 4B). Each protein displayed a localization pattern consistent with its proposed function. p28 DLC localized to flagella. Centrin localized to the basal bodies and around a portion of the internal axonemes of the posterolateral flagella. ARP7, TIP49, and ERK2 localized to the nuclei with various amounts of non-nuclear localization. HSP70 and 14-3-3 were found throughout the cell with slight enrichment at the cell anterior. None of these conserved proteins colocalized with prominent filamentous actin structures (see Fig. 1), which is consistent with the idea that they complex with G-actin (discussed below). It should be noted that standard tools such as fluorescent phalloidin and DNase I typically used to distinguish between monomeric and filamentous actin do not work in *Giardia* (13).

DISCUSSION

In this study, we undertook a biochemical approach to identify actin interactors in Giardia. Our easily adopted method for growing large-scale cultures and the use of the TwinStrep tag have the potential to make the process of defining interactomes routine in Giardia. During the course of our study, Svard and coworkers published a TAP-tagging approach for proteomics in Giardia (37). Similar to our approach, these researchers used two tandem Strep II tags but also included a Flag tag, the entirety of which is known as the SF-TAP tag. They overcame the surface area issues by distributing 2 liters of medium among 40 50-ml conical tubes. Our straw method simplifies cell culture, and our ability to identify actin-interacting proteins in a single purification step suggests that tandem purification is not generally required. This is significant because single-step purifications are able to isolate weaker interactors commonly lost in two-step purifications (22). Indeed, our laboratories have already performed proteomic analysis on two additional proteins using this approach. In all cases, 1 liter of medium was sufficient for isolating protein-protein interactors. Although analysis is still under way, in each case a unique set of high-frequency hits were identified. Conversely, several low-frequency hits are common to our data sets. One data set is for Polo-

TABLE 1 Identified interactors

$ \begin{array}{c} \mbox{Accentral dynch heavy chain} & 570,19 & 900 & Precedents & 2, 59 & ST 4476, 0.0 \\ (13.080, 11013) & Atonemal dynch heavy chain & 570,19 & 900 & Precedents & 2, 59 & ST 4476, 0.0 \\ (13.080, 12037) & P28 asomal dynch heavy chain & 570,19 & 948 & Precedents & 2, 59 & ST 4476, 1.0 \\ (13.080, 12037) & P28 asomal dynch heavy chain & 184,790 & 48 & Precedents & 2, 9 & ST 4476, 1.1 \\ (13.080, 12037) & Call asomal dynch heavy chain & 184,790 & 48 & Precedents & 2, 9 & ST 4478, 1.2 \\ (13.080, 12047) & Call asomal dynch heavy chain & 184,790 & 48 & Precedents & 2, 9 & ST 44418, 1.2 \\ (13.080, 14242 & Myeold levenis factor Hile & 25,702 & 14 & Pre- & 3 & ST 4418, 1.5 \\ (13.080, 14242 & Myeold levenis factor Hile & 25,702 & 14 & Pre- & ST 44444, 1.0 \\ (13.080, 11424 & Myeold levenis factor Hile & 25,702 & 14 & Precedents & 29 & ST 444444, 1.0 \\ (13.080, 11424 & Myeold levenis factor Hile & 25,702 & 14 & Precedents & 29 & ST 444444, 1.0 \\ (13.080, 11424 & Myeold levenis factor Hile & 25,702 & 14 & Precedents & 29 & ST 444491, 0.0 \\ (13.080, 11437 & TCP-1 chaperonin submit thet & 66,64 & 187 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1137 & TCP-1 chaperonin submit thet & 66,64 & 187 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 11347 & TCP-1 chaperonin submit factor & 66,64 & 187 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1713 & TCP-1 chaperonin submit factor & 66,64 & 187 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 6,592 & 174 & TCP-1 chaperonin submit factor & 6,592 & 174 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 6,592 & 174 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 6,592 & 174 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 16,599 & 174 & Precedents & 29 & ST 444871, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 16,599 & 174 & Precedents & 51 & ST 444444, 10, 0.0 \\ (13.080, 1741 & TCP-1 chaperonin submit factor & 16,599 & 174 & P$	Protein identification no. ^a	Name and/or description	Mol wt	No. of peptides	Interactor ^b	Reference(s)	<i>S. salmonicida</i> GenBank no. ^c
$ \begin{array}{c} cl.3803, 1 1950 \\ cl.3803, 1 1950 \\ cl.3803, 1 2951 \\ cl.3803, 1 2952 \\ cl.3803, 1 295 \\ cl.3803, 1 292 \\ cl.3803, 1 292 \\ cl.3803, 1 292 \\ cl.3803, 1 292 \\ cl.3803, 1 213 \\ cl.3803, 1 21 \\ cl.3803, 1 21$	Axonemal/cytoskeleton						
$ \begin{array}{c} L12803, 210136 \\ c12803, 21237 \\ c1280$	GL50803_111950	Axonemal dynein heavy chain	570,319	900	Precedents	32, 39	EST41976, 0.0
	GL50803_101138	Axonemal dynein heavy chain	578,219	849	Precedents	32, 39	EST46166.1, 0.0
$ \begin{array}{c} 10.3003, 2223 \\ 10.3003, 2233 \\ 10.3003, 2734 \\ 10.3003, 2734 \\ 10.3003, 2734 \\ 10.3003, 2734 \\ 10.3003, 2734 \\ 10.4000 \\ 10.3003, 3705 \\ 10.3003, 3705 \\ 10.3003, 3705 \\ 10.3003, 3705 \\ 10.3003, 10.12 \\$	GL50803_40496 CL50803_12273*	Axonemal dynein heavy chain	555,424	//8	Precedents $ID \perp \perp \perp$	32, 39	ES144588.1, 0.0
$ \begin{array}{c} \mbox{cl} Classible 3, 2749 \\ \mbox{cl} Classible 3, 2741 $	GL50803_13275	Axonemal dynein heavy chain 11	20,095	294 48	Precedents	32, 39	EST45975.1, 0E-150 EST41750 1 2E 61
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	GL50803_42285 GL50803_6744*	Centrin	18 687	40	IP+	32, 39	EST41730.1, 2E-01 EST41812 1 2E-98
Classibility Automene associated protein CASP-160 174,722 24 Free control No No Classibility Classibility Optimin kery chain cytoplasmic 03,353 11 Dr 55 FST42131,19-17 Classibility TCP-1 chapteronis submit byta 61,033 223 Precedents 29 57144631,10.0 Classibility TCP-1 chapteronis submit byta 61,033 223 Precedents 29 55144691,10.0 Classibility TCP-1 chapteronis submit that 64,646 187 Precedents 29 55144973,10.0 Classibility TCP-1 chapteronis submit that 64,646 187 Precedents 29 55144973,10.0 Classibility TCP-1 chapteronis submit that 64,659 71 Precedents 29 55144973,10.0 Classibility TCP-1 chapteronis submit data 64,591 71 Precedents 29 55144973,10.0 Classibility TCP-1 chapteronis submit data 65,391 71 Precedents 29 55144581,0.0 Classibility	GL50803_37985	Dynein heavy chain	118 679	26	Precedents	32, 39	FST48250 1 0 0
$ \begin{array}{c} \mbox{CLS005.1} ({\rm bl} 22^4) & \mbox{Multi} ps-17 & \mbox{CLS005.1} ({\rm bl} 224) & \mbox{Precedents} & Prec$	GL50803_37716	Axoneme-associated protein GASP-180	174,782	20	Treccucints	52, 59	No
$ \begin{array}{c} \text{GL3003}_1422 & \text{Dyncin heavy chain-cytoplasmic} & 633,335 & 11 \\ \hline \text{Chaperone} & \text{CP-1} & \text{Chaperonin subunit peakon } & 61,93 & 223 & \text{Precedents } 29 & \text{EST46693.1, 0.0} \\ \text{GL3003}_1197 & \text{TCP-1} & \text{Chaperonin subunit peakon } & 60,646 & 107 & \text{Precedents } 29 & \text{EST46493.1, 0.0} \\ \text{GL3003}_1037 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST46493.1, 0.0} \\ \text{GL3003}_1037 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST46993.1, 0.0} \\ \text{GL3003}_17438 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST4693.1, 0.0} \\ \text{GL3003}_17411 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST4693.1, 0.0} \\ \text{GL3003}_17411 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST47029.1, 0.0} \\ \text{GL3003}_17411 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 118 & \text{Precedents } 29 & \text{EST47039.1, 0.0} \\ \text{GL3003}_17411 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 12 & \text{Precedents } 29 & \text{EST47039.1, 0.0} \\ \text{GL3003}_17411 & \text{TCP-1} & \text{chaperonin subunit eta} & 60,941 & 12 & \text{Precedents } 54 & 56 & \text{EST4936.1, 0.0} \\ \text{GL3003}_17435 & \text{TPT-1} & \text{faperonin subunit eta} & 54,465 & 90 & \text{IP+} & 54 & 56 & \text{EST4936.1, 0.0} \\ \text{GL3003}_1763 & \text{TBP-interacting protein TIP49} & 51,418 & 76 & \text{IP++++} & \text{S4-56} & \text{EST4936.1, 1, 0.0} \\ \text{GL3003}_2783 & \text{Hypothetial/PKTS domain & 30,469 & 12 & \text{Precedents } 54 & 56 & \text{EST4936.1, 1, 0.2} \\ \text{GL3003_2876 & \text{Prokarvite DUB} & 102,336 & 11 & \text{Precedents } 54 & 56 & \text{EST4936.1, 1, 0.2} \\ \text{GL3003_2784 & \text{Hypothetial/PKTS domain protein & 102,336 & 11 & \text{Precedents } 54 & 56 & \text{EST4936.1, 1, 0.2} \\ \text{GL3003_2787 & \text{Hypothetial/PKTS domain protein & 102,336 & 11 & \text{Precedents } 57 & \text{EST4936.1, 1, 54 & \text{Precedents } 57 & ES$	GL50803 16424*	Myeloid leukemia factor like	29,702	14	IP-	35	EST47319.1, 9E-17
$ \begin{array}{c} \mbox{Caperon} & \mbox{Caperon} $	GL50803_14242	Dynein heavy chain-cytoplasmic	633,335	11			EST46283.1, 1E-27
$ \begin{array}{c} GL5003_11992 & TCP+ (daperonin subunit periodin 6), 193 & 223 & Precedents 29 & EST46466, 0.0 \\ GL5003_11023 & TCP+ (daperonin subunit that 4), 64, 752 & 196 & Precedents 29 & EST46466, 100 \\ GL5003_11023 & TCP+ (daperonin subunit that 4), 64, 752 & 196 & Precedents 29 & EST46466, 100 \\ GL5003_11023 & TCP+ (daperonin subunit data 56, 304 & 188 & Precedents 29 & EST4693, 1, 0.0 \\ GL5003_17482 & TCP+ (daperonin subunit data 56, 314 & 86 & Precedents 29 & EST4693, 1, 0.0 \\ GL5003_17482 & TCP+ (daperonin subunit agmma 6), 559 & 71 & Precedents 29 & EST4639, 0, 0 \\ GL5003_1741 & TCP+ (daperonin subunit agmma 6), 559 & 71 & Precedents 29 & EST4639, 0, 0 \\ GL5003_1712 & Bip & 74, 30 & 12 & 53 & EST462441, 0.0 \\ GL5003_1755 & TBP+interacting protein TP49 & 51, 418 & 276 & Precedents 39 & EST46394, 0, 0 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 162 & Precedents 34-56 & EST09851, 100 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 162 & Precedents 54-56 & EST09851, 0.0 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 162 & Precedents 54-56 & EST09851, 10, 0 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 162 & Precedents 54-56 & EST09851, 10, 0 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 12 & Precedents 54-56 & EST09851, 10, 0 \\ GL5003_1755 & TBP+interacting protein TP49 & 52, 616 & 12 & Precedents 54-56 & EST09851, 10, 0 \\ GL5003_1761 & AR7^{-1}AR$	Chaperone						
$ \begin{array}{c} \text{GLS003,11397} & \text{TCP-1 chaperon is subunit beta} & 56,064 & 204 & \text{Precedents} & 29 & \text{EST4446,1,0.0} \\ \text{GLS003,11301} & \text{TCP-1 chaperon is subunit that} & 66,046 & 187 & \text{Precedents} & 29 & \text{EST4652,1,0.0-175} \\ \text{GLS003,11301} & \text{TCP-1 chaperon is subunit that} & 60,046 & 187 & \text{Precedents} & 29 & \text{EST4653,1,0.0-175} \\ \text{GLS003,11301} & \text{TCP-1 chaperon is subunit that} & 60,046 & 187 & \text{Precedents} & 29 & \text{EST4703,1,0.0-175} \\ \text{GLS003,11211} & \text{TCP-1 chaperon is subunit apha} & 50,281 & 82 & \text{Precedents} & 29 & \text{EST4703,1,0.0-175} \\ \text{GLS003,17411} & \text{TCP-1 chaperon is subunit apha} & 50,281 & 82 & \text{Precedents} & 29 & \text{EST4703,1,0.0} \\ \text{GLS003,2751} & \text{TBP-interacting protein TP49} & 51,418 & 276 & \text{IP+++} & 54-56 & \text{EST49365,1,0.0} \\ \text{GLS003,2755} & \text{TBP-interacting protein TP49} & 51,418 & 276 & \text{IP+++} & 54-56 & \text{EST49365,1,0.0} \\ \text{GLS003,2755} & \text{TBP-interacting protein TP49} & 52,616 & 162 & \text{Precedents} & 54-56 & \text{EST49365,1,0.0} \\ \text{GLS003,2755} & \text{TBP-interacting protein TP49} & 52,616 & 162 & \text{Precedents} & 54-56 & \text{EST49365,1,0.0} \\ \text{GLS003,2755} & \text{TBP-interacting protein TP49} & 52,616 & 162 & \text{Precedents} & 54-56 & \text{EST49362,1,1E-27} \\ \text{GLS003,2868} & \text{Prokaryotic SMC domain protein } & 102,836 & 11 & \text{Precedents} & 54-56 & \text{EST49362,1,1E-27} \\ \text{GLS003,2680} & \text{Prokaryotic SMC domain protein } & 124,241 & 7 & \text{EST49521,1,0-15} \\ \text{GLS003,2764} & \text{Precedents} & 54-56 & \text{EST49521,1,0-15} & \text{No} & \text{EST49521,1,0-15} \\ \text{GLS003,2766} & \text{Precedents} & 54-56 & \text{EST49521,1,0-15} & \text{No} & \text{EST49521,1,0-15} \\ \text{GLS003,2764} & \text{Precedents} & 54-56 & \text{EST49521,1,0-15} & \text{No} & \text{EST49521,1,0-15} & \text{ST49521,1,0-15} & \text{ST49521,1,0-15} & \text{ST49521,1,0-15} \\ \text{GLS003,2767} & \text{Precedents} & 54-56 & \text{EST49521,1,0-15} & \text{ST49521,1,0-15} & ST49521$	GL50803_11992	TCP-1 chaperonin subunit epsilon	61,193	223	Precedents	29	EST46493.1, 0.0
	GL50803_11397	TCP-1 chaperonin subunit beta	56,604	204	Precedents	29	EST41446.1, 0.0
$ \begin{array}{c} \text{GL3083,} 1500 & \text{TCP} + 1 chaperon in subunit that be 0.646 & 187 & \text{Precedents 29} & \text{EST46624,} 0.0 & \text{GL3083,} 12431 & \text{TCP} + 1 chaperon in subunit 241 & 53.31 & 86 & \text{Precedents 29} & \text{EST4678,} 1.0 & \text{DTG13,} 1.45 & \text{DTG13,} 1.5 & \text{DTG13,} 1.5 & DTG1$	GL50803_16124	TCP-1 chaperonin subunit eta	64,752	196	Precedents	29	EST48572.1, 0.0
	GL50803_13500	TCP-1 chaperonin subunit theta	60,646	187	Precedents	29	EST46649.1, 0.0
$ \begin{array}{c} \mbox{L13003} \mbox{L142} \mbox{L1} \mbox{L2} \m$	GL50803_10231	TCP-1 chaperonin subunit zeta	60,941	118	Precedents	29	ES148976.1, 0.0
$\begin{array}{c} \mbox{L1:8003_1919} & \mbox{L2:1} (L-1) (Laperonin albumi appa b) (2.6) & 3.2 \\ \mbox{L2:1} (L-1) (Laperonin albumi appa b) (2.6) & 3.2 \\ \mbox{L2:1} (L-1) (Laperonin abbmi abbmi appa b) (2.6) & 1.2 \\ \mbox{L2:1} (L-1) (Laperonin abbmi abbmi appa b) (2.6) & 1.2 \\ \mbox{L2:1} (L-1) (L-1) (L-1) & 1.6 \\ \mbox{L2:1} (L-1) & 1.6 \\ \mb$	GL50803_1/482	TCP-1 chaperonin subunit delta	56,324	86	Precedents	29	ES141630.1, 4E-1/3
$ \begin{array}{c} L13003_1/41 & L1-4 \\ C12003_5/676^* & Crosselic HSP/0 \\ C12003_5/765^* & TBP-interacting protein TIP49 \\ C12003_2/815^* & TBP-interacting protein TIP49 \\ C12003_2/825^* & TBP-interacting protein TIP49 \\ T1452, T1452, T1452, T1452, T1452 \\ T1452, T1452, T1452, T1452 \\ T1452, T1452, T1452, T1452 \\ T1452, T1452, T1452 \\ T1452, T1452, T1452 \\ T1452, T1452, T1452 \\ T1452, T1$	GL50803_91919	TCP-1 chaperonin subunit alpha	59,281	82	Precedents	29	ES14/029.1, 0.0
$ \begin{array}{c} \mbox{Classons} 20.633 \\ Classon$	GL50803_1/411	Criteralia USD70	01,559	/1	Precedents	29	ES141555.1, 0.0
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	GL50803_17121	Bip	71,055	12	IP+	52	EST45859, 0.0 EST46254 1 0.0
Nuckar IP+++ 54-56 EST49365.1, 0.0 GL50803_9255* TBP-interacting protein TIP49 52,616 162 Precedents 54-56 EST4914.1, 0.6-31 GL50803_1255 HAP7-tikk 1,465 90 IP+ 54-56 EST49316.1, 66-31 GL50803_2155 SMARCCI 47,004 17 Precedents 54-56 EST49316.22, 19,9-78 GL50803_21761 Histone acetyltransferase MYST2 49,916 12 Precedents 54-56 No GL50803_686 Prokaryotic SMC domain protein 102,836 11 Precedents 54-56 No EST47350.1, 18-27 GL50803_6407 H4-3-3 protein 128,576 8 IP++ EST43282.7, 10.0 EST44224.1, 65-115 GL50803_12795 Phosducin-like 26,919 6 Precedents 57 EST44318.1, 15-14 GL50803_0430*/ Protein disulfide isomerase PDI2 50,408 6 Precedents 57 EST44353.1, 15-14 GL50803_04373* Dynamin 79,513 13 IP- 36 EST44318.1,	GL50005_17121	ыр	74,500	12		55	15140254.1, 0.0
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Nuclear						
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	GL50803_9825*	TBP-interacting protein TIP49	51,418	276	IP+++	54-56	EST49365.1, 0.0
	GL50803_17565	TBP-interacting protein TIP49	52,616	162	Precedents	54–56	EST48144.1, 0.0
	GL50803_15113*	ARP7-like	51,465	90	IP+	54–56	EST44310.1, 6E–31
$ \begin{array}{c} \text{GL20803} \ 8125 & \text{SMARCCI} & 47,004 & 17 & \text{Precedents} & 54-56 & \text{No} \\ \text{GL20803} \ 2851 & \text{Histone acetyltransferase MYST2} & 49,916 & 12 & \text{Precedents} & 54-56 & \text{EST45122,1, 9E-78} \\ \text{No} & \text{GL20803} \ 77461 & \text{SWIRM domain protein} & 102,836 & 11 & & & & & & & \\ \text{GL20803} \ 77461 & \text{SWIRM domain protein} & 124,241 & 7 & & & & & & & \\ \text{Signaling} & & & & & & & & \\ \text{GL20803} \ 6317 & \text{Putative DUB} & & & & & & & & & \\ \text{GL20803} \ 6317 & \text{Putative DUB} & & & & & & & & & \\ \text{GL20803} \ 77461 & \text{SWIRM domain protein} & & & & & & & & & \\ \text{GL20803} \ 22850^* & \text{EKT-like/giEK2} & & & & & & & & & \\ \text{GL20803} \ 22850^* & \text{IEK7-like/giEK2} & & & & & & & & \\ \text{GL20803} \ 22850^* & \text{IEK7-like/giEK2} & & & & & & & \\ \text{GL20803} \ 212795 & \text{Phosducin-like} & & & & & & & \\ \text{GL20803} \ 213795 & \text{Phosducin-like} & & & & & & \\ \text{GL20803} \ 213775 & \text{Phosducin-like} & & & & & & \\ \text{GL20803} \ 14373^* & \text{Dynamin} & & & & & & \\ \text{GL20803} \ 15264 & & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15264 & & & & & & & \\ \text{Hypothetical protein} & & & & & & & \\ \text{GL20803} \ 15264 & & & & & & & \\ \text{Hypothetical protein} & & & & & & & \\ \text{GL20803} \ 15264 & & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15246 & & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15246 & & & & & & & \\ \text{Hypothetical protein} & & & & & & & \\ \text{GL20803} \ 15246 & & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15246 & & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15246 & & & & & & \\ \text{Hypothetical protein} & & & & & & \\ \text{GL20803} \ 15246 & & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{GL20803} \ 15246 & & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{GL20803} \ 15726 & & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{GL20803} \ 15726 & & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{GL20803} \ 23577 & & & & & & \\ \text{Hypothetical protein} & & & & & \\ \text{GL20803} \ 23577 & & & & & & \\ Hypoth$	GL50803_9705	Hypothetical/YEATS domain	30,449	23	D		EST47502.1, 1E–27
GL30803_2831 Filsione acceptinalisense MTS12 49.910 12 Precedents 54-50 EST431221, 9E-76 GL50803_G886 Prokaryotic SMC domain protein 102,835 11 No EST43221, 9E-76 Signaling GL50803_686 Filsione acceptione 124,241 7 No EST43821, 10,61-15 GL50803_617 Putative DUB 150,389 16 EST43827, 10,0 EST443824, 1,1E-14 GL50803_6430° ERX7-like/giERX2 41,096 15 IP+ EST443827, 10,0 GL50803_6430° IA-53 protein 28,576 8 IP++ EST44321, 16-14 GL50803_15430 Protein disulfide isomerase PDI2 50,408 6 Precedents 57 EST43831, 15-43 Trafficking GL50803_15264 Hypothetical protein 404,245 153 No EST440221, 19E-76 GL50803_15454 Hypothetical protein 716,474 95 EST440221, 19E-76 No GL50803_15454 Hypothetical protein 75,198 S EST440221, 19E-76 No GL50803_15455 Hypothetical protein 716,474 95 EST440221, 19E-76 No	GL50803_8125	SMARCCI	47,004	17	Precedents	54-56	N0
GL3080_5080 F100H170ft SMC 600Hall protein 102,355 11 No GL50803_17461 SWIRM domain protein 124,241 7 EST43894.1,1E-14 Signaling GL50803_617 Putative DUB 150,389 16 P+ GL50803_122850* ERK7-like/giERX2 41,096 15 IP+ EST43894.1,1E-14 GL50803_12795 Phosducin-like 26,919 6 EST44531, 1,E-14 GL50803_12795 Phosducin-like 26,919 6 EST44531, 1,E-14 GL50803_12795 Phosducin-like 26,919 6 EST44521, 1,E-14 GL50803_1373* Dynamin 79,513 13 IP- 36 EST46023.1, 0.0 Unknown/Giardia specific GG50803_15264 Hypothetical protein 716,474 95 EST45121.1, 3E-25 GL50803_15264 Hypothetical protein 71,987 55 EST49821.1, 8E-39 GL50803_1742 Hypothetical protein 75,987 EST44942.1, 9E-40 No GL50803_1756 Zintyrin and WD repeat protein 59,785 EST49832.1, 2E-45	GL50803_2851	Histone acetyltransferase MYS12	49,916	12	Precedents	54-56	E5145122.1, 9E-78
	GL50803_17461	SWIRM domain protein	102,850	7			EST47570.1, 3E–51
	Signaling						
	GL50803_6317	Putative DUB	150,389	16			EST43894.1,1E-14
	GL50803_22850*	ERK7-like/giERK2	41,096	15	IP+		EST48827.1, 0.0
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	GL50803_6430*	14-3-3 protein	28,576	8	IP++		EST46224.1, 6E-115
GL 50803_9413 Protein disulfide isomerase PDI2 50,408 6 Precedents 57 EST43183.1, 5E-43 Trafficking GL 50803_14373* Dynamin 79,513 13 IP- 36 EST4023.1, 0.0 Unknown/Giardia specific GL 50803_15264 Hypothetical protein 404,245 153 No GL 50803_15264 Hypothetical protein 66,334 57 EST41505.1, 4E-25 GL 50803_17532 Hypothetical protein 716,474 95 EST44949.1, 9E-26 GL 50803_15485 Hypothetical protein 75,987 55 EST4394.1, 3E-19 GL 50803_15485 Hypothetical protein 56,506 53 No GL 50803_17807 WD-40 repeat protein 14,462 47 No GL 50803_17266 Ankyrin and WD repeat protein 90,538 29 EST41949.1, 2E-49 GL 50803_17394 Hypothetical protein 92,642 28 No No GL 50803_23897 Hypothetical protein 11,183 25 No GL 50803_3559 No GL 50803_3559 No GL 50803_3559<	GL50803_12795	Phosducin-like	26,919	6			EST41453.1, 1E-14
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	GL50803_9413	Protein disulfide isomerase PDI2	50,408	6	Precedents	57	EST43183.1, 5E–43
GL30805_143/3*Dynamin $79,513$ 13 $12^{}$ 36ES146023.1,0.0Unknown/Giardia specificGL50803_15264Hypothetical protein $404,245$ 153NoGL50803_39938Hypothetical protein $716,474$ 95EST41505.1,4E-25GL50803_17532Hypothetical protein $63,34$ 57EST445241.1,3E-19GL50803_13942Hypothetical Protein $56,506$ 53NoGL50803_13942Hypothetical Protein $56,506$ 53NoGL50803_17266Ankyrin and WD repeat protein $14,462$ 47 NoGL50803_17266Zinc finger domain protein $92,642$ 28NoGL50803_17596Zinc finger domain protein $92,642$ 28NoGL50803_23897Hypothetical protein $11,183$ 25NoGL50803_13726Hypothetical protein $20,414$ 22NoGL50803_20601Hypothetical protein $24,801$ 18NoGL50803_2359Hypothetical protein $24,801$ 18NoGL50803_37350Hypothetical protein $54,020$ 15NoGL50803_37350Hypothetical protein $76,898$ 11Sr43354.1,0.0GL50803_37350Hypothetical protein $76,898$ 11EST4354.1,0.0GL50803_37350Hypothetical protein $76,898$ 11EST4354.1,0.0GL50803_37350Hypothetical protein $76,898$ 11EST44629.1,6E-73GL50803_37350Hypothetical protein $76,898$ 11EST435	Trafficking			10			
Unknown/Giardia specific Vo GL50803_15264 Hypothetical protein 716,474 95 EST41505.1, 4E-25 GL50803_17532 Hypothetical protein 716,474 95 EST41505.1, 4E-25 GL50803_17532 Hypothetical protein 66,334 57 EST45241.1, 3E-19 GL50803_13942 Hypothetical Protein 55,056 53 No GL50803_13942 Hypothetical Protein 59,785 51 EST44362.1, 2E-65 GL50803_137711 Hypothetical Protein 719,629 41 EST41949.1, 2E-49 GL50803_137716 Hypothetical Protein 719,629 41 EST41949.1, 2E-49 GL50803_137711 Hypothetical Protein 719,629 41 EST41949.1, 2E-49 GL50803_137716 Hypothetical Protein 719,629 41 EST41949.1, 2E-49 GL50803_13792 Hypothetical Protein 11,83 25 No GL50803_13592 Hypothetical Protein 11,136 22 No GL50803_3540 Hypothetical Protein 11,136 22 No	GL50803_14373*	Dynamin	79,513	13	IP-	36	EST46023.1, 0.0
GL50803_15264 Hypothetical protein 404,245 153 No GL50803_39938 Hypothetical protein 716,474 95 EST41505.1, 4E-25 GL50803_15322 Hypothetical protein 66,334 57 EST41949.1, 9E-20 GL50803_1342 Hypothetical protein 55,066 53 No GL50803_1842 Hypothetical protein 59,785 51 EST4196.2.1, 2E-65 GL50803_17266 Ankyrin and WD repeat protein 719,629 41 EST41949.1, 2E-49 GL50803_17596 Zinc finger domain protein 90,538 29 EST41949.1, 2E-49 GL50803_3137711 Hypothetical protein 11,183 25 No GL50803_99726 Hypothetical protein 11,183 25 No GL50803_3590 Hypothetical protein 11,136 22 No GL50803_3559 Hypothetical protein 11,136 22 No GL50803_3559 Hypothetical protein 14,612 No No GL50803_3559 Hypothetical protein 11,36 22	Unknown/Giardia specific		101.015	150			
GL50803_37532 Hypothetical protein 716,474 95 ES145201,1,3E-19 GL50803_17532 Hypothetical protein 66,334 57 EST4521,1,3E-19 GL50803_15485 Hypothetical protein 751,987 55 EST41949,1,9E-20 GL50803_13942 Hypothetical Protein 59,785 51 EST44362,1,2E-65 GL50803_17266 Ankyrin and WD repeat protein 14,462 47 No GL50803_17596 Zinc finger domain protein 90,538 29 EST44949,1,2E-49 GL50803_17596 Zinc finger domain protein 92,642 28 No GL50803_2877 Hypothetical protein 11,183 25 No GL50803_9726 Hypothetical protein 489,165 24 No GL50803_13759 Hypothetical protein 11,136 22 No GL50803_31492 Hypothetical protein 24,801 18 No GL50803_14492 Hypothetical protein 51,753 11 No GL50803_37350 Hypothetical protein 51,753 11 No GL50803_37350 Hypothetical protein 76,898	GL50803_15264	Hypothetical protein	404,245	153			NO
GL50805_15485 Hypothetical protein 75,1987 55 55 5574194241,1,9E-20 GL50803_13942 Hypothetical protein 56,506 53 No GL50803_7807 WD-40 repeat protein 59,785 51 55 GL50803_17266 Ankyrin and WD repeat protein 719,629 41 55 51 GL50803_137711 Hypothetical protein 719,629 41 55 51 55 GL50803_137710 Hypothetical protein 719,629 41 55 51	GL50805_59958	Hypothetical protein	/16,4/4	95			ES141505.1, 4E-25
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	GL50803_92602	Hypothetical protein	342,328	6			EST44629.1, 6E-73

^{*a* *}, This protein was chosen for testing interaction with actin, as described in Results.

^b +++, strong interactor; ++, moderate interactor; +, weak interactor; -, no interaction detected.

^c The exponential values are BLAST Expect values, indicating the level of conservation between the homologs.

like kinase (S. Gourguechon and W. Z. Cande, unpublished data); because we did not find Polo in the actin data set, nor did we find actin in the Polo data set, we believe the low-abundance hits are likely false positives. The identity of these low-abundance hits may be useful for others using our same approach; therefore, we have identified the overlapping hits in Table S2 in the supplemental material.

Although once controversial, it is now clear that actin is part of the nucleoskeleton responsible for many nuclear processes, including transcriptional regulation, chromatin remodeling, and



FIG 4 Validation of identified interacting proteins. (A) Immunoprecipitation from *Giardia* extracts of C-terminally HA-tagged interactors, followed by anti-actin Western blotting demonstrates that these proteins interact with actin. (B) Colocalization of actin and HA-tagged interacting proteins in *Giardia* trophozoites. Actin is green, HA tagged proteins are red, DNA is blue. The first three columns are maximal projections, and the last column is a single slice through the middle of the cell. Arrowhead marks centrin localization associated with the posterolateral flagella. Scale bar, 5 μm.

general maintenance of genome organization and integrity (reviewed in reference 38). In contrast to localization studies performed in model eukaryotes, where it is difficult to detect actin in the nucleus, giActin is readily detectable in the nuclei, suggesting

that it has an important role in nuclear function (see Fig. 3B). Although we validated complex formation with actin for ARP7 and TIP49 (the second most abundant hit in terms of peptides/ molecular weight), we identified six other proteins containing domains that are consistent with a role in actin-based chromatin remodeling. It has been put forth that several proteins known to function in the cytoskeleton have roles in the nucleus; thus, they may have originally evolved to serve the genome (38). Our identification of conserved nuclear proteins and the lack of core cytoskeletal regulators are consistent with this notion.

Actin's role in the flagella is well established but largely ignored. Biochemical fractionation of flagella has shown that six of the seven inner dynein arm complexes are associated with actin (39). A conventional actin mutant of Chlamydomonas, ida5, lacks four of the inner-arm dynein complexes and, in this mutant, an actin-like protein, NAP, is upregulated to substitute for actin in the remaining two inner-arm dynein complexes, thus demonstrating the importance of actin to axonemal structure and function (39). Within the inner-arm dynein complexes, actin is thought to exist as a monomer in a complex with either a dimer of p28 or a monomer of centrin (32). Using super-resolution microscopy, we observed a regular repeating pattern for actin within all eight of the *Giardia* flagella (13). The precise molecular role of actin in the inner-arm dynein complexes remains enigmatic, but if actin simply acts as an adapter, it is perplexing to consider that Giardia may have lost proteins such as myosin, formin, and cofilin while maintaining actin's role in the flagella. Alternatively, some of the earliest functions of actin may include flagellar and nuclear processes.

Of the conserved proteins identified, 14-3-3 and ERK2 may be the most intriguing since they are likely regulators of actin dynamics or actin-related processes. 14-3-3 is known to play an important role in cytoskeletal regulation in other eukaryotes (40); however, the relationship between 14-3-3 and actin is complicated by multiple isoforms of 14-3-3 and conflicting results about 14-3-3 interactions with actin (reviewed in reference 41). In addition to our identification of 14-3-3 as an actin interactor, several efforts to define the 14-3-3 interactome in both Giardia and mammalian cells corroborate an actin-14-3-3 interaction (42-44). However, the current view is that 14-3-3 regulates actin through phosphodependent interaction with the actin-depolymerizing protein cofilin and does not bind to actin directly (45, 46). Notably, Giardia lacks both cofilin and its regulatory kinase LIM. Perhaps a more direct regulation of actin by 14-3-3 underlies the well-characterized cofilin-14-3-3 interaction. Our analysis of 14-3-3's role in actin regulation is in preparation (J. Krtková, J. W. Xu, and A. R. Paredez, unpublished data).

Giardial ERK2 is a potential regulator of actin-related processes. *Giardia* contains two ERK homologs: a prototypical ERK, giERK1, and the ERK7-like protein giERK2 (47, 48). Although ERK stands for extracellular signal-regulated kinase, ERK7, unlike other mitogen-activated proteins, is atypical in that it is thought to be auto-activated rather than responding to external signals (49, 50). giERK2 lacks the extended C-terminal domain found in ERK7 and may not be regulated in the same manner, and yet *in vitro* kinase assays have shown that giERK2 is much more active than giERK1 (48). In other eukaryotes, ERK7 kinases have been shown to regulate protein secretion and cell proliferation (50, 51). Our identification of an actin-ERK2 complex in *Giardia* and the localization of this kinase in the nucleus and throughout the cell do not exclude potentially conserved function.

This initial characterization of actin-associated proteins in *Giardia* has focused on validating interaction with conserved proteins (see above), both because of the evolutionary implications

and because these conserved proteins serve as proof of principle for our proteomic strategy. The largest group of identified proteins is, however, the novel/Giardia-specific category (see Table 1). We have begun to analyze the Giardia-specific interactors with the same endogenous-tagging approach used to study the conserved actin-interacting proteins. We have tagged 13 of the Giardia-specific proteins listed in Table 1 and have been able to immunoprecipitate giActin with 10 of these proteins (in preparation). Notably, five of these proteins localize to the nuclei, one localizes to the flagella, and the remaining proteins show a punctate pattern. None of the proteins identified thus far appear to colocalize with filamentous actin structures. Most canonical actin-binding proteins exclusively recognize globular or filamentous actin. The buffer conditions used in our affinity purification approach contained Ca²⁺ and ATP but lacked the Mg²⁺ and KCl typically found in buffers that support actin filament formation. Therefore, additional interactors remain to be discovered, and the set of interactors described here likely represents the globular giActin interactome. We did, however, test the ability of TS-Actin to polymerize, as assayed by ultracentrifugation (see Fig. S2 in the supplemental material). We observed that TS-Actin has some ability to pellet under filament-forming conditions that are consistent with the partial rescue we observed in our morpholino-knockdown experiments (Fig. 2). Future work will explore the identification of actin interactors in buffer conditions that support actin filament formation

In this first exploration of the giActin interactome, we found both conserved and *Giardia*-specific interactors. The subset of canonical actin-binding proteins in *S. salmonicida* suggests loss of actin-binding proteins from *Giardia*. Therefore, the retention of actin-interacting proteins in the nucleus and flagella suggest these processes are the most constrained of any actin processes. In any case, the role of actin in the nucleoskeleton and flagella are likely some of actin's most ancient functions and remain relatively unexplored compared to the role of actin in the cytoskeleton. The set of novel/*Giardia*-specific proteins remain intriguing. Many of these proteins have no recognizable domains; therefore, elucidation of their function will be a challenge. Simultaneously, if functional experiments demonstrate these proteins to be essential, they will become potential therapeutic targets to treat giardiasis.

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