Ycf93 (Orf105), a Small Apicoplast-Encoded Membrane Protein in the Relict Plastid of the Malaria Parasite Plasmodium falciparum That Is Conserved in Apicomplexa

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Abstract

Malaria parasites retain a relict plastid (apicoplast) from a photosynthetic ancestor shared with dinoflagellate algae. The apicoplast is a useful drug target; blocking housekeeping pathways such as genome replication and translation in the organelle kills parasites and protects against malaria. The apicoplast of Plasmodium falciparum encodes 30 proteins and a suite of rRNAs and tRNAs that facilitate their expression. orf105 is a hypothetical apicoplast gene that would encode a small protein (PfOrf105) with a predicted C-terminal transmembrane domain. We produced antisera to a predicted peptide within PfOrf105. Western blot analysis confirmed expression of orf105 and immunofluorescence localised the gene product to the apicoplast. PfOrf105 encodes a membrane protein that has an apparent mass of 17.5 kDa and undergoes substantial turnover during the 48-hour asexual life cycle of the parasite in blood stages. The effect of actinonin, an antimalarial with a putative impact on post-translational modification of apicoplast proteins like PfOrf105, was examined. Unlike other drugs perturbing apicoplast housekeeping that induce delayed death, actinonin kills parasites immediately and has an identical drug exposure phenotype to the isopentenyl diphosphate synthesis blocker fosmidomycin. Open reading frames of similar size to PfOrf105, which also have predicted C-terminal transmembrane domains, occur in syntenic positions in all sequenced apicoplast genomes from Phylum Apicomplexa. We therefore propose to name these genes ycf93 (hypothetical chloroplast reading frame 93) according to plastid gene nomenclature convention for conserved proteins of unknown function.

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Introduction

The widely used malaria prophylactic, doxycycline, was originally developed as an antibacterial that blocks protein translation by prokaryotic ribosomes [1]. Efficacy against malaria is believed to result from inhibition of translation in the reticuloplasts (apicoplast) of malaria parasites [2]. Other inhibitors of prokaryotic translation, plus inhibitors of apicoplast DNA replication (fluoroquinolones like ciprofloxacin) are also parasiticidal [2,3]. Indeed, any perturbation of apicoplast housekeeping appears to kill malaria parasites [2,3]. This highlights the importance of the maintenance of the apicoplast for parasite survival.

Apicoplasts arose by secondary endosymbiosis of a red alga within the common ancestor of Apicomplexa and dinoflagellates [4]. Most dinoflagellates remain photosynthetic, but Apicomplexa adopted a parasitic lifestyle and dispensed with photosynthesis [5,6]. The endosymbiotic organelle, however, persists and is indispensable [7]. The apicoplast of the human malaria parasite Plasmodium falciparum has a small (35 kb), circular genome similar to that of most plastids; the major difference being that the apicoplast genome encodes no genes for photosynthesis [8]. Thirty putative protein-encoding genes have been identified on the P. falciparum apicoplast genome, and seven of these are hypothetical proteins (open reading frames >50 amino acids) [8]. With the exception of SufB/ORF470, which has a role in the formation of iron-sulphur clusters, and ClpC, a protein chaperone subunit with a possible role in protein import [9], all apicoplast genes with known roles are devoted to expression of the genome (housekeeping) [8]. Further information of the roles of these genes, particularly the hypothetical ones, could thus be valuable in targeting apicoplasts in the battle against malaria.

To explore the roles of hypothetical apicoplast proteins we attempted to generate antisera to predicted peptides from a selection of the hypothetical apicoplast genes (ORFs) that possibly encode proteins. Here we describe the characterization of one previously hypothetical apicoplast protein – Orf105/PfYcf93 - to which we generated a successful antiserum allowing us to localize the protein to the apicoplast membranes, document its turnover during the cell cycle, and explore effects of a post-translation inhibitor on the apicoplast and PfYcf93. We also searched for homologues of PfYcf93 in the apicoplast genomes of other apicomplexan parasites and related photosynthetic symbionts of
corals such as the chromerids *Chromera velia* and *Vitrella brassicaformis*.

**Results and Discussion**

**PfYcf93** is an apicoplast-encoded protein

orf105 is an open reading frame (366 nucleotides) located on the IR-B section of the 35 kb circular genome of the apicoplast of *P. falciparum* (GenBank: X95276.2) that encodes a hypothetical protein of 121 amino acids [8]. The name orf105 stems from a previous annotation for a shorter version of the protein that invokes a downstream Met based on the presence of a transcribed and processed *trnS* gene within the N-terminus of the 121 amino acid open reading frame [10]. In keeping with plastid gene nomenclature convention for gene products conserved in multiple species [11] we designated the gene *ycf93* (hypothetical chloroplast open reading frame 88) because orthologues occur in numerous Apicomplexa (see below).

Goat antisera raised against a 14-residue peptide (NILLKSKNSNNYIY) were used for immunofluorescence on *in vitro* cultured blood stage *P. falciparum* parasites (Fig. 1A). Antigen (red) was localized to a small punctum (Fig. 1A). Co staining (green) with antisera to the apicoplast marker acyl carrier protein (ACP) [12] showed the antigen recognised by our goat antiserum co localized with the apicoplast adjacent to the nucleus (Fig. 1A). We conclude that the peptide antiserum recognises an antigen located in the apicoplast.

**PfYcf93** is an apicoplast membrane protein and may form dimers

Western blotting using the peptide antiserum identified a single band of 17.5 kDa (Fig. 1B), which is in reasonable agreement with the predicted mass for *PfOrf105/PfYcf93* of 15 kDa. Controls with pre-immune sera did not decorate any bands (Fig. 1B). Specificity of the antiserum for *PfYcf93* was verified by pre-incubation of the antiserum with 10 mg/ml peptide, which abrogated binding to the 17.5 kDa band (Fig. 1C). We conclude that the antiserum is specific for the gene product of the apicoplast gene *Pforf105*, confirming it as a protein-encoding gene, which we rename *Pfycf93*.

*PfYcf93* protein partitions into the detergent phase during Triton X-114 phase partitioning [13], indicating that it is a membrane protein (Fig. 1D). This membrane localization confirms the bioinformatic predictions of a single C-terminal transmembrane domain. In detergent partitioned non-denaturing Western blots, a second major band of apparent molecular mass 37 kDa was observed (Fig. 1D), indicating that *PfYcf93* may exist as a dimer. Based on the immunofluorescence, Western blots, and detergent partitioning we conclude that *Pfycf93* encodes a small apicoplast membrane protein. Considering the apicoplast appears to lack any systems such as YidC/Oxa/Alb3 chaperone family proteins involved in insertion, export or assembly of membrane proteins [14,15], it seems most likely that *PfYcf93* is located in the innermost of the four bounding membranes and self inserts.

![Figure 1. Characterization of Pfycf93 gene product in Plasmodium falciparum. A) Localization in the apicoplast of an antigen recognised by PfYcf93 peptide antiserum. PfYcf93 (red) localises to a small punctum that overlaps with the apicoplast marker acyl carrier protein (green) and sits beside the nucleus (blue) in a small, ring-stage parasite within a red blood cell. Bar = 1 μm. B) Western blot of SDS-PAGE separated proteins labelled with either pre-immune or immune sera demonstrates specific labelling of a 17.5 kDa protein. C) Labelling of the 17.5 kDa protein with the peptide antiserum is abrogated by pre-incubation of the membrane with 10 mg/ml of peptide. D) PfYcf93 protein partitions into the pellet during Triton X-114 partitioning, and a second band of 35 kDa is apparent. E) Western blotting of the 17.5 kDa band across the 48 hour asexual red blood cell cycle of synchronized parasites demonstrates that PfYcf93 is not detectable at 0, 4, or 12 hours but begins to be detectable at 16 hours and then increases in abundance from 20 to 40 hours and then decreases again at 44 and 48 hours. The change in abundance of the 17.5 kDa band is similar to published fluctuations of merozoite surface protein 1 (MSP-1), but contrasts to MSP-8 and apical merozoite antigen 1 (AMA-1), which have very different turnover patterns. Giemsa stained images of parasite development at different time points are shown. Loading was normalised by protein amount and equal loading confirmed by Ponceau staining (data not shown).](https://doi.org/10.1371/journal.pone.0091178.g001)
**PfYcf93 undergoes turnover**

During the 48-hour life cycle of malaria parasites the apicoplast undergoes dramatic growth, branching, and division resulting in the distribution of an apicoplast into each daughter merozoite [16]. We monitored the steady state levels of *PfYcf93* protein across the life cycle by Western blot on a synchronous culture of *P. falciparum* (Fig. 1E & 2A). In the initial part of the life cycle just following invasion (0 hours), the so-called ring stage parasites did not have any detectable *PfYcf93* (Fig. 1E). After 4 hours (early rings) and 12 hours (early trophozoites) *PfYcf93* was still not detectable (Fig. 1E). At 16 hours a faint *PfYcf93* band had developed, and by 20 hours (late trophozoites) there was a robust *PfYcf93* band, which persisted right through the trophozoite phase and into schizogony but then decreased at 44 hours and still further at 48 hours (Fig. 1E). By comparison, probing of the same samples with antibodies to the merozoite surface protein 1 (MSP1) confirms previous reports that MSP1 only begins to be expressed at 32 hours and persists through to early rings (0 hours and 4 hours). Conversely, merozoite surface protein 8 (MSP8) shows a restricted window of expression between 4 and 24 hours, as previously described [17], and the apical merozoite antigen (AMA1) has maximal expression in the shizont/merozoite phase (36 – 0 hours) as described [18].

The drastic disappearance of *PfYcf93* protein toward the latter part of schizogony (Fig. 1E & 2A) suggests that *PfYcf93* undergoes degradation, probably during the late stage of formation of merozoites. This is the first clear example of protein turnover within apicoplasts. Plastids of plants undergo substantial protein turnover due to activity of a range of proteases, most of which are inherited from the cyanobacterial endosymbiotic ancestor of plastids [19]. Three apicoplast proteases (signal processing peptidase [SPP] [20], falcilysin/PreP [21], and caseinolytic protease (Clp) [9,22,23]) have been identified in *Plasmodium*. SPP and PreP are believed to have roles in processing of leader peptides that mediate import of nucleus-encoded proteins into the apicoplast [20,21], so Clp is the prime candidate for turnover of *PfYcf93*.

Is *PfYcf93* subject to post-translational modification?

Like bacteria, plastids use a formyl methionine as the initiator Met for proteins encoded by the plastid genome and synthesised within the plastid [24]. The formyl group is added to the initiator methionine by formyl methionine transferase (FMT) and is later removed from the methionine by peptide deformylase (PDF) after translation [24]. In select instances (depending on the identity of the second amino acid) the deformylated methionine itself is then removed by methionine aminopeptidase (MAP) [24], which influences protein stability [25]. Genes for likely apicoplast FMT, PDF and MAP have been identified in *Plasmodium falciparum* [14,26] and *PfPDF* is indeed targeted to the apicoplast [27] suggesting homologous N-terminal processing occurs in the apicoplast. Several drugs inhibit PDF [28–32] and one, actinonin, has been noted to have antimalarial activity [33]. Recombinant *PfPDF* has been expressed and its structure determined [34].

![Figure 2](https://example.com/figure2.png)

**Figure 2. The effects of the prokaryotic post-translational modification inhibiting drug actinonin on *PfYcf93* turnover and apicoplast development and cytokinesis.** A) Western blot showing the effect of actinonin and the isoprenoid precursor synthesis inhibitor fosmidomycin on steady state levels of *PfYcf93* over one life cycle. Treatment with these drugs abrogates the normal turnover of *PfYcf93* observed during schizont (40 hour) to ring (56 hour) transition. Loading controls at each life cycle stage using anti-BiP are shown in the lower panels. B) Effect of the putative post-translational inhibitor actinonin on biogenesis of the apicoplast, mitochondria and nuclei. Live cell imaging of untreated parasite with red fluorescent apicoplast, green fluorescent mitochondrion and blue fluorescent nuclei shows the normal branching/division of the apicoplast, elaboration of the mitochondrion and multiplication of the nucleus at the schizont stage. Actinonin-treated parasite at the same stage shows a branched mitochondrion and multiple nuclei but an undeveloped, stunted apicoplast (arrow). C) Effect of fosmidomycin and actinonin on the apicoplast and parasitophorous vacuole (P.V.) at the schizont stage. Immunofluorescence assay of untreated parasite with red fluorescent apicoplast (anti-ACP), green P.V. and blue fluorescent nuclei shows the segregation of the apicoplast, nuclei into distinct merozoites during late schizont stage. Actinonin or fosmidomycin treated parasites at the same stage show multiple nuclei but an undeveloped apicoplast and severe disruption of internal membrane structure in the parasite.

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Activity of recombinant \( \text{PPDF} \) is inhibited by actinonin in in vitro assays [34]. Four apicoplast-encoded proteins from \( \text{P. falciparum} \) (Rps3, Rpl16, Orf91 and Orf105/Ycf93) contain the correct amino acid at residue two [8] to potentially undergo removal of the N-terminal methionine after deformylation by \( \text{PPDF} \). We therefore decided to explore the effect of actinonin on parasite growth, the N-terminus of \( \text{PPDF} \), and the turnover of \( \text{Ycf93} \) during the asexual red blood cell cycle.

In vitro culture drug trials of \( \text{P. falciparum} \) 3D7 confirmed reports [33] that the PDF inhibitor actinonin is parasitocidal with an IC\(_{50}\) of 4.9 (±1.05) \( \mu \)M for a 48 hour growth assay. Drugs targeting apicoplast housekeeping activities (DNA replication, RNA transcription, and protein translation) typically result in the curious phenomenon known as delayed death in apicomplexan parasites [3,7]. Delayed death alludes to the fact that the drugs have maximal effect during the cell cycle in which they are applied, but exhibit maximal lethality during the subsequent cell cycle, even if the drugs have been removed after exposure in the initial cycle [3,7,35–37]. How this phenomenon works remains uncertain, but the leading hypothesis is that a stockpile of some factor essential to the cell cycle is required to sustain the cell cycle. We exposed parasites with red fluorescent labeling antibodies to \( \text{P. falciparum} \) Ycf93 during the asexual red blood cell cycle. We examined the impact of fosmidomycin on \( \text{PPDF} \) turnover and observed the same effect as actinonin - abrogation of turnover (Fig. 2B). Similarly, fosmidomycin also blocked apicoplast growth, perturbed schizogony and the appropriate segmentation of the parasitophorous vacuole (Fig. 2C). The remarkable similarity of the drug response phenotypes for actinonin and fosmidomycin perhaps suggests that actinonin perturbs synthesis of isopentenyl diphasphate, but this remains to be established.

Whether or not actinonin impacts directly on \( \text{Ycf93} \) turnover is difficult to conclude since impairment of apicoplast development by either actinonin or fosmidomycin might indirectly prevent stage specific turnover simply by blocking stage progression. We attempted to determine the actual N-terminus of \( \text{Ycf93} \) with and without actinonin by immunoprecipitating the protein and N-terminal sequencing by Edman degradation to explore any N-terminal processing by PDF/MAP, but found the N-terminus to be blocked (data not shown). N-terminal blockage for Edman degradation occurs frequently when attempting to sequence proteins [41] and cannot be inferred to result from PDF modification or lack thereof.

### Ycf93 protein family in Apicomplexa

Apicoplast genomes are available for 14 apicomplexan parasites, namely nine species of the Haemosporida genus \( \text{Plasmodium} \) [8,42,43], one bird Haemosporida (\( \text{Leucocytozoon caulleri} \)) [44], two Coccidial (\( \text{Eimeria tenella} \) [45] and \( \text{Toxoplasma gondii} \) [46]), and two Piroplasms (\( \text{Babesia bovis} \) [47,48] and \( \text{Theileria parva} \) [49]). We examined the IR-B region (or equivalent) of these genomes plus that of the related chromerid algae \( \text{Vitrella brassicaeformis} \) [50] and \( \text{Chromera velia} \) [51] for possible orthologues of \( \text{ycf93} \) orthologues of \( \text{Plasmodium} \) and \( \text{Toxoplasma gondii} \) (Figs. 3 and 4). The Plasmodium spp. and \( \text{L. caulleri} \) \( \text{Ycf93} \) proteins all contain a predicted C-terminal transmembrane domain (Fig. 4A).

Amino acid alignment shows a 16 amino acid N-terminal extension in the \( \text{P. falciparum} \) \( \text{Ycf93} \) and a 17 amino acid extension in the \( \text{Ycf93} \) orthologue of another primate malaria parasite, \( \text{P. cynomolgi} \) (Fig. 4A). In both cases this extension is predicted to create a second, N-terminal transmembrane domain in addition to the single, predicted C-terminal transmembrane domain. We note that internal methionines in \( \text{PPDF} \) and \( \text{Ycf93} \) occur at the same site as the initiator methionines in the \( \text{Ycf93} \) proteins from \( \text{L. caulleri} \) and the other eight malaria parasite species (Fig. 4A), which includes human, bird and rodent infecting species. This internal methionine was initially invoked for \( \text{PPDF} \) due to the presence of a transcribed and processed \( \text{trnS} \) gene within the N-terminus of the longer version of \( \text{PPDF} \) [10]. A \( \text{trnS} \) gene also occurs in the currently annotated version of \( \text{PPDF} \) [43] so we consider it most likely that both \( \text{PPDF} \) and \( \text{Ycf93} \) use this internal methionine for initiation. If \( \text{Ycf93} \) were in fact initiated from the internal methionine, it would not be a substrate for MAP according to the prediction rules [8]. The apparent mass of \( \text{Ycf93} \) (slightly greater than predicted) might suggest that the longer version rather than the shorter version is translated, but the addition of a second transmembrane domain in just these two species seems unlikely, and (as mentioned above) confirmation of the N-terminus and of any processing remains to be made.

The apicoplast genomes of the coccidian apicomplexan parasites \( \text{Toxoplasma gondii} \) and \( \text{Eimeria tenella} \) have almost identical architecture and gene content to \( \text{Plasmodium} \) spp. [46] and likely orthologues of \( \text{ycf93} \) occur in these parasites (Fig. 3), though the sequence identity with \( \text{Ycf93} \) proteins is very low (not shown).
Figure 3. Gene maps (not to scale) for a portion of the IR-B section of the apicoplast genomes of sequenced apicomplexan parasites and the closely related chromerid alga Vitrella brassicaformis. Genomic accession numbers and species names are provided above each map. A small open reading frame (orf105/ycf93) occurs in syntenic positions in each genome. All the photosynthesis related genes (petD, petA, psbC and chlB) from Vitrella brassicaformis have disappeared from the apicoplast genomes and reorientation of the rpo/rps operon has occurred in select species.

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Figure 4. A) Alignment of PfYcf93 homologues from the 10 sequenced Plasmodium spp. and the related bird parasite Leucocytozoon caulleri reveals obvious sequence identity and a possible N-terminal extension or an internal initiator methionine in P. falciparum and P. cynomolgi. Putative transmembrane domains are underlined. B) The PFyc93 homologues of the two Coccidia, Toxoplasma gondii and Eimeria tenella, are well conserved. C) The PFyc93 homologues of two Piroplasmorida, Theileria parva and Babesia bovis also show clear sequence identity. D) Sequence identity between the putative Vitrella brassicaformis Ycf93 homologue and the apicomplexan proteins is far less obvious.

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Nevertheless, TgYcf93 and EYcf93 are similar size proteins, have a predicted C-terminal trans membrane domain, and are well conserved between these two Coccidia (Fig. 4B). Possible ycf93 orthologues also exist in two members of the Protoplastomorpha, namely B. bovis [47] and T. parva [49] (Fig. 3). The proteins encoded by these small open reading frames lack much overall sequence identity but are relatively similar within lineages (Fig. 4C). Moreover, a predicted C-terminal trans membrane domain exists in TgYcf93 (Fig. 4C), and the genes are situated in syntetic position to other members of the family (Fig. 3) so we believe them to be orthologous.

It is also possible that a ycf93 orthologue exists in the plastids of the related photosynthetic coral symbiont Vitrella brassicaformis [50], a chromerid alga related to Chromera velia [51]. These algae share a common origin of the plastid with Apicomplexa, but remain photosynthetic [4]. In V. brassicaformis good conservation of the gene order in the plastid DNA compared to the apicoplast genome is evident, particularly in comparison with the apicoplast of B. bovis, the key difference in this region being the obvious loss of the photosynthesis genes petD, petH, petB and chlB in the parasite genome (Fig. 3). A small open reading frame, situated between rpl20 and rpoB in V. brassicaformis [4], might be an orthologue of apicoplast ycf93 (Fig. 3), but it shares only modest sequence identity (Fig. 4D), and, unlike all the orthologues in Apicomplexa, is not predicted to contain any trans membrane domain. Drastic rearrangement of the plastid genome in C. velia [4] prevents any identification of putative Ycf93 homologues.

Synteny comparison between the apicoplast genomes and the plastid genome of V. brassicaformis in the ycf93 region indicate that the rps4/rps2 operon underwent reorientation(s) (Fig. 3). The alga V. brassicaformis and the two Protoplastomorpha (B. bovis and T. parva) share the same orientation of this operon, but the Haemosporida (Plasmodium and Leucocytozoon) and the Coccidia (Toxoplasma and Eimeria), which share an almost identical architecture [44], have the reverse orientation (Fig. 3). Rearrangements in plastid genomes are common, and it is difficult to interpret a single reorientation in isolation, especially because the Protoplastomorpha apicoplast genomes are unusual in being transcribed off a single strand and presumably underwent substantial rearrangements to achieve this.

Conclusions

We have demonstrated that of105/ycf93 from the apicoplast genome of P. falciparum encodes a small membrane protein that likely resides in the inner membrane of the apicoplast and undergoes dramatic turnover during the asexual red blood cell life cycle of the parasite. The complete disappearance of this protein after schizogony, and its putative location in the inner apicoplast membrane, might suggest a role in apicoplast biogenesis, but sequence comparison does not give any clues to function. Orthologues of of105/ycf93 occur in all apicoplasts, and perhaps the plastid of the related photosynthetic coral symbiont V. brassicaformis. In keeping with plastid gene nomenclature convention [11] we elevate the gene from of status to ycf (hypothetical chloroplast open reading frame) status as a protein encoded in plastids of multiple organisms, and we occupy the next available number ycf93.

Materials and Methods

Parasite Culture and Drug Treatment

Plasmodium falciparum strains 3D7, D10, mitoYFP/apicoRFP [16], and KAHRP-GFP (Wickham et al., EMBO J 2001) were in vitro cultured using standard techniques [52]. We thank the Australian Red Cross for red blood cells. Parasite synchronization was done using sequential sorbitol lysis as previously described [3]. Parasites used were isolated from red blood cells by incubating with 0.15% saponin and 0.1% BSA in PBS at 4°C followed by extensive washing with PBS.

Drug trials were carried out on tightly synchronized parasites using a modified version of the SYBRgreen assay [3,53]. IC50 values were calculated using the four-component inhibition model in Graphpad Prism.

For microscopy and Western analysis of drug treated parasites, inhibitors were added to double synchronised parasites within 4 hours after invasion of a new host cell.

Peptide Antisera

The peptide NILLKSKNSNNY was selected based on an analysis of the antigenicity and hydrophobicity of the amino acid sequence of PYYcf93 using VectorNTI (Invitrogen). The peptide was compared to the P. falciparum genome and found to be unique for sequences >6 amino acids in length. The peptide was conjugated to Diphtheria toxin (Mimotopes) and used to generate polyclonal goat serum (Antibodies Australia). Antibodies were affinity-purified from serum using immobilized peptide as described [54].

Western Blotting

Protein extraction and Western blotting were done as previously described [54]. For the competition assay, unconjugated PYYcf93 peptide (Mimotopes) was added to affinity-purified antibodies immediately prior to probing the membrane. Triton X-114 membrane partitioning and subsequent Western blot analysis was carried out as previously described [55]. Affinity purified anti-PYYcf93, anti-MSP1-19 [56], anti-AMA1 [18], anti-MSP8 [17], and anti-BiP [57] antibodies were used at 5000:1, 2000:1, 2000:1, 1000:1 and 1000:1 dilutions, respectively.

Immunofluorescence Assay

Parasite fixation and labelling was carried out as previously described [27] except for the PYYcf93 assay in which cell permeabilization and subsequent washing was carried out at 60°C. Affinity purified goat anti-PYYcf93, rabbit anti-ACP [12] antibodies were used at 250:1 and 500:1 respectively. Alexafluor conjugated anti-goat 488 and anti-rabbit 543 (Invitrogen) were use at 250:1 and 500:1 respectively. Triton X-114 antibody enrichment and subsequent Western blot analysis was carried out as previously described [55]. Affinity purified anti-PYYcf93, anti-MSP1-19 [56], anti-AMA1 [18], anti-MSP8 [17], and anti-BiP [57] antibodies were used at 5000:1, 2000:1, 2000:1, 1000:1 and 1000:1 dilutions, respectively. Images were collected using a Leica SP2 confocal microscope and assembled for publication using Adobe Photoshop.

Bioinformatics

Protein sequences were aligned with ClustalW2 using default parameters [58]. Transmembrane domain predictions were made with THHMM version 2.0 [59].

Author Contributions

Conceived and designed the experiments: CDG GIM. Performed the experiments: CDG GIM. Analyzed the data: CDG GIM. Wrote the paper: GIM.
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