A Toxoplasma gondii Class XIV Myosin, Expressed in Sf9 Cells with a Parasite Co-chaperone, Requires Two Light Chains for Fast Motility*§

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Background: TgMyoA, a Toxoplasma gondii class XIV myosin, is an important parasite virulence factor.

Results: Expression of functional TgMyoA in Sf9 cells requires co-expression with a T. gondii myosin co-chaperone.

Conclusion: TgMyoA can be heterologously expressed and needs two bound light chains to propel actin at fast speeds.

Significance: High yield expression of TgMyoA will enable structure-function studies and drug screening.

Many diverse myosin classes can be expressed using the baculovirus/Sf9 insect cell expression system, whereas others have been recalcitrant. We hypothesized that most myosins utilize Sf9 cell chaperones, but others require an organism-specific co-chaperone. TgMyoA, a class XIVa myosin from the parasite Toxoplasma gondii, is required for the parasite to efficiently move and invade host cells. The T. gondii genome contains one UCS family myosin co-chaperone (TgUNC). TgMyoA expressed in Sf9 cells was soluble and functional only if the heavy and light chain(s) were co-expressed with TgUNC. The tetratricopeptide repeat domain of TgUNC was not essential to obtain functional myosin, implying that there are other mechanisms to recruit Hsp90. Purified TgMyoA heavy chain complexed with its regulatory light chain (TgMLC1) moved actin in a motility assay at a speed of ~1.5 μm/s. When a putative essential light chain (TgLCL1) was also bound, TgMyoA moved actin at more than twice that speed (~3.4 μm/s). This result implies that two light chains bind to and stabilize the lever arm, the domain that amplifies small motions at the active site into the larger motions that propel actin at fast speeds. Our results show that the TgMyoA domain structure is more similar to other myosins than previously appreciated and provide a molecular explanation for how it moves actin at fast speeds. The ability to express milligram quantities of a class XIV myosin in a heterologous system paves the way for detailed structure-function analysis of TgMyoA and identification of small molecule inhibitors.

Toxoplasma gondii, a member of the phylum Apicomplexa, is a common infectious agent of humans that can cause severe disease in immunocompromised individuals and the developing fetus. This obligate intracellular parasite must invade a host cell and replicate to survive. The invasive stage of the parasite relies on a unique form of substrate-dependent motility called gliding motility, which is driven by a class XIVa myosin motor, TgMyoA.4 The TgMyoA heavy chain, one of 11 myosin heavy chains found in T. gondii (1), is important for efficient parasite motility, invasion, and egress from the host. Parasites lacking TgMyoA are avirulent in a mouse model of infection (2).

The TgMyoA motor is located between the plasma membrane and the inner membrane complex (IMC), a double membrane that is continuous around most of the cell (3) (see Fig. 1). TgGAP50 (a 50-kDa gliding-associated protein), an integral membrane glycoprotein of the IMC, acts as a membrane receptor for the motor (4); TgMyoA is linked indirectly to TgGAP50 through an apicomplexan-specific N-terminal extension of its regulatory light chain, TgMLC1, and TgGAP45 (a 45-kDa gliding-associated protein). Two other membrane-associated proteins, TgGAP40 (a 40-kDa gliding-associated protein) and TgGAP70 (a 70-kDa gliding-associated protein), have recently been identified as additional components of this myosin motor complex (5). The precise mechanism by which the motor complex generates motility is the subject of intense investigation (see Fig. 1 for one model), but there is little debate about the central importance of TgMyoA in the process.

TgMyoA is a single-headed motor with several unusual features that distinguish it from other myosin motors. It lacks a highly conserved amino acid in a key actin-binding surface loop that is typically either an acidic residue or a phosphorylatable residue but is a Gln in TgMyoA. It also lacks a conserved Gly in a helix in the motor domain that is thought to act as a pivot point for motion of the light chain binding lever arm (6, 7). The lever arm amplifies small motions at the active site into the larger motions needed to move actin. The light chain-binding region of myosin heavy chains also typically contains one or

The abbreviations used are: TgMyoA, T. gondii myosin A (TgMyoA heavy chain with or without bound light chain(s)); IMC, inner membrane complex; MTIP, myosin tail-interacting protein; TgLCL1, T. gondii essential light chain-1; TgGAP40, 45, 50, and 70 kDa, respectively; TgMLC1, T. gondii myosin light chain-1; TPR, tetratricopeptide repeat; TgUNC, T. gondii myosin co-chaperone of the UCS family; UCS, a domain conserved in UNC-45/CRO1/She4p proteins.

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more α-helical “IQ motifs” (consensus sequence IQXXRXGXXX), which bind specific myosin light chains and/or calmodulin. The sequence of the IQ motif in TgMyoA is degenerate (6, 7), and thus the number of bound light chains is unclear.

Isolation of a functional motor complex from *T. gondii* showed that despite these unique features TgMyoA was a bona fide motor able to propel actin at fast speeds of ~5 μm/s in an *in vitro* motility assay (7–9). The speed observed *in vitro* is similar to the speed of parasite motility on two-dimensional surfaces (1–3 μm/s) (10) and in three-dimensional Matrigel (maximum speed, 2 μm/s) (11). The *in vitro* biophysical characterization of TgMyoA was a tour de force given the limited amounts of protein that could be isolated from the parasite (7). However, it raised some interesting questions. For example, how can a single-headed myosin with only one bound light chain propel actin at such fast speeds? The speed of actin movement by myosin depends on the kinetics of nucleotide binding and release from the motor domain as well as the length of the lever arm, which is determined by the number of bound light chains (12, 13). The unitary step size of TgMyoA was measured to be ~5.3 nm (7), which is strikingly similar to the step size measured for single-headed subfragment-1 from either fast skeletal muscle myosin or smooth muscle myosin, both of whose lever arms are known to be stabilized by two bound light chains (for a review, see Ref. 14).

A recent mass spectrometry-based study, which analyzed proteins that bound to the motor complex in the presence and absence of calcium, identified a smaller ~15-kDa second calmodulin-like light chain in the motor complex immunoprecipitates, which they called the essential light chain (TgELC1) (15). The functional consequences of TgMLC1 and TgELC1 binding to the TgMyoA heavy chain are unknown. To address structure-function questions like these, it is essential to have an expression system that produces sufficient amounts of pure, functional motor protein.

Unfortunately, our previous efforts to express soluble and functional TgMyoA using the baculovirus/Sf9 insect cell expression system were unsuccessful. This was surprising given that many classes of myosin are successfully expressed and properly folded in Sf9 cells including class II smooth (16) and non-muscle myosins (17) and many classes of unconventional myosins including classes V (18–20), VI (21), and VII (22). There is another exception: functional class II striated muscle myosins have not been expressed in Sf9 cells but can be expressed in the C2C12 muscle cell line, implying that proper folding of striated muscle myosins requires factors that are present in muscle cells but absent from Sf9 cells (23–25). Similarly, although active TgMyoA can be purified from *T. gondii* (7–9), we speculated that a parasite-specific factor was needed for proper folding of TgMyoA in Sf9 cells.

Myosin folding and assembly into filaments in striated muscle is mediated by the general chaperones Hsc70 and Hsp90 and a striated myosin-specific co-chaperone of the UCS protein family, Unc45b (26–28). A second myosin chaperone called Unc45a is expressed in all vertebrate tissues (29). Unc45b forms a stable complex with Hsp90, binds only to the unfolded striated muscle myosin motor domain, and promotes motor domain folding (26–28). The N-terminal region of Unc45 interacts with Hsp90 via three tetratricopeptide (TPR) repeats, a degenerate 34-amino acid motif that is involved in protein-protein interactions (30). The central domain has no known function, and the C-terminal UCS domain of ~450 amino acids interacts with myosin (26). The UCS domain is named for the three founding members of this family: UNC-45 from *Caenorhabditis elegans*, CRO1 from the fungus *Podospora anserina*, and She4p from *Saccharomyces cerevisiae*.

In the present study, we identified a single unc-45 homolog in *T. gondii*, referred to as TgUNC, that encodes a protein containing all three domains found in the canonical Unc45 protein. We showed that isolation of functional TgMyoA from Sf9 cells strictly requires co-expression with TgUNC. Using this novel co-expression strategy, we showed that recombinant TgMyoA simultaneously binds both TgMLC1 and TgELC1, similar to the domain structure of conventional class II myosins. The functional consequence of having two bound light chains was a doubling of the speed of actin movement in motility assays compared with that observed when TgMyoA has only a single bound light chain (TgMLC1). The successful expression of TgMyoA will enable structure-function analysis of this unusual class of myosin motor proteins as well as identification of small
molecule inhibitors that can serve as leads for new drugs to prevent or treat toxoplasmosis.

**EXPERIMENTAL PROCEDURES**

**Expression Constructs**—Full-length (FL) TgUNC (GenBank™ accession number XM_002367241.1) was amplified from *T. gondii* cosmids number p559; (cosmid generously provided by Dr. M. J. Gubbels) using the primer pair TPRForANcol (AGCACATGGAGATTGGTCAAGCGC) and TPRRevAGFull MycStpKnl (CGCCGGATCTCAGGTCTCTTTCAGAGATCAGTATCTGGTTGAGTTG). A PCR product coding for TgUNC followed by a Myc tag was cloned into the baculovirus transfer vector pAcSG2 (BD Biosciences). TgUNC pAcSG2 was truncated to make two other constructs. TgUNCΔTPR starts at residue Leu-161, eliminating the TPR region but keeping the central and UCS domains along with the C-terminal Myc tag. The TgUNC U5-TPR consists starts with residue Glu-682, thus eliminating both the TPR and central domains.

The plasmid pEB2-FLAG-MyaO⁵ was used as a template to amplify a product coding for N-terminally FLAG-tagged TgMyoA heavy chain (accession number AAC47724.1) using primer pair EcoR1-FLAG-MyaF (GGGGAATTCATGGACATTGCAGACATCATGACATCGAGATGCATCACTACAGGATGCAAGGCGACGATGATGACGACAAGATGGC) and c-TgMyoA-EcoR1 (GGGGGAATTCATGGACATTGCAGACATCATGACATCGAGATGCATCACTACAGGATGCAAGGCGACGATGATGACGACAAGATGGC) and cloned into the bacterial expression vector pET3a (Novagen).

A dual light chain plasmid was constructed for immunoprecipitation studies. DNA coding for TgELC1 with a C-terminal 3xHA tag was cloned into vector pAcUW51 downstream of the p10 promoter, whereas DNA coding for TgMLC1 with an N-terminal 3xMyc tag was cloned downstream of the polH promoter. All constructs were sequenced prior to transfection.

**Co-immunoprecipitations**—Sf9 cells were harvested 72 h after infection with recombinant baculoviruses (TgMyoA heavy chain and tagged light chains) and lysed with 10 mM imidazole, pH 7.4, 150 mM NaCl, 1 mM EGTA, 5 mM MgCl₂, 7% (w/v) sucrose, 3 mM Na₂3P, 1% (v/v) Nonidet P-40, 1 mM DTT, and 1 μM protease inhibitor mixture (Sigma-Aldrich, catalogue number P8340). Following addition of 5 mM MgATP, the lysate was spun at 350,000 × g for 20 min at 4 °C. The supernatant was incubated with either rabbit anti-TgMLC1 (a generous gift from Dr. Con Beckers) or rat anti-HA (Roche catalogue number 11867423001) overnight at 4 °C. Rec-Protein A-Sepharose beads (Invitrogen) were added, and the samples were incubated at 4 °C for 60 min. The beads were washed four times with lysis buffer. Bound proteins were eluted by boiling in SDS-PAGE sample buffer, centrifuged at 100 × g for 2 min, and resolved on 4–12% gradient gels. The protein was transferred to Immobilon-FIL (Millipore, Bedford, MA) and probed with mouse anti-Myc 9E10 (1:6,000; Developmental Studies Hybridoma Bank, University of Iowa), rat anti-HA (1:4,000), or rabbit anti-HA (1:2,000; Abcam catalogue number 9110) and mouse anti-FLAG (1:7,500; Sigma-Aldrich), LI-COR Biosciences (Lincoln, NE) secondary antibodies (anti-mouse IRDye680RD, anti-rabbit IRDye800CW, and anti-rat IRDye800CW) were used according to manufacturer’s instructions, and blots were scanned using an Odyssey CLx Infrared Imaging System (LI-COR Biosciences).

**Protein Expression and Purification**—Sf9 cells were co-infected with recombinant baculovirus coding for TgMyoA heavy chain (tagged at the C terminus with a Bio tag and FLAG tag), untagged light chain(s), and the co-chaperone TgUNC. The cells were grown in medium supplemented with 0.2 mg/ml biotin. After 72 h, the cells were lysed by sonication in 10 mM imidazole, pH 7.4, 0.2 mM NaCl, 1 mM EGTA, 5 mM MgCl₂, 7% (w/v) sucrose, 2 mM DTT, 0.5 mM 4-(2-aminoethyl)benzenesulfonyl fluoride, 5 μg/ml leupeptin, and 5 mM benzamidine. To determine TgMyoA heavy chain solubility, the extracts were centrifuged at 350,000 × g for 20 min. For motor purification, bacterially expressed TgMLC1 and TgELC1 at 25 μg/ml (final concentration) each and 5 mM MgATP were added to the lysate, which was then clarified at 200,000 × g for 30 min. The supernatant was applied to a FLAG affinity resin column. The cultures were induced with 0.4 mM isopropyl-β-D-thiogalactopyranoside and grown overnight at 27 °C.

⁵ L. Tilley and G. E. Ward, unpublished data.
⁶ A. Heaslip and G. E. Ward, unpublished data.
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before being pelleted and frozen. The pellets were lysed by sonication in 10 mM sodium phosphate, pH 7.4, 0.3 mM NaCl, 0.5% (v/v) glycerol, 7% (w/v) sucrose, 7 mM β-mercaptoethanol, 0.5 mM 4-(2-aminoethyl)benzenesulfonyl fluoride, and 5 μg/ml leupeptin. The cell lysate was clarified at 200,000 × g for 30 min. TgELC1, which is found in the supernatant, was boiled for 10 min in a double boiler and then clarified at 26,000 × g for 30 min. Soluble protein was applied to a HIS-Select ρnickel affinity column (Sigma-Aldrich). Nonspecifically bound protein was removed by washing the resin with buffer A (10 mM sodium phosphate, pH 7.4 and 0.3 mM NaCl). TgELC1 was then eluted from the column with buffer A containing 200 mM imidazole. The protein was dialyzed overnight against 10 mM imidazole, pH 7.4, 150 mM NaCl, 1 mM EGTA, 1 mM MgCl₂, and 50% (v/v) glycerol.

Bacterially expressed TgMLC1 is found in the insoluble inclusion bodies. The cell lysate was clarified at 26,000 × g for 10 min. The pellet was dissolved in 20 ml of 8 M guanidine, 150 mM NaCl, 10 mM NaPO₄, pH 7.5, and 10 mM DTT and stirred at room temperature until dissolved. It was then clarified at 200,000 × g for 30 min and dialyzed overnight against 2 × 1 liter of buffer A containing 7 mM β-mercaptoethanol and 1 μg/ml leupeptin. The next day the sample was clarified at 26,000 × g for 30 min, and the supernatant was applied to a HIS-Select ρnickel affinity column. The column was washed with 15 ml of dialysis buffer and TgMLC1 was eluted with buffer A containing 200 mM imidazole. The protein was dialyzed overnight against 10 mM imidazole, pH 7.4, 150 mM NaCl, 1 mM EGTA, 1 mM MgCl₂, and 50% (v/v) glycerol. Both purified light chains were stored at −20 °C.

Gels—Proteins were separated on a 4–12% Bis-Tris NuPAGE gel (Invitrogen) run in MES buffer according to the NuPAGE technical guide.

In Vitro Motility—To prepare the flow cell, 0.2 mg/ml biotinylated BSA in buffer B (150 mM KCl, 25 mM imidazole, pH 7.5, 1 mM EGTA, 4 mM MgCl₂, and 10 mM DTT) was added to the nitrocellulose-coated flow cells for 1 min followed by three rinses with 0.5 mg/ml BSA in buffer B. Neutravidin (50 μg/ml; Thermo Scientific) in buffer B was applied for 1 min followed by three rinses with buffer B. Before introduction into the flow cell, TgMyoA was mixed with a 2-fold molar excess of F-actin and 10 mM MgATP in buffer B and centrifuged for 25 min at 350,000 × g to remove ATP-insensitive myosin heads. TgMyoA (0.5 μM) was then introduced into the flow cell. To further block any ATP-insensitive heads, 1 μM vortexed F-actin in buffer C (50 mM KCl, 25 mM imidazole, pH 7.5, 1 mM EGTA, 4 mM MgCl₂, and 10 mM DTT) was added for 60 s followed by a 10 mM MgATP wash. Rhodamine-phalloidin-labeled actin was then introduced for 1 min followed by one rinse with buffer C. Three volumes of buffer C, which also contained 5 mM MgATP (unless stated otherwise), 0.5–0.7% methylcellulose, 1 μM TgMLC1, 1 μM TgELC1, 3 mg/ml glucose, 0.125 mg/ml glucose oxidase (Sigma-Aldrich), and 0.05 mg/ml catalase (Sigma-Aldrich) were then flowed into the cell. When assayed with calcium, 1.2 mM calcium was added to this buffer.

Actin movement was observed at 30 °C using an inverted microscope (Zeiss Axiovert 10) equipped with epifluorescence, a Rolera MGi Plus digital camera, and dedicated computer with the Nikon NIS Elements software package. Data were analyzed using a semiautomated filament tracking program described previously (31). The velocities of >600 filaments were determined. Speeds were fit to a Gaussian curve.

Actin-activated ATPase Activity—Assays were performed in 10 mM imidazole, pH 7.0, 5 mM NaCl, 1 mM MgCl₂, 1 mM NaN₃, and 1 mM DTT at 30 °C. Purified TgMyoA (52 nm) was incubated with various concentrations of skeletal actin. Activity was initiated by the addition of 5 mM MgATP and stopped with SDS every 2 min for 8 min. Inorganic phosphate was determined colorimetrically (32). A low salt concentration was needed to keep the Kₘ values as low as possible. Data were fit to the Michaelis-Menten equation.

Sedimentation Velocity—Sedimentation velocity runs were performed at 20 °C in an Optima XL-I analytical ultracentrifuge (Beckman Coulter) using an An60Ti rotor at 30,000 rpm. The solvent was 20 mM HEPES, pH 7.4, 0.1 M NaCl, and 2 mM DTT. An N-terminally FLAG-tagged TgMyoA heavy chain (no Bio tag) bound to TgMLC1 was used for this experiment. The sedimentation coefficient was determined by curve fitting to one species using the dc/dt program (33).

RESULTS

TgMyoA Heavy Chain Is Insoluble When Expressed in Sf9 Cells—Recombinant baculoviruses encoding TgMyoA heavy chain and its regulatory light chain, TgMLC1, were used to co-infect Sf9 cells. The C terminus of the TgMyoA heavy chain contained two tags: the FLAG tag facilitates purification by affinity chromatography, and the Bio tag, which becomes biotinylated within the Sf9 cells (34), allows the motor to be specifically attached via its C terminus to a streptavidin-coated coverslip for in vitro motility assays (Fig. 2A). Unfortunately, although both TgMyoA heavy chain and TgMLC1 were expressed following infection as detected by Western blotting of the total Sf9 cell lysate, none of the TgMyoA heavy chain was present in the soluble fraction (Fig. 3).

Identification of a UCS Family Gene in the T. gondii Genome—We speculated that the endogenous Sf9 protein folding machinery was not sufficient to fold this unusual myosin and that a parasite-specific cofactor was required. Potential candi-
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Production of Soluble TgMyoA in Sf9 Cells Requires Co-expression with TgUNC—In marked contrast to what was observed in the absence of TgUNC, co-expression of TgMyoA heavy chain and TgMLC1 with TgUNC yielded soluble protein (Fig. 3). We then set up a large scale infection for protein purification, whereas TgMyoA heavy chain was detected with anti-FLAG antibody.

To determine whether calcium regulates the speed at which TgMyoA moves actin, an in vitro motility assay was performed in the presence or absence of free calcium. TgMyoA was preincubated in buffer containing either 0.2 mM free calcium or 1 mM EGTA for 1 h before the assay. Whether the myosin contained TgMLC1 only or both light chains, calcium did not affect the speed of actin movement (Fig. 5, open symbols).

Steady-state Actin-activated ATPase Activity—The speed of actin filament movement increased with MgATP concentration in the in vitro motility assay (Fig. 7A). The fit to a rectangular hyperbola defined a maximal speed of 4.6 ± 0.3 μm/s and a K_m of 1.3 ± 0.3 mM MgATP. Steady-state actin-activated ATPase assays were thus performed with 5 mM MgATP. Rates of ATP hydrolysis were determined as a function of skeletal actin concentration, and data were fit to the Michaelis-Menten equation. TgMyoA with both light chains bound showed a V_max of 84 ± 9.5 s^{-1} and a K_m for actin of 136 ± 22 μM at 30 °C (Fig. 7B).

In a separate set of experiments, we showed that the actin-activated ATPase activity of TgMyoA with only TgMLC1 bound was indistinguishable from that obtained with TgMyoA containing both light chains (p > 0.1, Student's t test). The values obtained were 55.4 ± 5.4 s^{-1} (n = 5) for TgMyoA with expressed TgMyoA was slower than that obtained with myosin isolated from parasites (7, 9), which led us to question the reason for the difference.

Binding Both TgMLC1 and TgELC1 to the TgMyoA Heavy Chain Doubles the Speed of Actin Movement—In addition to TgMLC1, an essential light chain called TgELC1 was recently identified as part of the myosin motor complex (15). To test whether TgMLC1 and TgELC1 bind simultaneously to the same heavy chain, we prepared a recombinant baculovirus containing both light chains, each with a different tag. TgMLC1 had an N-terminal 3xMyc tag, whereas TgELC1 had a C-terminal 3xHA tag. Following infection with heavy and light chains, TgMyoA was immunoprecipitated from the Sf9 cell lysate with either an anti-HA antibody (TgELC1) or an anti-TgMLC1 antibody (Fig. 6). The Myc antibody could not be used for TgMLC1 immunoprecipitation because TgUNC was also tagged with Myc. The eluates were analyzed by Western blotting with antibodies to detect TgUNC (anti-Myc), TgMyoA heavy chain (anti-FLAG), TgMLC1 (anti-Myc), and TgELC1 (anti-HA).

The results showed that the proteins co-immunoprecipitating with TgMLC1 included TgELC1 and conversely that the proteins co-immunoprecipitating with TgELC1 included TgMLC1. Earlier work showed that individual motor complexes do not physically associate with each other (9), and thus we conclude that both light chains simultaneously bind to the TgMyoA heavy chain.

When TgELC1 was co-expressed with TgMyoA heavy chain, TgMLC1, and TgUNC, both light chains co-purified with the heavy chain, indicating that both TgMLC1 and TgELC1 are tightly bound subunits of TgMyoA (Fig. 4A, lane 3). When TgMyoA containing both light chains was assayed in the in vitro motility assay, it moved actin at 3.4 ± 0.7 μm/s, which is more than twice the speed seen with TgMyoA containing only TgMLC1 (Fig. 5, solid red circles).

Calcium Does Not Regulate Motility Speed in Vitro—To determine whether calcium regulates the speed at which TgMyoA moves actin, an in vitro motility assay was performed in the absence of free calcium. TgMyoA was preincubated in buffer containing either 0.2 mM free calcium or 1 mM EGTA for 1 h before the assay. Whether the myosin contained TgMLC1 only or both light chains, calcium did not affect the speed of actin movement (Fig. 5, open symbols).
only TgMLC1 bound and 55.6 ± 10.4 s⁻¹ (n = 6) for TgMyoA with both light chains (at 60 μM actin). These values were obtained at 2 mM MgATP to decrease the ionic strength (relative to Fig. 7B, which was done at 5 mM MgATP). The lower ionic strength decreases the $K_m$ and allows the measured values to be nearer $V_{max}$.

The TPR Domain of TgUNC Is Not Required to Express Functional Myosin—We next determined the minimal domain of TgUNC that produces an active motor. In addition to full-length TgUNC, two shorter constructs were cloned (Fig. 2B): one contained the central and UCS domains (ΔTPR), whereas the other contained the UCS domain only. Small scale infections were performed using these three TgUNC constructs, and expression and solubility of TgMyoA were determined by Western blotting (Fig. 8). Total and soluble fractions were probed with either anti-FLAG (top panel) for TgMyoA heavy chain or anti-Myc (bottom panel) for TgUNC and its truncations. The results showed that the TPR domain was dispensable to obtain soluble myosin, whereas the UCS domain alone produced very little soluble myosin.

To determine the functionality of the TgMyoA co-expressed with these truncated chaperones, large scale co-infections and purifications were done with TgMyoA heavy chain, the two light chains, and each of the TgUNC constructs. All three infections yielded protein, but the yields were reduced for the two shorter TgUNC constructs. Per 10⁹ Sf9 cells, full-length TgUNC yielded ~1.5 mg of TgMyoA, ΔTPR yielded ~0.35 mg of TgMyoA, and UCS yielded only ~0.1 mg of TgMyoA. In the
in vitro motility assay, the motor expressed with either full-length TgUNC or ∆TPR generated similar actin sliding speeds (3.4 ± 0.7 and 3.1 ± 0.5 μm/s, respectively; see Fig. 9). In contrast, speeds decreased more than 4-fold to 0.7 ± 0.2 μm/s when TgMyoA was expressed in the presence of the UCS domain alone (Fig. 9).

DISCUSSION

The first major finding of this study is that when the class XIVa myosin TgMyoA simultaneously binds two light chains, the well established TgMLC1, and a second light chain called TgELC1 (7, 15) the speed at which TgMyoA moves actin in an in vitro motility assay is more than twice that obtained when only TgMLC1 is bound (1.5 versus 3.4 μm/s). TgMyoA had previously been described as containing a motor domain that binds actin and MgATP followed by a short tail. Our data imply that TgMyoA has a fairly conventional lever arm with two bound light chains formed by TgELC1 and TgMLC1 binding to adjacent sites near the C terminus of the TgMyoA heavy chain as suggested previously (15) (Fig. 10). The lever arm amplifies small changes at the active site into the larger motions necessary to propel actin at fast speeds, and the length of the lever arm dictates the speed (for a review, see Ref. 14). Consistent with a number of other studies with skeletal muscle myosin (35), smooth muscle myosin (13), Dictyostelium myosin (36), and myosin V (12), the light chains affect the in vitro motility speed by changing the lever arm length, not by changing the kinetics of the motor domain.

The faster speed we obtained when both light chains were bound (~3.4 μm/s) is close to the speed of the motor complex isolated from parasites (~5 μm/s) (7, 9). The difference in the two values may be due to the fact that we used a semiautomated tracking program that calculates speeds of hundreds of moving filaments without user bias, whereas prior studies tracked a smaller number of individual filaments manually. A more interesting possibility would be that accessory proteins such as GAP45, which were not included in our study, can influence motor speed.

Role of TgMyoA Light Chains—TgELC1 was only recently identified as a component of the TgMyoA motor (15) because with a molecular mass of only ~15 kDa it can be easily missed depending on how the motor complex components purified from parasites are resolved by SDS-PAGE. Our data and those of Nebel et al. (15) establish that TgELC1 is a bona fide subunit of TgMyoA. In principle, sequence analysis could have predicted that a second light chain would bind, but the region of the heavy
residue at position 7 of the IQ motif in T. gondii heavy chain in a compact conformation (39, 40). Presumably the Arg instead of the consensus Gly of the IQ motif (44) might regulate motor function. However, our data show no calcium-dependent changes in the speed of actin movement, and thus calcium binding does not directly regulate this motor. Because the motility assay is performed in the presence of MgATP as expected for a single-headed motor. The extrapolated maximal ATPase rate of ~80 s⁻¹ gives a total time per cycle of MgATP hydrolysis of ~12 ms. From the measured unitary step size (d_unit) of 5.3 nm (7) and a speed of actin movement of ~3.4 µm/s (v), the time the motor spends strongly attached to actin is ~1.5 ms (t_on = d_unit/v). This is a small percentage of the total cycle time, and thus this motor has a low duty cycle as do most motors designed for speed. Earlier studies also concluded that TgMyoA has a low duty cycle based either on kinetic (7) or in vitro motility data (9). An unusual feature of TgMyoA is the high MgATP concentration (1.3 mM) required to obtain half-maximal speed in the motility assay. This apparent low affinity for substrate is consistent with the K₉ of ~800 µM measured for the dissociation constant of ADP from the actomyosin complex (7).

Folding of TgMyoA into a Functional Motor Requires a Par- site-specific Myosin Co-chaperone—A second major finding of this work is that a T. gondii myosin co-chaperone is required to properly fold TgMyoA heavy chain in the baculovirus/Sf9 insect cell expression system. This is the first demonstration that functional class XIV myosin from an apicomplexan parasite can be expressed in a heterologous system. Although TgMyoA can be purified directly from the parasite, the yields are low, making it difficult to rigorously characterize the parasite-derived motor complex. In contrast, yields of 1 mg of TgMyoA were readily attainable from 1 x 10⁶ infected Sf9 cells (200-ml culture), which is comparable with yields obtained with myosins that do not require co-expression with an exogenous chaperone.

TgUNC has all three domains found in the canonical Unc45 protein, i.e. an N-terminal TPR domain, a central domain, and a C-terminal UCS domain that binds to myosin. The crystal structure of UCS family myosin chaperones from S. cerevisiae (43), Drosophila melanogaster (44), and C. elegans (45) have been solved. She4p, the UCS family protein from budding yeast, lacks a TPR domain, whereas DmUnc45 from Drosophila con-
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tains one, but it was disordered and not seen in the crystal structure. The structure of the C. elegans UNC-45 was the first to show the structure of the full-length protein. The UNC-45 structure resembles a checkmark with the shorter arm consisting of the TPR and central domain and the longer arm consisting of the UCS domain (43–45). The structure is composed almost entirely of stacked α-helical motifs known as arginine-rich motif repeats. Forty percent of the arginine-rich motif consensus residues identified in DmUnc45 (44) are identical in TgUNC (supplemental Fig. S1), making it highly likely that their overall structures are similar. Based on sequence homology within the UCS domains of functionally diverse UCS proteins, there is a strikingly high degree of identity at positions that map to a hydrophobic groove near the C terminus of the UCS domain. Based on this, it was proposed that the last five arginine-rich motifs (numbered 17–21) are the sites of interaction with myosin (44).

A novel feature observed with the C. elegans UNC-45 was its ability to form a scaffold of linear protein chains, a desirable feature in that it offers multiple adjacent binding sites for Hsp70, Hsp90, and the client protein myosin (45). It would be interesting to know whether this seemingly advantageous oligomerization is a feature common to all three-domain UCS family proteins including TgUNC.

The N-terminal TPR domain of TgUNC was dispensable for expression of functional TgMyoA in S9 cells. Similarly, the TPR domain of C. elegans UNC-45 is not required to rescue lethal unc-45-null mutants arrested in embryonic muscle development (46) even though this domain is typically thought to be essential for recruiting the chaperone Hsp90 to the client protein myosin. The comparable UCS family proteins in both budding (Sh4p4) and fission (rng3p) yeast also lack a TPR domain, but Rng3p still interacts with fission yeast Hsp90 (Swo1p) by yeast two-hybrid and co-immunoprecipitation analyses (47). This result implies that in fission yeast Hsp90 can bind to another domain of the UCS protein either directly or via another protein. Perhaps TgUNC also has alternative binding sites for Hsp90 that are utilized in the absence of the TPR domain. Alternatively, Hsp90 and TgUNC may act sequentially through the TPR and central domain and the longer arm of the UCS domain (43–45). The structure is composed almost entirely of stacked α-helical motifs known as arginine-rich motif repeats. Forty percent of the arginine-rich motif consensus residues identified in DmUnc45 (44) are identical in TgUNC (supplemental Fig. S1), making it highly likely that their overall structures are similar. Based on sequence homology within the UCS domains of functionally diverse UCS proteins, there is a strikingly high degree of identity at positions that map to a hydrophobic groove near the C terminus of the UCS domain. Based on this, it was proposed that the last five arginine-rich motifs (numbered 17–21) are the sites of interaction with myosin (44).

Although we have clearly established a requirement for TgUNC in TgMyoA heavy chain folding, many questions still remain about the folding pathway of myosins in general. Why can so many myosins utilize the endogenous folding machinery of S9 cells, whereas others such as TgMyoA and vertebrate skeletal and cardiac myosins require specific cofactors? Do other T. gondii myosins (48) require TgUNC for folding? While this paper was under review, it was shown that co-expression with Unc45b and Hsp90 led to the successful purification of mouse myosin 15 from S9 cells (49), demonstrating that chaperone-assisted expression may be a useful approach for other myosins. Regardless of the answers to these and other basic mechanistic questions about myosin folding, we have accomplished here the long sought after goal of obtaining milligram quantities of TgMyoA that will for the first time make high-throughput screening for drugs against this unusual, virulence-associated motor possible. Furthermore, the genomes of other apicomplexan parasites encode homologs of TgUNC, suggesting that this approach may be similarly useful for the expression of functional class XIV myosins from other apicomplexan parasites of medical or veterinary importance.

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