Isolation and Identification of Actin-binding Proteins in *Plasmodium falciparum* by Affinity Chromatography

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The invasion of the erythrocyte by *Plasmodium falciparum* depends on the ability of the merozoite to move through the membrane invagination. This ability is probably mediated by actin dependent motors. Using affinity columns with G-actin and F-actin we isolated actin binding proteins from the parasite. By immunoblotting and immunoprecipitation with specific antibodies we identified the presence of tropomyosin, myosin, α-actinin, and two different acts in the eluate corresponding to F-actin binding proteins. In addition to these, a 240-260 kDa doublet, different in size from the erythrocyte spectrin, reacted with an antibody against human spectrin. All the above mentioned proteins were metabolically radiolabeled when the parasite was cultured with 35S-methionine. The presence of these proteins in *P. falciparum* is indicative of a complex cytoskeleton and supports the proposed role for an actin-myosin motor during invasion.

Key words: actin - α-actinin - malaria - myosin - spectrin - tropomyosin - *Plasmodium falciparum* - affinity chromatography

The invasion of erythrocytes by parasites of the genus *Plasmodium* is a phenomenon well-described from a morphological and ultrastructural point of view (Aikawa et al. 1978, Bannister & Dluzewski 1990). However, the machinery and the molecular mechanisms involved in the process are practically unknown. The penetration of the erythrocyte depends on the ability of the merozoite to move through the membrane invagination. This ability is probably mediated by actin dependent motors since it is inhibited by Cytochalasin B (Miller et al. 1979, Field et al. 1993). Furthermore, in a related parasite *Toxoplasma* the involvement of actin fibers has been directly demonstrated (Dobrowolski & Sibley 1996).

The cytoskeleton of *Plasmodium* is poorly known. Two actin genes have been characterized: one, coding for a 376-aminoacid residues protein, is expressed during the sexual and the asexual cycles and its sequence is highly divergent from other actin genes found in lower eukaryotes. The other one is expressed only in sexual forms (Wesseling et al. 1988a,b, 1989). With a DNase-I inhibition assay it has been possible to show the simultaneous presence of actin fibers (F-actin) and monomers (G-actin) in the merozoite cytoplasm. Two proteins, 43 and 55 kDa were detected with anti-actin antibodies in immunoblots. The bigger band was ubiquitinated (Field et al. 1993). The distribution of actin in the cytoplasm of the parasite has also been shown by immunofluorescence microscopy (Webb et al. 1996).

One β-tubuline and two different α-tubuline genes have been cloned (Delves et al. 1989, Holloway et al. 1989, 1990). The microtubule organization during schizogony has been visualized by fluorescent microscopy (Read et al. 1993). Short filaments of unknown composition were shown by electron microscopy to link the plasma membrane of the merozoite with the pellicular cisternae (Bannister & Mitchell 1995). A myosin was detected in *P. berghei* and its distribution was studied using confocal laser scanning microscopy (Suhrbier et al. 1993); recently, a myosin was also detected in *P. falciparum* (Pinder et al. 1998, Forero 1998). The apparent molecular weights of the former myosins are very different compared with each other and somehow strange compared with most myosins reported (Titus 1993, Mooseker & Cheney 1995). Using F-actin affinity columns and ATP-agarose chromatography Tardieux et al. (1998) identified a 70 kDa protein (HSC70) in *P. knowlesi*
merozoites, characterized as a *Plasmodium* heat shock protein and a doublet of 32/34 kDa. The HSC70/32/34 complex inhibited the polymerization of rabbit skeletal muscle actin *in vitro*.

In our laboratory we have studied the role of calcium during invasion and we have shown that its depletion in the medium causes an inhibition of invasion partly due to the inability of the merozoite to reorientate and penetrate the erythrocytes (Wasserman et al. 1982, 1990). It has been well established that calcium is an important mediator of assembly and disassembly of cytoskeleton (Bennet & Weeds 1986) and of molecular motors (Barkalow & Hartwig 1995, Wolenski 1995). The study and characterization of actin binding proteins in *Plasmodium* should contribute to the understanding of the force events required for the invasion. The purpose of the present work was the detection and isolation by affinity chromatography of actin binding proteins in the parasite. Some of the purified proteins were identified with antibodies against well-known and conserved F-actin binding proteins and a myosin was immunoprecipitated from metabolic radiolabelled extracts.

**MATERIALS AND METHODS**

**Parasite growth conditions and obtention of protein extracts** - The Colombian strain FCB1 of *P. falciparum* was cultured in O(+)-erythrocytes at 5% hematocrit in medium RPMI 1640 (Sigma Chem. Co. St. Louis, Mo) supplemented with 10% human serum (Trager & Jensen 1976). Cultures were synchronized with sorbitol 5% (Lambros & Vanderberg 1979) and parasitemias of 5-10% schizonts were obtained after 47 h incubation. For 35S-methionine labeling, 100 ml of a 7% parasitemia culture of 18-h-age rings were washed twice with HBS (Hepes 20 mM pH 7.4, NaCl 160 mM) and resuspended in the same volume of methionine free RPMI-1640 medium. A volume of 24 µl of 35S-methionine (9.8 µCi/ml, 1.18 Ci/mmol - Amersham International plc, UK) was added and the culture was incubated for 30 h more. The cells were washed twice with HBS and erythrocytes were lysed with saponine 0.15% in HBS. Erythrocyte membranes were eliminated after centrifugation over a Percoll cushion (Wiser & Lanners 1979) and parasitemias of 5-10% schizonts were obtained after 47 h incubation. For 35S-methionine labeling, 100 ml of a 7% parasitemia culture of 18-h-age rings were washed twice with HBS (Hepes 20 mM pH 7.4, NaCl 160 mM) and resuspended in the same volume of methionine free RPMI-1640 medium. A volume of 24 µl of 35S-methionine (9.8 µCi/ml, 1.18 Ci/mmol - Amersham International plc, UK) was added and the culture was incubated for 30 h more. The cells were washed twice with HBS and erythrocytes were lysed with saponine 0.15% in HBS. Erythrocyte membranes were eliminated after centrifugation over a Percoll cushion (Wiser & Lanners 1992). Free parasites were washed three times with HBS; the weight of the pellet was evaluated and the parasites were resuspended 1:10 w/v in extraction buffer (Hepes 5 mM pH 7.5, EDTA 0.5 mM, EGTA 0.5 mM, Nonidet P40 0.05%, supplemented with the following mixture of protease inhibitors: PMSF 1 mM, Benzamidine 0.01 mM, Phenantroline 1 µg/ml, Aprotinin 10 µg/ml, Leupeptin 10 mg/ml and Peptatin A 10 mg/ml) and frozen at -70°C. Immediately before chromatography one volume of the extraction buffer with protease inhibitors was added to the parasite suspension, and it was homogenized with a teflon Potter-Elvehjem homogenizer with ten strokes at maximum speed. The suspension was centrifuged at 10,000xg during 20 min; the supernatant was adjusted to a concentration of 50 mM Hepes pH 7.5, DTT 2 mM and centrifuged again at 100,000xg during 1 h to remove ribosome, membranes and residual polymeryzed cytoskeleton proteins. The clear supernatant was loaded to the affinity columns for chromatography. The same process was followed with uninfected erythrocytes as a control to verify the parasitic source of the radiolabeled proteins.

**Affinity chromatography - F-actin columns** - Bovine muscle actin (Sigma) was resuspended in a low ionic strength buffer -buffer G- (Hepes 5 mM pH 7.5, CaCl₂ 0.2 mM and ATP 0.2 mM) and was polymerized by adding Hepes to reach 50 mM, KCl to 0.1 M and MgCl₂ to 5 mM (buffer F). The mixture was incubated at 20°C for 1 h and Phalloidin was added to the mixture to reach an equimolar concentration with the actin. Polymerization was checked out by electrophoresis in 5% polyacrilamide gels under denaturing and non-denaturing conditions. The F-actin columns were prepared as described (Miller & Alberts 1989, Miller et al. 1991) with slight modifications. All procedures were performed at 4°C. Two ml of Affigel-10 (BioRad, Richmond, CA) and 2 ml of Sepharose-Cl6B (Pharmacia Biotech, Piscataway, NJ) were carefully mixed in a 1-cm-diam and 10-cm-long glass column. The mixture was washed three times with deionized water and twice with buffer F containing 10 µg/ml Phalloidin. The buffer was drained out and 2 ml of previously polymerized actin were added to a protein concentration of 2.5 mg/ml. The mixture was left overnight with very gentle agitation on a rotatory mixer (Taiyo RT50). After the former step, the column was left in a vertical position for 1 h, the supernatant was carefully drained out and the gel was washed with 3 vol of buffer F with a flow rate of 3 ml/h. The column was stored with buffer F supplemented with 10 µg/ml Phalloidin and 0.02% NaN₃.

The F-actin columns were checked out for their binding specificity by doing a chromatography of a mixture of commercial pure bovine myosin (Sigma) and BSA (Bovine Serum Albumin). The mixture, 2 ml containing 1 mg/ml of each protein in sample buffer A (Hepes 50 mM pH 7.5, DTT 2 mM, EGTA 0.5 mM, EDTA 0.5 mM, KCl 50 mM, Nonidet P40 0.05%, glicerol 10% and the above described mixture of protease inhibitors), was applied at a flow rate of column vol/h. Unbound proteins were eluted with the same buffer. After extensive washing, the bound proteins were eluted
in two steps. The first with 5 ml of buffer A plus ATP 1 mM and MgCl₂ 3 mM and the second with 5 ml of the same solution added with KCl 1M. Fractions were analyzed by SDS-PAGE. The column was used five times with identical results.

**G-actin columns** - The resin and the column were prepared as formerly described. The affigel-10 was preincubated 90 min with buffer G and washed twice with the same buffer. Two ml of bovine G-actin (3 mg/ml) in buffer G were added and mixed as described. Unbound protein was drained out and the mixture was left during 3 h in 2 ml of buffer G containing 50 mM ethanolamine pH 7.8. The column was washed again and used only once during the week following its preparation.

**BSA column** - The control column was prepared as described for the F-actin column, (but this time no Phalloidin was added). Four ml of a 4 mg/ml BSA solution in buffer F were used.

**Affinity chromatography** - Chromatography was carried out simultaneously on the three columns using a multichannel peristaltic pump (WIZISCO). After stabilizing the columns with buffer A, 1 ml of the extract was loaded at a flow rate of 0.5 column-ml/h (the extracts contained 10⁸ DPM/ ml or 4 mg protein/ml). Four 1 ml fractions containing the unbound proteins were collected. The columns were washed with four column-volumes of buffer A, and the bound proteins were eluted in two steps as formerly described. Fractions were analyzed on 8% SDS polyacrilamide gels that were silver stained (Oakley et al. 1980), dried, and exposed to Hyperpaper-35S (Amersham) for autoradiography.

**Electrophoresis, immunoblot and slot blot** - The samples were electrophoresed as described (Laemmli 1970) and transferred to Immobilon P (Millipore, Bedford, MA) at 0.8 mA/cm² (Towbin et al. 1979). Membranes were blocked by overnight incubation with a solution containing Tris-HCl 20 mM pH 7.5, NaCl 150 mM, Tween20 0.1% and fat-free dry milk 5%. As secondary antibodies we used biotinylated anti-rabbit-IgG (BRL Life Technologies Inc, Gaithersburg, MD), biotinylated anti-mouse-IgG (Amersham), biotinylated anti-goat-IgG (Sigma), peroxidase conjugated anti-mouse IgG (Dako, Denmark), and a streptavidine conjugated with alkaline-phosphatase (Amersham). The final conjugate was incubated at the previously established dilution for 1 h and washed as described. For signal development, 5bromo-4chloro-3indoyl-phosphate (BCIP) and nitroblue tetrazolium were used with alkaline phosphatase and the ECL Western Blotting System (Amersham) with peroxidase.

The slot blot was done as follows: samples containing 1 and 2 μg of rabbit muscle myosin or of column protein eluates were filtered on a nitrocellulose membrane using a Bio-Dot SF (BioRad) equipment. The membranes were dried between two Whatman 3MM filters and the signal was developed with HRP-conjugated secondary antibodies using the ECL Western Blotting System (Amersham).

**Immunoprecipitation** - The immunoprecipitation was carried out both with the radiolabelled crude extracts and with the fractions eluted from the affinity chromatography. We used the Firestone and Winguth (1990) method as follows: to avoid inespecific reactions, the antigen-antibody reaction took place in the presence of BSA, in HENN 250 buffer (25 mM Hepes KOH pH 7.5/5 mM EDTA pH 7.5/250 mm NaCl/1% Nonidet P40). We used 20 μl of the monoclonal antibody (anti-pan myosin Amersham RPN-1169) at a 1:10 dilution and 300 μl of the radiolabelled fractions. The positive control used was bovine muscle myosin (Sigma M-6643). To demonstrate the specificity of the immunoprecipitation assay we performed a control using the same samples eluted from the F actin column but without the addition of the primary anti-myosin antibody. The samples were left overnight in low agitation at 4°C, and after adding 20 μl of the secondary antibody (rabbit anti-mouse IgGs) at 1:50 dilution, they were incubated for 1h at
20°C. Afterwards, we added 20 µl of Sepharose 4B Protein G (Sigma P 3296) to each tube and we incubated them with agitation for 5 min. Then the samples were layered over a 1M Sucrose cushion in HENN 250 buffer and were centrifuged for 3 min at 10,000g and 20°C. The unadsorbed proteins were eliminated carefully from the top of the tube and the walls of the tube in contact with the sample were carefully washed with 2 M urea in 500 HENN buffer (500 mM NaCl and other components as in 250 HENN) for 2 min. Finally, the urea and the sucrose were discarded and the pellet was washed twice with 250 HENN buffer and once with the same buffer without detergent. The pellet was resuspended in Laemmli buffer for SDS-PAGE. The gel was electrotransferred to a PVDF (Millipore) membrane and was exposed to Hyperpaper-35S (Amersham) at -70°C.

RESULTS

Affinity chromatography - We tried to construct affinity columns for filamentous actin (F-actin), stabilizing the fibers with chemical crosslinkers (Miller et al. 1991) or with Phalloidin (Miller & Alberts 1989). When we used the chemical crosslinker dimethylsuberimidate we found it was not efficient to construct the columns. At the reported concentration of 10 µg/ml, Phalloidin was not enough to maintain long fibers. We tested higher concentrations, and we found that when they were equivalent to one or two times the molar concentration of actin, the polymerization and the stability of the filament were efficient and the binding of the fibers to the gel improved. Six F-actin columns were used through the present work and the binding of actin to the support varied between 85.2 to 96%. The specificity of the F-actin columns was verified doing a chromatography of a sample prepared by mixing BSA and myosin (from bovine skeletal muscle). Fig. 1 shows that BSA did not bind to the column and eluted with the void volume and the initial washing. The myosin bound strongly to the F-actin. It eluted only with a solution that contained ATPMg2+ and 1M KCl.

The G-actin columns were prepared similarly, but in a buffer that prevented polymerization. Around 54% of the G-actin was incorporated to the gel support. The BSA bound to the gel in the control columns with an efficiency of 75%.

Similar 1 ml samples containing 108 DPM of a 35S-methionine-labeled parasite-protein extract were loaded on F-actin, G-actin and BSA columns. The chromatography was performed as described in Materials and Methods, under identical conditions in the three columns. Radioactivity was quantified and the eluted fractions were analyzed by SDS-PAGE and autoradiography. A 2.9% of the radioactivity bound to the F-actin column. Ten proteins were eluted from the F-actin column with the addition of ATP and Mg2+, and six more when the concentration of KCl was raised to 1M (Fig. 2A). Only 0.9% of the radioactivity bound to the G-actin column. Two proteins were eluted with ATP and Mg2+, and two more with the addition of KCl to 1M (Fig. 2B). No protein bound to the BSA control column (Fig. 2C).

Identification of F-actin binding proteins - The 16 proteins that eluted from the F-actin column with the addition of ATP and Mg2+, and six more when the concentration of KCl was raised to 1M (Fig. 2A). Only 0.9% of the radioactivity bound to the G-actin column. Two proteins were eluted with ATP and Mg2+, and two more with the addition of KCl to 1M (Fig. 2B). No protein bound to the BSA control column (Fig. 2C).

Identification of F-actin binding proteins - The 16 proteins that eluted from the F-actin columns with ATP, Mg2+ and 1M KCl were separated by SDS-PAGE and transferred to PVDF-cellulose membranes. Several different antibodies against known and conserved actin binding proteins were used to assay strips of the membrane containing the electrophoresed proteins. One polyclonal antibody against tropomyosin (chicken gizzard) recognized three proteins. One sized 45 kDa produced a very strong signal, while the other two of 33 and 34 kDa produced mild signals (Fig. 3A). A polyclonal antibody against chicken actin recognized two defined proteins of 43 and 55 kDa (Fig. 3B). Those proteins probably correspond to the reported normal and ubiquitinated forms of the parasite actin (Field el al. 1993). An anti α-actinin (chicken gizzard) polyclonal antibody detected two bands of 100 and 56 kDa (Fig. 3C).

Two proteins were recognized with two different antibodies against human erythrocyte spectrin (Fig. 3D). They seem to be of parasitic origin, since
they were labelled with 35S-methionine (Fig. 1A, lane 7). They show a degradation pattern different from that of the human spectrin (not shown), and they apparently have a molecular weight slightly higher than that of human spectrins.

Using two different antibodies against myosin, a polyclonal prepared against bovine skeletal muscle myosin and a monoclonal against human myosins I and II, we identified two bands of 120 and 72 kDa (Fig. 3E, lanes 4, 5). An antibody against a *P. falciparum* myosin peptide (kindly provided by Dr Walter Gratzer from MRC London) recognized myosin in a slot-blot (Fig. 3F) of the mixture eluted from the F-actin column, but not a bovine muscle myosin. Moreover, a myosin was immunoprecipitated from a crude radiolabeled extract with the anti-pan-myosin antibody, the molecular weight of the protein being 120 kDa (Fig. 2).
Actin-binding Proteins in *P. falciparum* • C Forero, M Wasserman

4), which coincides with the band identified by immuno-blots. Negative results were obtained with antibodies against vinculin, gelsolin and filamin.

**DISCUSSION**

We isolated several proteins of *P. falciparum* that bind specifically to G-actin and to F-actin. The parasite proteins were metabolically radiolabeled and some of them were identified using antibodies raised against the conserved homologous proteins of other organisms. These proteins were not detectable in the erythrocyte extract (data not shown), demonstrating their parasite origin.

Four proteins bound to the G-actin column (Fig. 2 B). Two of them with molecular weights of about 100 and 97 kDa eluted with ATP and Mg\(^{2+}\), and were present in the same fraction of the F-actin column. The other two eluted when the salt concentration was raised to 1M KCl: one, 55 kDa, was also isolated in the F-actin column and has the same
molecular weight of the reported ubiquitinated actin (Field et al. 1993). The other one, a protein of 13 kDa, could be a profilin, although we were not able to confirm its identity. Profilin has been found in most eukaryotic cells playing an important role in the regulation of the polymerization kinetics of actin fibers (Goldschmidt-Clermont et al. 1990, Machesky & Pollard 1993).

Several proteins were isolated from the F-actin columns. Most eluted when besides the 1 mM ATP and 3 mM Mg2+, the salt concentration was 1M (KCl). A similar result was reported when F-actin binding proteins of chicken gizzard were purified (Miller & Alberts 1989). It is interesting to point out a high molecular weight doublet (approx. 260-240 KDa), that migrates in a slightly different way from the human host spectrin, and was recognized by anti-spectrin antibodies. The proteins are undoubtedly from parasite origin since they were metabolically radiolabeled. In several unicellular organisms, non-erythroid spectrins have been described. In

...Plasmodium... leads to the obvious assumption of the existence of actin binding proteins in the parasite. We present here direct evidence of the existence in Plasmodium of a myosin, and also of a spectrin-like protein, an α-actinin, and a tropomyosin. This preliminary finding opens the way to further studies on the proteins and their possible role in the biological process of the invasion. The recent finding that cytochalasin B and E apparently do not cause a significant depolymerization of actin fibers in the parasite (Webb et al. 1996) may imply that different proteins are acting by capping and crosslinking, thus stabilizing the actin filament network. Force events and cytoskeletal rearrangement seem to be crucial during the invasion of Plasmodium, thereby stressing the importance of a better knowledge of the proteins involved in those processes and of their interactions.
REFERENCES


