Infection with *Plasmodium berghei* ookinetes alters protein expression in the brain of *Anopheles albimanus* mosquitoes

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**Abstract**

**Background:** The behaviour of *Anopheles* spp. mosquitoes, vectors for *Plasmodium* parasites, plays a crucial role in the propagation of malaria to humans. Consequently, it is important to understand how the behaviour of these mosquitoes is influenced by the interaction between the brain and immunological status. The nervous system is intimately linked to the immune and endocrine systems. There is evidence that the malaria parasite alters the function of these systems upon infecting the mosquito. Although there is a complex molecular interplay between the *Plasmodium* parasite and *Anopheles* mosquito, little is known about the neuronal alteration triggered by the parasite invasion. The aim of this study was to analyse the modification of the proteomic profile in the *An. albimanus* brain during the early phase of the *Plasmodium berghei* invasion.

**Results:** At 24 hours of the *P. berghei* invasion, the mosquito brain showed an increase in the concentration of proteins involved in the cellular metabolic pathway, such as ATP synthase complex alpha and beta, malate dehydrogenase, alanine transaminase, enolase and vacuolar ATP synthase. There was also a rise in the levels of proteins with neuronal function, such as calreticulin, mitofilin and creatine kinase. Concomitantly, the parasite invasion repressed the expression of synapse-associated proteins, including enolyl CoA hydratase, HSP70 and ribosomal S60 proteins.

**Conclusions:** Identification of upregulated and downregulated protein expression in the mosquito brain 24 hours after *Plasmodium* invaded the insect midgut paves the way to better understanding the regulation of the neuro-endocrine-immune system in an insect model during parasite infection.

**Keywords:** *Plasmodium*, Ookinet, *Anopheles albimanus*, Neuron, Proteomic, Brain

**Background**

Anopheline mosquitoes represent the principal vectors for transmission of protozoan parasites of the genus *Plasmodium* to humans. In 2015, 215 million malaria cases were reported worldwide, causing 438,000 deaths [1]. To control this public health problem, it is necessary to deepen the understanding of the interaction of this insect species with the parasite.

The molecular interplay between *Anopheles* and *Plasmodium* has been extensively studied in regard to the immune response of the mosquito when invaded by the parasite [2–4] and the molecular mechanisms of evasion developed by the latter [5]. Previous reports have shown that *Plasmodium falciparum* and *P. yoelii nigeriensis* alter the behaviour of their hosts, *Anopheles gambiae* and *An. stephensi*, respectively [6, 7]. Specifically, infection of mosquitoes causes a greater insect contact rate with the vertebrate host [8, 9] because of the increase in the number of blood-feedings [9]. At the same time, enzymatic activity in the mosquito salivary glands diminishes [10].
Since the mosquito brain and nervous system dictate feeding behaviour, muscle activity, neuropeptide and hormonal secretion, as well as others functions [11], they are expected to play a crucial role in malaria dissemination. Moreover, there is cross-talk between the immune and nervous systems, which occurs through a complex set of neurotransmitters, cytokines, hormones and neuropeptides [12]. The neural response to a parasite invasion could be generic and stress dependent or pathogen-specific. The pathway of infection-related signal transmission from midgut cells to the brain is still unknown. The effects of midgut cell invasion by *Plasmodium ookinete*, and the consecutive shedding reaction may generate stress signals that are perceived by mosquito brain cells. Signals related to energy deprivation as well as oxidative and immune stress are transmitted throughout the organism. Our group has demonstrated that when midguts of *An. albimanus* are invaded by *P. berghei* ookinetes, they produce and release nitric oxide (NO) and hydrogen peroxide (H$_2$O$_2$) at 24 h post-infection, compounds that are able to activate a systemic immune response [13]. Furthermore, the presence of the parasite in the *An. albimanus* midgut induces changes in the protein profile of midgut cells [14].

In the early stage of the ookinet invasion of mosquito midgut tissues (24 h post-intake of infected blood meal), there is an alteration in the expression of immunity/defence, mitochondrial redox, metabolism, and transcription/regulation of proteins in these tissues [14]. However, the changes occurring in the mosquito neural system during the early phases of *Plasmodium* infection are as yet scarcely explored. Considering that the mosquito brain regulates feeding conduct as well as muscular activity, the analysis of its proteome may help unveil the mechanisms underlying the behavioural changes of mosquitoes that have been observed during a *Plasmodium* infection. A differential head proteome of sporozoite-infected versus uninfected anopheline mosquitoes revealed that various proteins associated with metabolism, synaptic transmission, heat-shock response, signalling, and cytoskeleton function were altered by the presence of the parasite 20 days post-infection [15]. We herein used a proteomic approach to analyse protein expression in the mosquito brain 24 h after feeding on blood infected with *P. berghei* ookinetes. Compared to the brain of control mosquitoes, we observed modifications in the protein expression of different physiological processes at 24 h post-invasion of midgut cells by ookinetes. The current results shed light on the early processes taking place in the mosquito brain during an ookinet invasion of the midgut.

**Methods**

**Mosquitoes and infection with *P. berghei***

White stripe strain *An. albimanus* females [16] were obtained from the insectary of the National Institute of Public Health (INSP) in Cuernavaca, Mexico. Mosquitoes were bred under a 12:12 photoperiod at 28 °C and 70–80 % relative humidity. To determine that the changes in the proteins expression were mainly due to the presence of *Plasmodium* in the midgut, mosquitoes were treated with antibiotics [17]. They had *ad libitum* access to food, which was 8 % sucrose with PSN 1x (5,000 U of penicillin, Streptomycin at 5 mg/ml, and Neomycin at 10 mg/ml) and gentamicin (50 μg/ml) (Thermo Fisher Scientific, Waltham, Massachusetts, USA) absorbed in cotton pads and provided during the 72 h before infection. Cotton pads were changed daily. This antibiotic treatment almost eliminates all bacteria in the midgut of *An. albimanus* [17]. At 4 days post-emergence, mosquitoes were infected with *P. berghei* ANKA expressing the green fluorescent protein (GFP) [18] (kindly donated by Robert E. Sinden, Imperial College, UK).

Ookinetes were produced by culturing gametocyte-infected mouse blood, as described previously [19]. Groups of 300 female mosquitoes were fed for 1 h using artificial membrane feeders with: (i) mouse blood + GFP ookinetes (infected group, with approximately 800 ookinetes per μl), or (ii) mouse blood only (control group). Unfed mosquitoes were removed and the engorged ones incubated at 21 °C to allow for parasite invasion and interaction with the mosquito midgut. Ookinetes recognize, interact with and invade the midgut cells.

**Brain samples**

Brains were extracted from mosquitoes at 24 h post-feeding. Mosquitoes were cold anesthetized for 10 min at 4 °C and maintained on ice. Three hundred brains were obtained from each group of mosquitoes. Tissues were kept in 200 μl PBS 1x with protease inhibitors (2 mM PMSF, 2 mM TLCK, 0.1 mg/ml leupeptin and 1 mM EDTA) (Sigma-Aldrich, St. Louis, Missouri, USA) and stored at -70 °C to await further use. Midguts were analysed under a fluorescence microscope to confirm the presence (infected group) or absence (control group) of ookinetes. In the former case, only tissues from mosquitoes infected with *P. berghei* were used for the analysis of expression. Prior to first dimension IEF, the brains were solubilised in a solution containing 7 M urea, 100 mM DTT, 2 % CHAPS and 0.2 % ampholine (ampholine lysis buffer; Sigma-Aldrich). The protein content was determined using the BCA method (Thermo Fisher Scientific). Triplicates of 125 μg of control and experimental samples were separated on individual IEF strips.

**2D electrophoresis and image analysis**

First dimension IEF was carried out in an Ettan IPGphor 3 unit as described by the manufacturer (Amersham Biosciences, Uppsala, Sweden). For the isoelectric focusing separation, 13 cm, pH 3–10 NL (nonlinear) precast
IPG strips were submitted to 2000 V tension for 17 h at 20 °C, as per the manufacturer’s instructions. Prior to SDS-page second dimension separation (2-D), the IPG strips were placed in the rehydration tray and the proteins in the strip were reduced and alkylated by sequential incubation in the following solutions: 0.04 M Tris-HCl, pH 6.8, 1 % SDS; 30 % glycerol (equilibration buffer-EB) supplemented with 4 mg/ml DTT in EB; and then 40 mg/ml iodoacetamide in EB. After isoelectric focusing, the IPG strips were applied directly to 13 % SDS-polyacrylamide gels for second-dimension electrophoresis at 150 V in an Owl electrophoretic chamber (Thermo Fisher Scientific). The gels were fixed and then stained with PhastGel Blue (Amersham Biosciences), according to the manufacturer's instructions. The stained gels were digitalized using a GS-800 densitometer (Bio-Rad Hercules, CA, USA). Images from 2-D gels of three biologically independent protein extracts from control and experimental group were processed using the PD-Quest 8.0.1 software (Bio-Rad). Protein spots in all replicates were detected automatically by the software, and the detection was improved by the manual addition of omitted spots and the elimination of incorrectly detected spots. Normalization of gel images was performed using the local regression model normalization method supply by PD-Quest software. To appropriately compare the samples, the gel images were adjusted to fit a common distortion model; this was done by matching spots that were common to all the gel images. The gel images from the different protein samples were compared to each other in order to generate a robust data set containing all the spots represented in the samples with 98 % statistical confidence (P < 0.01) in a Student’s t-test. Lastly, the protein spots in the statistical data set displaying ± 2-fold abundance change were selected and then submitted to mass spectrometry (MS) identification.

Protein proteolysis, MS identification and functional annotation
Selected spots from Coomassie blue-stained two-dimensional gels were excised manually and frozen at -70 °C until further use. Samples were prepared for mass spectrum analysis using a slight modification of a previously described procedure [20, 21]. Protein spots were destained, reduced, alkylated and digested with trypsin (Promega, Madison, WI, USA). Before the mass spectra of the peptide mixtures were obtained, these solutions were desalted using a C18 Zip Tip (Millipore, Bedford, MA, USA), according to the manufacturer’s recommendations.

Mass spectra were determined using a Bruker Daltonics Autoflex (Bruker Daltonics, Billerica, MA, USA) operated in the delayed extraction and reflectron mode. Spectra were externally calibrated using a peptide calibration standard (Bruker Daltonics). Peptide mixtures were analysed using a saturated solution of alpha-cyano-4-hydroxycinnamic acid in 50 % acetonitrile-0.1 % trifluoroacetic acid. Lists of the peaks of tryptic peptide masses were generated and searched against the NCBI nr (NCBI, Maryland, USA; https://www.ncbi.nlm.nih.gov/nuccore/about/nonredundantproteins/#reference) [22] and a anopheine protein data bank (https://www.vectorbase.org/organisms) [23] using the Mascot search program (Matrix Science, London, UK; http://www.matrixscience.com) [24], with the following parameters: one missed cleavage allowed, carbamidomethylcysteine as the fixed modification, and oxidation of methionine as the variable modification. We accepted proteins with scores > 50 and P < 0.05.

Differential brain protein expression between control and infected mosquitoes
The average protein levels in the spots were quantified, and those showing relative changes in abundance of > 2-fold between conditions (increase or decrease) at the 99 % confidence level (P < 0.01) were considered significantly different (for details see Methods). A total of the 188 differential spots were observed there were 165 in the control (88 %) and 23 (12 %) in the infected group, evidencing that the Plasmodium midgut tissue invasion does trigger a brain response (Fig. 1).

Identification
Nineteen differentially expressed proteins were identified with an e-value, score, identity and coverage that are compatible with correct identification and assignment (Fig. 2 and Additional file 1: Table S1). These proteins correspond to 19 genes previously registered in the An.
albimanus genome database (https://www.vectorbase.org/taxonomy/anopheles) [23]. The 19 proteins that presented an altered expression level were classified into five categories: ATP synthesis coupled proton transport, cytoskeleton rearrangements and synaptic plasticity, oxidation-reduction process, signal transduction, and miscellaneous (Tables 1 and 2).

**ATP synthesis coupled proton transport**

During the ookinet invasion of the mosquito midgut, the following proteins were upregulated in the mosquito brain: alpha (AALB010020-PA) and beta (AALB005889-PA) proteins (spots 10 and 19, respectively) of the ATP synthase complex, the vacuolar ATP synthase subunit f (AALB009730-PA) and a voltage-dependent anion-selective channel protein (AALB000444-PA) (spots 13 and 7, respectively).

**Cytoskeleton rearrangements and synaptic plasticity**

The synapse-associated protein (AALB008424-PA), which contains a BTF-2 like transcription factor and a Dos2-like domain (BSD), was downregulated in the brains of infected mosquitoes (spot 4). This protein was found specifically in neurons and is an important molecular element of the nervous system [27, 28].

The tubulin \( \beta \) chain protein (AALB000046-PA) (spot 16) is a key element of microtubule assembly in the cell cytoskeleton [29]. Cofilin (AALB010134-PA) (spot 14) is
a protein implicated in regulation of neuronal polarity [30]. Both proteins were upregulated.

Calreticulin (AALB003666-PA) (spot 8), a multifunctional protein mainly involved in directing the proper conformation of proteins, controlling the calcium level, and resisting infection [31], was upregulated during the parasite invasion, potentially as a consequence of mitochondrial stress signalling.

The oxidation-reduction process
Enoyl-CoA hydratase (AALB010381-PA) was downregulated during infection (spot 1). This mitochondrial enzyme catalyses the second step in the beta-oxidation pathway of fatty acid metabolism [32]. Another four enzymes of this process were upregulated during infection: (i) malate dehydrogenase (AALB008565-PA) (spot 3), an enzyme that participates in the citric acid cycle, catalysing the oxidation reaction of the malate to oxaloacetate; (ii) enolase (AALB006338-PA) (spot 6), a protein involved in carbohydrate metabolism and energy production; (iii) an alanine aminotransferase (AALB006338-PA) (ALAT, EC 2.6.1.2) (spot 12) that catalyses the transfer of an amino group from L-alanine to $\alpha$-ketoglutarate and is considered as a marker of tissue damage in various organisms; and (iv) mitofilin (AALB002066-PA) (spot 12), an enzyme that has been reported to counteract sudden oxidative stress.

Signal transduction
Downregulation was found for the HSP70 proteins (spots 2 and 15) (AALB008255-PA), which are molecular chaperones that minimize unfolded protein aggregation through degradation or removal [33].

### Table 1

| Spot | pI/MW/AA          | An. albimanus | Protein family                  | Gene ontology                                                                 | Fold change |
|------|-------------------|--------------|---------------------------------|-------------------------------------------------------------------------------|-------------|
| 13   | 6.1/14.1/127      | AALB009730-PA| Vacuolar ATP synthase subunit f  | GO:0015991: ATP hydrolysis coupled proton transport                            | 383 Up      |
| 19   | 9.5/107.7/551     | AALB005889-PA| F-type H+-transporting ATPase   | GO:0015986: ATP synthesis coupled proton transport                            | 6.6 Up      |
| 10   | 4.7/53.7/503      | AALB010020-PA| ATP synthase beta subunit       | GO:0015986: ATP synthesis coupled proton transporter                           | 2.9 Up      |
| 7    | 6.4/76.2/675      | AALB000444-PA| Voltage-dependent anion-selective channel | GO:0055085: Transmembrane transport                                           | 2.4 Up      |

### Table 2

| Spot | pI/MW/AA          | An. albimanus | Protein family | Gene ontology                              | Spot intensity |
|------|-------------------|--------------|---------------|--------------------------------------------|----------------|
| 8    | 4.1/46.5/409      | AALB003666-PA| Calreticulin  | GO:0005509: Calcium ion binding            | 1.1            |
| 12   | 8.6/198.3/1,771   | AALB002066-PA| Mitofilin     | GO:0003676: Nucleic acid binding           | 2.5            |
The guanine nucleotide binding protein beta 3 (AALB004444-PA) was upregulated during infection (spot 17). This protein has been shown to be part of the protein complexes allowing for the integration of signals between receptor and effector proteins.

**Miscellaneous**

The 60S acidic P0 ribosomal protein that binds to 25S rRNA (AALB008188-PA) (spot 18) was downregulated during infection, potentially reflecting a higher turnover of the proteins involved in the translation processes.

Two of the proteins that were upregulated during early infection were the enzyme arginine kinase (EC:2.7.3.3) (AALB004623-PA) (spot 9), which catalyses the transfer of phosphate from ATP to arginine, and the enzyme peptidyl-prolyl cis-trans isomerase (spot 5), which is related to high protein folding demand (AALB007271-PA).

Three proteins identified in the *An. albimanus* database represent “hybrid proteins” and potentially correspond to more than one protein (Additional file 2: Figure S1a−c). The annotation of the AALB008255-PA transcript reports a potential protein that presents domains of heat-shock protein 70 in its COOH terminal and arrestin homology in its amino terminal. Since the deduced protein showed a molecular weight of 149.08 kDa and had 1,374 amino acid residues, and the protein herein identified showed a molecular weight of 71.67 kDa, had 656 amino acid residues and was identical to HSP70 of *An. gambiae*, we propose the reannotation of this transcript (Additional file 2: Figure S1a). The AALB000444-PA transcript translation presents porin domains in the amino (NH) terminal as well as proteasome component domains (PCI) in the carboxyl (COOH) terminal. The newly annotated protein would have a molecular weight of 76.21 kDa and had 282 amino acid residues. The voltage-dependent anion-selective channel protein presently identified showed a molecular weight of 30.85 kDa and had 282 amino acid residues, identical to the *An. darlingi* protein (Additional file 2: Figure S1b). AALB010381-PA, with a molecular weight of 66.9 kDa and 620 amino acid residues, had domains of the enoyl-CoA hydratase/isomerase, the HSP10−like chaperonin and alcohol dehydrogenase. The Enoyl CoA hydratase herein identified had a molecular weight of 32.28 kDa and 295 amino acid residues. The sequence is identical to that reported for *An. darlingi* (Additional file 2: Figure S1c).

**String analysis**

Interaction analysis showed that Enolase (Eno) and F-type H+ transporting ATPase α (blw) were molecules with the most interactions (eleven), while ATP synthase β (FBpp0305828) and HSP70 (Hsc70-4) had nine and seven interactions, respectively (Fig. 3).

**Discussion**

We herein employed a proteomic approach to analysing the alterations in the protein profile in brains from infected and non-infected mosquitoes. We identified 19 differentially expressed proteins, of which 14 were upregulated and 5 were downregulated. These proteins were mainly associated with the ATP synthesis coupled proton transport, cellular metabolism, cytoskeleton rearrangements, synaptic plasticity and signal transduction. For an overall understanding, the intracellular distribution was defined for the aforementioned 19 proteins (Fig. 4).

In insects, response to stressors are regulated and coordinated by a few hormonal compounds as neuropeptides, ecdysteroids and juvenile hormones. The metabolic response of insects to stress is linked to secretion of biogenic amines in the brain. Within a few minutes after a stress stimulus, reserve energy substrates are mobilized from the fat body. A previous study of *An. gambiae* infected by *P. berghei* showed an alteration in the mosquito head proteome during the process of infection. Specifically, proteins related to energy metabolism presented differential expression in the brain of sporozoite-infected versus uninfected mosquitoes [15]. Other proteins also showed a differential concentration between infected and uninfected mosquito head extracts, including the synapse-associated protein, the 14-3-3 protein, calmodulin, the stress response protein HSP20 and proteins (e.g. tropomyosin) involved in the cell structure [15]. The proteomic approach presently employed provides a direct portrait of the functional status of the mosquito brain during the early phase of infection by *Plasmodium*. A large number of proteins found to be related to the stress response. The signalling in brain cells (an uninvaded tissue) involved in this response may result from the liberation of intracellular proteins in the haemolymph during the crossing of ookinetyes to the basal membrane.

Upregulation of heat-shock proteins (HSPs) during cellular stress, leads to the presence of inducible HSPs such as HSP70 and HSP90 in the haemolymph. These proteins serve as danger signals [34]. Indeed, HSP 70 has been defined as a powerful stress elicitor in many insects. Since these secondary messengers transit through the haemolymph towards all insect tissues, it is expected that brain cells would respond to this danger signal. HSP70 is a family of proteins very strongly upregulated by heat stress and toxic chemicals, particularly heavy metals exposure. However, in anophelines, these proteins also have been associated with dehydration stress [35] and ageing [36]. Surprisingly, during the *Plasmodium* invasion, HSP70 was herein downregulated in the brain of *An. albimanus*.

Several proteins affecting nerve cell function, such as arginine kinase, which were upregulated during
*Plasmodium* infection. This protein is involved in energy production and might modulate immune processes via changes in the concentration of nitric oxide (NO) and the activity of iNOS, as reported in the scallop *Chlamys farreri* [37]. NO is a well-known immune stress mediator, involved in systemic immune response signalling [13]. Furthermore, NO is liberated during immunological reactions and it is also a neuromodulator coordinating many neuronal activities in insects [38]. Interestingly, phosphoarginine, the end product of the arginine kinase activity, alters the Na$^+$ - Ca$_2^+$ exchanger balance [39]. This signalling has been linked to stress signal transduction, affecting the neural function of the squid Na$^+$ - Ca$_2^+$ exchange [40]. Other proteomic results have shown that arginine kinase is differentially expressed in the brain of infected versus uninfected of *Gammarus insensibilis* and *G. pulex* [41].

Besides being a structural protein, mitofilin is also involved in protein transport into the mitochondrion and promotes protein import via the mitochondrial intermembrane space assembly pathway [42]. Since the mitochondria may be modified in the neurons of infected mosquitoes, mitofilin may be critical for maintaining the mitochondrial morphology.

The presence of tubulin $\beta$ chain and cofilin might reflect the reorganization of neurons during infection. Cofilin, is related to the growth of neurons and depolymerization of actin filaments, specifically in neurons and dendritic

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**Fig. 3** Differentially expressed proteins in the infected versus uninfected mosquito brain. Their interaction networks are depicted by STRING and visualized by Medusa. Each node represents the upregulated and downregulated proteins. The edges represent putative protein interactions recorded or predicted by STRING. The graph was obtained using the spectral clustering option, with five clusters. Node rectangles are coloured and the edges predicted functional links. Neighborhood (green lines), Gene Fusion (red lines), Co-occurrence (blue lines), Co-expression (grey lines), Experiments (purple lines), Databases (light blue lines), Textmining (light green lines) and Homology (light grey lines). Abbreviations: CG6543, enolyl-CoA hydratase; CG32683, HSP70; Men-b, malate dehydrogenase; Sap47, synapse-associated protein; CG14715, peptidyl-prolyl cis-trans isomerase; Eno, enolase; Porin, VDAC porin; Crc, calreticulin; Argk, arginine kinase; FBpp0305828, ATP synthase $\beta$; CG1640, alanine transaminase; CG6455, mitofilin; Vha14-1, V-type proton ATPase $\alpha$; Tsr, cofilin; Hsc70-4, HSP70; CG3157, tubulin; GTPase, Gbeta13F protein G $\beta$-1; RplP0, 60s Ribosomal LPO; Blw, F-type H+ transporting ATPase $\alpha$. 
spines [43]. This protein points to an early neuronal response to Plasmodium infection since it has been linked to synapse remodelling [44]. Therefore, cofilin could potentially be a protein responsible or associate for the Plasmodium-induced alteration in mosquito behaviour observed in the later stages of the infection.

In Anopheles gambiae, the vacuolar ATP synthase subunit f and a voltage-dependent anion-selective channel protein interact with Plasmodium enolase mimetic peptide SM-1 [45], a porin whose function in brain cells is still unclear. In our model, Plasmodium enolase could has induced upregulation of these proteins. Also, could arise from its interaction with prohibitin, a protein involved in the stress response [46].

The identification of the alpha and beta subunits of the mitochondrial ATP synthase complex might reflect a surge in energy requirements, potentially resulting from the progression of the Plasmodium infection.

The protein identified that has a BSD domain related to synapse-associated proteins was previously found in the head of An. gambiae infected with P. berghei sporozoites [15]. During synaptogenesis of the mouse brain, orthologous synapsis-associated protein 102 (SAP102) is implicated in the transport of ionotropic glutamate receptors to the cell membrane [27]. In D. melanogaster, this protein is required for proper behavioural and synaptic plasticity [28]. In anopheline mosquitoes, it may be associated with the same processes described in those organisms.

Calreticulin upregulation could potentially be a consequence of mitochondrial stress signalling during infection. Nevertheless, this protein is also involved in the olfactory system [47], anaesthetic sensitivity [48], and phagocytosis of apoptotic cells [49]. In the innate immune system, calreticulin regulates phagocytosis [50] and encapsulation [51]. In An. albimanus, the interaction of this protein with parasite surface molecules has been reported [52].

Enoyl-CoA hydratase also known as crotonase, is essential for metabolizing fatty acids to produce both acetyl CoA and energy. The enoyl-CoA hydratase
presently found suggests that the mosquito brain reduces the use of fatty acids as substrates for oxidative energy metabolism during the Plasmodium infection.

The malate dehydrogenase reflects a higher energetic demand. Interestingly, this enzyme has been studied as a potential target for species-specific insecticidal compounds [53].

The presence of two spots attributed to the enolase protein, showing different isoelectric points (8.5 and 6.5, respectively) and masses (47.7 0 and 50 kDa), implies that this protein exists in two distinct isoforms.

Alanine aminotransferase (ALAT, EC 2.6.1.2), which participates in maintaining the alanine-proline cycle between flight muscles and fat body during the flight of Aedes aegypti [54], is upregulated during injury and toxic substance exposure [55]. Many insects yield lactate and alanine as anaerobic end products, but other species have been known to produce a wide array of other products during hypoxia, including sorbitol, succinate, glycerol, α-glycerol-3-phosphate, pyruvic acid and fatty acids [56]. In An. stephensi, infection with P. yoelii diminishes the expression of transcripts of two different guanine nucleotide-binding protein genes at 4 hours post-feeding [57]. Hence, it is not surprising that this enzyme is present in the brain cells of An. albimanus infected with P. berghei. The absence of this protein decreases cellular proliferation and increases apoptosis, due to abnormal mitochondrial function [58].

In the army worm Spodoptera exigua, it has been shown that small G proteins regulate part of hemocyte phagocytosis, nodulation and encapsulation through cytoskeletal alteration [59]. In the case of Anopheles, guanine nucleotide-binding protein beta 3 (belonging to the small G-protein family) might be related to immune dampening processes.

The diminution of the ribosomal 60S acidic protein would hinder protein synthesis in the mosquito brain. This decrease could limit the amount of energy consumption of the mosquito in order to withstand the energy impounding imposed by Plasmodium infection.

Peptidyl-prolyl cis-trans isomerase has been related to high protein folding demand and is upregulated in the midgut of An. gambiae (Giles) during the invasion of the o’nyong-nyong virus. Therefore, this protein might constitute a general infection-related responsiveness marker [60] in anophelines during infection.

The STRING program was used to analyse the interactions between the set of 19 brain proteins whose levels were altered during the early stage of a Plasmodium infection. Given the scant data available in the anopheline database, we used the orthologous protein from Drosophila melanogaster, having defined UniProtKB accession numbers. When the Drosophila orthologues of the nineteen regulated proteins were mapped, there was a significant interaction between them. The enolase was determined to be the alpha protein of this interactome. This indicates that the metabolism of neuronal cells is the first cellular process altered by Plasmodium infection (Fig. 3). It is noteworthy that the Drosophila ortholog proteins identified show up to 60 % identity with the proteins identified in An. albimanus (Additional file 3: Table S2). Therefore, it is reasonable to consider that the functionality and interaction between them may be conserved. On the other hand, it is important to perform studies to correlating the proteins expression with behavioural changes in mosquitoes. Several aspects must be considered in further experiments: (i) perform experiments in several stages of the parasite life cycle into mosquitoes; (ii) use other Plasmodium-Anopheles models; and (iii) analyse the effect of the microbiota in the parasite interaction and the changes in the brain proteome. Finally, our results provide support for future studies using Plasmodium vivax, a natural parasite for Anopheles albimanus.

Conclusions
The present results indicate that ookinetes promoted the modification of protein expression in the mosquito brain at 24 hours post-infection. The pattern of differential protein expression in infected versus uninfected mosquitoes indicates that the invasion of midgut tissue by Plasmodium triggered a brain response. Whether this change is due to a stress or immune response remains unclear. Nevertheless, we hereby demonstrate that as soon as ookinetes penetrated the intestine, the protein composition of the mosquito brain was altered, indicating potential functional changes. These findings provide novel information about the mechanisms underlying the behaviour and immune response of Anopheles during a Plasmodium invasion, but there must be continuing study to explore the functional analysis and specific involvement of each of these proteins.
Abbreviations
fluorescent protein; H2O2: Hydrogen peroxide; HSP: Heat-shock protein; BSD: Dos2-like domain; COOH: Carboxyl terminal; Eno: Enolase; GFP: Green

Acknowledgments
This study was supported by the Consejo Nacional de Ciencia y Tecnología de México (CONACYT) (Grant 243171). Alejandro Alvarado-Delgado, a doctoral student in PROGRAMA DE DOCTORADO EN CIENCIAS BIOMÉDICAS-UNAM, appreciates the support given by CONACYT (Beca-98230). We thank R. E. Sinden for kindly supplying the P. berghei ANKA strain clone 2.34. The authors would like to thank Mario Henry Rodriguez Lopez, PhD, for his critical review of the manuscript and Guillermo Romero Tecua for his help in the preparation of Fig. 4. We are also grateful to Laura Salgado Albarran and Patricia Munguia Vega for technical assistance with mosquito and parasite/ookinete cultures, respectively.

Funding
The research described herein is supported by Conacyt, Grant 243171 to HLM and Beca-98230 to AAD.

Availability of data and materials
The datasets supporting the conclusions of this article are included within the article and its additional files.

Authors' contributions
AAD: Experimental design, mosquito infection, mosquito tissue extraction, manuscript writing. GPO: String design and analysis, mosquito tissue extraction, manuscript writing. ATTL: Gel electrophoresis, manuscript writing. SE: Mass spectrometry analysis. RC: Manuscript writing. AGMB: Mass spectrometry experiments and analysis. KMF: Mosquito tissue extraction and mosquito breeding. HLM: Experimental design, manuscript writing. All authors read and approved the final version of the manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
Not applicable.

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Received: 19 July 2016 Accepted: 2 October 2016 Published online: 11 October 2016

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Additional file 3: Table S2. Identification of differentially displayed proteins in Anopheles albimanus and orthologs in Anopheles darlingi and Drosophila melanogaster. (XLSX 18 kb)
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