**ORIGINAL ARTICLES**

**Characterization of plasma membrane bound inorganic pyrophosphatase from *Leishmania donovani* promastigotes and amastigotes**

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**Abstract**

**Background:** Currently, a major problem in the management of visceral leishmaniasis or kala-azar, especially in the Indian subcontinent, is the growing unresponsiveness to conventional antimonials therapy. Membrane bound pyrophosphatase (PPases) do not exist in plasma membrane from mammals. Thus, H⁺-PPases from *Leishmania* plasma membrane might be potential target in rational chemotherapy of the disease caused by *Leishmania* parasites.

**Objective:** To characterize the activities of inorganic pyrophosphatase (PPase) in the plasma membrane of *Leishmania donovani* promastigote and amastigote.

**Methods:** Culture method of promastigote and amastigote were developed. We assayed PPase activity in isolated plasma membrane of *L. donovani*.

**Results:** We characterized K⁺-PPase present in the plasma membrane of *Leishmania donovani* and investigated its possible role in the survival of promastigote and amastigote. PPase activity was stimulated by K⁺ ions and sodium orthovanadate, inhibited by pyrophosphate analogs imidodiphosphate and alendronate, KF, DCCD, thiol reagent parachloromercuribenzenesulfonate (PCMBS), N-ethylmaliemide (NEM), phenylarsineoxide, ABC superfamily transport modulator verapamil and was also by F₁Fₒ-ATPase inhibitor quercetin.

**Conclusion:** We conclude that there are significant differences within promastigote, amastigote and mammalian host in cytosolic pH homeostasis to merit the inclusion of PPase transporter as putative targets for rational drug design.

**Keywords:** *Leishmania donovani*; pyrophosphatase; plasma membrane

**Abbreviations used:** PPase- Pyrophosphatase, NEM- N-ethylmaliemide, PAO- Phenylarsineoxide, KF- Potassium fluoride, DCCD- N,N'-dicclohexylcarbodiimide, PCMBS-Parachloromer curibenzenesulfonate, SA- Sodium azide, SOV- Sodium orthovanadate, IDP-Imidodiphosphate, QTN- Quercetin, LDC- *Leishmania donovani* cell, DPC- Digitonin permeabilized cell, PM- Plasma membrane, PPᵢ- Pyrophosphate, gp- Glycoprotein, Pro- Promastigote, Am- Amastigote

**Introduction**

Protozoa parasites are responsible for important diseases that threaten the lives of nearly one-quarter of the human population world-wide. Among them, leishmaniasis has become the a leading cause of death, mainly due to the emergence of parasite resistance to conventional drugs1. *Leishmania donovani*, the causative agent of visceral leishmaniasis, encounters a wide range of pH values in its life cycle. The gut of the phlebotamine insect vector is extremely alkaline, whereas promastigotes of *L. donovani* invade host cells via acidic lysosomes2. Mechanisms to cope with this varied environmental pH and maintain cytosolic pH homeostasis might involve the use of proton pumps (H⁺-ATPases and H⁺-PPases) on both plasma membrane and internal membranes. These transport proteins specifically and actively mobilize ions, generating chemical gradients across a membrane. This movement of ions is vital for numerous cellular functions ranging from energy production, motility, nutrient uptake, ionic homeostasis, intracellular signaling, and differentiation, to name a few. Membrane-bound proton-translocating inorganic pyrophosphatases (H⁺-PPase; EC 3.6.1.1) belong to a new category of proton pumps, distinct from F-, P-, and V-ATPases, which utilize pyrophosphate hydrolysis as the driving force for H⁺ movement across the biological membranes3. The membrane-bound, proton-pumping inorganic pyrophosphatase was first described4 in chromatophores from the photosynthetic bacterium *Rhodospirillum rubrum*. Under physiological conditions this enzyme can both
synthesize and hydrolyze PP$_i$. It was also demonstrated\textsuperscript{8} that PP$_i$ could drive a number of energy requiring reactions in chromatophores including ATP synthesis in the dark\textsuperscript{7}. H$^+$-PP synthase is the only known alternative to the well-known ATP synthase in biological electron transport phosphorylation\textsuperscript{8}. H$^+$-PPase gene have been reported to occur in \textit{Leishmania} parasites\textsuperscript{8}. Eukaryotic cells also possess soluble inorganic pyrophosphatases. Pyrophosphate (PP) is formed as a byproduct in several metabolic reactions, for example, DNA and RNA synthesis. It has to be hydrolyzed in order not to stop these reactions. This is a major function of the soluble, cytoplasmic PPases. Membrane bound PPases do not exist in plasma membrane from mammals\textsuperscript{10}, thus, H$^+$-PPases from \textit{Leishmania} plasma membrane might be potential target in relation to rational chemotherapy of the disease caused by \textit{Leishmania} parasites\textsuperscript{11}.

In the present work, we demonstrate that \textit{L. donovani} promastigote and amastigote plasma membrane possesses PPase activity with features in common with the trypanosomatid and plant enzymes\textsuperscript{4}. Our results also indicate that PP$_i$ analogues inhibit the PPase of \textit{Leishmania} parasite.

\section*{Methods}

\textit{Culture methods}\textsuperscript{12, 13, 14}

\textit{L. donovani} promastigote strain MHOM/IN/1978/UR6, a clinical isolate from a confirmed kala-azar patient, was grown at 24$^\circ$C on blood agar medium, pH 7.5. The cells were washed at 500x g twice in cold Tris-sucrose-salt solution (250 mM sucrose, 50 mM NaCl, 20 mM KCl, 1 mM ethylenediamine tetra acetic acid, 20 mM Tris, pH 7.2) and kept it at 4$^\circ$C until use. Viability of harvested cells was monitored microscopically by trypan blue exclusion.

\textit{Amastigote culture method}\textsuperscript{15, 16, 17}

\textit{L. donovani} amastigote strain MHOM/IN/1978/UR6 was grown and maintained as described by Debrabant A et al. Axenically grown amastigotes of \textit{L. donovani} are maintained at 37$^\circ$C with 5\% CO$_2$ by weekly subpassages in MAA/20 (medium for axenically grown amastigotes) at pH 5.5 in 177 cm$^2$ Petridishes. Under these conditions, promastigotes differentiated to amastigotes within 120 hr. Cultures were maintained by 1:3 dilution once a week. The axenic amastigotes remained stable in culture for a long time. Axenic amastigotes were routinely recycled every 10 weeks by differentiation back to promastigotes, and in parallel, initiating a new line of promastigotes. Transformation of amastigotes to promastigotes was performed by centrifugation of amastigotes (1,000 x g at room temperature for 10 min), suspension in promastigote medium, and incubation at 25$^\circ$C. Under these conditions amastigotes differentiate to promastigotes within 72 hr.

\textit{Assay of pyrophosphatase activity}\textsuperscript{18}

Pyrophosphatase activity, in terms of $P_i$ release, was assayed according to the method of Katewa and Katyare. The reaction mixture contains buffer B [Sucrose 300mM, KCl 50mM, Tris 50mM, pH 7, EGTA 2mM pH 7], 0.5 mM Magnesium acetate, 0.25 mM pyrophosphate disodium together with 25 $\mu$g \textit{L. donovani} plasma membrane preparation in a final volume of 1 ml. The mixture was incubated at 25$^\circ$C for 15 min, and the reaction was terminated by adding 0.14 ml of 2.5 M HClO$_4$. In the blank experiments, the reaction mixture was incubated, and the protein solution was added after reaction termination. The amount of enzyme catalyzing hydrolysis of 1 $\mu$mol of pyrophosphate per minute is taken to the unity. Activity was calibrated with a phosphate standard solution. For enzyme inhibition study, inhibitor was added to the reaction mixture containing enzyme but lacking pyrophosphate. The mixture was preincubated for 10 min at 25$^\circ$C before starting the reaction with pyrophosphate. Activity values shown below represent means $\pm$S.E. of three independent experiment.

\textit{Plasma membrane preparation}\textsuperscript{19}

Plasma membrane from \textit{L. donovani} promastigotes was prepared according to the method as described before. Parasites were harvested at a concentration of 10$^8$ cells/ml (5000 x g, 10 min, 4$^\circ$C) washed twice in PBS, and pooled with $^{125}$I cells. All subsequent steps were performed on ice or in refrigerated centrifuges. The cell pellet was suspended to 2 x 10$^7$ cells/ml in PBS buffer, pH 7.2 that contained 10 mM MgCl$_2$ and rapidly mixed with equal volume of 1 mg/ml concanavalin A in the same buffer. Cell aggregation was apparent within 1 min. After 5 min, cells were gently spun at 1,000 x g for 1 min to remove excess concanavalin A. The supernate was discarded, and the cell pellet was re-suspended in 12 ml of 10 mM Tris-HCl buffer, pH 7.5, that contained 10$\mu$g leupeptin/ml and 1mM MgCl$_2$. After swelling for 10min in that hypotonic buffer, cells were homogenized by 18-20 strokes in tight...
fitting Dounce-type homogenizer. Cell lysates and formation of membrane sheets were verified by phase-contrast microscopy. The homogenate was layered over a two-step gradient consisting of 8 ml of 0.5 M mannitol over 4 ml 0.58 M sucrose, both in Tris buffer, and spin at 1000 x g for 20 min. For analysis, material remaining at the top of the 0.5 M mannitol was saved. Large crude plasma membrane fragments were separated as a tight pellet at the bottom of the gradient. This pellet was re-suspended in Tris buffer that contained 1M α-methylmannoside and left on ice for 40 min with occasional mixing.

This plasma membrane, free from bulk concanavalin A, were diluted into three volumes of Tris buffer and homogenized by 80 strokes with a glass Dounce-type homogenizer. This second homogenate was layered on a single-step gradient that consisted of 20% sucrose in Tris buffer and spun for 30 min at 500 x g. Scrolls and large plasma membrane sheets above the 20% (w/v) sucrose layer was collected by centrifugation at 40,000 x g for 1 hr. The pellet containing the enriched plasma membranes, was re-suspended in Tris buffer. All samples were either assayed immediately or frozen at -20°C for further use.

**Preparation of digitonin permeabilized Leishmania cell**

*L. donovani* promastigote and/or amastigote cells were collected, washed once by buffer A (140mM NaCl, 20mM KCl, 20mM Tris, 1 mM EDTA, pH 7.5) and resuspended in isolation buffer (20 mM MOPS, pH 7.0; 0.3% bovine serum albumin; 350 mM sucrose; 20 mM potassium acetate; 5 mM magnesium acetate and 1 mM EGTA). Cells were permeabilized in separate tube with 50µg digitonin/ mg protein and incubated on ice for 10 min. After incubation, the cells were centrifuged at 6,000×g for 7 min. Pellets were re-suspended in assay buffer.

**PPase activity in plasma membrane**

Activity of PPase was optimum at pH 6.8 (data not shown). KCl greatly stimulated PPase activity. Replacing 50 mM KCl with 50 mM NaCl or 50 mM choline chloride in the buffer resulted in substantial loss of PPase activity as shown in figure 1.

**Figure 1: Effect of salt concentration on plasma membrane PPase activity.** The values represent the means ± S. D. of three independent experiments.

Panmede was purchased from Paines and Byrne (Greenford, Middlesex, UK).

**Statistical analysis**

All experiments were performed in triplicate, with similar results obtained in at least three experiments. Statistical significance was determined by student’s t test. Significance was considered as P < 0.05.

**Results**

**Isolation of plasma membrane**

Development of a method based on cell disruption by osmotic swelling in presence of a plasma membrane specific lectin concanavalin A and differential centrifugation has proved to be ideal for obtaining a plasma membrane fraction from *L. donovani* promastigote. This membrane fraction consists of open membrane sheets with microtubules attached to the membranes. The utility of the method was assessed by assaying marker enzymes. The activity of tartarate-resistant acid phosphatase, a plasma membrane marker of *L. donovani* was enriched in this fraction. An enrichment of 28 fold was obtained when cell surface plasma membrane was labeled with 125I.

**Materials**

All the biochemicals unless otherwise mentioned were from Sigma-Aldrich (St. Louis, MO, USA).
Table 1 shows that activity of PPase in plasma membrane is located at endoplasmic face of plasma membrane. Permeabilized *L. donovani* cell showed substantial PPase activity, whereas nonpermeabilized cell did not. However, very little ecto-PPase activity have been observed at exoplasmic face of the plasma membrane.

**Table 1: Substrate specificity of pyrophosphatase in *L. donovani* promastigote membrane**

<table>
<thead>
<tr>
<th>Addition (Substrate)</th>
<th>Enzyme activity in DPC</th>
<th>Enzyme activity in LDC</th>
<th>Enzyme activity in PM</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP_i (0.25 mM)</td>
<td>47 ± 6</td>
<td>10 ± 0.8</td>
<td>117 ± 18</td>
</tr>
</tbody>
</table>

*Assay of pyrophosphatase activity have been described in Materials and methods. Values represent mean ± S.E.M. of three independent experiments. Enzyme activity (nmol phosphate/min/mg protein)*

Figure 2 shows PPase activity and the amount of plasma membrane proteins added was linear. Linearity of PPase activity was also observed for digitonin permeabilized cell. It is also evident from Figure 3 that in digitonin permeabilized amastigote cell, PPase activity was lesser than promastigote.

**Figure 2: Effect of plasma membrane protein concentration on the rate of PP_i hydrolysis by PPase. Each point represents the average value of three experiments.**

**Figure 3: Effect of digitonin permeabilized Leishmania cell protein concentration on the rate of PP_i ([●]), hydrolysis by promastigote PPase, and amastigote PPase ([▲]). Each point represents the average value of three experiments.**

**Specific inhibitors and stimulators of PPase activity**

PP hydrolysis in the plasma membrane and digitonin permeabilized *L. donovani* promastigote cell was inhibited, in a dose-dependent manner by KF as shown in figure 4. PP_i hydrolysis was inhibited by the PP_analogue, IDP and alendronate as shown in figure 5B. The sulphhydryl group inhibitors p-chloromercuribenzenesulphonate (PCMB) and phenylarsineoxide (PAO) showed potent inhibition on PPase activity. Some compounds used to inhibit H^+^-ATPase activity, such as N-ethylmaleimide (NEM) and DCCD\(^{35}\), are also able to inhibit the plasma membrane PPase. F_F^i^-ATPase and F_F^i^-ATPase inhibitor sodium azide\(^{37}\) did not inhibited the PPase activity as shown in figure 5B. The bioflavonoid quercetin which can inhibit F_F^i^-ATPase strongly at low concentration and F_F^i^-ATPase moderately at high concentration\(^{39}\) had inhibited plasma membrane bound PPase moderately at high concentration. Sodium orthovanadate, a P-type H^+^-ATPase inhibitor\(^{38}\) and P-type ABC superfamily of transporters, such as P-glycoprotein involved in the multidrug resistance\(^{39}\), stimulated PPase activity. Effects of stimulators and inhibitors on PPase in digitonin permeabilized amastigotes were very similar to the effect on PPase of promastigote plasma membrane (data not shown). Verapamil, a P-gp ATPase stimulator\(^{30}\), inhibited PPase activity.
Figure 5: Concentration-response curve describing the effect of effectors on promastigote plasma membrane PPase. Each point represents the average value of three experiments. The effectors were added to plasma membrane 10 min before the addition of substrates.

5A: PCMBS ( ), SOV ( ), QTN ( ), Verapamil ( ), DCDD ( ), NEM ( )

5B: Aledronate ( ), SA ( ), PAO ( ), IDP ( )

Discussion
In the present study, we have identified and characterized PPase activity in the plasma membrane of L. donovani promastigote. This is the first report demonstrating biochemically the presence of a PPase in the plasma membrane of L. donovani promastigote. Our results suggest that K⁺ was necessary for PPase activity, but substantial amount of K⁺ and Na⁺ independent PPase was also present in the plasma membrane. Since IDP is a bisphosphonate, we tested other bisphosphonate (alendronate) used clinically in the treatment of bone resorption disorders for its inhibitory effect on PP hydrolysis. The chemical modification with the trivalent arsenical reagent PAO indicates the involvement in PPase activity of a viscinal sulfhydryl group.

The significance of the presence of a PPase in the amastigote plasma membrane may be proton pumping to maintain neutral cytosolic pH in an environment of acidic pH of phagolysosome. Data provided evidence that pH formation in promastigote and amastigote was partially inhibited by H⁺-ATPase inhibitor DCCD. This fact suggested that mechanisms in addition to that inhibited by DCCD may also be involved in controlling homeostasis of pH. It appears from our observations that PPase may also play an important role in the extrusion of proton from the cytosol.

Conclusion
In conclusion, this is the first report of a PPase in plasma membrane of an organism different from plants. Since membrane bound PPases are apparently absent from vertebrates, analysis of the role of PPase in pH regulatory mechanisms of L. donovani amastigote might serve as potential target for putative therapeutic intervention.

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