**Leishmania donovani** Secretory Mevalonate Kinase Regulates Host Immune Response and Facilitates Phagocytosis

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**Summary Statement:** *Leishmania* secretes over 151 proteins during *in vitro* cultivation. Cellular functions of one such novel protein: mevalonate kinase is discussed here; signifying its importance in *Leishmania* infection. Visceral *Leishmania* infection, caused by *Leishmania donovani* in Indian subcontinent. This persistence is partly due to phagocytosis and evasion of host immune response. The underlying mechanism involves secretory proteins of *Leishmania* parasite; however, related studies are meagre. We have identified a novel secretory *Leishmania donovani* glycoprotein, Mevalonate kinase (MVK), and shown its importance in parasite internalization and immuno-modulation. In our studies, MVK was found to be secreted maximum after 1 h temperature stress at 37°C. Its secretion was increased by 6.5-fold in phagolysosome-like condition (pH ~5.5, 37°C) than at pH ~7.4 and 25°C. Treatment with MVK modulated host immune system by inducing interleukin-10 and interleukin-4 secretion, suppressing host’s ability to kill the parasite. Peripheral blood mononuclear cell (PBMC)-derived macrophages infected with mevalonate kinase-overexpressing parasites showed an increase in intracellular parasite burden in comparison to infection with vector control parasites. Mechanism behind the increase in phagocytosis and immunosuppression was found to be phosphorylation of mitogen-activated protein (MAP) kinase pathway protein, Extracellular signal-regulated kinases-1/2, and actin scaffold protein, cortactin. Thus, we conclude that *Leishmania donovani* Mevalonate kinase aids in parasite engulfment and subvert the immune system by interfering with signal transduction pathways in host cells, which causes suppression of the protective response and facilitates their persistence in the host. Our work elucidates the involvement of *Leishmania* in the process of phagocytosis which is thought to be dependent largely on macrophages and contributes towards better understanding of host pathogen interactions.

**Keywords:** *Leishmania*, mevalonate kinase, immune response, phosphorylation, infection, secretion, invasion, phagocytosis
INTRODUCTION

The Leishmaniases are a set of neglected tropical diseases caused by 20 species of *Leishmania* parasites and is transmitted to humans by infected female sandfly bite. Among different types of the disease caused by *Leishmania*, Visceral Leishmaniasis (VL) is the most severe one which is often fatal if not treated in time. In India, VL is caused by *Leishmania donovani* species. Progresses in controlling the Leishmaniases will require better understanding of pathogenesis to recognize novel drug targets or vaccine candidates.

*Leishmania* exists in two forms; the extracellular, flagellated, motile form is promastigote, that resides in the alimentary canal of the sandfly. Blood feeding activity of the vector results in the transmission of the parasite to the human where it is phagocyted and transformed into intracellular, non-flagellated, non-motile amastigote form. *Leishmania* is an enormously successful organism considering its two structural variants and ability to exist in the harsh host environment. Some of these survival mechanisms may be attributed to a large repertoire of proteins secreted by the parasite. *L. donovani* releases a total of 151 proteins in abundance to the extracellular media (Silverman et al., 2008). These are not a set of unrelated proteins, rather, these are functionally related group of proteins (Geiger et al., 2010). The exoproteome is known to assist the entry of parasite into host cells which is a prerequisite for infection (Nandan et al., 2002; Choudhury et al., 2010a; Zylbersztejn et al., 2015). In addition, *Leishmania* exosome treatment induces immune suppression of macrophage prior to infection and creates an environment to support early infection (Silverman et al., 2010). The secretory proteins and surface molecules on parasites form an interface between the parasite and host. Though, the major cell surface molecules of parasites are well characterized (Connell et al., 1993; McConville et al., 1993; Winter et al., 1994), very less information is available regarding secretory proteins/antigens for their role in host infection.

Mevalonate kinase, one such secreted protein has been reported with different organisms. However, Mevalonate kinase in *Leishmania donovani* is not known. Mevalonate pathway, present in most of the eukaryotic cells, is essential for various cellular functions, such as, cell cycle regulation, control of cell growth and size, autophagy, and protein glycosylation (Fu et al., 2002; Miettinen and Björklund, 2016). The mevalonate pathway also provides precursors for cholesterol biosynthesis. Mevalonate kinase (MVK) is an important enzyme of this pathway catalyzing Mg^{2+}-ATP dependent phosphorylation of mevalonic acid to mevalonate-5-phosphate. This step is regulated by feedback inhibition (Dorsey and Porter, 1968; Henneman et al., 2011). *L. major* MVK crystal structure was elucidated and its ATP binding site was found to be structurally distinct (Sgraja et al., 2007). In *Trypanosoma cruzi*, MVK is secreted outside the cell and it modulates the host cell signaling (Ferreira et al., 2016). In *L. major* and *T. cruzi*, dimeric MVK has a high enzymatic activity (Sgraja et al., 2007; Ferreira et al., 2016). However, the mevalonate pathway in *L. donovani* has not been studied till date. Here, we have demonstrated that MVK protein is present in *L. donovani* and is secreted. It was observed that it regulates host immune response and induce parasite entry through phosphorylation of ERK-1/2, p-38, and cortactin. Altogether, our work sheds light on the involvement of *Leishmania* in the process of phagocytosis and contributes towards better understanding of host pathogen interactions.

MATERIAL AND METHODS

Ethics

All experiments were assessed and approved by the Institutional Animal Ethical Committee (AH/RMRI/IAEC/09/33-37), Indian Council of Medical Research, Rajendra Memorial Research Institute of Medical Sciences, Patna, and are managed by CPCSEA (The Committee for the Purpose of Control and Supervision of Experiments on Animals), Government of India, New Delhi. ICMR-RMRRMS (Indian Council of Medical Research-Rajendra Memorial Research Institute of Medical Sciences) follows “The Guide for the Care and Use of Laboratory Animals,” were followed as per 8th edition by the Institute for Laboratory Animal Research. For the use of human sample too, approval from ethical committee was taken (RMRI/EC/02/20).

Parasites

Ag83 strain of *L. donovani* originally obtained from an Indian VL patient was maintained routinely in mice as earlier described (Das et al., 2012). Promastigotes were maintained in Gibco™ Medium-199 (ThermoFisher Scientific; #10063372) containing 10% (v/v) heat-inactivated Gibco™ fetal bovine serum (FBS; ThermoFisher Scientific; #10082139), 25 mM 4-(2-hydroxyethyl)-1-piperazinenuethanesulfonic acid (HEPES) (Calbiochem), 4 mM sodium bicarbonate (NaHCO₃) (Sigma-Aldrich), 100 μg/ml streptomycin (Sigma-Aldrich), and 100 units/ml penicillin G-sodium (Sigma-Aldrich) at 24°C as earlier described (Kumar et al., 2018).

Generation of Axenic Amastigotes

Axenic amastigotes were generated as described previously (Kaul et al., 2000) with minor modifications. Promastigote culture grown in Medium199 supplemented with 20% FBS was incubated at 37°C. After 24 h, cells were maintained in Medium199 (20% fetal bovine serum) with pH 5.5 under same temperature conditions for 5 days. Obtained axenic amastigote culture were used for the experiments.

Signal Peptide and Transmembrane Protein Prediction

Protein sequence of LdMVK was obtained from sequenced MVK gene in FASTA format and uploaded in SecretomeP 2.0 server to study the possibility of its secretion by non-classical i.e. non-signal peptide triggered protein secretory route. Due to the limited organism group in list, mammalian was selected as the organism group and the sequence was submitted.

Polymerase Chain Reaction Amplification

Full length LdMVK gene coding region was amplified from the genomic deoxy-ribonucleic acid (DNA) of Ag83 strain using following primer pair: Forward- 5' TTTTGGATCCATGCCA
GTTTGACGCGGTGG3 and Reverse- 5’TTTTAAGCTTCAG GTTTGACGCGGTGG3. Polymerase Chain Reaction (PCR) conditions were: 60 s at 94°C, 45 s at 56°C, 60 s at 72°C (35 cycles), and 10 min at 72°C for final extension. Full length MVK gene sequence was obtained through di-deoxy sequencing of MVK-PCR product.

**Soluble Leishmania Antigen Preparation**

To prepare Soluble *Leishmania* antigen (SLA), promastigotes harvested from 3 to 4 days of culture was washed two times in PBS and resuspended in PBS containing protease inhibitors cocktail (Sigma). Ultra-sonication was performed followed by centrifugation (20,000g for 20 min) and protein quantification of the supernatant was done by Bicinchoninic acid method and SLA was stored at −20°C for further use.

**Supernatant Processing and Determination of Mevalonate Kinase Activity**

*Leishmania* was inoculated at a concentration of 1 × 10⁶ cells/ml. Promastigotes and amastigotes of 4th day culture were washed three times with 1X PBS and incubated in pre-warmed (37°C) Medium199 (without FBS) containing Halt protease inhibitor cocktail (Thermo Scientific) at a concentration of 1 × 10⁵ cells/ml for required time. The viability of parasites was assessed by trypan blue dye exclusion test to ensure over 98% viability. The culture was centrifuged (3,000 g, 10 min, 4°C) and the obtained supernatant was passed through 0.45 µm pore size syringe filters (Millipore) to remove any remaining parasites. It was then concentrated at 4°C to 100 µl volume by 10 kDa MWCO (Molecular weight cut off) centricron (Millipore) according to the manufacturer’s protocol. Distilled water was added to the concentrated sample to reduce the salt and phenol red content and sample was again concentrated to required volume. Wherein required, the obtained supernatant consisting of secretory proteins was quantified by bichinchoninic acid assay and then stored at −20°C until use. Once obtained, the supernatant was kept on ice at all times.

For time kinetics experiments and heat and pH based relative studies, normalization of the detected MVK amount could not be done with other secretory protein since there are no known proteins that are constitutively secreted from parasites in a stable manner. Therefore, the parasites were carefully counted for each sample before supernatant preparation and soluble *Leishmania* antigen was prepared by already described method. For normalizing blots, both cell lysate and supernatants for the same population was run and the protein of interest was normalized against cell number control (β-actin).

To check the presence of mevalonate kinase activity in culture supernatant, *L. donovani* promastigotes were incubated in serum-free Medium199 for 6 h at 37°C. Culture supernatant was collected, concentrated, and mevalonate kinase assay was performed. Reaction mixture was prepared consisting of 100 mM glycine, 25 mM sodium chloride (NaCl) pH 9.0, 4 U lactic dehydrogenase, 4 U pyruvate kinase, 1 mM phosphoenolpyruvate, 5 mM adenosine triphosphate (ATP), 5 mM Magnesium chloride (MgCl₂), 4 mM mevalonate, and 30 µM β-nicotinamide adenine dinucleotide hydrogen (β-NADH). It was incubated at 25°C for 10 min and assay was initiated by concentrated supernatant addition (0.6 µg/µl; 260/280: 0.63). MVK activity was examined by coupling ADP release with oxidation of NADH by pyruvate kinase and lactate dehydrogenase, which was measured at 340 nm for 600 s (time scan) using Shimadzu UV-visible spectrophotometer. For blank assays, Medium199 processed using centrific was used. The reactions were performed in triplicates and culture supernatants from different batches were used for each assay.

**Cloning, Expression, Purification, and Polyclonal Antisera Generation Against LdMVK**

The PCR product was ligated to pET-28a expression vector (Novagen). pET28a is a bacterial expression vector that expresses protein in fusion with amino terminal and carboxy terminal His₆ tag. Recombinant vector and PCR fragment was sequenced by dideoxynucleotide chain termination method for confirmation of cloning. *E. coli* pET-28a-LdMVK vector were transformed in BL-21(DE3) cells, grown at 37°C, 150 rpm and 1 mM IPTG induction was performed for 4 h. Cells were resuspended in buffer containing 50 mM Tris-HCl, 300 mM sodium chloride (NaCl), 1X protease inhibitor cocktail (Roche), and 0.1 mg/ml of lysozyme (Sigma) and incubated in ice bath for 30 min. Sonication disrupted cells and debris was removed by centrifugation (20,000 g, 15 min). Expression of protein was evaluated by western blot using anti-his antibody (1:1,000 dilution; ThermoFisher Scientific; #MA1-135).

For affinity chromatography purification, cell lysate was passed through Nickel-nitriotriacetic acid (Ni-NTA) resin column (Qiagen). Protein was eluted from the nickel column with an increasing buffer gradient consisting of 150 mM Tris pH-7.8, 300 mM NaCl, and 250 mM imidazole. Anti-LdMVK antibodies were generated by one subcutaneous immunizations with pure rLdMVK along with complete Freund’s adjuvant (Sigma-Aldrich, St. Louis, CA, USA), following three subcutaneous immunizations along with incomplete Freund’s adjuvant (Sigma-Aldrich, St. Louis, CA, USA) in rabbit at 14-day intervals. Serum was isolated 10 days after last immunization and western blot of rLdMVK and Ag83 soluble *Leishmania* antigen were performed to verify antibody specificity.

**Ouchterlony Test**

Gel slides were prepared using 1.2% agarose in assay buffer and poured 4 ml/slide. Wells were bored on gel. Then 10 µl rLdMVK protein was placed on the well at the center and 10 µl each of post immune sera, distilled water, and pre-immune sera was placed the other wells. The slide was incubated overnight in a moist chamber at 37°C and the next day it was observed.

**Western Blotting**

To confirm the expression of recombinant MVK (containing his-tag in N and C-terminal), expressed bacterial cell lysate was separated on 12% SDS-PAGE gel and probed with anti-his antibody at 1:1,000 dilution for 1h and then with anti-rabbit antibody (1:5,000 dilution; Jackson ImmunoResearch laboratories; AB_2307391) for 45 mins.
For secretion experiments, culture supernatant of *L. donovani* was concentrated and run in SDS-PAGE. Secretory proteins were transferred onto PVDF membrane and processed for western blots analysis with anti-MVK antibody raised in rabbit at 1:1,000 dilutions for 1 h and then with anti-rabbit antibody (1:5,000 dilution; Jackson ImmunoResearch laboratories; AB_2307391) at 1:5,000 dilutions for 45 mins. Antibody antigen complexes were detected by enhanced chemiluminescence (GE Healthcare; #RPN2209) and bands were visualized using Image J software. Band densitometry was performed using ImageJ software. To validate that the supernatant containing secretory proteins was collected from equal number of cells, cell lysate was prepared for each condition and probed with anti-actin antibody which was kindly provided by Dr Anuradha Dube.

**Characterization of MVK**

**Enzymatic Activity of r-LdMVK**

Specific activity of purified r-LdMVK was tested. Reaction mixture consisting of 100 mM glycine, 25 mM sodium chloride (NaCl) pH 9.0, 4 U lactic dehydrogenase, 4 U pyruvate kinase, 1 mM phosphoenolpyruvate, 5 mM magnesium chloride (MgCl2), 4 mM mevalonate, and 30 µM β-nicotinamide adenine dinucleotide hydrogen (β-NADH) were incubated at 25°C for 10 min. To start the reaction, r-LdMVK was added, cuvette with lid was inverted for mixing, and time scan was performed immediately at 340 nm for 600 s to monitor NADH oxidation. Mevalonate kinase activity converts ATP to ADP which is used by pyruvate kinase to form pyruvate. Pyruvate, in the presence of pyruvate kinase, is converted to lactate, oxidizing NADH. Thus, NADH oxidation is linked to mevalonate kinase activity. Blank assays were performed in r-LdMVK absence. Enzyme activity of one unit corresponds to production of 1 mol NADH/min. Specific activity was expressed as µmole product formed/min/mg protein. The reactions were done in triplicates and purified proteins from different batches were used for each assay. Also, different concentrations of rLdMVK were used. To determine heat stability, the reaction mixture was incubated for 15 min at temperatures between 25 and 60°C, brought to room temperature, assay was performed and readings were taken. The temperature with the highest activity were considered as 100% and from this the residual kinase activity was determined.

**Glycoprotein Staining**

LdMVK protein was electrophoresed (SDS-PAGE) and staining of glycoprotein was carried out as stated in Pro-Q Emerald glycoprotein gel and blot stain kit manufacturer’s protocol (Molecular Probes, Eugene, OR). Protein in gel was fixed in a solution of 50% methanol and 5% acetic acid and washed in 3% acetic acid solution. Gel was incubated in oxidizing solution provided in the kit for 30 min and washed two times. Incubation in Pro-Q Emerald 300 staining solution was done for 2 h and stained gel was then visualized by illuminating in UV light in BioRad transilluminator. CandyCane™ glycoprotein molecular weight standards (Molecular Probes, Eugene, OR, USA) containing a mixture of alternate glycosylated and non-glycosylated proteins was separated by SDS-PAGE and stained by Pro-Q Emerald glycoprotein gel and blot stain kit, and it served as positive and negative control for the experiment. The staining was performed at least three times and protein from different batches were used each time.

**Immunofluorescence Assay**

Immunofluorescence assay was carried out to know the localization of MVK within parasite using anti-LdMVK antibody and anti-Ld-pyruvate phosphate dikinase antibody (glycosomal marker, raised in mice, 1:1,000 dilution). Briefly, exponentially grown *L. donovani* (5 × 10⁶ cells/ml) of Ag83 strain were collected by centrifugation, washed three times in PBS. Cells were incubated in fixation/permeabilization solution (BD Biosciences) for 30 min at 4°C. Blocking (3% BSA in PBS) was performed for 30 min and cells were incubated for 45 min with anti-MVK antibody (1:200 dilution) prepared in 1% BSA and Triton X-100 containing PBS. Cells were then incubated with FITC conjugated anti-rabbit antibodies (1:200 dilution; Jackson ImmunoResearch; #AB_2337972) in dark for 45 min. Further, cells were incubated with anti-pyruvate phosphate dikinase antibody (1:200 dilution; glycosomal marker) for 30 min, following incubation with TRITC conjugated anti-mouse antibodies (1:200 dilution; Jackson ImmunoResearch; #AB_2337972). Washing of cells were carried out with permeabilization/wash buffer (BD Biosciences; #554723) three times. Finally, cells were resuspended in PBS, mixed with prolong anti-fade solution (ThermoFisher Scientific; #P10144) and photographs were obtained from confocal laser scanning microscope (LSM880 zeiss) using a 100× Numerical aperture (NA) 1.44 PlanApo oil immersion objective. The assay was performed more than three times.

**Studies on Host Immune Response to MVK**

**Peripheral Blood Mononuclear Cells and Macrophages Isolation**

To estimate the effect of r-LdMVK on immune response of host and parasite entry, venous blood from a non-endemic healthy control was collected in heparinized tube and mixed with phosphate buffered saline (PBS) (1:1). This suspension was layered on Ficoll-Hypaque (Sigma; #17144003) in 1:1 ratio and density gradient centrifugation was performed. Obtained peripheral blood mononuclear cells (PBMC) were washed thrice with Gibco™ RPMI-1640 medium (ThermoFisher Scientific; #31800022). For infection study, monocytes were segregated from PBMC in plastic culture flask since they have a property of plastic adherence. Five times washing with pre-warmed PBS was performed to remove non-adherent cells. Cell viability was evaluated by trypan blue dye (Ferreira et al., 2016). Adherent cells were cultured in complete RPMI-1640 medium (10% FBS) for up to 6 days and the medium was replaced after every 3 days. Accutase (Sigma; #A6964) was used to obtain adherent macrophages, cells were counted and re-adhered on chamber slides or six-well plate.

**Enzyme Linked Immunosorbent Assay**

The level of Interferon-γ (IFN-γ), Interleukin-12 (IL-12), Interleukin-2 (IL-2), Tumor necrosis factor-α (TNFα) in addition to Interleukin-10 (IL-10) and Interleukin-4 (IL-4) were calculated with commercial ELISA kits. PBMC’s from
healthy person (1 × 10^6 cells/ml) were plated in 24-well culture plates for 24 h. Soluble *Leishmania* antigen (50 µg/ml), r-LdMVK (1 µg/ml), or lipopolysaccharide (LPS) (100 ng/ml) (Sigma; #54881) were added in triplicate wells, incubated for 16 h and cytokine response was observed according to the manufacturer’s protocol (BD OptEIA kit, USA). The results were obtained as picograms of cytokine/ml, based on the standard curves generated using a recombinant cytokine provided in the kit. Anti-inflammatory to pro-inflammatory cytokine ratios associated with different antigens-induced PBMC were compared. The experiment was carried out with three biological replicates, each performed in triplicates.

### Infection Studies

#### Cell Binding Assay

**Method 1**

PBMC-derived macrophages (5 × 10^4) were seeded in 96-well microtiter plates and incubated overnight in RPMI 1640 supplemented with 10% FBS at 37°C in 5% CO2. Cells were fixed with 3.7% paraformaldehyde in PBS, washed three times with PBS, and blocked with 10% FBS diluted in PBS for 1 h at room temperature. Cells were washed three times with 0.05% Tween 20 containing PBS (T-PBS). Increasing amount of purified recombinant MVK was added to the wells (0.1 to 15 µg/ml) and incubated for 1 h at room temperature. Cells were washed with T-PBS and incubated with anti-MVK antibody (1:1,000 dilution) for 30 min and then with HRP tagged anti-rabbit antibody (1:1,000 dilution) for 30 min. After three washes with T-PBS, 3,3′,5,5′-Tetramethylbenzidine (TMB) (BD Biosciences; #555214) was added, reaction was stopped and absorbance was measured at 495 nm. There were three biological replicates under our study and for each condition ELISA were performed in triplicates.

**Method 2**

MVK binding to host cells was confirmed as previously described (Huynh and Carruthers, 2016). Precisely, r-LdMKV (1 µg/ml) or soluble *Leishmania* antigen was incubated with PBMC-derived macrophages adhered to six-well plate for 1 h. Monolayer cells were then washed with PBS-CM (1X PBS, 1 mM CaCl_2, and 1 mM MgCl_2) four times and the last wash was stored to check for r-LdMVK presence. Cells were lysed with RIPA buffer supplemented with protease inhibitor cocktail (Roche; #11697498001). The obtained cell bound fraction was separated by 12% SDS-PAGE and immunoblotted with anti-MVK antibody.

#### Generation of MVK-Overexpression Parasites

For the generation of MVK-OE (MVK-Overexpression) parasites, the LdMVK coding sequence was amplified using following primers: FP: 5′-TTTATAGCTTATGCAAGCCGGTCAAG-3′ and RP: 5′-TTTCTGATCCACGGTTGACCCCGTGG-3′. The amplified products were ligated within HindIII and BamHI restriction sites and sense cloning was done in the same vector. The resulting recombinant pLGFPN vector was cloned in DH5α cells for stability. For transfection, late log phase promastigotes (2 × 10^8 cells/ml) were washed with electroporation buffer containing 6 mM glucose, 21 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), pH 7.5, 5 mM Potassium chloride (KCl), 137 mM NaCl, and 0.7 mM Disodium phosphate (Na₂HPO₄) and transfected with recombinant pLGFPN (10 µg) by electroporation in 4 mm electroporation cuvette using a Gene Pulsar (Bio-Rad). Electroporation was carried out according to the high-voltage protocol: 25 µF, 1,500 V (3.75 kV/cm) pausing 10 s between two pulses. Parasites were allowed to recover for 24 h and *Leishmania* that stably incorporated pLGFPN and pLGFPN-LdMVK vectors were chosen by culturing parasites for 4 weeks in the presence of increasing antibiotic Geneticin (G418) concentration (5 µg/ml to 50 µg/ml).

### Quantitative PCR

Fourth-day culture of MVK-overexpression and vector control strains were harvested at 800 g, 5 min and washed with PBS twice. Pellet was collected, resuspended, cells were counted, and 10^7 cells were used for each strain. RNA was extracted from the cells using the TRIzol™ Plus RNA purification kit (ThermoFisher Scientific, #12183555) following manufacturer’s protocol. Amount of extracted RNA was quantified by Nanodrop Spectrophotometer. Exon of sequences were identified from the NCBI database and MVK and 18S rRNA qPCR primers were designed using IDT (Integrated DNA Technologies, USA). MVK FP sequence was: 5′ CGGATGAAGGTTGATACGAG 3′; and MVK RP sequence was: 5′ GTATGGCGGACTCATTGCC 3′. 18S rRNA was used for normalization and its FP and RP sequences were: 5′ GGGCCGATTAATGGAGATGC 3′; and MVK RP sequence was: 5′ CCAGATTCACCTAGGCTGG 3′ respectively. Quantitative Polymerase Chain Reaction (qPCR) conditions were: 30 s at 95°C, 45 s at 58°C, 30 s at 72°C (45 cycles), and 5 min at 95°C for initial denaturation. cDNA was prepared using High capacity cDNA reverse transcription kit (ThermoFisher Scientific, #4368814) and qPCR was performed using SYBR green PCR master mix (ThermoFisher Scientific, #4309155). Negative template controls were made for each qPCR analysis. Three biological replicates were used and each time the reaction was carried out in triplicates.

### Infection of Macrophages

PBMC-derived macrophages (5 × 10^5 cells/well) were prepared as previously described, seeded onto four-well chamber slides (Nunc Lab-Tek) and incubated overnight in RPMI 1640 supplemented with 10% FBS at 37°C in 5% CO2. Cells were washed with PBS and infected with metacyclic *L. donovani* promastigotes in 0.4 ml RPMI 1640 at 1:10 ratio (macrophage: parasite). After 4 h at 37°C in 5% CO2, parasites were removed by three PBS washes and chamber slides were incubated for required periods of time. This was followed by methanol fixation and either Giemsa staining or immunofluorescence. For MVK treatment, r-LdMVK (1 µg/ml) was added to the macrophages along with the parasites. For antibody inhibition assay, promastigotes were pre-incubated with anti-MVK antibody for 30 min at 25°C. The measurement of intracellular parasites was carried out in 400 macrophages per well and the data was shown as the total number of intracellular parasites per 100 macrophages. Infection was repeated with mutant strains of parasites: MVK-Overexpression/Vector control strains (4 h duration) as already discussed. For each condition, experiment was carried out in triplicates and the data were analyzed using an
unpaired Student’s t test and indicated as mean ± SE of three independent experiments. P value <0.05 was considered to be significant.

**Phosphoprotein Assay**

PBMC-derived macrophages (4 × 10⁶) were seeded and grown for 24 h. Cells were incubated for another 24 h with serum-free RPMI to reduce constitutive signaling. Following starvation, cells were incubated with r-LdMVK (1 ng/ml) from 5 to 90 min. Cells were then washed with PBS containing 2 mM sodium orthovandate (Na₃VO₄) and 5 mM sodium fluoride (NaF) to minimize phosphatase activity. Cell lysis was performed with mammalian lysis buffer (Cell Signaling Technology; #8003) supplemented with phosphatase and protease inhibitors. Protein was quantified by Bicinchoninic acid method and probed with antibodies against Phospho-Extracellular signal-regulated kinase (ERK-1/2) (1:1,000 dilution; Cell Signaling Technology, #9101), ERK-1/2 (1:1,000 dilution; Cell Signaling Technology, #9102), Phospho-p38 MAP kinase (1:1,000 dilution; Cell Signaling Technology, #9211), p38 MAP kinase (1:1,000 dilution; Cell Signaling Technology, #9212), Phospho-Cortactin (1:1,000 dilution; Merck, # AB3795), Cortactin (1:1,000 dilution; Cell Signaling Technology, #3502), and housekeeping gene, Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (1:1,000 dilution; Santa Cruz Biotechnology, #sc-32233). Each experiment was carried out three times.

**Expression and Characterization of Recombinant Mevalonate Kinase**

Full length amino acid sequence of LdMVK has a molecular mass of 35.61 kD and a theoretical isoelectric point of 9.13 as predicted by Expasy. Protein sequence similarity between human MVK and L. donovani MVK was only 26%. LdMVK was cloned into E. coli (Figure 2A) and confirmed by DNA sequencing that showed 99% homology with putative MVK sequence. Recombinant protein (r-LdMVK) tagged with poly-histidine was expressed (Figure 2B) and MVK expression was confirmed through western blot technique using anti-his antibody (Figure 2D). The expressed protein was purified using Ni-NTA affinity chromatography and a single band of expected size was obtained (Figure 2C). The purified r-MVK was then used for polyclonal antibody production. Obtained anti-MVK antisera was validated for its specificity by western blot (Figure 2E) and Ouchterlony test (Figure 2F). Western blot of whole cell lysate (WCL) of L. donovani promastigotes and axenic amastigotes probed with anti-MVK antibody demonstrated a single band of expected molecular weight (35 kD) that further confirmed the presence of MVK in both forms of L. donovani (Figures 2G, H). The presence of MVK in the culture supernatant was confirmed by western blot. Both promastigotes and axenic amastigotes of Ag83 strain released MVK into the extracellular medium (Figures 2G, H). The size of secreted LdMVK was found to be the same as that of the cytosolic LdMVK, hence, the released forms are not proteolytically processed. WCL (whole cell lysate) was prepared from only 3% of the pellet and loaded. Supernatants can be compared to the diluted lysates of cell (3% of pellet). Natural death during the course of Leishmania culture does not include the release of intracellular content. It is earlier reported that glycosylation inhibitors inhibit Leishmanial infectivity (Nolan and Farrell, 1985; Kink and Chang, 1987), so we also attempted to verify if LdMVK is a glycoprotein using glycoprotein specific staining procedure as discussed in methods section. Glycoprotein specific staining successfully stained r-MVK confirming its glycoprotein nature (Figure 2K). Also, analysis of LdMVK model indicated the presence of three possible N-linked glycosylation regions and one O-linked glycosylation regions (Supplementary Figure 1).

**Enzymatic activity of r-LdMVK** was measured by methods described earlier (Ferreira et al., 2016; Duarte et al., 2018); by relating ADP release with oxidation of NADH by pyruvate kinase and lactate dehydrogenase. With increasing concentration of MVK, change in absorbance with respect to time increased linearly (Figure 2I). The specific activity of r-LdMVK was found to be 0.83 micromoles NADH/min/mg. Thermal stability was also studied and r-LdMVK was functional over broad range of temperature: 25–60°C and exhibited 33% of its optimal activity at 60°C. It was found to be temperature resistant as it retains its activity...
at 60°C (Figure 2J). All the enzymatic activity studies were carried out in glycine buffer. These results represent first step toward understanding of properties of MVK in *Leishmania*.

Subcellular Localization of MVK in Promastigote and Amastigote Forms

To check the localization of MVK in both promastigotes and amastigotes, confocal laser scanning microscope (LSM880 zeiss) was used. Immunofluorescence detection of MVK probed with FITC-conjugated secondary antibody (1:200 dilution; Jackson ImmunoResearch; #AB_2337972) showed a glycosome-like punctuate pattern which co-localized with glycosome specific pyruvate phosphate dikinase in promastigotes (Bringaud et al., 1998) which was probed with TRITC-conjugated anti-mice antibodies (1:200 dilution; Jackson ImmunoResearch; #AB_2337972). In amastigotes, MVK was observed to be present in glycosome, nucleus as well as concentrated on the cell membrane (Figure 2L). Three spots selected on promastigote and amastigote were shown to have overlapping histogram of fluorescent signal from PPDK and MVK. This showed partial co-localization of MVK and PPDK.

MVK Secretion Is Time, Temperature, and pH Dependent

The exoproteome of parasite was examined for the level of MVK release after temperature stimulation (37°C) to mimic initial infection condition. Sixty min post heat stress (37°C), maximum amount of MVK was observed in the extracellular medium, which declined with time (Figures 3A, B), suggesting its probable involvement in initial stage of infection. To examine if the release pattern is related to protein function, time kinetics of enolase release was studied. *Leishmania* enolase facilitates invasiveness (Vanegas et al., 2007) and its release was maximum at 30 min post temperature stimulation which declined with time (Figures 3E, F). On the contrary, the release of pyruvate phosphate dikinase (PPDK), a gluconeogenesis pathway enzyme, increased linearly with time (Figures 3G, H). To our knowledge, this is the first report on the release of PPDK by *L. donovani*. *Leishmania* encounters heat shock and acidic environment as they enter mammalian host (37°C) through sand fly bite (26°C). Therefore, it was of interest to examine changes in release of MVK in response to such stimulus. Conditioned medium from *L. donovani* promastigotes incubated at different conditions were collected after 1 h incubation. Temperature stimulation at 37°C induced the release of MVK by ~4 times compared to parasites incubated at 25°C (Figures 3C, D). On the other hand, phagolysosome-like conditions (37°C and pH 5.5) increased the secretion by ~6.5 times in comparison to MVK secretion at 25°C (Figures 3C, D). For these experiments, intraacellular actin was used as control to ensure that the exoproteome was from equivalent number of cells (Figure 3I). Each sample was normalized against β-actin of parasite lysate.

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**FIGURE 1** | Presence of mevalonate kinase in *L. donovani* and its release in extracellular media. (A) PCR amplification was carried out using MVK gene specific primers and *L. donovani* Ag83 strain genomic DNA as template to reveal its presence in the parasite. Kb, kilobase; M, DNA ladder; 1, MVK-PCR product. Similar results were obtained from at least two independent experiments. (B) The PCR fragment amplified using MVK gene specific primers was sequenced using di-deoxy sequencing and full length MVK gene sequence was obtained. (C) Parasite was incubated in serum-free medium for 6 h at 37°C and culture supernatant was collected and concentrated. The concentrated supernatant consisting of released proteins was checked for the presence of MVK by mevalonate kinase assay. Decrease in absorbance due to NADH oxidation relates to increased MVK activity. Representative image of at least three independent experiments is shown. Blank assays were performed with centrifric concentrated incomplete Medium199.
Characterization of L. donovani mevalonate kinase.

(A) Figure showing confirmation of cloning after HindIII+BamHI double digestion of pET28a-MVK. HindIII+BamHI double digested product was run on agarose gel electrophoresis. M, DNA ladder; uncut, uncut pET28a-MVK; HindIII, HindIII digested recombinant plasmid; BamHI, BamHI digested recombinant plasmid; HindIII+BamHI, double digested HindIII+BamHI pET28a-MVK product. Representative of at least three independent experiments is shown.

(B) SDS-PAGE showing the expression of rLdMVK by induced and transformed BL21 cells. M, Protein marker; BL21, bacterial cell lysate; BL21-pET28a, pET28a transformed bacterial lysate; BL21-pET28a-MVK, pET28a-MVK transformed bacterial lysate; Ind. BL21-pET28a-MVK, pET28a-MVK transformed and induced bacterial lysate.

(C) Ni-NTA purified sample stained with Coomassie brilliant blue. This purified rLdMVK was used for all functional characterization experiments. rMVK: Sample from final purification step of Ni-NTA purified extract.

(D) Western blot confirming the expression of rMVK. MVK expression by induced and transformed BL21(DE3) cells was verified by western blot of transformed cell lysate using anti-his antibody. Ind. BL21, transformed induced bacterial extracts expressing r-LdMVK; unind. BL21, transformed uninduced bacterial lysate; BL21, untransformed BL21 cells.

(E) Validation of generated anti-MVK antisera. SLA and rMVK were probed with anti-MVK antisera; and pre-immune sera. M, protein marker; SLA, soluble Leishmania antigen; rMVK, recombinant mevalonate kinase; WB, western blot.

(F) Oucherlony test validating anti-MVK antisera. Line of precipitation denotes the line where rMVK and anti-MVK antisera meet and interacts. Line of precipitation was not observed with pre-immune sera.

(G) Western blot analysis of promastigotes conditioned medium (Cm) confirming secretion of MVK. Stationary phase promastigotes were washed three times in PBS, incubated for 6 h in serum-free M199 medium (1 x 10^8 parasites/ml), culture supernatant was concentrated 25 times using 10MWCO centricon, and the 6 h conditioned medium was loaded on SDS-PAGE gel. Protein transferred on PVDF membrane was probed with anti-MVK antibody at 1:1,000 dilution. 6 h Cm, Ld promastigotes 6 h conditioned medium; WCL, 3% whole cell lysate of promastigotes; Media, concentrated serum-free M199 medium; Media, concentrated serum-free M199 medium.

(H) Western blot analysis of amastigotes conditioned medium (Cm) confirming secretion of MVK. Stationary phase axenic amastigotes were incubated for 6 h in serum-free M199 medium (1 x 10^8 parasites/ml), culture supernatant was concentrated 25 times by 10MWCO centricon, and the 6 h conditioned medium was loaded on SDS-PAGE gel. Protein transferred to PVDF membrane was examined with anti-MVK antibody. Fat 1:1,000 dilution. 6 h Cm, Ld promastigotes 6 h conditioned medium; WCL, 3% whole cell lysate of promastigotes; Media, concentrated serum-free M199 medium.

(I) Specific activity of rLdMVK was determined with increasing MVK concentrations. Graph depicts a linear increase in MVK enzymatic activity with increasing concentration of rLdMVK.

(J) Graph depicting a decline in enzymatic activity of LdMVK on treatment at different temperatures (25 to 60°C).

(K) Candycane glycoprotein molecular weight standards (M1 and M2) were run on 12% SDS PAGE. Candycane glycoprotein molecular weight standards (Molecular Probes, Eugene, OR, USA) containing a mixture of alternate glycosylated and non-glycosylated proteins was separated by SDS-PAGE and stained by Pro-Q Emerald glycoprotein gel and blot stain kit, and it served as positive and negative control for the experiment. Staining with Coomassie Brilliant blue reveals all eight band; glycoprotein stainings shows four glycosylated proteins bands. LdMVK was run on 12% SDS-PAGE, oxidized using perchloric acid, and stained for carbohydrates (Emerald 300 Q glycoprotein stain; M1, Coomassie brilliant blue stained Candycane glycoprotein marker; M2, glycoprotein stained Candycane glycoprotein marker; SLA, soluble Leishmania antigen; rMVK, recombinant LdMVK; CBB, Coomassie Brilliant blue. LdMVK is co-localized with glycosomal marker, PPDK. Double immunofluorescence pictures taken using confocal microscope depicts that LdMVK is present in glycosomes of L. donovani promastigotes and glycosomes, membrane and nucleus of L. donovani amastigotes. Differential interference contrast (DIC); DAPI (blue); rabbit anti-LdMVK (green); rabbit anti-LmPPDK (red); merged image (merged); histogram: overlapping histogram of PPDK and MVK shows co-localization of both the protein. Images are representative of three independent experiments.
**FIGURE 3** | MVK secretion in *L. donovani* promastigotes is time, temperature, and pH dependent. Western blot of promastigote conditioned medium collected at different time points or different environment conditions and normalized with housekeeping gene (*L. donovani* actin). Stationary phase parasites (1 × 10⁹ cells) were washed three times in PBS and resuspended in pre-warmed serum free medium to final density of 1 × 10⁸ cells/ml. Parasites were stimulated for secretion at 37°C for 5 to 240 min. Conditioned medium consisting of extracellular proteins were probed with anti-MVK antibody (A), anti-enolase antibody (E), anti-PPDK antibody (G), and compared. Parasite lysate for each sample was probed with anti-Ld actin antibody that served as cell number control. Graphical representation of densitometry values of extracellular MVK / intracellular β-actin (B), extracellular enolase / intracellular β-actin (F), extracellular PPDK / intracellular β-actin (H) with respect to time. (C) Release of MVK was observed at different temperatures and pH conditions: 25°C and 7.4 pH; 37°C and 7.4 pH; and 37°C and 5.5 pH. Serum-free M199 medium was maintained at different temperatures and pH and parasites were incubated in it for 1 h. Conditioned medium was obtained, concentrated, and western blot was performed using anti-MVK antibody. (D) Graphical representation of densitometry values of extracellular MVK obtained from different conditions normalized with β-actin. (I) 3% whole cell lysates from all the conditions were probed with anti-actin antibody. Error bars in all graphs represent standard error from three independent experiments performed in triplicates. * p value < 0.05 is denoted by *, ** p value ≤ 0.01 is denoted by ** and *** p value ≤ 0.001 is denoted by ***.
r-LdMVK Stimulates PBMCs to Express Th2 Cytokine Profile

*Leishmania* manipulates host defense system for their own survival and targeting these immune components is a reliable method to monitor the disease. In the same queue we studied the potential of LdMVK in shifting immune axis in support of parasite survival and disease progression (Table 1). We have observed that the IL-10/IL-12 ratio was 6.4 ± 0.41 in the LdMVK treated PBMC, 0.5 ± 0.02 in soluble *Leishmania* antigens (SLA) treated PBMC, and 1.6 ± 1.1 in untreated PBMC obtained from healthy person (Figure 4A). Further, IL-4/IL-12 ratio was found to be 7.33 ± 0.8 in LdMVK treated cells, 0.6 ± 0.01 in SLA treated cells, and 1.4 ± 0.9 in untreated cells of a healthy person (Figure 4B). IL-4/IL-2 cytokine ratio in LdMVK treated cells was found to be 12.2 ± 2.6 compared to 4 ± 0.2 in untreated cells of healthy person (Figure 4C). These results suggest that MVK has an immunosuppressive function.

**LdMVK Has a Role in Phagocytosis**

Cell membrane adhesion assay using r-LdMVK and fixed PBMC-derived macrophages showed an increase in enzyme bound to cell surface of macrophage with increasing concentration up to 1 µg/ml (Figure 5Aa, b). No increase in bound enzyme was observed on increasing the concentration further. Hence this concentration of protein was used later in experiments. To validate the association of r-MVK with macrophage, r-MVK was incubated with PBMC-derived macrophages for 1 h and cell lysate was immunoblotted against anti-LdMVK antibody. LdMVK specific staining was observed in the macrophages confirming the association of LdMVK with host macrophages (Figure 5Ac). As discussed earlier, maximum MVK is released 1 h post temperature stimulation (Figures 3A, B), when most of the parasite enters macrophage. This led us to speculate that MVK may have a role in parasite internalization. PBMC-derived macrophages from healthy person were treated with 1 µg/ml r-LdMVK followed by incubation with promastigotes (Ag83 strain) for 4 h. It was observed that LdMVK treatment increased phagocytosis by ~1.8-fold (Figure 5B). Antibody inhibition using anti-LdMVK antibody prevented MVK induced parasite entry, affirming that it has a role in phagocytosis. For validation, we generated MVK-Overexpression and Vector control strains of *L. donovani* parasites (Figure 5C). Western blot of MVK-OE and VC parasite’s lysate with anti-MVK antibody showed two bands indicating that exogenous protein (GFP and MVK-GFP) was expressed. MVK was overexpressed by ~2.3-fold in MVK-OE parasites as shown by western blot experiments (Figure 5Cc). qPCR results indicated mean normalized Ct value of 1.06 and 1.18 for MVK-overexpressed and vector control strains respectively and a difference in mean Ct value of 3 (Figure 5Cd). Infection assay wherein PBMC-derived macrophages were treated with mutant strains of parasites for 4 h, demonstrated significant increase in number of internalized parasites (~1.4-fold) in MVK-OE strain compared to VC strain (Figure 5Ce).

**LdMVK Treatment Leads to Host Cell Signaling Proteins’ Phosphorylation**

To understand the mechanism behind the increased parasite entry and immunomodulation induced by LdMVK, its effect on phosphorylation status of key host cell signaling components was observed with respect to time (0 to 90 min). Phosphorylated ERK

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**TABLE 1 | Secreted cytokine values (pg/ml) of PBMC treated with r-LdMV, soluble *Leishmania* antigen (SLA), or lipopolysaccharide (LPS).**

|        | IFN-γ | IL-12 | TNFα | IL-2 | IL-10 | IL-4 |
|--------|-------|-------|------|------|-------|------|
| UN     | 112 ± 8 | 41 ± 8 | 69 ± 7 | 31 ± 1 | 126 ± 32 | 128 ± 4 |
| SLA    | 310 ± 19 | 96 ± 21 | 81 ± 24 | 34 ± 5 | 152 ± 13 | 182 ± 9 |
| LPS    | 109 ± 46 | 33 ± 9 | 161 ± 14 | 4 ± 4 | 195 ± 35 | 165 ± 21 |
| MVK    | 155 ± 61 | 29 ± 5 | 108 ± 5 | 18 ± 5 | 193 ± 44 | 213 ± 26 |

Unstimulated culture was used as a control.

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**FIGURE 4 | Secreted level of cytokines in macrophages after r-LdMVK stimulation.** (A) IL-10/IL-12, (B) IL-4/IL-12, and (C) IL-4/IL-2 cytokine ratio of PBMC of healthy control, cultured in the presence of r-LdMVK, soluble *Leishmania* antigen (SLA), or lipopolysaccharide (LPS). PBMC was assessed for their effect on various cytokines release: IFN, IL-12p70, TNFα, IL-10, IL-2, and IL-4. Levels of cytokines were measured by ELISA after 16 h treatment. Unstimulated culture was used as a control. The data represents mean±SE of three independent biological replicates (including three technical replicates). p value < 0.05 is denoted by *, p value ≤ 0.01 is denoted by ** and p value ≤ 0.001 is denoted by ***.
1/2 was shown to be upregulated more than two times after 15 min MVK treatment (Figures 6A, B). On the contrary, no significant change was observed in the phosphorylation status of p38 MAP kinase (Figures 6A, C). Cortactin, a key actin scaffold protein, was phosphorylated three times more at 90 min on MVK treatment (Figures 6A, D).

**DISCUSSION**

The characterization and role of individual secretory proteins of *Leishmania* has remained elusive since many years. Here we describe first time one such novel protein from *L. donovani*, the mevalonate kinase (MVK) in *Leishmania* infection. Our investigation demonstrated that LdMVK is secreted by the parasite during in vitro cultivation and plays an important role in VL pathogenesis by assisting in the initial phase of infection and contributing to the efficient entry of the parasites in the host cells. Our results also suggest that LdMVK is capable of inducing interleukin-4 and interleukin-10 secretion through subversion of host cell. Our work sheds some light on the role of parasite in the entry process which is thought to largely dependent on macrophage in *Leishmania* infection.

The present study confirmed the presence of MVK protein in both promastigote and axenic amastigote of *L. donovani*. Cloned, expressed, and purified recombinant LdMVK protein was found to be functional (0.83 µmol/min/mg), as assessed by in vitro assays using its synthetic substrate, mevalonic acid. On the contrary,
L. major MVK activity was found to be significantly low (20 pmol/min/mg) in another study (Sgraja et al., 2007). LdMVK modelled structure is predicted to have similar α/β-folding patterns as that of LmMVK. The reliability of the proposed model was also supported by the presence of both N- and O-glycosylation sites. The glycosylation sites that were predicted, recommended that glycans will fit into these sites without steric clashes, thus signifying LdMVK glycosylation probability. Further, r-LdMVK expressed in a bacterial system was studied and revealed the glycosylated nature of protein. It is now established that bacteria express glycoproteins and carry out both N-linked and O-linked glycosylation pathways with many commonalities with their eukaryotic counterpart (Nothaft and Szymanski, 2010) and both the N-linked and O-linked glycosylation pathways can modify multiple proteins. Considering the significance of glycosylation in Leishmania virulence and extracellular vesicle pathophysiology (Kielian et al., 1982; Wang et al., 1996; Escrevente et al., 2011; Staubach et al., 2012; Gerlach et al., 2013), it was thought that the glycosylation of MVK might assist it in performing cellular functions.

Partial co-localization of MVK with the glycosomal protein, PDPK in typical punctuate structures showed its glycosomal compartmentalization. This was in compliance with the previous studies that showed MVK in Leishmania donovani is glycosome residing (Jardim et al., 2018). Other work done in L. major, T. cruzi, and T. brucei also displayed colocalization with glycosomes (Carrero-Lerida et al., 2009; Ferreira et al., 2016). The compartmentation of environment-sensitive parts of metabolism, like MVK, within glycosome could help parasites to acclimatize to acidic pH in phagolysosome. In contrast to promastigotes, it was observed that MVK in amastigotes were concentrated at the periphery of the cells, indicating that probably secretion of MVK in amastigotes is higher than that in promastigotes. Similar result was observed while mimicking phagolysosome-like condition that led to significant increase in MVK secretion. However, this was not confirmed in our study.

Silverman et al. have suggested the release of whole glycosome or glycosomal cargo by parasites outside cells (Silverman et al., 2008). Also, metabolic enzymes are shown to be post-
transitionally modified, changing their location and allowing to phosphorylate protein substrates. MVK’s glycosylated nature, specific compartmentalization in glyosome, and its secretory nature in Trypanosoma cruzi (Kielian et al., 1982; Wang et al., 1996; Ferreira et al., 2016) embarked us to investigate if it is secreted by L. donovani too. Going through the published secretome data we found that mevalonate kinase protein is not listed in the L. donovani and L. infantum secretome profiles (Silverman et al., 2008; Douanne et al., 2020). But, the list also does not include some other known Leishmania secreted proteins like serine protease, histidine acid phosphatase, and glycoprotein 63 (Jaffe and Dwyer, 2003; Joshi et al., 2004; Choudhary et al., 2010a). This could be because of the stringency in data collection method to involve abundantly secreted proteins.

The secretion of MVK into extracellular medium was confirmed by mevalonate kinase enzymatic assay and western blot of culture supernatant obtained from both promastigote and axenic amastigote form. Obtaining amastigotes from infected macrophages led to death of more 5% of amastigotes. Death of amastigotes due to harsh processing conditions could release their intracellular contents outside the cell interfering with the secretory protein profile. Hence, axenic amastigotes were used in all the experiments. The size of secreted LdMVK was found to be the same as that of the cytosolic LdMVK, suggesting that the released forms are not proteolytically processed as in the case of MIC2 (Carruthers et al., 2000), MIC5 (Brydges et al., 2000), and glycosylphosphatidylinositol-anchored micronemal antigen (TgGAMA) (Huynh and Carruthers, 2016). It is well known that eukaryotes use classical secretion pathway for the distribution of proteins throughout cell and outside the cell. But in Leishmania, most reports assert that majority of secretory proteins lack a signal peptide (Silverman et al., 2008; Kima et al., 2010; Hassani et al., 2011). Though the secretory pathway in this parasite is not well studied, present understanding exhibit non-classically secreted modes: exosome, apototic bodies, and plasma membrane blebs, all involving microvesicles (Silverman et al., 2010; Hassani et al., 2011; Forrest et al., 2020). The absence of signal sequence in MVK and prediction as non-classically secreted protein by Secretome P 2.0 server, suggested that its release is by non-classical secretion pathway.

Most of the L. donovani promastigotes are internalized by 60 min (Chakrabarty et al., 1996). Maximum release of LdMVK, 60 min after heat treatment could be because of its involvement in initial phase of infection (host cell colonization). L. donovani enolase, that supports parasite entry in host cell also shows similar release pattern in our study, whereas the release of PPDK (unknown secondary function) was shown to increase with time. This is one of the few studies that described the dynamics of protein release with respect to time. It indicated that the amount of protein released outside cell is not just dependent on the environment but also on the time of infection. If the maximum release time of a protein is dependent on its function is yet to be studied.

It is known that Leishmania protein release involves functional enrichments based on temperature and pH variations (Silverman et al., 2010; Hassani et al., 2011). Heat shock for 24 h increase vesicle release by 3-fold and lowering the pH to 5.5 (phagolysosome condition), increase or decrease the secretion depending on the protein (Silverman et al., 2010). In present study, MVK release was found to be increased by ~4.2-fold on heat stress (37°C) after 1 h stress treatment in comparison to secretion at 25°C. The temperature induced increase in MVK level must have also been contributed by the global increase in protein level. But, the fold increase value of 4.2 is high and suggests that all of this increment must not have been a part of global increase. Moreover, though acidic pH does not change the bulk of protein release, it led to 6.5-fold change in MVK expression. In extreme acidophiles, an adapted mevalonate pathway is followed, signifying MVK’s involvement in sustaining life in extremely acidic environments (Vinokur et al., 2016) which could be the case in Leishmania as well.

A protein can have one function within the cell and another outside it (Jeffery, 1999). A growing number of secretory proteins have been shown to be involved in virulence (Table 2). MVK’s partial localization outside glyosome and glycosylated nature could assist in performing an extra function beyond ergosterol synthesis. Besides, it’s different oligomeric state in L. major could be used to switch between functions. Thus, to study if MVK performs additional roles apart from ergosterol synthesis in parasite, its role in disease pathogenesis was studied.

Cytokine response can give a broad prospect of the complicated interaction between host and parasite. Lipopolysaccharide (LPS) induction leads to reduced TH1 response (Lauw et al., 2000). Its exposure increases TNF-ß and interleukin-10 production significantly; and the release of interferon-γ and interleukin-12 remains unaffected (Jansky et al., 2003). Soluble Leishmania antigen (SLA) treatment on the other hand, is considered as good control for Leishmania specific TH1 response (Hailu et al., 2005) and triggers interferon-γ and TNF-α producing cells. Similar results were observed in our study when PBMC were treated with LPS and SLA. Treatment with MVK triggered interleukin-10 and interleukin-4 secretion which is indicative of disease susceptibility (Bhattacharya et al., 2001). Interleukin-10 plays an important role in pathogenesis by inhibiting TH1 mediated response, activation of macrophages, and antigen presentation (De Medeiros et al., 1998; Nylen and Sacks, 2007). Increase in the level of interleukin-4 trigger macrophage in a different manner; and assist in polynyme synthesis, assisting in growth and survival of parasites (Bhattacharya and Ali, 2013). Unexpectedly, there was an increase in the level of interferon-γ release also, though insignificant. It has been shown by Murphy et al. that abundance of interleukin-10 is more important than the level of interferon-γ (Murphy et al., 2001); and interleukin-10 resists interferon-γ induced macrophage activation. Reduced IL-12 secretion in response to MVK treatment would delay TH1 cells development allowing parasite to transform to withstand adverse macrophage conditions. The TH1/TH2 balance is a determining factor of the consequence of leishmaniiasis. Anti-inflammatory to pro-inflammatory cytokines ratios (IL10/IL-12 ratio; IL4/IL-12 and IL-4/IL2 ratio) were found to be significantly higher in LdMVK-treated macrophage compared to untreated and SLA-treated cells. Such high ratios during Leishmania
infection, specify Th2 dominance, host immunosuppression, and disease progression. Our results are concurrent with the previous studies where mice treated with \textit{L. major} excretory secretory proteins released more IL-4 and IL-10 thereby enhancing parasite survival and disease progression (Tonui et al., 2004).

Hence, LdMVK can be among the proteins responsible for transient immunosuppression, favoring parasite internalization inside the host cells.

Earlier observations indicates that proteins involved in parasite host interactions usually bind to the host cell surface.
To survive inside macrophages, parasite modulates host cell signaling pathways and its antagonistic and synergistic molecular actions decides the fate of parasite. *Leishmania* causes reciprocal regulation of ERK-1/2 and p38 (Mathur et al., 2004). ERK-1/2 pathway activation causes increased IL-10 production, induces Th-2 type immune response, and parasite survives (Bhardwaj et al., 2010). p38 MAPK activation leads to IL-12 production, guides Th-cell differentiation into Th1-type cell in the favor of host (Dong et al., 2002; Arthur and Ley, 2013). Since, protein phosphorylation is the major regulatory mechanism (Ptacek et al., 2005), phosphorylation kinetics of ERK-1/2 in rLdMVK exposed macrophages was measured and was found to be upregulated transiently post-MVK incubation. Similar results were seen in LPG (Balaraman et al., 2005) and *L. donovani* (Soares-Silva et al., 2016) incubated macrophages. The MKV-induced change in ERK1&2/p38 expression reflects in the high IL-10/IL-12 ratio post MKV treatment, skewing the T cell response towards Th-2 type and in the favor of parasites. ERK also regulates the activation of proteins involved in microfilament remodeling like Cortactin (Navratil et al., 2014) and have also been linked with increased invasion in *T. cruzi* infection (Magdesian et al., 2007). Furthermore, increase in phosphorylation status of cortactin on MKV induction indicates possible involvement of MKV in actin polymerization, since cortactin is the key scaffold for actin regulation (Han et al., 2014). This gives an idea behind the mechanism involved in MKV-mediated parasite internalization and immunosuppression.

We have not shown direct activation of ERK by rLdMVK. MKV mediated indirect phosphorylation of ERK through ERK kinase is also possible. Hence, MKV, apart from phosphorylating the metabolite: mevalonate, might also acts like a protein kinase and phosphorylate ERK-1/2 and cortactin either directly or indirectly. There are several reports of metabolic kinases moonlighting as protein kinases (Lu and Hunter, 2018). Current studies cannot explain how the active site of a metabolic kinase responsible for recognizing a small metabolite and ATP/ADP, can also recognize Ser, Thr, or Tyr residues of protein and phosphorylate it and it needs further investigation. A co-crystal structure of mevalonate kinase bound to its protein substrate could define how an amino acid in the protein substrate can fit in the active site.

Extracellular secretion of proteins is one of the mechanisms of virulence in *Leishmania*. MKV has to cross two membranes through their journey into phagolysosome. Its secretion from *Leishmania* glycosome occur through non-classical mode of secretion in the form of microvesicles or whole glycosome. It then binds to the membrane of macrophage and is internalized before the internalization of parasite. Once in cytosol, MKV mediates the phosphorylation of host cell signaling pathway proteins. This could lead to cortical actin polymerization, formation of actin rich structure: pseudopodia that would internalize parasite through phagocytosis. MKV-mediated activation of host cell proteins increases and decreases the secretion of interleukin-10 and interleukin-12 respectively. This could help in evasion of the immune system by parasite, facilitating their persistence in host (Figure 7).

Together, in this study, we have identified and characterized a novel *Leishmania* secretory protein, MKV, that is involved in internalization of parasite and immunosuppression. It can be predicted that once *Leishmania* sense the changes in environment, the release of MKV is increased that facilitates parasite entry and induce immune suppression through MKV-mediated phosphorylation of host cell components. Overall, these results will improve our understanding on the capabilities of secretory proteins in internalization of parasite and host immunomodulation which may be further explored for their potential importance in diagnosis and immunoprophylaxis.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Institutional Human Ethics Committee (IHEC). The patients/participants provided their written informed consent to participate in this study. The animal study was reviewed and approved by Institutional Animal Ethical Committee, affiliated by
The Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA).

AUTHOR CONTRIBUTIONS

Conceptualization: TB and PD. Methodology: TB and AS. Validation: MK and AjK. Formal analysis: TB, MD, and KA. Investigation: TB, TS, and AsK. Writing—original draft preparation: TB. Writing—review and editing: TB, SD, AS, and PD. Supervision: PD. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2021.641985/full#supplementary-material

Supplementary Figure 1 | Various parameters for structure validation. (A) Structure of LdMVK depicting α-helix, β-sheet, and turn regions illustrated in distinct colors, and representing three predicted N-linked glycosylation sites (N130, N137, N222), and one O-linked glycosylation site (S228). (B) Stereochemical properties of the homology model were investigated in Ramachandran plot using PROCHECK analysis. (C) ProSA was employed to assess the 3D structure model of protein for errors. Graph represents score plot acquired through ProSA web server. (D) The refined model was confirmed by Verify-3D profile analysis method.

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