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Characterization of Trichomonas vaginalis Survival Factor under Nutrient Starvation and A Protein That Mediates Parasite Host Cell Binding and Killing

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Characterization of *Trichomonas vaginalis* Survival Factor under Nutrient Starvation and A Protein That Mediates Parasite Host Cell Binding and Killing

A dissertation submitted in partial satisfaction of the requirements for the degree Doctor of Philosophy in Molecular Biology

by

Yi-Pei Chen

2018
Characterization of *Trichomonas vaginalis* Survival Factor under Nutrient Starvation and A Protein That Mediates Parasite Host Cell Binding and Killing

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Yi-Pei Chen

Doctor of Philosophy in Molecular Biology

University of California, Los Angeles, 2018

Professor Patricia J. Johnson, Chair

*Trichomonas vaginalis* is a unicellular extracellular sexually transmitted parasite. While the infection may be cleared by nitroimidazole drugs, in cases where re-infection occurs or the treatment fails, the infected individual may live with inflammation, soreness, pain or itch surrounding urogenital areas without an alternative solution. To establish an infection, *T. vaginalis* attaches to the host and acquires nutrients from the host. Several parasite surface molecules and its secreted exosomes have been shown to be important for the parasite attachment to the host. However, none of the factors identified is solely responsible for host cell binding. In this study, we characterize a parasite surface protein TVAG_393390 (or cadherin-like protein) that significantly increases host binding and killing when it is overexpressed. In addition, to also understand how the parasite survives in the host under nutrient-deficient
conditions, we unravel the role of a survival protein in the parasite (TvMIF) that is homologous to human macrophage migration inhibitory factor (huMIF). We found that TvMIF-overexpressing parasites gain a strong survival advantage under nutrient stress. To obtain further insights into functions of parasite proteins, we adapted gene knockout method CRISPR (cluster regularly interspaced palindromic repeat)-Cas9 (CRISPR-associated protein 9) in T. vaginalis to efficiently knock out genes of interest. We then successfully knocked out TvMIF gene using CRISPR-Cas9 and observed a significant reduction in the survival of TvMIF knockout cells compared to the wild-type when these parasites were serum-starved. Last, we examine the human innate immune responses to T. vaginalis encounter. Human macrophages and dendritic cells produce strong inflammatory responses when T. vaginalis and Mycoplasma hominis are both present but remarkably less when M. hominis is absent. Together, these data reveal several independent mechanisms which allow T. vaginalis to successfully establish infection in human bodies and provide a framework for future studies on the “trichy” parasite.
The dissertation of Yi-Pei Chen is approved.

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2018
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In Chapter 3, the project involves an important T. vaginalis protein selected by Dr. Angelica Riestra from her former studies. She identified the key sites for the protein function and established its importance in parasite binding and host killing. Without her, the project would not have developed.
Throughout my PhD career, Dr. Brian Janssen has given countless ideas and suggestions that guided me through obstacles in my projects. Furthermore, his dedication to CRISPR-Cas9 development significantly advanced the knockout method in the *T. vaginalis* field which is described in Chapter 4. In Chapter 5, Dr. Frances Mercer has offered her expertise for working with immune cells and examining cytokine production.

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Chapter 1:

Introduction
*Trichomonas vaginalis* is the causative agent of the most common non-viral sexually transmitted infection, trichomoniasis, both in the United States and the world [1]. Trichomoniasis has been classified as a neglected parasitic infection for being underestimated for its public health impact and disproportionate effects on minorities [2]. In fact, *T. vaginalis* infection outnumbers the total infection numbers of *Neisseria gonorrhoeae, Chlamydia trachomatis* and syphilis combined [1]. Poor recognition of the disease has been contributed to 70-85% asymptomatic nature of the infection [3]. However, in the symptomatic cases, mild to moderate levels of inflammation in the men and women’s urogenital tracts is observed [2][4]. The symptoms include pain during sex and urination, lesions of urogenital epithelium, vaginal discharge, unpleasant smell, itching and irritation [2][3]. In worse scenarios, preterm labor [5], increased risks of other sexually transmitted diseases including human immunodeficiency virus (HIV) [6], and higher risks of cervical cancer [7] and extraprostatic cancer development have also been reported [8].

The current treatments for trichomoniasis are two different nitroimidazole drugs, metronidazole and tinidazole. Metronidazole is prescribed first and if the treatment fails, tinidazole with longer half-life is then prescribed [9][10]. Clinical data have recorded failed treatment using both nitroimidazole drugs [11]. In addition, drug-resistant parasites are reported to be on the rise [12]. The mode of actions of both drugs are not fully understood but evidence suggests that the nitroimidazoles form adducts with proteins and damage of the proteins induced by drug toxicity results in *T. vaginalis* killing [13]. The nitro functional group for drug activation of the two 5-nitroimidazoles is identical so it is likely that the parasite strains resistant to metronidazole are also resistant to tinidazole. Successful treatment in patients does not prevent
them from being re-infected. As a result, there is a need to develop novel treatments for trichomoniasis.

**Biology of *T. vaginalis* survival mechanisms under stress**

Understanding the parasite biology is crucial to drug development. For example, how the parasite survives in the constantly changing vaginal environment is poorly understood [10]. Targeting parasite survival factors can be used to prevent trichomoniasis. As an obligatory human parasite, *T. vaginalis* acquires nutrients from human host in order to survive and grow [10]. For example, *T. vaginalis* has high demand for iron uptake [14]. Iron-containing proteins from humans such as hemoglobin provide an iron source for *T. vaginalis* growth and its iron-containing enzymes [15][16]. Under iron-deficient condition, *T. vaginalis* accumulates nitric oxide to maintain its survival [17]. Glucose levels of vaginal secretions from *T. vaginalis*-infected patients reveal that the glucose concentrations can vary between 0.3 and 36.65 mM [18]. Under glucose restriction, *T. vaginalis* displays autophagic behavior and increases expressions of anti-oxidant genes which result in induced resistance to hydrogen peroxide [19]. Programmed cell death (apoptosis) in a protozoan parasite such as *T. vaginalis* has been speculated to be a mechanism to enhance population fitness under these stressful conditions [20] or reduce a potential risk of causing fatal effects on the host if the parasites overgrow [21]. Regulated death may prevent release of toxic elements from dying parasites [20] as we know in mammalian systems, necrosis releases a protein that can trigger inflammation [22]. Apoptosis has been described in *T. vaginalis* when treated with apoptotic inducers [23]. Drugs-induced apoptosis shows DNA condensation and fragmentation, externalization of phosphatidylserine on cell surface, and dissipation of hydrogenosomal (mitochondria-related organelle) membrane potential [23]. A closely related species *Tritrichomonas foetus* also demonstrates DNA condensation and
fragmentation when treated with hydrogen peroxide [24]. In addition, *T. foetus* exhibits vesicular formation and reaction with anti-human/mouse-caspase-3 antibody with immunocytochemical staining [24]. However, there is no homologous caspase 3 present in *T. foetus* or *T. vaginalis* [24]. This could mean the active sites of the caspase are conserved in *T. vaginalis* and can be recognized by the human/mouse antibody but the rest of the protein is too dissimilar to be identified by analyzing the primary sequence with BLAST. It is notable that metacaspases which are believed to have the same role as mammalian caspases in yeast are also present in protozoan parasites but their specific roles in parasites are not entirely understood [25][26]. Other conserved apoptotic players such as endonuclease G, caspase-activated DNase or inhibitors of caspase-activated DNase in other eukaryotic systems are not identifiable by searching the *T. vaginalis* genome [21]. Apoptosis in *T. vaginalis* presents some similar features as in multicellular eukaryotes but the mechanism and the players involved are likely to have diverged.

**Biology of *T. vaginalis* adherence mechanisms**

*T. vaginalis* is an extracellular parasite. The biology of *T. vaginalis* factors involved in parasite binding to its host is critical for understanding how the parasite establishes the infection. Killing of host epithelial cells by *T. vaginalis* is contact-dependent [27]. The differences between 26 examined *T. vaginalis* strains in host cell adherence and killing are up to 45-fold and 96-fold, respectively [27]. The drastic differences between *T. vaginalis* strains allow us to study the parasite factors involved in host binding and killing. A *T. vaginalis* surface proteome comparing 3 adherent strains and 3 less adherent strains revealed proteins that are more abundant in the adherent strains relative to the less adherent strains that may be involved in attachment [28]. The main polysaccharide lipoglycan (LG) found on *T. vaginalis* surface was found to be involved in host binding [29][30]. LG mutant parasites created by chemical mutagenesis significantly
reduced adherence to host and LG isolated from the parasites competes with wild-type parasite binding to host [30]. Other T. vaginalis factors such as laminin-binding proteins [31], adhesins [32][33][34] and cysteine and serine proteinases [35][36][37][38] have also been found to play a role in host binding. Recently, our laboratory demonstrated that parasite-derived small, secreted vesicles called exosomes mediate host attachment via both host:host and host:parasite interactions [39][40][41]. By pre-incubating either the parasites or the host, or both with exosomes from a highly adherent strain increases parasite adherence to host in less adherent strain [39]. In this dissertation, another T. vaginalis surface protein that plays an important role in adherence and host killing will be described in Chapter 3.

**Host immune responses to T. vaginalis**

Immune responses from human cells to T. vaginalis have been characterized to understand why chronic infection and re-infection can occur. Antibodies against T. vaginalis have been reported to be present in infected patients [42][43]. Our laboratory has found that primary human T cells and B cells are killed by a clinical T. vaginalis strain but much less so by a laboratory adapted strain and T. vaginalis [44]. Phagocytosis of leukocytes by T. vaginalis [45] and cysteine proteinases that can degrade host antibodies have been described [46]. Killing of leukocytes by T. vaginalis could be one of the reasons why the parasite is able to maintain a chronic infection. We and others found that only low levels of IL-8 are secreted by human monocytes by T. vaginalis alone, either laboratory adapted or clinical strain, but the presence of T. vaginalis with its symbiotic bacteria Mycoplasma hominis results in significant increase in IL-8 production and induction of IL-6 and IL-1β that are not detectable in M. hominis-free samples [44][47]. T. vaginalis and M. hominis co-infection occurs at 5-90% rates. The dramatic differences in co-infection rate depends largely on the sources of the samples [48] and the
dissimilarity between *M. hominis*-free and *hominis*-containing cytokine responses could be responsible, at least in part, for the diverse infection outcomes.

*T. vaginalis* exosomes also induce IL-6 and IL-8 responses from host ectocervical cells (Ects) [39]. In addition, IL-8 cytokine induction is lessened when Ects are pre-incubated with *T. vaginalis* exosomes, a potential mechanism that the parasite employs to minimize the host immune response by priming the host with the parasite exosomes [39]. Olmos-Ortiz et al examined other immunomodulatory roles of *T. vaginalis* exosomes [49]. They found that *T. vaginalis* exosomes induced IL-10, IL-6 and TNF-α from mouse macrophages and in their mouse model, pretreatment with the exosomes significantly reduced the inflammatory responses [49], reminiscent of our in vitro study [39]. Identification of a homologue of human cytokine macrophage migration inhibitory factor in *T. vaginalis* (TvMIF) reveals a secretory factor which induces IL-8 production from human monocytes [50]. Isolated LG from the parasite surface is also found to induce a large amount of IL-8 and macrophage inflammatory protein 3α, and mild IL-6 responses from Ects [51]. However, whether *M. hominis* is present or not in the LG preparation was not addressed. Revisiting the story with the *M. hominis*-containing and *M. hominis*-free controls may be necessary to parse the true immunomodulatory roles of LG.

Additionally, *T. vaginalis* contains endosymbionts that are double-stranded RNA (dsRNA) viruses called *T. vaginalis* viruses (TVVs) [52]. A TVVs-containing strain upregulates Toll-like receptor 3, and also induces IL-8, IFNβ, Regulated upon Activation, Normal T-cell Expressed, and Secreted (RANTES), IL-1 and MIP-3α production in host cells while a TVVs-free strain does not [53]. *T. vaginalis* and its symbionts have multiple factors that could elicit host immune responses and more may yet to be discovered.

**Lacking genetic tools to study *T. vaginalis* biology**
Despite the interesting findings revealed previously by using molecular biology, biochemical and other tools, there are numerous technical limitations that restrict researchers in the field from better understanding *T. vaginalis* biology. Introducing DNA to *T. vaginalis* using electroporation was demonstrated more than two decades ago [54] but only ~3% of the parasites obtain the plasmids using this method and thus, selection with a drug-resistant marker is required [55]. In addition, using immunofluorescent assays to detect multiple overexpressed proteins in the parasites in separate experiments in our laboratory suggests that the existing transfection method results in highly variable protein expression levels within a population (Fig. 1-1). This renders characterization of protein function ambiguous. Large gene family amplification, abundant repetitive elements in the genome [56] and low transfection efficiency has impeded the development of gene knockout tools in *T. vaginalis*. Prior to the development of CRISPR (cluster regularly interspaced palindromic repeat)-Cas9 (CRISPR-associated protein 9) in *T. vaginalis*, only two separate knockouts have been accomplished using traditional homologous recombination method [57][58]. In this dissertation, adaptation and employment of CRISPR-Cas9 in our laboratory to knock out *T. vaginalis* genes will be described as a much more efficient method than traditional homologous recombination [55].

Due to low success rate of gene depletion in *T. vaginalis*, knockdown of transcripts of gene in the parasites was used as an imperfect alternative. MicroRNAs (miRNAs) and its machinery have been identified in *T. vaginalis* [59]. An endogenous miRNA and introduction of its mimics reduced *T. vaginalis* malate dehydrogenase protein expression by 60% [60]. Use of synthetic small interfering RNAs (siRNAs) reduced the transcripts of two genes by 48-67% and 33-72% [61]. Antisense targeting a transcription factor Myb3 [62], a metabolic enzyme glyceraldehyde-3-phosphate (GAPDH) [63], and two surface proteins involved in parasite
adherence, AP33 [33] and AP65 [34], and a major serine/threonine protein phosphatase called protein phosphatase 1 gamma [64] have been reported separately. Despite multiple attempts on knocking down genes in *T. vaginalis* were described, experiences in our laboratory revealed that knocking down genes is unreliable and the results are inconsistent (unpublished data). The presence of residual gene expression in knockdown parasites also prevents full characterization of a protein. Hence, efforts towards development of a reliable knockout system in *T. vaginalis* became essential.

The goals of my dissertation are to focus on understanding *T. vaginalis* proteins and host responses in shaping how the parasite becomes a “successful” parasite. First, a *T. vaginalis* protein called TvMIF is a survival factor for the parasite to survive in the constantly changing host environment [10]. How TvMIF enhances the parasite survival is described in Chapter 2. Chapter 3 focuses on a different protein called TVAG_393390 (or cadherin-like protein) which plays a critical role in parasite binding to and killing of host cells. To better understand parasite proteins and their function, we adapted CRISPR-Cas9 for knocking out genes in *T. vaginalis*, as discussed in Chapter 4. In Chapter 5, we shifted our focus from parasite factors to host immune responses to the infection to understand the other side of the story in this host: pathogen relationship.
Figure 1-1: Immunofluorescent assay on TvMIF-overexpressing cells with anti-HA and DAPI staining to show variations of exogenous protein levels between cells. Standard transfection of *T. vaginalis* using G418-selection to exogenously express an epitope-tagged protein shows expressions of the protein are highly variable. Parasites shown here are expressing TvMIF with HA tags. The white arrow denotes an example of a parasite nucleus stained with DAPI (blue) but has almost no exogenous expression of the TvMIF protein (red) within a transfection population. The yellow arrow denotes a parasite with high expression of exogenous TvMIF protein. Red: anti-HA. Blue: DAPI.
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protozoan parasite Trichomonas vaginalis is involved in proliferation and cell attachment to the host cell. *Int J Parasitol* (42) 715–727.
Chapter 2:

*Trichomonas vaginalis* macrophage migration inhibitory factor mediates parasite survival during nutrient stress
Trichomonas vaginalis Macrophage Migration Inhibitory Factor Mediates Parasite Survival during Nutrient Stress

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ABSTRACT Trichomonas vaginalis is responsible for the most prevalent non-viral sexually transmitted disease worldwide, and yet the mechanisms used by this parasite to establish and maintain infection are poorly understood. We previously identified a T. vaginalis homologue (TvMIF) of a human cytokine, human macrophage migration inhibitory factor (hMIF). TvMIF mimics hMIF’s role in increasing cell growth and inhibiting apoptosis in human host cells. To interrogate a role of TvMIF in parasite survival during infection, we asked whether overexpression of TvMIF (TvMIF-OE) confers an advantage to the parasite under nutrient stress conditions by comparing the survival of TvMIF-OE parasites to that of empty vector (EV) parasites. We found that under conditions of serum starvation, overexpression of TvMIF resulted in increased parasite survival. Serum-starved parasites secrete 2.5-fold more intrinsic TvMIF than unstarved parasites, stimulating autocrine and paracrine signaling. Similarly, we observed that addition of recombinant TvMIF increased the survival of the parasites in the absence of serum. Recombinant hMIF likewise increased the parasite survival in the absence of serum, indicating that the parasite may use this host survival factor to resist its own death. Moreover, TvMIF-OE parasites were found to undergo significantly less apoptosis and reactive oxygen species (ROS) generation under conditions of serum starvation, consistent with increased survival being the result of blocking ROS-induced apoptosis. These studies demonstrated that a parasitic MIF enhances survival under adverse conditions and defined TvMIF and hMIF as conserved survival factors that exhibit cross talk in host-pathogen interactions.

IMPORTANCE Macrophage migration inhibitory factor (MIF) is a conserved protein found in most eukaryotes which has been well characterized in mammals but poorly studied in other eukaryotes. The limited analyses of MIF proteins found in unicellular eukaryotes have focused exclusively on the effect of parasitic MIF on the mammalian host. This was the first study to assess the function of a parasite MIF in parasite biology. We demonstrate that the Trichomonas vaginalis MIF functions to suppress cell death induced by apoptosis, thereby enhancing parasite survival under adverse conditions. Our research reveals a conserved survival mechanism, shared by a parasite and its host, and indicates a role for a conserved protein in mediating cross talk in host-pathogen interactions.

KEYWORDS Trichomonas vaginalis, apoptosis, macrophage migration inhibitory factor, nutrient starvation

Trichomonas vaginalis, an extracellular unicellular parasite, causes the most common non-viral sexually transmitted infection in the world (1) but has been long neglected. Thus, the mechanisms that drive parasite pathogenesis and the disease epidemiology are poorly understood (2). However, this is changing as more molecular and genetic tools for analyses are developed (3, 4). The majority of T. vaginalis
infections are asymptomatic. When they are symptomatic, the manifestations of infection vary greatly and may include inflammation of the urogenital tract, preterm delivery, and increased chances of HIV co-infection (5, 6). The factors that determine how the parasite maintains the infection in the vaginal microenvironment, where the nutrients, hormones, and pH are constantly changing, remain largely unknown (7–9).

Macrophage migration inhibitory factor (MIF) is a highly conserved eukaryotic protein found across unicellular protists, plants, arthropods, and mammals (10–13). Human MIF (humMIF) has been widely studied and is known to play essential roles in cell growth, survival and in cancer growth in humans (14–16). In our previous study, we reported that T. vaginalis shares a homologous protein (TvMIF) with the human host and that TvMIF can activate the same survival pathways as huMIF in human cells (17). MIF homologues found in other eukaryotic parasites are known to modulate the host immune system and to activate huMIF pathways (10, 18–21). Although the effects of parasite-derived MIFs from parasites on host cells have been studied, the role of MIF in parasites and non-mammalian systems is poorly understood.

T. vaginalis and other parasitic protists share certain apoptotic phenotypes with mammalian systems such as DNA fragmentation and phosphatidylserine exposure (22–25). However, no factor that either stimulates or suppresses apoptosis in these divergent, unicellular eukaryotes has yet been identified. In this report, we describe the anti-apoptotic effect caused by TvMIF, a conserved eukaryotic protein, and reveal similarities between this protein and its mammalian homologue. We also provide evidence that T. vaginalis is able to exploit huMIF to enhance its survival during nutrient starvation. These studies uncovered a highly conserved eukaryotic protein used by a parasite and its host to enhance survival.

RESULTS

T. vaginalis MIF (TvMIF) enhances survival of the parasites under conditions of nutrient stress. We have previously shown that the homologue of huMIF in the parasite T. vaginalis (TvMIF) can induce the growth and activation of anti-apoptotic pathways in human host cells (17). To investigate whether TvMIF can perform a similar function and hence enhance the survival of the parasite under adverse conditions, we have studied the role of TvMIF in parasite survival under conditions of nutrient and density stress. T. vaginalis is typically grown in Diamond’s media supplemented with serum (26). The parasite will not grow in the absence of serum as it provides lipids, precursors of nucleotides, and amino acids required for parasite survival (26, 27). Parasites also cease to swim and die within hours after reaching a density of $5 \times 10^7$ cells/ml in Diamond’s media supplemented with serum (27). As a first step toward determining whether TvMIF plays a role in parasite survival, we compared the survival rates of parasites transfected with (28) and overexpressing TvMIF (TvMIF-OE) with those transfected with an empty vector (EV). Immunoblotting confirmed that expression of TvMIF in TvMIF-OE parasites is approximately 17-fold greater than in EV control parasites (see Fig. S1 in the supplemental material).

Fluorescence-activated cell sorter (FACS) analyses of living cells assessed using Zombie Red viability dye to compare TvMIF-OE and EV parasites were conducted using parasites grown in serum-free media or at a high density. We found that TvMIF-OE parasites survived significantly better than EV parasites in serum-free media after 8 h of incubation and that this survival phenotype became more pronounced after 24 to 32 h of starvation (Fig. 1A; see also Fig. S2). Comparison of the survival rates of TvMIF-OE and EV parasites at densities as high as $10^7$ cells/ml or $2 \times 10^7$ cells/ml also revealed the death of a significantly higher proportion of EV parasites than of TvMIF-OE parasites at 4 and 8 h after they were subjected to density stress (Fig. 1B). When grown in regular media supplemented with serum, EV and TvMIF-OE had similar growth rates (Fig. S3), demonstrating that TvMIF promotes parasite survival under adverse conditions.

TvMIF is secreted as both a free soluble protein and an exosomal protein. We have previously demonstrated that T. vaginalis secretes small vesicles called exosomes (29, 30) and that TvMIF is found in the exosomal proteome (31). As the survival
pathways activated by huMIF require a secreted soluble form of the protein (16, 32), we examined both the exosomal fractions (Exo) and the non-exosomal soluble fractions (NESF) secreted by the parasite for TvMIF using a Vivaflow crossflow cassette and an ultracentrifugation-based method (Fig. 2A). Parasites grown overnight were collected and then incubated in phosphate-buffered saline (PBS)–5% sucrose at either 16°C (negative control for secretion) or 37°C. After 2 h of incubation, the cells were pelleted and lysed to make whole-cell lysates (Wcl). The supernatant was passed through a Vivaflow crossflow cassette using a 100-kDa molecular weight cutoff (MWCO) to separate the heavier exosomes (Exo > 100 kDa) from the lighter non-exosomal secreted fraction (NESF < 100 kDa). The level of NESF TvMIF was found to be 1.4-fold higher than that of Exo TvMIF by immunoblotting using an anti-TvMIF antibody (4) (Fig. 2B). As the Exo fraction had much less protein than the Wcl fraction and NESF from equal numbers of cells, equal (20 μg) amounts of protein were loaded in each fraction. At 37°C, 20 μg of protein is equal to 2.87% of the total NESF and 17.2% of the Exo fraction. As a result, we can reason that approximately 8.4-fold-more TvMIF is secreted in the NESF than in the Exo (17.2% divided by 2.87% and then multiplied by 1.4-fold).
FIG 3. Endogenous TvMIF protein is induced under conditions of serum starvation. (A) Anti-TvMIF immunoblot showing that the TvMIF level was induced during serum starvation. Detection of GAPDH served as the loading control. One representative immunoblot of three independent experiments is shown. (B) Quantification of TVMIF was determined by normalizing the TvMIF signal to GAPDH for each sample; the data were compared to the 0-h time point value (set at 1) in each experiment, so no error bar was made for 0 h. Data are means ± standard errors. (C) TvMIF secretion was induced in the absence of serum. + Serum, cells grown in complete media; − Serum, cells grown without serum. Secreterd TvMIF was induced ~2-fold at 3 h and ~2.5-fold at 6 h after serum starvation comparing − serum and + serum signals. Neo was used as a negative control for cell lysis.

These data show that the majority of total secreted TvMIF is present as a soluble protein.

Both intracellular TvMIF and secreted TvMIF are induced during serum starvation. As shown in Fig. 1, overexpression of TvMIF increased the survival of the parasites during nutrient stress. Thus, we asked if induction of TvMIF expression and its secretion play a role in the enhanced survival. First, to test if intracellular TvMIF expression is induced under conditions of serum starvation, we collected the whole-cell lysates (Wcl) from cells starved for 16 or 24 h and TvMIF levels were quantified by immunoblotting. Intracellular TvMIF was found to be induced 1.6-fold at 16 h and 2.2-fold at 24 h after starvation (Fig. 3A and B). For determining TvMIF secretion levels under conditions of serum starvation, parasites grown overnight in complete Diamond’s media were collected, resuspended, and incubated in serum-free media for 3 h or 6 h at 37°C. Cells were then spun, resuspended in PBS–5% sucrose, and incubated for an additional 2 h at 37°C. Cells were then pelleted and lysed to make Wcl, and the supernatant was collected to assess secretion. Immunoblot analyses showed that the secreted TvMIF were induced ~2-fold and 2.5-fold higher from cells incubated in serum-free media for 3 and 6 h, respectively, than from parasites grown in serum-containing media (Fig. 3C). Neomycin phosphotransferase (Neo), a non-secreted protein, was used as a control to monitor cell lysis. The presence of Neo signal only in the Wcl confirms that the TvMIF signal detected was the result of secretion and not cell lysis.

Addition of recombinant MIF increases survival of the parasite during serum starvation. As survival and anti-apoptotic pathways are activated by huMIF in an autocrine manner (15, 33, 34) and TvMIF secretion is induced during serum starvation (Fig. 3), we generated recombinant MIF and added it to parasites to directly test whether soluble TvMIF plays a role in enhancing parasite survival during serum starvation. As shown in Fig. 4A, 50 ng/ml of recombinant TvMIF (rTvMIF) increased
parasite survival 1.2-fold at 4 h and 1.3-fold at 8 h after serum starvation. Human MIF has been reported to be secreted during parasitic infection (35, 36). Thus, we tested whether recombinant huMIF (rhMIF) can also increase parasite survival. We found that 50 ng/ml of rhMIF induced parasite survival 1.2-fold at 8 h (Fig. 4B). These results support the idea of ability of the parasite to respond to both TvMIF and huMIF, increasing parasite survival, and indicate that T. vaginalis can hijack huMIF to enhance its survival.

Parasites overexpressing TvMIF can enhance survival of neighboring parasites in co-cultures. Having established that secreted TvMIF is involved in signaling to parasites, we next tested whether this occurs in an autocrine manner only or can operate in trans. To test if secreted TvMIF can enhance the survival of neighboring parasites, we set up a co-culture of EV and TvMIF-OE parasites in a transwell apparatus using complete Diamond’s media. The bottom wells contained EV, and the top wells, separated from the bottom wells by membranes with 0.4-μm pores, contained either EV or TvMIF-OE. Parasites in both the top and bottom wells were passaged into new media daily for 7, 14, and 21 days (Fig. 5A) to maximize exposure of EV to abundant TvMIF secreted from TvMIF-OE. Then, the cultures were switched to serum-free media to induce nutrient stress and we measured the survival of the EV parasites in the bottom wells under conditions of co-culture with either EV or TvMIF-OE parasites. We found that EV parasites co-cultured with TvMIF-OE parasites for 7 days had a 1.4-fold-higher survival rate than those co-cultured with EV control parasites, that those co-cultured for 14 days had a 1.7-fold-higher survival rate, and that those co-cultured for 21 days had a 2.3-fold-higher survival rate and that the survival advantage became more dramatic with increasing numbers of days of co-culturing (Fig. 5B). These data indicate that the abundance of TvMIF secretion from TvMIF-OE parasites enhanced the survival of neighboring EV parasites, possibly exerting this effect via a positive-feedback loop.

TvMIF inhibits parasite apoptosis. HuMIF is known to activate anti-apoptotic pathways in human cells (14, 15, 37–39). To test whether TvMIF inhibits the apoptosis of the parasites during serum starvation, we employed a double-staining method similar to the commonly used annexin V and propidium iodide (PI) method (40) except that we replaced PI with Zombie Red as the viability dye to exclude dead cells in the
FIG 5 Increased secretion of TviMF by TviMF-OE parasites increases survival of EV parasites after co-culture with TviMF-OE parasites. (A) Scheme of co-culture transwell assay. EV parasites in the wells indicated at the bottom were separated from EV or TviMF-OE in the wells indicated at the top by a membrane with 0.4-μm pores. (B) Co-culturing EV parasites with TviMF-OE parasites confers a survival advantage. EV parasites (bottom well in Fig. 5A) were co-cultured with TviMF-OE parasites (black bars) or EV parasites (white bars) for the number of days indicated on the x-axis. The EV parasites in the bottom well were then transferred to serum-free media for 24 h, and survival was measured. Error bars represent means ± standard errors. *, P value = 0.05; **, P value = 0.01. The data represent results from 3 independent experiments, each done in triplicate.

population. Annexin V, which labels externalized phosphatidylserine, is used as an indicator of the number of parasites undergoing apoptosis (40). Using this assay, we observed that EV parasites had slightly more apoptosis than TviMF-OE parasites at 16 h (P value = 0.07). Moreover, after 24 h of serum starvation, EV parasites were significantly more apoptotic than TviMF-OE parasites (P value = 0.01) (Fig. 6A). To further

FIG 6 TviMF inhibits parasite apoptosis. (A) Double staining with annexin V and Zombie Red was used to assess apoptosis. TviMF-OE parasites (black bars) underwent less apoptosis at 16 h and 24 h after serum starvation than EV parasites (white bars). MFI, mean fluorescence intensity; Healthy, DMSO-treated control; STS, staurosporine (apoptosis inducer positive control). Error bars represent standard errors. *, P value = 0.05. (B) At 0 h, both EV and TviMF-OE nuclei were intact with little damage. (C) Nuclei were stained with DAPI. At 16 h after serum starvation, DNA damage in EV parasites was clearly visible. White arrows indicate examples of DNA fragmentation. (D) DNA damage in EV parasites was significantly greater than that observed in TviMF-OE parasites after 16 h of serum starvation. The imaging data shown in panels B and C were quantified by counting the number of fragmented DNA in a total of 300 nuclei from both EV and TviMF-OE parasites. *, P value = 0.05. Error bars represent means ± standard errors. All data represent results from 3 independent triplicated experiments.
validate the apoptotic phenotypes of the serum-starved parasites, we examined the nuclei of EV and TVMIF-OE parasites grown in serum-free media for 16 h, as the nuclei of *T. vaginalis* are reported to become condensed and fragmented when treated with apoptosis inducers (24, 25). As shown in Fig. 6B and quantified in Fig. 6D, both EV and TVMIF-OE parasites had low levels of DNA fragmentation at 0 h. In contrast, 16 h after serum starvation, ~55% of EV parasites exhibited apoptotic-like DNA fragmentation, whereas only ~38% of TVMIF-OE parasites displayed this phenotype (Fig. 6B to D). These data demonstrate that TVMIF enhancement of parasite survival is associated with a decrease in apoptosis. It is notable that the observed difference between TVMIF-OE and EV parasites with respect to the anti-apoptotic phenotypes seen at 16 and 24 h is not as dramatic as that observed for the survival phenotype (Fig. 1A), indicating that other mechanisms may contribute to survival during serum starvation.

**TVMIF inhibits parasite apoptosis during serum starvation via ROS suppression.** Reactive oxygen species (ROS) activate apoptosis signaling in mammalian systems (41), and huMIF is known to inhibit apoptosis by suppressing ROS production (42, 43). To test whether TVMIF-induced survival under conditions of serum starvation is dependent on ROS inhibition, we measured levels of superoxide, a known by-product of ROS signaling, by staining EV and TVMIF-OE parasites with dihydroethidium (DHE). Upon oxidation by superoxide, DHE is converted to fluorescent 2-hydroxyethidium (44) and the intensity can then be used as a measure of ROS levels. We found that TVMIF-OE parasites produced significantly less superoxide than EV parasites when grown without serum for 16 or 24 h (Fig. 7A). To validate the specificity of the DHE signal, we treated EV and TVMIF-OE parasites with the superoxide dismutase (SOD) mimetic manganese(III) tetrakis(1-methyl-4-pyridyl)porphyrin (MnTmPyP) during serum starvation. MnTmPyP significantly decreased the superoxide signal in EV parasites at both 250 μM and 500 μM and in TVMIF-OE parasites at 500 μM (Fig. 7B). The less dramatic effect that MnTmPyP had on TVMIF-OE parasites is consistent with the ability of TVMIF-OE parasites to suppress ROS production prior to the treatment. In addition, the survival of EV parasites
FIG 8 Parasites with the TvMIF gene knocked out have significantly less survival than the wild-type (WT) parasites. (A) Immunoblot using anti-TvMIF antibody to confirm the loss of TvMIF in knockout (KO) parasites and its presence in addback parasites. GAPDH is the loading control. (B) Knockout of TvMIF in the parasite severely reduces the survival of the parasite under conditions of serum starvation. Adding back of TvMIF in knockout parasites that exogenously overexpress the protein restores and enhances the survival phenotype, reminiscent of the increased survival seen with TvMIF-OE. All time points are normalized to time point 0 h for each parasite. Data are means ± standard errors of results from triplicates, and data from 1 of 3 independent experiments are shown. *, P value < 0.05; **, P value < 0.01; ***, P value < 0.001.

was strongly enhanced by MnTmPyP treatment, whereas TvMIF-OE parasite survival was increased only slightly by the addition of 500 μM MnTmPyP (Fig. 7C). The low increase in survival of TvMIF-OE parasites resulted from normalizing the percentages of living cells of treated TvMIF-OE and untreated TvMIF-OE parasites (Fig. 7C) as the untreated TvMIF-OE parasites had higher survival rates than EV. Together, these data indicate that inhibition of superoxide by TvMIF contributes to parasite survival under conditions of serum starvation.

Gene knockout (KO) of TvMIF severely reduced parasite survival in serum-free media. After completing the work described above, using TvMIF-OE and EV parasites to study the role of TvMIF in parasite survival under nutrient stress, we succeeded in developing CRISPR-Cas9 methods to knock out (KO) genes in T. vaginalis (4). These methods allowed us to KO TvMIF in the parasite (4). As shown in Fig. 8A, immunoblot analyses using the anti-TvMIF antibody confirmed that TvMIF was depleted in the TvMIF KO parasites. To test whether the effects observed in comparisons of wild-type and KO parasites were specific to the loss of TvMIF, we restored TvMIF to the KO parasites (referred to here as “addback parasites”) to test whether this rescued the KO phenotype and caused reversion to that of the wild-type parasites. Immunoblot analysis of addback parasites where TvMIF was overexpressed on a plasmid showed that this resulted in greater expression of TvMIF than was observed in the wild-type parasites (Fig. 8A). Testing the wild-type, TvMIF KO, and addback parasites for survival under conditions of serum starvation, we found that TvMIF KO parasites had 16-fold less and 9-fold less survival than the wild-type parasites and 23.7-fold and 27.7-fold less survival than the addback parasites after 16 h and 24 h under conditions of serum starvation, respectively (Fig. 8B). The increase in survival of addback parasites relative to wild-type parasites is consistent with the higher levels of TvMIF expressed in addback parasites, due to the gene’s presence on a multicopy plasmid. To further validate the survival effect of TvMIF, we added rTvMIF to TvMIF KO parasites in serum-free media and found that 100 ng/ml rTvMIF increased the survival 1.7-fold after 24 h, compared to vehicle control (Fig. S4). Together, these results provide definitive evidence that TvMIF plays a crucial role in resistance to death of the parasite under conditions of nutrient stress.

DISCUSSION

We have established a role for TvMIF in the survival of the parasite under adverse conditions. Although MIF has been widely studied in mammals, the function of this conserved protein in other eukaryotes is largely unknown. We demonstrated that overexpression of TvMIF increases parasite survival under conditions of nutrient starvation, which may be important for the parasite to maintain chronic infection in the constantly changing vaginal environment (5). Nutrient starvation was also shown to
induce the expression and secretion of TpMIF by the parasite. By co-culturing parasites that overexpress TpMIF (TpMIF-OE) with empty vector (EV) parasites that do not, secreted TpMIF was shown to enhance the survival of EV parasites upon prolonged co-culturing, suggesting the presence of an intracellular positive-feedback pathway by which uptake of TpMIF triggers an increase in TpMIF expression and secretion. We were ultimately able to knock out (KO) TpMIF and to show definitively that the loss of TpMIF severely altered the ability of T. vaginalis to survive under conditions of nutrient stress.

We found that TpMIF inhibits parasite apoptosis during serum starvation, consistent with the increased survival phenotype conferred on TpMIF-OE parasites. These data imply that nutrient deprivation during infection induces parasite apoptotic pathways and that increased production of TpMIF may allow the parasite to survive, awaiting more favorable conditions within the urogenital tract. The ability of huMIF to inhibit apoptosis and to contribute to oncogenesis is well established (15, 38, 45–48). This functional conservation between huMIF and TpMIF is remarkable given their evolutionary divergence. Our findings are consistent with host-parasite interactions providing an environment that allowed co-evolution of the two proteins.

Previous studies indicated that T. vaginalis undergoes apoptosis when treated with drugs known to induce apoptosis in mammalian cells. Treatment with staurosporine (STS) led to DNA condensation and staining with annexin V, both indicators of apoptosis (24). The survival of T. vaginalis under conditions of iron depletion, glucose restriction, or serum starvation has also been described, followed by examination of the resulting differential gene expression (49–51). Re-evaluation of these data revealed that under conditions of glucose restriction, TpMIF mRNA is upregulated by 3.8-fold at 12 h and 1.3-fold at 24 h, indicating that TpMIF may be involved in enhancing parasite survival under conditions of glucose starvation (50). In contrast, RNA analyses of parasites grown under conditions of iron depletion revealed that TpMIF mRNA levels were 2.5 times lower under iron-deficient conditions than under iron-rich conditions (49). However, the time point used to extract RNA for the transcriptome sequencing (RNA-seq) analysis was not stated. Therefore, it is possible that TpMIF was upregulated to exert its survival effect at a different time during the analyses.

Although the studies on RNA regulation under glucose-deficient and iron-deficient conditions (49, 50) provide insights into potential pathways that induce parasite survival, prior to this study, no survival factor had been characterized or manipulated by genetic approaches to directly address mechanisms underlying the survival phenotype. Our study results show that the induction of parasite survival by TpMIF via the inhibition of apoptosis is accompanied by reactive oxygen species (ROS) suppression. These data confirm the evolutionary conservation of apoptosis inhibition pathways, as ROS has been shown to activate apoptosis signaling in mammalian systems (41) and huMIF is known to inhibit apoptosis by suppressing ROS production (42, 43).

Induction of apoptosis in the parasitic protists Leishmania spp., Trypanosoma cruzi, and Trypanosoma brucei is accompanied by an increase in ROS (52–57). However, whether the modulation of apoptosis by MIF has been generally conserved between protists and mammals, as appears to be the case for ROS, is yet to be determined. In this regard, it is notable that the known roles of MIF in Xenopus laevis and Caenorhabditis elegans are not directly related to survival under conditions of stress (58, 59). The main functions of other parasite MIFs have been shown to be focused on their effects on hosts. Leishmania major MIF (LmMIF), Plasmodium species MIF, Toxoplasma gondii MIF, and Entamoeba histolytica MIF can bind huMIF receptor CD74 and modulate host immune responses, activate huMIF pathways such as the extracellular signal-regulated kinase/mitogen-activated protein kinase (ERK/ MAPK) pathways, and affect the infection outcome (10, 18, 20, 60, 61). LmMIF and Plasmodium yoelii MIF also inhibit apoptosis of host macrophages by activating mammalian survival pathways (10, 61, 62). However, the possible anti-apoptotic effect of parasite MIFs on these parasites is yet to be examined.

We found that both TpMIF and huMIF confer cell survival, which may be the result of host-parasite co-evolution. This notion is supported by our previous studies showing
that TvMIF and huMIF induce the same signaling pathways in human cells, conferring an anti-apoptotic and increased-growth phenotype (17), as well as by the data reported here indicating that addition of rhuMIF induces parasite survival under conditions of serum starvation. The latter observation implies that secreted huMIF could also directly affect parasite survival under adverse conditions during infection.

TvMIF was originally identified in the T. vaginalis exosomal proteome (31). HuMIF is also present in the exosomal proteome derived from a variety of human cell types (63–71). The effect of huMIF on cell survival has been shown to be mostly focused on soluble huMIF (14, 72, 73). However, Costa-Silva et al. showed that pancreatic ductal adenocarcinoma-derived exosomes with abundant huMIF play a role in inducing liver metastasis, although they did not test whether exosomal huMIF directly triggers anti-apoptotic pathways (71). Further analyses will be required to determine whether exosomal huMIF or TvMIF is capable of triggering anti-apoptotic pathways when exosomes fuse and deliver their protein cargo into human and/or trichomonad cells. Likewise, future transcriptomic and/or proteomic analyses should assist in identifying the anti-apoptotic pathway(s) triggered by TvMIF.

MATERIALS AND METHODS

T. vaginalis cell culture and transfection. T. vaginalis strain B7RC2 (ATCC 50167) was cultured in Diamond’s medium supplemented with 100 U/ml penicillin and 100 µg/ml streptomycin (Thermo Fisher Scientific), 180 µM ferrous ammonium sulfate (Fisher), 28 µM sulfasalicylic acid (Fisher), and 10% horse serum (Sigma) in complete Diamond’s media (26). TVAg219770 (TvMIF) was overexpressed in Master-Neo-HA (hemagglutinin), plasmid and transfected as previously described (17). The plasmid was maintained with 100 µg/ml of G418 (Gibco) for selection. Parasites were cultured at 37°C and passaged daily for 2 weeks or less.

Serum starvation assay. T. vaginalis was cultured overnight in complete Diamond’s media (26). The parasites were then pelleted by centrifugation and resuspended at 10^6 cells/ml in the same media, with the exception that no horse serum was added (serum-free Diamond’s media), and incubated at 37°C at the time points indicated. After each time point, 500 µl of culture was taken from each sample and read on a FACS instrument. The survival rates of the parasites were determined by excluding dead cells using Zombie Red viability dye at a 1:1,000 dilution in PBS (BioLegend) and quantitated using CountBright Absolute Counting Beads (Thermo Fisher Scientific) with a BD LSRFortessa cell analyzer (see Fig. S2 in the supplemental material). The percentages of living cells at all of the time points were normalized to the level at time point 0 h, which was set as 100%.

Density stress assay. EV and TvMIF-EC parasites were grown to mid-log phase (10^6 cells/ml) overnight and concentrated to 10^7 cells/ml or 2 × 10^7 cells/ml in complete Diamond’s media. These parasites were then incubated for 4 h or 8 h. The cells were subjected to FACS analyses using a BD LSRFortessa cell analyzer. The percentages of dead cells were determined by gating the Zombie Red-positive population and determining its percentage relative to the total cell population.

Growth assay. EV and TvMIF-EC parasites (4 × 10^6 cells/ml) were cultured in complete Diamond’s media. At 12 h and 24 h, 500 µl of each sample was taken for FACS analyses to determine the numbers of living cells. Numbers of living cells were determined as described for the serum starvation assay.

Secretion assay. The parasites were grown to mid-log phase overnight in complete Diamond’s media, collected, and resuspended in PBS–5% sucrose at 10^6 cells/ml. The parasites (5 × 10^6) were incubated at either 16°C or 37°C for 2 h. The 16°C condition was used for the secretion inhibition control. After the incubation, the parasites were spun at 3,200 rpm for 10 min and the cell pellets were collected and lysed in lysis buffer containing 50 mM Tris·HCl (pH 7.5), 2% SDS, and 1× Halt protease inhibitor cocktail (Thermo Fisher Scientific) to make whole-cell lysates (WCL). The supernatant was passed through a PES Vivaflow crossflow cassette (Sartorius) (MWCO, 100 kDa) to separate the exosomes (Exo) from the non-exosomal soluble fraction (NESF). The NESF was concentrated by using Amicon Ultra Centrifugal filters (EMD Millipore) (MWCO, 10 K). The Exo were pelleted at 100,000 × g for 70 min. Anti-neomycin phosphotransferase (Neo) (Jackson Laboratory) (1:2,500) and anti-TvMIF antibodies (polyclonal rabbit antiserum raised against TvMIF) (1:500) (14) were used as the primary antibodies and anti-rabbit antibody (Jackson Laboratory) (1:25,000) was used as the secondary antibody for both Neo and TvMIF probing.

Intracellular and secreted TvMIF quantitation. Anti-TvMIF polyclonal antibody, anti-glyceraldehyde-3-phosphate dehydrogenase (anti-GAPDH) antibody (Cocalico Biologicals) (1:10,000), and anti-Neo antibody (Jackson Laboratory) (1:2,500) were used as the primary antibodies, and anti-rabbit (Jackson Laboratory) was used as the secondary antibody. For intracellular TvMIF quantitation, mid-log-phase parasites were spun down and washed with PBS–5% sucrose–1× Halt protease inhibitor cocktail (Thermo Fisher Scientific). The cells were then lysed in 50 mM Tris·HCl (pH 7.5)–2% SDS–1× Halt protease inhibitor cocktail lysis buffer. Equal total proteins were loaded from whole-cell lysates from 0 h, 16 h, and 24 h. For secreted TvMIF quantitation, T. vaginalis parasites cultured overnight in complete Diamond’s media were collected and resuspended in serum-free or complete Diamond’s media for 3 h or 6 h of incubation. After the incubation, the parasites were resuspended in PBS–5% sucrose for 2 h of incubation for secretion collection. The cells were spun down and collected using the method described for intracellular TvMIF quantitation as the whole-cell lysate (WCL) control. The supernatant was then concentrated with
Amicon Ultra Centrifugal Filters (MWCO = 10 k). Equal total protein amounts were loaded for the WC1 control and secreted fractions for all samples.

Production and purification of *rTvMIF* and *rhuMIF*.
TvMIF and huMIF was cloned into the pET SUMO expression vector with an N-terminal SUMO domain with 6× His tag separately. Both constructs were transformed into BL21(DE3) E. coli (Thermo Fisher Scientific) for protein expression (74). *TvMIF* and *rhuMIF* were purified using a HisPur nickel-nitrilotriacetic acid (Ni-NTA) Spin column (Thermo Fisher Scientific) and dialyzed into 20 mM Tris-HCl (pH 8.0)–150 mM NaCl–1 mM dithiothreitol (DTT). His-tagged SUMO protease (mclab) in combination with SUMO protease buffer (50 mM Tris (pH 8.0), 1 mM DTT) was used to cleave off N-terminal SUMO and the 6×His tag to produce rTvMIF and rhuMIF with their native sequences, SUMO protease and the SUMO domain with the His tag were removed by the use of a HisPur Ni-NTA Spin column. rTvMIF and rhuMIF were then purified by the use of Econo-Pac 10DG desalting prepacked gravity flow columns (Bio-Rad).

Exogenous *rTvMIF* and *rhuMIF* addition and survival assay. A 50 ng/ml volume of *rTvMIF* or rhuMIF or PBS vehicle control was added to TvMIF-OE grown in serum-free Diamond’s media. The survival rates were determined 4 h and 8 h after starvation. *rTvMIF* (100 ng/ml) or PBS was added to TvMIF KO parasites in serum-free Diamond’s media. The survival rates were determined 24 h after starvation.

Co-culture in transwell. Transwell (Corning) with a 0.4-μm-pore-size polycarbonate membrane insert was used. EV or TvMIF-OE (1.8 × 10⁴) was plated in 300 μl of complete Diamond’s media in each individual top insert, and EV (2 × 10⁴) was plated in 1 ml of media in each individual bottom well. The top and bottom cells were passaged and rediluted daily to the concentrations described above in complete Diamond’s media. Every 7 days, the cells were resuspended in serum-free Diamond’s media at the concentration described above and incubated for 24 h for survival tests. The survival rates were determined as described for the serum starvation assays.

Apoptosis assays. Zombie Red (1:1,000) and 4,5 μg/ml of fluorescein isothiocyanate (FITC)-annexin V (BioLegend) in annexin V binding buffer (BioLegend) were used to stain parasites grown without serum at 0 h, 16 h, and 24 h, and the cells were subjected to FACS analysis using a BD LSRII Fortessa cell analyzer. The live parasites were gated by excluding positive Zombie Red signal. Apoptotic levels were determined by analysis of the mean fluorescence intensity (MFI) of annexin V of live parasites. Parasites treated with dimethyl sulfoxide (DMSO) were used as the healthy control, and parasites treated with 4 μM staurosporine (STS) for 16 h composed the apoptotic control.

DNA fragmentation was done by incubating EV and TvMIF-OE in serum-free Diamond’s media and fixing parasites in 4% formaldehyde-PBS. Parasite nuclei were stained with ProLong Gold antifade mountant with 4,6-diamidino-2-phenylindole (DAPI) (Thermo Fisher Scientific). The slides were then imaged using an Axioskop 2 epifluorescence microscope (Zeiss). The percentage of fragmented DNA was determined by the number of fragmented DNA in a total of 300 nuclei counted in each sample by using ZEN lite software.

Reactive oxygen species (ROS) detection and superoxide dismutase (SOD) mimic treatment.
EV and TvMIF-OE parasites were grown in serum-free Diamond’s media as previously described. At 0 h, 16 h, and 24 h, cells were stained with 2 μM dihydroethidium (DHE) (Thermo Fisher Scientific) mixed with PBS at 37°C for 30 min in the dark. The MFI of ROS signal was determined by the MFI of phycoerythrin-Texas Red (PE-Texas Red)-treated live cells gated by the use of a flow cytometer.

For SOD mimic treatment, a 250 μM or 500 μM concentration of manganese(II) tetraakis(1-methyl-4-pyridyl)pyrophosphin (MnTMPyP) (Sigma) or DMSO was used to treat parasites grown in serum-free Diamond’s media at time point 0 h. At 16 h, parasites were stained with DHE and CountBright absolute counting beads (Thermo Fisher Scientific) were added to determine the number of live cells. The ROS signal and the percentage of living cells were measured by the use of a flow cytometer.

Gene knockout and adding back of *TvMIF* in *T. vaginalis* B7RC2. The constructs and reagents used to create the TvMIF knockout were as described by Jannsen et al. (4). Adding back of *TvMIF* to TvMIF KO cells was done by transfection of TVAG_219770 (TvMIF) in Master-Nec-{|HA}p plasmid with selection of G418 at the concentration of 900 μg/ml in order to select for the plasmid in the presence of a neomycin phosphotransferase (Neo) gene knock-in at the TvMIF gene locus, replacing the TvMIF gene.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00910-18.

**FIG S1.** TIF file, 2.3 MB.
**FIG S2.** TIF file, 2.2 MB.
**FIG S3.** TIF file, 1.8 MB.
**FIG S4.** TIF file, 1.3 MB.

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REFERENCES


Chapter 3:

A *Trichomonas vaginalis* cadherin-like protein mediates adherence to and killing of host cells
Abstract

Trichomonas vaginalis, a worldwide prevalent sexually-transmitted parasite, adheres to and induces cytolysis of human mucosal epithelial cells. We have functionally characterized a hypothetical protein, TVAG_393390, whose tertiary structure modeling using Phyre2 revealed similarity to cadherin proteins. TVAG_393390 contains four predicted calcium-binding sites at structurally similar locations to cadherin proteins; thus we renamed it cadherin-like protein (CLP). CLP was found to be surface localized and CLP mRNA was significantly up-regulated when parasites contacted host cells. To test the roles of CLP and its calcium-binding domains in host cell adherence, wild-type CLP (CLP) and a calcium-binding site mutant (CLP-mut) were overexpressed in T. vaginalis. We found that CLP parasites have ~3.5-fold greater adherence to host cells relative to the empty vector control (EV), and this increased adherence is ablated by mutating the 4th calcium-binding domain. Additionally, competition with recombinant CLP decreased parasite binding to host cells. As cadherin proteins help mediate cell-cell interactions and adherent T. vaginalis strains also display parasite-parasite interactions, we tested the contribution of CLP and CLP-mut on parasite aggregation in the presence and absence of both calcium and host cells. Overexpression of CLP induced parasite aggregation whereas CLP-mut overexpression did not, and CLP-induced parasite aggregation was significantly increased in the presence of calcium, further establishing a Ca^{2+}-binding dependency for CLP’s function. Lastly, parasites overexpressing wild-type CLP increases killing of host cells by ~3.3 and ~2.35-fold, compared to parasites overexpressing EV and CLP-mut, respectively. Our data identify the first CLP characterized in a unicellular eukaryote that contributes to both parasite-parasite and host-parasite interactions. CLP may represent convergent evolution of a parasite cadherin protein that
is structurally similar to the mammalian cell adhesion protein cadherin and whose function also contributes to pathogenesis.

**Introduction**

*Trichomonas vaginalis* is an extracellular eukaryotic parasite that causes trichomoniasis, the most common non-viral sexually transmitted infection, which affects more than 275 million people worldwide annually [1]. In the United States, trichomoniasis is classified as a neglected disease due to limited knowledge of the consequence of infection and its disproportionate affliction of low-income populations and minorities [2][3]. Although the majority of *T. vaginalis* infections are asymptomatic, trichomoniasis can result in inflammation of the urogenital tract of both men and women, resulting in vaginitis, prostatitis, pruritus, dysuria and discharge [4]. Furthermore, *T. vaginalis* is also associated with adverse pregnancy outcomes and HIV co-infection [5][6][7].

As a parasite that does not invade host cells, it is critical for *T. vaginalis* to attach to urogenital epithelial cells in order to establish an infection and acquire nutrients from host cells. *T. vaginalis* can also attach to, lyse, and phagocytose leukocytes and red blood cells [8][9]. *T. vaginalis* strains display different abilities to bind host cells in vitro, with up to a 45-fold difference in attachment observed between different strains [10]. Therefore, understanding the molecular mechanisms of how *T. vaginalis* attaches to host cells is the key to understanding how the parasite establishes infection. In an attempt to determine what factors play a role in adherence, we previously compared the plasma membrane surface proteome of 3 adherent and 3 less adherent *T. vaginalis* strains, identifying proteins that are significantly more abundant in the adherent strains relative to the less adherent strains [11]. Mining this surface proteomics data
revealed a hypothetical protein, TVAG_393390, that is more abundant in 2 out of 3 adherent strains relative to less adherent strains by 1.7 to 3.4-fold [11]. TVAG_393390 was also identified as a putative substrate of the *T. vaginalis* rhomboid protease 1, TvROM1, a membrane serine protease that we have shown is involved in parasite attachment and host cytolysis [12]. Together, these findings highlight a potential role of the TVAG_393390 protein contributing to *T. vaginalis*-host cell interactions.

*T. vaginalis* cell-cell interactions may also be an important phenotype contributing to pathogenesis. Groups of parasites are readily visible when attached to ectocervical cells and prostate cells [13][14]. The ability of *T. vaginalis* to aggregate also correlates with a strain having higher host cell adherence and cytolytic properties [15]. In metazoans, the strongest forms of cell-cell attachment are mediated by cadherin proteins that bind to each other on apposing cells forming adherens junctions [16]. While single-celled eukaryotes, such as *T. vaginalis*, do not contain cell junctions, it has been hypothesized that protein precursors found in protozoans may have given rise to the complexes that allowed cell-cell interactions leading to multicellularity [17][18]. Bioinformatic analysis revealed that TVAG_393390 is predicted to be structurally similar to metazoan cadherin proteins, indicating its potential role in mediating cell-cell adhesion in the protozoan *T. vaginalis*.

Classic cadherin proteins are large single-pass, transmembrane proteins with extracellular cadherin (EC) repeats, calcium-binding sites, and a cytosolic tail that is involved in intracellular signaling and interactions with p120 catenin, β-catenin, α-catenin and, indirectly, with f-actin [19][20]. Cadherin proteins are involved in both homophilic and heterophilic interactions [21]. Homophilic interactions occur when cadherin protein binds to the same type of cadherin protein *in trans* on another cell. A classic example is the mouse epithelial cadherin (E-cadherin), which
mediates cell-cell interactions in epithelial cells [22][23][24]. Heterophilic interaction involves cadherin proteins binding to a different type of cadherin protein on a different cell type [25]. An example of a heterophilic interaction that has a role in pathogenesis, is the binding of the human E-cadherin protein to a bacterial *Listeria monocytogenes* surface protein, an interaction which helps to mediate invasion of the bacteria [25].

Here we characterize TVAG_393390 and show that it is structurally and functionally similar to cadherin proteins. This cadherin-like protein (CLP) was also found to play significant roles in parasite attachment to and lysis of host cells, as well as parasite aggregation. CLP may thus represent an evolutionary relic of cadherin-like proteins. To our knowledge, this is the first report of a cadherin-like protein in protozoans contributing to host-pathogen and parasite-parasite interactions.

**Results**

**Bioinformatic analysis reveals that TVAG_393390 is structurally similar to cadherin proteins**

The *T. vaginalis* protein encoded by TVAG_393390 is listed as a conserved hypothetical protein in the genome sequence database for *T. vaginalis* [trichdb.org/trichdb](http://trichdb.org/trichdb). This protein was found to be more abundant in the surface proteome of adherent *T. vaginalis* strains relative to less adherent strains by 1.7 to 3.4-fold [11]. We also identified TVAG_393390 as a putative substrate of a *T. vaginalis* rhomboid protease demonstrated to be involved in parasite attachment and host cytolysis [12]. To further investigate a potential role of TVAG_393390 in the *T. vaginalis* adherence, we searched both the gene sequence and the protein sequence using National Center for Biotechnology Information (NCBI) Basic Local Alignment Search Tool
(BLAST) and found no homologues in other organisms. We also used InterPro [26] and Pfam [27] analyses to identify functional domains and none was identified. Next, we employed a secondary structure prediction program called Phyre2 which predicts protein structure by comparing a large database of known protein secondary structures and building threedimensional models based on the identified homology [28]. Using this approach, we found that the most common modeling for TVAG_393390 to a particular type of protein was to cadherin proteins, with each model having at least 97.9% confidence scores (probability that a template is homologous to our sequence) and 7% identity (Fig. 3-1A). Fig. 3-1B shows a TVAG_393390 model derived using one of the highest ranking tertiary structures, mouse E-cadherin protein. Similar to cadherin proteins, TVAG_393390 is predicted to have 5 extracellular domains with the classical β-sandwich domains and Greek-key folding of cadherin proteins [29][30] (Fig. 3-1B).

Another defining characteristic of cadherin proteins is their binding to extracellular Ca\(^{2+}\) via calcium-binding pockets located in between the extracellular domains [31]. Visual inspection of the TVAG_393390 protein sequence for the highly conserved Ca\(^{2+}\)-binding sites LDRE, DXD, DXXD, \(x = \) any amino acid [29], located at the interfaces of the extracellular domains predicted by Phyre2, identified four candidate Ca\(^{2+}\)-binding sites (Fig. 3-1C). TVAG_393390 is also predicted to have one transmembrane domain, near the C-terminus of the protein, based on the transmembrane protein topology prediction software TMHMM [32]. Thus, the predicted orientation of TVAG_393390 would result in the cadherin-like modeled region being exposed to the outside of the cell with a small C-terminal tail located intracellularly (Fig. 3-1B). Due to TVAG_393390’s high confidence modeling to cadherin proteins and the presence of key cadherin-like features, we renamed TVAG_393390 as cadherin-like protein (CLP).
CLP is expressed on the surface of the parasite

To determine whether CLP was surface-localized and displays the predicted orientation, we cloned the gene in our standard *T. vaginalis* expression vector, MasterNeo [33] under the control of *T. vaginalis* α-succinyl Co-A synthetase promoter and fused with two C-terminal hemagglutinin (HA) tags. The construct was then introduced into *T. vaginalis* by transfection and parasites were selected with G418 as previously described [34]. Using indirect immunofluorescence assay with an anti-HA antibody, exogenously overexpressed CLP was shown to localize to the surface of the parasite (Fig. 3-2A & B). Since it is also predicted that the C-terminal tail of the protein with the fused HA tags will be located inside the cell, to further probe the orientation of CLP, we performed indirect immunofluorescence with and without permeabilization. We found that the fluorescent signal from CLP is significantly stronger when the parasites are treated with a permeabilizing agent (Fig. S3-1A & C) as permeabilization allows anti-HA antibody access to the intracellular HA tags. These results provided additional support that the C-terminal domain of CLP is located inside the cells. The data are consistent with the predicted CLP structure generated with the Phyre2 analysis (Fig. 3-1B) and demonstrated that overexpression of the protein resulted in the predicted membrane localization with the cadherin-like domains being exposed on the outer surface of the parasite. Therefore, the predicted CLP protein topology is graphically depicted in Fig 3-2C.

CLP mRNA is up-regulated during host contact

Upon parasite contact with human ectocervical cells (Ects), *T. vaginalis* up-regulates expression of a variety of proteins including actin, actin-binding proteins [35] and tetraspanin proteins (TvTSP3, TvTSP5, TvTSP6, and TvTSP8), the latter of which are transmembrane proteins that modulate their own expression and subcellular localizations during host contact.
Thus, we hypothesized that surface localized CLP might be up-regulated upon interaction of *T. vaginalis* with host cells. To test this, wild-type parasites were incubated with human Ects for 30 min, 1 h, 2 h or 6 h, followed by removal of unbound parasites in suspension and extraction of RNA from only the parasites adhered to Ects. We found that CLP mRNA is up-regulated by ~5-fold, 19- and 20-fold at 1h, 2h and 6h, respectively, relative to 30 min after host contact (Fig. 3-3). As reduced temperature might affect the ability of the parasites to sense the environment initially and modulate surface protein expression, we used 30 min instead of 0 min for baseline comparison because immediately before co-culturing the parasite with Ects, the parasites are exposed to 4°C to collect and concentrate them. These data support a role for CLP in host cell sensing and/or binding.

**Calcium binding is predicted to be important for the function of CLP**

Calcium binding plays a critical role in the adhesive role of cadherin proteins by rigidifying the extracellular domains and mediating binding between cadherin proteins on apposing cells [37][38]. Specifically, aspartate residues that help coordinate calcium ions at the base of the cadherin extracellular domains constitute one of the most conserved domains of cadherin proteins across different species [29]. Therefore, to understand how CLP functions, we used Phyre2 and SuSPect mutational analysis to help identify which aspartate residues of the 4 predicted Ca\(^{2+}\)-binding domains [28][39] (shown in Fig. 3-1C) would lead to the strongest phenotypic effects if mutated (Fig. 3-4A and Fig. S3-2). The two aspartate residues D443 and D445 in the putative fourth Ca\(^{2+}\)-binding domain are predicted to be the most sensitive to mutation (Fig. 3-4A and Fig. S3-2) so we proceeded to mutate both of these residues to alanine as this type of mutation is standardly performed to study the function of calcium-binding domains in cadherin proteins [40]. We then overexpressed both the wild-type CLP (CLP) and the
D443A D445A CLP mutant (CLP-mut) with an N-terminal GFP tag in *T. vaginalis* and an empty vector (EV) as a negative control. Next, we compared the expression of CLP and CLP-mut proteins in the selected transfectants and found that expression of the wild-type CLP and CLP-mut was similar as determined by immunoblotting using an anti-GFP antibody (Fig. 3-4B).

**CLP protein contributes to increased host cell binding which is dependent on the CLP calcium-binding domain**

To test whether CLP is important for host binding, we compared the ability of CLP vs CLP-mut overexpressing cells to bind host Ects. We found that CLP increases the host attachment by 3.5-fold compared to EV (Fig. 3-5A). In contrast, CLP-mut has a significantly decreased attachment phenotype that is similar to that of EV (Fig. 3-5A). This data strongly indicate a role for CLP in host cell binding.

To further support the role of CLP in host cell binding, we competed parasite binding to Ects with addition of recombinant CLP. The extracellular domain (EC) of CLP (rCLP EC) was used instead of the entire protein for the ease of expressing the protein in *E. coli* periplasm and since its EC is predicted to be responsible for the cadherin protein-protein interactions [31]. We added rCLP EC to Ects for 30 min prior to the addition of parasites. We found that as the concentration of rCLP EC was increased, the percentage of parasite binding to the host cell decreased (Fig. 3-5B). These results further support a role of CLP in host binding.

**CLP contributes to parasite-parasite clumping and the effect is abolished in CLP-mut parasites**

We and others have observed that more adherent *T. vaginalis* strains appear to clump with each other more readily than poorly adherent strains. In addition, overexpression of the TvTSP8 surface protein which increases parasite adherence to host cells also increased parasite
clumping compared to the EV control parasites [15][10]. Furthermore, cadherin proteins are known to mediate homophilic interactions where cadherin proteins interact with the same type of cadherin on another cell [21][41][42]. We thus compared the clumping behavior of EV, CLP and CLP-mut parasites with or without Ca$^{2+}$ and in the presence or absence of the host. CLP displayed a ~7.5-240 fold increase in parasite clumping compared to the EV control and CLP-mut had almost the same levels of clumping as EV under all conditions tested (Fig. 3-6A & B). Addition of Ca$^{2+}$ but not the presence of host cells significantly increased the clumping of CLP parasites (Fig. 3-6A & B). These data together suggest that the CLP-mediated clumping phenotype is calcium-dependent and the ability does not necessitate a host signal.

**CLP contributes to increased killing of host cells**

Since CLP is involved in host cell binding and parasite clumping, we hypothesized that these properties could also contribute to increasing host cell death as epithelial host cell killing by *T. vaginalis* is contact-dependent [10] and the increased parasite-parasite association may further increase the number of parasites attacking the host. We found that CLP overexpression increased host death by 3.3-fold compared to EV (Fig. 3-7). In contrast, CLP-mut only displayed a 1.4-fold increase in host killing compared to EV (Fig. 3-7). The slight increase in cytotoxicity observed with the CLP-mut may be due to the fact that the CLP-mut has one of the four hypothesized Ca$^{2+}$-binding domains mutated so it may not completely abolish the enhanced killing effect of CLP.

**Discussion**

We have identified and characterized a surface protein of *T. vaginalis* called cadherin-like protein (CLP) and showed that it plays a significant role in host binding, parasite clumping
and host cell killing. This is the first functional analyses of a cadherin-like protein in a unicellular eukaryote. Mammalian cadherin proteins are known to be co-opted for pathogen invasion or colonization by several bacteria and the pathogenic yeast *Candida albicans* [43][44][45]. However, the identification of CLPs or a role for unicellular pathogen CLPs in the adherent to and killing of mammalian cells have not been previously described to our knowledge. We show that this *T. vaginalis* CLP mimics the structure and the function of host cadherin proteins, raising the possibility that other parasites may also use CLPs with low sequence identity but with structural similarity to mammalian cadherins to interact with host cells. The presence of a CLP in *T. vaginalis* that is structurally and functionally similar to mammalian cadherins is likely an example of convergence evolution of 3D structures with similar properties, in the absence of strong primary sequence homology.

Cadherin proteins are known to be involved in homophilic interactions that cause the same cell types to adhere to each other [25][41]. We observed that CLP-overexpressing parasites clump significantly more than empty vector control and the CLP mutant-overexpressing parasites, demonstrating a role for parasite CLP in homophilic adherence of parasites to one another. We also demonstrated that CLP-induced parasite clumping is significantly increased in the presence of calcium, consistent with the calcium-dependent homophilic interaction mediated by cadherin proteins [41]. Highly adherent strains of *T. vaginalis* have been observed to clump (i.e. adhere to each other) significantly more than poorly adherent strains [15]; however, the role of clumping in infection is unclear. To our knowledge, only another family of surface proteins, tetraspanins, have been found to help mediate parasite aggregation [15]. In mammalian cells, integrin proteins are one of the predominant protein groups complexed by tetraspanins [46]. However, the *T. vaginalis* genome does not contain any proteins annotated as
integrin-like. On the other hand, it has been reported that human E-cadherin protein interacts with its tetraspanin protein in colon carcinoma [47]. Therefore, future investigation into whether CLPs and TSPs functionally interact is merited. It should also be noted that this *T. vaginalis* CLP is predicted to have a small C-terminal tail with only 2 amino acid residues (Fig. 3-1B). In classical cadherin proteins, the C-terminal is important for cadherin-mediated signaling [48]. It is therefore possible that if CLP represents an early evolutionary form of cadherin-like proteins, part of its functions such as the role we observed in sensing the presence of host cells and increasing its expression in response, may in part be mediated by associating with other proteins that help recruit the signaling proteins. Overall, we speculate that parasites clumping increases the number of parasites attaching to host cells, which in turn, increases the likelihood of parasites successfully colonizing the host. Additionally, human neutrophils have been shown to kill *T. vaginalis* by taking bites from the parasites, in a process called trogocytosis [49]. Aggregation of the parasite could potentially protect parasites in the center of an aggregate from being attacked by neutrophils. Future analyses will better define the functional importance of parasites adhering to one another in parasite survival and infection.

Pre-incubation of host cells with recombinant CLP (rCLP) reduced parasite binding to host cells, confirming a role for parasite CLP in host cell binding. While *T. vaginalis* attachment to host cells likely depends on multiple factors which have been identified to date [11][12][50][51][52][53][54][55] [56][57][58], the highest amounts of rCLP tested reduced parasite binding by a significant 21% (Fig. 3-7). Furthermore, the observed 3.3-fold increase in host cell killing is also, to our knowledge, one of the strongest phenotypic effects mediated by exogenous expression of a single protein factor. E-cadherin and N-cadherin are expressed in the human male and female urogenital and reproductive tracts [59][60]. E- and N-cadherin are also found on
spermatozoa [60] to which *T. vaginalis* can also attach and phagocytose [61]. It is therefore of interest to investigate in future studies whether CLP has a conserved role in attaching to and lysing multiple cell types and how it may structurally mediate molecular mimicry.

Previous functional studies on cadherin proteins have revealed that mutations at the first and second calcium-binding sites cause the strongest disruption in homophilic interactions of cadherin proteins that mediate cell-cell binding, whereas mutation at the latter two calcium-binding sites have a slight or almost no effect [40][62]. However, antibody blocking experiments and subsequent mapping of the epitope of the blocking antibody points to a direct interaction between the antibody and the fourth calcium-binding site in E-cadherin protein [63][64]. Here we show that mutating the 4th calcium-binding domain in CLP almost completely reverses the enhanced host attachment, parasite clumping and host cell killing observed with parasites overexpressing the wild-type CLP. It is possible that the overexpression of the *T. vaginalis* protein mutated in the 4th calcium-binding domain results in a dominant negative effect on the endogenous wild-type CLP. Alternatively, the low identity of *T. vaginalis* CLP to mammalian cadherins (10%) (Fig. 3-1A) could explain why the 4th calcium-binding domain of *T. vaginalis* plays a significant role in host cell interactions relative to the 1st or 2nd domain. Further analyses of the evolution and function of this, and perhaps other, CLPs in pathogens is warranted.

Although CLP surface expression levels are different in adherent vs less adherent *T. vaginalis* strains, the fact that CLP is expressed by all the six strains surveyed in our prior proteomics study [11] may indicate an important and conserved function. Our work has uncovered new roles for a family of previously undescribed proteins in *T. vaginalis*, helping us to further scratch the surface of the mechanistic interactions contributing to pathogenesis. Our
work also places *T. vaginalis* as a model organism for the broader study of cell-cell interactions and cell-cell adhesion.

**Material and Methods**

**Bioinformatic analyses**

To predict TVAG_393390 function, we used InterPro [26], Pfam [27] and Phyre2 [28] programs to analyze its protein sequence and BLAST with both the gene and protein sequences published on TrichDB [65]. For Phyre2 analysis, we chose the intensive modeling mode, then used the “Run Investigator” feature using the mouse E-cadherin template and performed SuSPect mutational analysis on the four calcium-binding domains/eight aspartate residues in TVAG_393390. The topology of TVAG_393390 was generated with the TOPO2 program [66].

**T. vaginalis and ectocervical cell line Ect1 culture**

*T. vaginalis* strain RU393 (ATCC 50142) was grown as previously described [67]. Parasites were grown at 37°C and sub-cultured daily for up to 2 weeks. The human ectocervical cell line Ect1 E6/E7 (ATCC CRL-2614) (Ects) was grown and passaged as previously described [68].

**CLP wild-type and mutant plasmid construction and T. vaginalis transfection**

TVAG_393390 (CLP) wild-type sequence was cloned into the Master-Neo-(HA)\textsubscript{2} plasmid [33] or the N-terminal enhanced green fluorescent protein (eGFP)-Master-Neo plasmid [12]. Two rounds of site-directed mutagenesis were performed using QuikChange kit (Stratagene) in order to introduce the D443A and D445A mutations sequentially using the following primer sets. To introduce D443A mutation: D443A Fwd:

CACAGCCGTAGTTGTTGcTCCAGATACTAACTTTG and D443A Rev:
CAAAGTTAGTATCTGGAgCAACAACTACGGCTGTG. To introduce the D445A mutation: 
D445A Fwd: GTAGTTGTGTcTCCAGcTACTAACTTTGATTCC and D445A Rev: 
GGAATCAAAGTTAGTAgCTGGAgCAACAACTAC. Introduction of the desired mutations 
was confirmed by sequencing (Genewiz). The constructs were transfected into *T. vaginalis* strain 
RU393 and parasites containing the constructs were selected using G418 as previously described 
[34].

**Indirect immunofluorescence assays**

Parasite transfectants overexpressing the wild-type (WT) CLP with C-terminal HAx2 
were plated on glass coverslips coated with 100 µg/ml of poly-l-lysine (Sigma) and then fixed in 
4% formaldehyde in phosphate-buffered saline (PBS) for 20 minutes. The cells were then 
permeabilized in 0.2% Triton X-100 in PBS or just PBS for non-permeabilized control for 15 
minutes and blocked in 3% BSA for 30 minutes. 1:1000 dilution of anti-HA mouse (BioLegend) 
and 1:5000 goat anti-mouse Alexa Fluor488-conjugated secondary antibody (Molecular Probes) 
were used for staining. The coverslips were then mounted using ProLong Gold Antifade 
Mountant with 4’,6-diamidino-2-pheylindole (DAPI) (Thermo Fisher Scientific). The images 
were taken by a Zeiss confocal microscope with Yokogawa spinning disc and analyzed with 
SlideBook 6 software.

**Real-time reverse transcription PCR (qRT-PCR)**

1 X 10^7 of non-transfected RU393 parasites were incubated with 80% confluent Ects for 
30 minutes, 1 hour, 2 hours or 6 hours. Unbound parasites were removed and RNA was collected 
by adding TriZol to attached parasites on Ects. Total RNA was purified by phenol-chloroform 
extraction and treated with TURBO DNase (Invitrogen). cDNA was prepared by using 
SuperScript III with oligo dT primers (Thermo Fisher Scientific). Platinum SYBR Green qPCR
SuperMix-UDG and the manufacturer’s protocol (Thermo Fisher Scientific) were used for real-time PCR. *T. vaginalis* beta-tubulin was used as the housekeeping gene control. The primer sequences for beta-tubulin are Tub-f: GGCTCGTAACACATCCTACTTC and Tub-r: CTGTTGTGTTGCCGATGAATG. The primer sequences for CLP are 393-f: GACGATGGTCTTTCACAGCC and 393-r: CCATCAGAGTTTGATCTTGAAATTGA.

**Immunoblot Analyses**

Mouse anti-GFP polyclonal antibody (1:1,000) (Clontech) and anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (1:10,000) (Cocalico Biologicals) were used as the primary antibodies and anti-mouse (1:25,000) and anti-rabbit (1:25,000) (Jackson Labs) were used as the secondary antibodies, respectively. 5 X 10^6 parasites were taken from *T. vaginalis* culture and washed with PBS + 5% sucrose + 1xHalt Protease Inhibitor Cocktail (Thermo Fisher Scientific). The cells were then lysed in lysis buffer (0.1% Nonidet P-40, 0.5% deoxycholate, 2% SDS, 50 mM Tris, pH 8, 5 mM EDTA, 150 mM NaCl) + 1xHalt Protease Inhibitor Cocktail (Thermo Fisher Scientific). Equal amounts of proteins were loaded from each sample.

**Production and purification of rCLP ECD**

Sequence encoding the CLP ECD was cloned into the pET22b(+) expression vector flanked by a N-terminal pelB periplasm signal sequence and C-terminal 6xHis tag. The primers for cloning were pET22b_393-f:

TAATTCGGATCCGATGATTTGGACTTTTTTATTGAGGATG and pET22b_393-r:

ACTAAGCTCGAGCTTCTTTTGTCTTTTTCTTTTG. The plasmid was transformed into C41(DE3) *E. coli* cells [69]. An overnight culture was inoculated into 1L LB medium and when this culture reached OD_{600} = 0.5 expression was induced with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) for 16h at 25°C. The culture was then spun at 5000 g for 10 min.
The cultured media was precipitated with 50% saturated ammonium sulfate at 4°C overnight and dialyzed against PBS overnight. Protein in the periplasm was extracted as previously described [70][71]. In brief, the cell pellet was washed with ice-cold 20% sucrose + 30 mM Tris pH8 + 1 mM EDTA and then the periplasm was extracted with ice-cold 5 mM MgSO4. The periplasmic fraction was then dialyzed against PBS at 4°C overnight. A small-scale protein production was done initially to determine that rCLP ECD was abundant in both the culture media and periplasm. As a result, the medium fraction and the periplasmic fraction were combined and loaded onto HisPur Ni-NTA Spin column (Thermo Fisher Scientific). Purified rCLP ECD was dialyzed into PBS and the protein concentration was determined by Pierce BCA Protein Assay (Thermo Fisher Scientific).

**T. vaginalis attachment to Ects and attachment with rCLP ECD competition**

Attachment of *T. vaginalis* to Ects was performed as described in [10]. Briefly, 5 X 10⁴ of CellTracker Blue CMAC (Thermo Fisher Scientific) labeled *T. vaginalis* were incubated with confluent Ects for 30 minutes and the coverslips were fixed in 4% formaldehyde in PBS and mounted on slides using Mowiol (Calbiochem). Fifteen images of each coverslip were acquired using an Axioscope 2 epifluorescence microscope (Zeiss) and cell counts were quantified using Zen lite and Image J software.

Attachment assays that included rCLP ECD to compete for Ect binding were performed using wild-type, non-transfected RU393 parasites and the difference in the procedure was addition of 0.25 µg, 1 µg or 4 µg of rCLP ECD or 4 µg of BSA as a negative control to Ects for 30 min. Media was then removed and replenished with new media containing CellTracker Blue labeled parasites.

**Parasite clumping assay**
The parasite clumping assay was done as described in [15] with modifications indicated below. CellTracker Blue CMAC (Thermo Fisher Scientific) pre-labeled *T. vaginalis* was plated at 1 X 10⁶/ml on glass coverslips covered with confluent Ects or coverslips alone and incubated in completed keratinocyte-SFM with no CaCl₂ or 1 mM of CaCl₂ for 30 minutes. Parasites were then fixed in 4% formaldehyde in PBS and mounted on slides using Mowiol (Calbiochem). Fifteen images of each coverslip were acquired using Axioscope 2 epifluorescence microscope (Zeiss) and analyzed by Zen lite software. A clump is defined as an aggregate of 10 or more parasites.

**T. vaginalis-induced cytotoxicity of ectocervical cells**

Cytotoxicity of Ects was measured as described [10]. The only modification was 3 X 10⁵ parasites was added to confluent Ects and incubated for 4 hours.
Figure 3-1: Tertiary structure modeling of TVAG_393390 predicts cadherin-like protein function. (A) The most common high-quality 3D models of CLP predicted by Phyre2 revealed homology modelling to cadherin proteins. Characteristics of the aligned regions of these models are shown. (B) The predicted structure of TVAG_393390 generated by Phyre2 using one of the highest confidence models, mouse E-cadherin, as the template. (C) Inspection of TVAG_393390 sequence for LDRE, DXD, DXXD, x = any amino acid revealed four predicted Ca\textsuperscript{2+}-binding sites.
Figure 3-2 Cadherin-like protein (CLP) is localized to the surface of the parasites. (A) The parasite exogenously expressing CLP with two C-terminal HA tags (CLP-2X-HA) were stained for immunofluorescence microscopy using an anti-HA antibody (green) and 4’-6’-diamidino-2-phenylindole (DAPI) for nuclear staining (blue). This image is representative of 60 parasite viewed. (B) Brightfield image of (A). (C) The predicted topology of CLP generated with the TOPO2 program [66]. The predicted orientation is based on results from immunofluorescent assays performed in the absence or presence of a permeabilizing reagent (Fig. S3-1) as well as from the structural prediction from Fig. 3-1B. Predicted transmembrane residues are shown in blue.
Figure 3-3 CLP mRNA is up-regulated during host contact. T. vaginalis was exposed to host ectocervical cells (Ects) for 1h, 2h and 6h and the amount of CLP mRNA was quantified by qRT-PCR. CLP mRNA levels relative to 30 min exposure time point are shown. CLP is up-regulated 5, 19, and 20-fold after contact with Ects for 1h, 2h and 6h, respectively. 1 representative experiment of 3 independent experiments is shown. Data shown are means of triplicates ± standard deviations. *** = p ≤ 0.001
Figure 3-4 Generation of a CLP calcium-binding mutant. (A) Phyre2 and SusPect analyses identify the predicted calcium-binding site composed of D443 and D445 as the most sensitive to mutation (see Fig. S3-2 for analysis and comparison of other predicted calcium-binding sites in CLP). The height and color of the bars shown in the legend indicate the predicted functional impact of mutating the aspartate residue to the amino acids shown at the bottom of the histogram. Long and red bars in the histogram indicate that introduction of that particular amino acid would lead to the greatest phenotypic change while short blue bars have the smallest predicted phenotypic effect. (B) A CLP mutant that has D443 and D445 mutated to alanines (CLP-mut) was generated to investigate the functional effects of calcium-binding in CLP. Wild-type CLP and CLP-mut were exogenously expressed with an N-terminal GFP tag. As a negative control, parasites were transfected with an empty vector plasmid (EV). Immunoblot using an anti-GFP antibody confirmed that there are approximately equal amounts of CLP wild-type and CLP-mut overexpression. GAPDH is shown as a loading control. Fold represents the CLP expression levels between CLP and CLP-mut relative to CLP (=1).
Figure 3-5 CLP contributes to *T. vaginalis* adherence to host cells Ects. (A) Empty vector (EV) and CLP or CLP-mut transfectants were fluorescently labeled and incubated with Ects for 30 min followed by quantification of adhered parasites. The average fold changes of CLP and CLP-mut relative to EV were 3.5- and 0.83-fold. (B) The ability of CLP’s extracellular domain to compete with *T. vaginalis* binding to host cells was tested by the addition of 0.25 µg (7.7 nM), 1 µg (30.8 nM) or 4 µg (123.2 nM) of rCLP. Exogenous rCLP decreased host binding by 14%, 19%, and 21% compared to bovine serum albumin (BSA)-treated control. Both data represent 3 independent experiments, each performed in triplicate. Error bars indicate ± standard deviations. * = p ≤ 0.05, ** = p ≤ 0.01, *** p ≤ 0.001
Figure 3-6 CLP increases parasite clumping and the CLP-mut reversed the enhanced clumping. (A) The clumping ability of the parasites was assessed by quantifying an aggregate of 10 or more parasites. The parasites were incubated in the absence or presence of the host and in the absence or presence of 1 mM CaCl$_2$. Images shown are with 100X magnification and each white dot is a single parasite. Yellow arrows denote clumps with 10 or more parasites. (B) Quantification of clumping behavior observed in (A). The fold changes shown are relative to the EV - Ca + host condition. The results represent 3 independent experiments, each performed in triplicate. Error bars indicate ± standard deviations. *** = p ≤ 0.001
Figure 3-7 Death of Ects induced by *T. vaginalis* is increased by the wild-type CLP overexpression by 3.3-fold relative to the EV parasites and the CLP-mut parasites reduces the cytotoxicity level to 1.4-fold of the EV parasites. Data shown are means of triplicates ± standard deviations and one representative set of 3 independent experiments. *** = p ≤ 0.001
Supplemental Figure 3-1 To determine the topology of overexpressed CLP, indirect immunofluorescent assay in the presence or absence of a permeabilizing agent on C-terminally HA-tagged CLP was performed. (A) Bright green signal from anti-HA staining on permeabilized parasites versus faint green in non-permeabilized parasites (C) suggests that C-terminally tagged HA is on the intracellular side of the parasites. (B) and (D) are the brightfield images of (A) and (C), respectively. Green: HA. Blue: DAPI. These images are representative of ~30 parasites viewed under each condition.
Supplemental Figure 3-2 Mutational analysis of the rest of predicted Ca\(^{2+}\)-binding sites in CLP. Phyre2 and SuSPect analyses of the other 3 predicted Ca\(^{2+}\)-binding sites (from Fig. 3-1C). See Fig. 3-4A for the Ca\(^{2+}\)-binding sites which is the most sensitive to mutation. The height and color of the bars shown in the legend indicate the predicted functional impact of mutating the aspartate residue to the amino acids shown at the bottom of the histogram. Long and red bars in the histogram indicate that introduction of that particular amino acid would lead to the greatest phenotypic change while short blue bars have the smallest predicted phenotypic effect.
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Chapter 4:

CRISPR/Cas9-mediated gene modification and gene knock out in the human-infective parasite *Trichomonas vaginalis*
CRISPR/Cas9-mediated gene modification and gene knock out in the human-infective parasite

Trichomonas vaginalis

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The sexually-transmitted parasite Trichomonas vaginalis infects ~1/4 billion people worldwide. Despite its prevalence and myriad adverse outcomes of infection, the mechanisms underlying T. vaginalis pathogenesis are poorly understood. Genetic manipulation of this single-celled eukaryote has been hindered by challenges presented by its complex, repetitive genome and inefficient methods for introducing DNA (i.e. transfection) into the parasite. Here, we have developed methods to increase transfection efficiency using nucleofection, with the goal of efficiently introducing multiple DNA elements into a single T. vaginalis cell. We then created DNA constructs required to express several components essential to drive CRISPR/Cas9-mediated DNA modification: guide RNA (gRNA), the Cas9 endonuclease, short oligonucleotides and large, linearized DNA templates. Using these technical advances, we have established CRISPR/Cas9-mediated repair of mutations in genes contained on circular DNA plasmids harbored by the parasite. We also engineered CRISPR/Cas9 directed homologous recombination to delete (i.e. knock out) two non-essential genes within the T. vaginalis genome. This first report of the use of the CRISPR/Cas9 system in T. vaginalis greatly expands the ability to manipulate the genome of this pathogen and sets the stage for testing the role of specific genes in many biological processes.

Trichomonas vaginalis is an obligate extracellular, unicellular flagellated protozoan parasite and the causative agent of trichomoniasis, an infection that affects ~1/4 billion people worldwide. Infection can be asymptomatic or result in a variety of negative outcomes in women and men including vaginitis, cervicitis, pelvic inflammatory disease, prostatitis and urethritis. Alarmingly, public health studies have reported an increased incidence of drug resistance to the drug used to treat infections, metronidazole. Despite being a very common sexually transmitted infection, the basic mechanisms underlying pathogenesis and drug resistance are poorly understood. This is in part due to the slow adoption of molecular tools to study the genes and proteins involved in these processes. The development and application of cutting-edge gene technologies to manipulate this medically-important parasite are needed to further advance our knowledge of disease mechanisms.

Studies aimed to test the function of T. vaginalis genes have been limited by inefficient methods of genomic manipulation. Most reports have relied on isolates naturally lacking expression of particular genes or those directed by lab-acquired mutations that result in changes in gene expression. Only two studies have reported gene knockouts of non-essential proteins by integrating a drug resistance selection cassette in the gene locus relying purely on homologous recombination. A few reports have also utilized knockdown-based strategies utilizing antisense RNA expression and modified oligonucleotides. Knockdown techniques do not result in elimination of gene expression, which is often required to effectively test gene function, limiting the usefulness of this technology.

Recent advances in in vivo gene modification techniques have increased the ability to test gene function in both model and non-model organisms. The CRISPR (clustered regularly interspaced short palindromic repeats) system has recently revolutionized the ability to specifically target genes for modification in many organisms. 

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The CRISPR system is composed of two components, the Cas9 double stranded DNA nuclease of *Streptococcus pyogenes* and a user-customized version of its cofactor (guide RNA) to selectively target genes and induce double stranded breaks. The double stranded breaks are repaired by non-homologous end joining (NHEJ), microhomology-mediated end joining (MMEJ) or homology-directed repair (HDR) pathways. In many organisms, Cas9 gRNA expression induces repair of the double stranded break by the NHEJ and MMEJ pathways, which can result in insertion (NHEJ) or deletions (NHEJ and MMEJ) of nucleotides that may cause gene disruption and loss-of-function mutations. A more directed approach utilizes Cas9 gRNA in combination with a provided DNA template containing homology to the repair site to direct specific user-defined gene modifications. This approach requires a small region of donor DNA homology around the site of modification. More recently, CRISPR-based techniques have been applied to modify genes in a variety of parasitic protozoa and have been proposed for multicellular parasites.

Here we report the first application of CRISPR/Cas9 to alter genes in *T. vaginalis*. We demonstrate Cas9-gRNA directed modification of a nanoluciferase reporter gene using homology directed repair. Cas9-gRNA mediated knockout of two endogenous *T. vaginalis* genes by replacement with a drug resistance gene cassette were also achieved. This report lays the foundation for further development and utilization of this CRISPR/Cas9 to rigorously test gene function in this understudied parasite.

**Results**

**Expression of Cas9 and gRNA in Trichomonas vaginalis.** To date only two genes are reported to have been deleted (i.e. knocked out) from the haploid genome of the human parasite *T. vaginalis*. In order to improve the ability to modify *T. vaginalis* genes, we have developed the use of the CRISPR/Cas9 system as a means to directly modify genes in this parasite. One technical problem encountered when developing CRISPR/Cas9 in *T. vaginalis* was to overcome the toxicity associated with Cas9 expression in *T. vaginalis*. In initial experiments in which *cas9* was constitutively expressed using our standard *T. vaginalis* expression vector (pMasterNEO) and selection with G418, we were unable to select viable parasites, consistent with possible Caas9 toxicity. To address whether Cas9 was toxic in *T. vaginalis* cells selected for stable expression, we cloned the *cas9* gene as a fusion protein with an N-terminal FKBP destabilization domain (FKBP-DD). The presence of this domain on a protein results in the degradation of the protein in the absence of the stabilizing ligand, Shield-1. We also added both a hemagglutinin (HA) tag and two SV40 C-terminal nuclear localization motifs at the C-terminus of Cas9, to allow detection of the protein by immunoblot and to direct it into the nucleus, respectively (pMNfklp-cas9; Fig. 1A). Using this construct, we were able to transfect and G418 select for parasites with the pMNfklp-Cas9 plasmid in the presence or absence of the stabilization ligand Shield-1. Regulating the expression of Cas9 using the FKBP-DD and Shield-1 greatly reduced, but did not eliminate, basal expression of the FKBP-Cas9 protein in the absence of Shield-1 (Fig. 1B). We found that a low concentration of Shield-1 (0.1 μM) resulted in ~4-fold increased stabilization of the FKBP-Cas9 protein compared to parasites lacking Shield-1. A greater than 6.5-fold increase was observed using 1 μM Shield-1 (Fig. 1B). Moreover, we noticed no growth difference in parasites containing the pMNfklp-cas9 plasmid compared to parasites transfected with an empty vector (pMN:EV) when grown in the absence of Shield-1 (data not shown). Upon addition of Shield-1 parasite numbers were reduced in the pMNfklp-cas9 transfectants compared to the pMN:EV parasites which did not exhibit growth rate reduction (data not shown), confirming the initial suspicion that Cas9 might exhibit toxicity in *T. vaginalis*.

We then made a construct that would allow the expression of gRNA(s) using the pMasterNEO plasmid backbone (Fig. 1C, see Methods for details). The gRNA construct (pMN-ggRNA) contains the 360 bp sequence upstream of the U6 transcription start site (TSS), and is flanked at the 3′ end by the original 37 bp downstream of the U6 transcription unit to ensure proper transcription termination. Between these 5′ and 3′ regulatory elements, we inserted the first 20 nt sequence of the U6 snRNA transcript (5′-AGCTGGAGATATGCCTAGT-3′) followed by the conserved 76 bp scaffold sequence necessary for Cas9 targeting specificity and gRNA interaction, respectively (Fig. 1C). To evaluate whether transcription of this small RNA (gRNA mimic) would be driven by the putative *T. vaginalis* U6 promoter, we performed RT-PCR on total RNA extracts from cells transfected with this construct (pMN-ggRNA). Transcription was only detected from cells transfected with this construct (Fig. 1D) and was absent in negative control samples, transfected with an empty vector (pMN:EV). Next, sequential mutagenesis was done to replace the first 20 nt of *T. vaginalis* U6 snRNA, except for the first adenosine. None of the nt substitutions had an effect on transcription by end-point RT-PCR (data not shown). While the first transcribed nucleotide in the U6 transcript of metazoans and plants is typically a guanosine, *T. vaginalis* uses an adenosine. Since the first transcribed nucleotide is necessary for U6 transcription in other systems, but dispensable for the DNA target specificity of the Cas9 gRNA, we did not alter the +1 adenosine in our gRNA construct. The ability of the 360 nt upstream of the *T. vaginalis* U6 gene to drive transcription of an internally altered U6 gene indicates that the *T. vaginalis* U6 gene uses an upstream promoter as is found in metazoans and plants, and does not rely on internal promoter sequences as described for some unicellular eukaryotes (e.g. yeast and trypanosomes). These data also demonstrate that the *T. vaginalis* U6 promoter can be used to drive transcription of a sequence-customizable gRNA. In metazoan and plants, the core promoter of the U6 snRNA gene sits within the first ~700 bp upstream of the TSS. In *T. vaginalis*, the core promoter of the U6 snRNA gene is unknown however we observed by end-point RT-PCR that there was no effect on transcription when the upstream region was reduced from 360 bp to 131 bp (data not shown). All additional gRNA plasmids produced contain the 360 nt sequence 5′ of the U6 gene (Fig. 1C) preserving the +1 adenosine followed immediately by a 19 nt seed region that is customized to target the gene of interest.

**Increasing transcription efficiency of Trichomonas vaginalis.** Previous reports on transfection of *T. vaginalis* have used electroporation to deliver plasmids, linearized DNA or small oligonucleotides into the cell. Although sufficient to introduce DNA, transfection efficiency using this method is low. With the need
to introduce multiple DNAs into the same cell to achieve CRISPR/Cas9 editing, we sought to improve the transfection efficiency using the Amaxa nucleofection system. We first compared electroporation versus nucleofection using the nanoluciferase reporter system. Nanoluciferase (also termed NanoLuc) is a newly developed luciferase gene from a deep sea shrimp that produces stronger bioluminescence than the firefly luciferase (Promega). After normalization of parasite numbers, we subjected cells to either electroporation using our previously described protocol or nucleofection to introduce the wild-type nanoluciferase gene with a duplicate hemagglutinin (HA) tag at the 3′ end (pMN:nluc) (Fig. 2A). To optimize nucleofection, multiple programs and buffers were tested. Two of the programs tested (D-023 and X-001) and buffer (Parasite-2) resulted in detection of nanoluciferase signal compared to controls, however the signal was less than that obtained using standard electroporation (Supplemental Figure 1 and Fig. 2B). We found that one program (U-033) yielded better nanoluciferase signal, although parasite survival was low (<25% of untransfected controls at +24 hours post transfection, comparable to electroporation). Increased nanoluciferase signal was observed for two nucleofection buffers tested, T-cell and Parasite-1. As predicted, cells transfected with the pMN::nluc(stop) plasmid containing a premature stop codon that eliminates nanoluciferase protein production (Fig. 2A and B) produced only background signals. In comparison to the signal intensity achieved using nucleofection we found that nanoluciferase signals measured from parasites electroporated with pMN::nluc barely increased over background signal (Fig. 2B). This is consistent with the low frequency of transfection of parasites by electroporation, 3% at best for surviving
Figure 2. Parameters for transfection of *Trichomonas vaginalis* for detection of nanoluciferase activity. (A) Cartoon representation of nanoluciferase plasmids and predicted outcome for protein production. (B) A total of 1 × 10⁶ parasites were either nucleofected using the U-033 (“U”) or D-023 (“D”) programs (Amaxa) or mock (no nucleofection) with 10 μg of pMN::nLuc or pMN::nLuc(stop) in buffers Parasite-1 or T-cell (Lonza) or electroporated (975 μF, 350 V, Bio-Rad) in completed TYM media. Plasmids are pMN::nLuc, denoted as “nLuc” and pMN::nLuc(stop), denoted as “nLuc(stop)” where indicated. Parasites were immediately recovered in completed TYM media. After 24 hours, a total of 1 × 10⁶ parasites were collected and nanoluciferase activity was measured by NanoGlo assay (Promega). Graph represents the average relative luminescence value +/− standard deviation of two transfections per condition. (C) Samples transfected with the pMN::nLuc or pMN::nLuc(stop) constructs and the U-033 program from Fig. 2B were subjected to G418-selection and allowed to grow for an additional 24 hours (+48 hours total, post-transfection). Equivalent numbers of parasites (1 × 10⁶) were assayed for nanoluciferase activity and the graph represents the average relative luminescence value +/− standard deviation of two attempts per condition.
cells. Nucleofection with the U-033 program increased nanoluminescence signal 18-fold using Parastat-1 buffer and 30-fold using T-cell buffer, relative to signals detected using electroporation.

To determine if signal was stable and detectable at later time points, we passaged the parasites in the presence or absence of 100 μg/ml G418 to select for the pMN:nLuc and pMN:nLuc(stop) plasmids (Fig. 2C). We noted that G418 selection for the pMN:nLuc plasmid resulted in high levels of detectable nanoluminescence activity compared to transient nucleofection (Figures S2 and S3). The overall nanoluminescence signal for the unselected pMN:nLuc parasites was reduced at +48 hours post-nucleofection compared to +24 hours. Selection for parasites harboring the pMN:nLuc plasmids using G418 resulted in a dramatic increase with greater than 20-fold higher nanoluminescence signal compared to the transiently nucleofected (Fig. 2C). Electroporation did not yield the same increase (average >1.2-fold) over the unselected counterparts (Fig. 2C and Fig. S3). Based on these data, we selected the U-033 program and the T-cell buffer as the optimal conditions to use for nucleofection of this strain of T. vaginalis. Together, these data demonstrate that nucleofection greatly improves transfection efficiency of T. vaginalis.

CRISPR repair of a dead-luciferase gene in Trichomonas vaginalis. To test the activity of Cas9, we used the highly-sensitive nanoluminescence repair assay originally described by Striepen and colleagues. To express the Dd-Cas9 and gRNAs together in T. vaginalis, we first constructed a plasmid containing both elements as well as a puromycin acetyltransferase (PAC) gene, pCas9-gRNA (nLuc) (Fig. 3A). As outlined in Fig. 2A, we then preselected parasites containing the plasmid that expresses a dead luciferase (i.e. pMN:nLuc(stop)) using G418. We subsequently co-nucleofected preselected parasites with the pCas9-gRNA (nLuc) plasmid expressing a gRNA targeting the premature stop codon (Y180cde) region of the gene and pre-annealed oligos capable of both repairing the stop codon (“repaired”; ochreY180cde) and introducing a mutation that eliminates the PAM (AGG→AGG). Mutation of the PAM requires the correction of two sites by oligo template-directed repair, increasing the confidence that repair is Cas9-gRNA directed, and not due to random point mutation that changed a stop codon back to a tyrosine codon. Mutation also prevents further targeting of the repair site by Cas9-gRNA (nLuc). Transient nucleofection was used, without drug selection for expression of the gRNA or Cas9. Using this approach, we were able to demonstrate that CRISPR/Cas9 is capable of repairing the mutation to allow detection of nanoluminescence signal above background levels in T. vaginalis (Fig. 3).

The nanoluminescence repair assays were done using different lengths of oligos (50 nt and 125 nt) and we found that although the 50 bp annealed oligos yielded similar or better efficiency of repair, more variability was also observed (Fig. 3B). We also tested the use of PCR products corresponding to the 125 bp annealed oligos and the full length nLuc gene. These also repaired pMN:nLuc(stop), restoring nanoluminescence activity (Fig. 3C) however, the efficiency was not increased.

To further explore the ability to utilize CRISPR/Cas9 in T. vaginalis, we co-nucleofected parasites with pCas9-gRNA (nLuc), pMN:nLuc(stop) and either the 50 bp or 125 bp annealed repair oligos and selected for parasites containing both the pCas9-gRNA (nLuc) (puromycin selection) and pMN:nLuc(stop) (G418 selection) plasmids (Fig. 4A). Assaying for nanoluminescence signal 24 hours post-nucleofection, prior to drug selection, yielded a low overall signal compared to previous nucleofections (Fig. 4B). This is likely the result of requiring the same parasite to acquire all three DNAs to allow Cas9-directed repair. We also found that the 30 bp annealed oligo treatment did not work effectively, compared to previous experiments using parasites preselected to contain pMN:nLuc(stop) target. Conversely, treatment with the 125 bp annealed oligos yielded a ~six-fold increase in nanoluminescence signal over background (water).

We then subjected the nucleofected parasites to selection with G418 and puromycin for either 7 or 14 days and tested for nanoluminescence activity. The amount of nanoluminescence signal observed was very high for parasites nucleofected with the 125 bp repair oligo (Fig. 4C). This signal did not increase between the 7 and 14 day measurements. To confirm expression of intact nanoluminescence protein at 14 days, total protein was extracted and analyzed by immunoblot analysis for the presence of repaired nanoluminescence-2xHA. We observed expression of the protein in parasites nucleofected with all three DNAs (pCas9-gRNA (nLuc), pMN:nLuc(stop) and the 125 bp annealed repair oligos), but not in parasites where the pCas9-gRNA (nLuc) was replaced with an empty vector (pMPAcEV) (Fig. 4C, inset), consistent with the nanoluminescence activity assay. These data show that the repair of the dead luciferase was dependent on the presence of both essential components of the CRISPR/Cas9 system: the Cas9 and gRNA.

To confirm the repair of nanoluminescence by directly examining the DNA sequence of the luciferase gene, we cloned parasites from a selected population demonstrated to express the enzyme by activity and immunoblot assays, using limiting dilution. We then assayed 79 clones for nanoluminescence activity and observed that 91% (72 clones) gave signal above background, while 9% (7 clones) were below a cutoff of 30,000 relative luminescence units. The high percentage of clones expressing nanoluminescence indicates that the majority of parasites in the culture were expressing a repaired nanoluminescence gene. To support this observation, we collected total protein from 8 clones and performed immunoblot analysis (Fig. 4D, upper). Although varied in amount of observable signal, we were able to detect nanoluminescence signal in all 8 clones. We further PCR amplified and then sequenced the nLuc gene from total gDNA extracts of 8 clones (Fig. 4D, lower). In all 8 clones, the codon 17 contained a mixed population of guanine (GGT, glycine) and cytosine (GCA, alanine) sequence. Additionally, without exception, codon 18 contained a mixed population of thymine (TA) tyrosine) and adenine (TAA, ochre/stop). The presence of a mixed population of nLuc(stop) (GCTAATG or GCTAAT) and nLuc (repaired) (GCTAAT or GCTAAT) sequences in the examined clones is expected. Our previous study indicates that ~15–20 episomes are maintained per T. vaginalis cell, hence CRISPR/Cas9-mediated repair may not reach every single plasmid copy within a cell. In fact, repair of a single modified nLuc(stop) gene would likely be sufficient to be detectable by DNA sequencing and to produce enough luciferase activity for detection using the sensitive NanoGlo assay (Figure S2). Together, the data presented clearly demonstrate that the expression of Cas9 & gRNAs in T. vaginalis can mediate gene repair via the CRISPR/Cas9 pathway.
Figure 3. Nanoluciferase repair assay. (A) Upper: cartoon representation of nanoluciferase repair assay. Below: Cas9-gRNA(nluc) editing sites of nluc(stop) with mutation to create a premature stop codon (red, underlined). Also represented are the wild-type (nluc) and repaired (nluc(repaired)) versions. (B) A total of $5 \times 10^7$ parasites pre-selected for the pMN:nluc(stop) plasmid were nucleofected in T-cell buffer using the U-033 program (Amaza) with 10 μg or 100 μg of annealed repair oligos and 10 μg of pCas9-gRNA(nluc). Oligos contained the repair sequence and either 50 bp or 125 bp annealed repair oligos. Parasites were immediately recovered in completed TYM media $+/−$ 1 μM Shield-1 and assayed for nanoluciferase activity after 24 hours. Equivalent numbers of parasites ($1 \times 10^7$) were assayed for nanoluciferase activity and the graph represents the average relative luminescence value $+/−$ standard deviation for each condition. (C) A total of $5 \times 10^7$ parasites pre-selected for pMN:nluc(stop) were nucleofected in T-cell buffer using the U-033 program (Amaza) with 10 μg pCas9-gRNA(nluc) and either 100 μg of PCR-amplified 125 bp repair sequence (equivalent to ds repair oligos) or the full length nluc (“Full nluc PCR”) sequence. Parasites were immediately recovered in completed TYM media $+1 \mu M$ Shield-1 and assayed for nanoluciferase activity after 24 hours. Equivalent numbers of parasites ($1 \times 10^7$) were assayed for nanoluciferase activity and the graph represents the average relative luminescence value $+/−$ standard deviation for each condition.
CRISPR knockout of the Ferredoxin-1 and Mif genes from the *T. vaginalis* genome. To test whether the CRISPR/Cas9 system we have established is capable of knocking out a gene in the complex *T. vaginalis* genome, we first attempted to knock out the ferredoxin-1 (Fd-1) gene previously shown to be non-essential to the parasite. A neomycin resistance (NeoR) gene flanked by ~1000 bp of the 5′ and 3′ UTRs of Fd-1 was constructed (Fig. 5A). Parasites were co-nucleofected with either the pMPAC:EV or the pCas9-gRNA(Fd) plasmid that expresses a gRNA targeting the 5′ end of the gene and an ~2800 bp PCR product containing the NeoR genes flanked by Fd-1 gene 5′ and 3′ UTRs (Fig. 5A). After selection for parasites resistant to G418, parasites were screened for the presence of the NeoR gene in the Fd-1 locus, using PCR and primers that sit just outside the Fd-1 UTRs included in the ~2800 bp PCR product. Positive amplions of expected size were observed in the pCas9-gRNA(Fd) nucleofected parasites, while the pMPAC:EV parasites did not grow as expected. We then obtained cloned parasites from a NeoR positive population using limited-dilution cloning. Two clones (2-2 G4 "ΔFd-1 clone 1" and 2-1 D9 "ΔFd-1 clone 2") were randomly selected and tested for the presence of the NeoR gene as well as the wild-type Fd-1 gene (Fig. 5B). Confirmation of the sequence of all PCR products clearly demonstrates that both clones contain the NeoR gene in the Fd-1 locus and to also lack the endogenous Fd-1 gene (Fig. 5B). Immunoblot analysis using an antibody that detects Fd-1 also confirmed the loss of Fd-1 protein in both clones (Fig. 5C).
Figure 5. CRISPR-directed KO of ferredoxin-1 and mif. (A) Cartoon representation of the KO scheme using nuclease of parasites with a linearized neo resistance gene flanked by the ferredoxin-1 UTRs and a plasmid containing Cas9 and Fd gRNA (pCas9-gRNA(Fd)) followed by selection for G418-resistance (image not to scale). The resulting parasites were then screened for the presence of the KO and wt alleles using sequence-specific primers (horizontal arrows). (B) PCR analysis of G418-resistant clones for the presence of the neo resistance gene in the ferredoxin-1 locus. PCR screens utilized primers specific to the neo resistance gene (yellow arrows in 5′ A) combined with primers specific to the ferredoxin-1 locus (blue arrows in 5′ A) in regions outside the region present in the original KO PCR introduced by nuclease. The 5′ end PCR screen (upper panel) predicts a product of 1665 bp and the 3′ PCR screen (middle panel) predicts a product of 1236 bp if the neo gene is present in the ferredoxin-1 locus. Lowest panel: PCR analysis of the ferredoxin-1 gene in clones. Amplification utilized primers specific to the wild-type ferredoxin-1 (yielding a 324 bp product, if present). The PCR products for the different panels were run in parallel from reactions on 100 ng of the indicated genomic DNA extracts and resolved on separate agarose gels then imaged using a Bio-Rad Gel Doc and ImageLab software. (C) Immunoblot protein analysis of the clones yielding positive PCR products for KO in Fig. 5B. Total protein extracts were resolved by SDS-PAGE and immunoblotted using anti-Ferredoxin-1 and TβGAPDH antibodies. All immunoblots contained equal amounts of the same experimental samples and were analyzed in parallel using a Bio-Rad Gel Doc and ImageLab software. (D) Cartoon representation of the KO scheme using nuclease of parasites with a linearized neo resistance (Neor) gene flanked by the mif UTRs and a plasmid containing Cas9 and mif gRNAs (pCas9-gRNA(mif)) followed by selection for G418- and puromycin-resistance (image not to scale). The resulting parasites were then screened for the presence of the KO and wt alleles using sequence-specific primers (horizontal arrows). (E) PCR analysis of G418 and puromycin-resistant clones for the presence of the neo resistance gene in the mif locus. PCR screens utilized primers specific to the neo resistance gene (yellow arrows in 5′ D) combined with primers specific to the mif locus (orange arrows in 5′ D) in regions outside the region present in the original KO PCR introduced by nuclease. The 5′ end PCR screen (upper panel) predicts a product of 1290 bp and the 3′ PCR screen (middle panel) predicts a product of 1220 bp if the neo gene is present in the mif locus. Lowest panel: PCR analysis of the mif gene in clones. Amplification utilized primers specific to the wild-type mif gene (yielding a 580 bp product, if present). The PCR products for the different panels were run in parallel from reactions on 100 ng of the indicated genomic DNA extracts and resolved on separate agarose gels then imaged using a Bio-Rad Gel Doc and ImageLab software. (F) Immunoblot protein analysis of the clones yielding positive PCR products for KO in Fig. 5E. Total protein extracts were resolved by SDS-PAGE and immunoblotted using anti-MIF and TβGAPDH antibodies. All immunoblots contained equal amounts of the same experimental samples and were analyzed in parallel using a Bio-Rad Gel Doc and ImageLab software. Full length blots/gels are presented in Supplementary Figure S5.
To further test the CRISPR/Cas9 system, we attempted to knock out the single copy \textit{mif} gene, which codes for a homologue of the human macrophage migration inhibitory factor (MIF) and was recently reported to affect host cell physiology. A neomycin resistance (NeoR) gene flanked by ~1000 bp of the 5' and 3' UTRs of \textit{mif} was constructed (Fig. 5D). To illustrate the ability to use CRISPR/Cas9 in multiple \textit{V. vulnificus} strains, the 5'RC2 strain was used. Parasites were co-nucleofected with the pCas9-gRNA(mif) plasmid expressing a gRNA targeting the 5' end of the gene and an ~2800 bp PCR product containing the NeoR gene flanked by \textit{mif} gene 5' and 3' UTRs (Fig. 5D). After selection for parasites resistant to G418 and paromycin, parasites were screened for the presence of the NeoR gene in the \textit{mif} locus, using PCR and primers that sit just outside the \textit{mif} UTRs included in the ~2800 bp PCR product. Two positive clones (one expected size were observed in the pCas9-gRNA(mif) nucleofected parasites (Fig. 5E). Control reactions wherein parasites were nucleofected with only the linear KO PCR product survived G418 selection, however no positive PCR signal was observed for the NeoR gene in the \textit{mif} locus (Supplementary Figure S4). We further examined these control parasites lacking the pCas9-gRNA(mif) plasmid by screening 800 cells, split into 40 subpopulations of 20 parasites each to directly test whether the NeoR gene was in the \textit{mif} locus. No PCR product was detected in any of the 40 subpopulations (Supplementary Figure S4). This is in contrast with obtaining positive PCR signal for NeoR integration at the \textit{mif} locus in 39/40 subpopulations of the parasites nucleofected with both the pCas9-gRNA(mif) plasmid and the linear KO PCR product. To confirm that the \textit{mif} gene was knocked out in these NeoR PCR positive populations, parasites were cloned using limited-dilution cloning. Two clones ("\textit{mif} clone 1" and "\textit{mif} clone 2") were randomly selected and tested for the presence of the NeoR gene in the \textit{mif} locus and the absence of the wild-type \textit{mif} gene (Fig. 5E). Confirmation of the sequence of all PCR products clearly demonstrated that both clones contain the NeoR gene in the \textit{mif} locus and also lack the endogenous \textit{mif} gene (Fig. 5E). Immunoblot analysis using an antibody that detects MIF likewise confirmed the loss of the MIF protein in both clones (Fig. 5F). Together, our findings demonstrate that CRISPR/Cas9 can be used to disrupt gene sequences in the \textit{V. vulnificus} genome by a homology-directed repair pathway.

**Discussion**

Here we describe the development of a system to achieve CRISPR/Cas9 gene modification and gene knockout in \textit{T. vaginalis}. Three technical advances made this possible. One was to decrease the toxicity of Cas9 when expressed in \textit{T. vaginalis} by using FKBP-DD/Shield-1 regulated expression of the endonuclease. The second was to increase transfection efficiency using nucleofection. These together, with the use of a sensitive nanoluciferase reporter assay to detect homology directed repair of a premature stop codon to wild-type firstly described by Streipen and colleagues, provided a sensitive assay to facilitate identifying optimal nucleofection parameters. Finally, we were able to utilize the strong RNA polymerase III \textit{T. vaginalis} U6 promoter to drive transcription of a sequence-customizable gRNA. This, the first report of CRISPR/Cas9 gene modification in \textit{T. vaginalis}, will greatly enhance the ability to test gene function in this pathogen.

It has been previously reported that expression of Cas9 in various organisms leads to cell growth issues. We found that initial attempts at plasmid-based Cas9 expression by transfection and selection of \textit{T. vaginalis} repeatedly failed. Different reports have addressed decreased cell growth by transient transfection of \textit{cas9} mRNA in plasmids or transfecting with \textit{Cas9} mRNA or homologous genes. We chose to regulate Cas9 expression at the protein through fusion to the FKBP destabilization domain (FKBP-DD), which by default selects proteins for proteolytic degradation. In the presence of a stabilizing ligand Shield-1, the protein accumulates. Using this method allowed the regulation of cellular Cas9 levels in \textit{T. vaginalis}, which in turn yielded viable parasites expressing sufficient, but not toxic, levels of Cas9.

Testing the biological role of genes in \textit{T. vaginalis} has previously been severely compromised by limited technologies to alter the highly, repetitive genome of this parasite. Only two previous reports have successfully knocked out \textit{T. vaginalis} genes and knockdown of gene expression is limited to four reports. Lacking the ability to knock out gene expression, researchers have primarily relied on episomal over-expression of gene variants to study protein function. One reason that homologous gene replacement has been hampered in \textit{T. vaginalis} is the poor transfection efficiency achieved with standard electroporation. Here, using nucleofection, we have greatly enhanced the transfection efficiency by 20-fold or greater as compared to that achieved using electroporation method. This enhancement of transfection efficiency has enhanced both the probability of successful modification of genes and the ability to detect the modifications by sensitive techniques such as nanoluciferase activity assays or PCR. Thus, developing successful nucleofection methods allowing sensitive screening and selection for gene modifications was key to our ability to achieve and monitor CRISPR/Cas9 modification of \textit{T. vaginalis} genes.

Our analyses of the \textit{T. vaginalis} genome for homologs of NHEJ pathway components indicated that this pathway is absent or highly divergent from other organisms. Based on this observation, we focused on homology-directed repair pathways to establish the use of CRISPR/Cas9 in this parasite. We were able to detect CRISPR activity using the sensitive technique of early termination mutation repair first used to demonstrate CRISPR in \textit{Cryptosporidium parvum}. Using a 125 bp repair template of annealed oligonucleotides with flanks on both sides of the CRISPR cut site, we were able to repair an early termination codon restoring nanoluciferase activity as demonstrated by enzyme activity assays, immunoblot analysis and DNA sequencing. The repair template makes two specific point mutations both changing the stop codon to a tyrosine and a proximal glycine to alanine. The nanoluciferase repair assay demonstrates that small oligonucleotides are sufficient to modify multi-target episomal maintained genes and thus the length of a homology template may not have to be large. Repair is due to homology-directed repair (HDR) using the oligonucleotide template, as opposed to NHEJ activity. Notably, in the absence of the HDR template, we did not observe signal of repaired nanoluciferase above background signal, consistent with the lack of NHEJ-mediated repair.
Nucleofection of pre-selected pMN:nlux(stop) parasites with pCas9-gRNA(nlux) and repair oligos resulted in an increase of nanoluciferase activity. This varied based on concentration of oligos with 100 μg working more effectively than 10 μg. Transient co-nucleofection of all DNA components (pMN:nlux(stop), pCas9-gRNA(nlux) and repair oligos) resulted in lower nanoluciferase signals, however, this could be overcome using drug selection to enrich for parasite populations containing DNAs required for CRISPR/Cas9-mediated repair. Using drug selection and cell cloning by limited dilution, we found that 91% of the clones had nanoluciferase activity above background. DNA sequencing of the clones confirmed the modification of the nlux(stop) to nlux(repaired), demonstrating CRISPR-Cas9 mediated repair of the nlux(stop)-Cas9 gene.

We also have demonstrated the ability to use CRISPR-Cas9 mediated homology-repair to eliminate two genes (ferredoxin-1 and mlf) from the genome of T. vaginalis. CRISPR-mediated gene knock-out was achieved in two different T. vaginalis strains. The methods we have developed to increase transfection efficiency for both plasmids and KO cassettes and to direct CRISPR/Cas9 gene modification will greatly increase the success rate of gene knockouts in this parasite. The expanded toolkit for molecular modification of the T. vaginalis genome will significantly enhance the ability to assess gene function in this medically-important pathogen.

Methods

Parasite culture. Parasites (T. vaginalis strain B7RC227 and MSA 1103425) were maintained by daily passage in modified TYM media28 supplemented with 10% heat-inactivated horse serum (Sigma), 10 U/ml penicillin/10 μg/ml streptomycin (Invitrogen), 180 μM ferrous ammonium sulfate and 28 μM sulfosuccinic acid (completed TYM media). To select for the pMasterPAC (pMPAC)53,76 or pMasterNEO (pMN)22 plasmids, puromycin (AG scientific) or G418 (Invitrogen) was added to cultures to a final concentration of 30 μg/ml and 100 μg/ml, respectively.

Plasmid constructs. Plasmid pMN:cas9 was created by PCR amplification of the human codon optimized cas9 gene of plasmid pM1920214 using primers For-hcas9-SacI and Rev-hcas9-BamH. All oligos used in this study are listed in Supplemental Table 1. The resulting hcas9 product was digested with SacI/BamHI and ligated to pMN:fkbp. The resulting construct was then PCR amplified using primers For-dS-Hucas9 and Rev-MNSUTR to identify mutate an internal SacI sequence. This product was then used in megaprimer PCR with primer For-MN-SacI and subsequently digested with SacI/BamHI and ligated into pMN:fkbp to produce pMN:fkbp-cas9. To create the pMPAC:fkbp-cas9 construct, pMN:fkbp-cas9 was digested with SacI/BamHI and the product cloned into the pMPAC-empty vector plasmid (pMPAC-EV). The gRNA construct was generated by PCR amplification of a synthetic gRNA construct (Table S1) containing the U6 seed region and gRNA scaffold flanked upstream by the 360 bp 5′ of the U6 start nucleotide and downstream with 37 bp of the 3′ UTR of the U6 gene using primer U6_Sacl_F1 and downstream 3′ UTR with primer U6_Sacl_R1. The resulting product was then digested with SacI and ligated to the unique SacI site in pMasterNEO53. The nlux gRNA was constructed by megaprimer amplification of the gRNA scaffold using primers For-Nluc-gRNA and U6_Sacl_R1 and then used in a megaprimer PCR with U6_Sacl_F1. The resulting PCR product was digested with SacI and ligated to pMPAC:fkbp-cas9, the final construct. The nlux gRNA was abbreviated as pCas9-gRNA(nlux).

Construction of nanoluciferase plasmids utilized codon-optimized plasmid templates of nlux (wt) and nlux (Y180chne, "dead"/stop) from the Nluc-Neo and Dead Nluc-Neo plasmids (a generous gift from the Boris Striepen laboratory, University of Georgia)64, respectively. For each nlux variant, a megaprimer PCR was performed utilizing primers For-MN-SacI and Rev-Nluc-Nde to amplify the 5′ UTR of a SCS on pMasterNEO with a portion of the 5′ end of the nlux gene. The resulting products were gel-purified and used with primer Rev-Nluc-Nde and ligated to pMasterNEO such that the nlux (or repaired) sequence would be fused to a 2×HA epitope tag when translated. These plasmids were termed pMN:nlux and pMN:nlux(stop).

For construction of the pCas9-gRNA construct targeting the ferredoxin-1 gene (TVAG_003900), the 20 nt seed region of pMPAC:fkbp-cas9-gRNA(nlux) was replaced after the adenines reside start with 19 nucleotides of the ferredoxin-1 gene (residues 5-23, 5′-TCT CTC AAG TTT GCC CCT T-3′) which lie 5′ upstream of a TGG PAM sequence. The product was constructed by PCR amplification with primers For-gRNA-Fer-1 and U6_Sacl_R1. The resulting PCR product was purified and used in a second PCR with primer U6_Sacl_F1, followed by digestion with SacI and cloning into pMPAC:fkbp-cas9-gRNA(nlux). The resulting plasmid was termed pCas9-gRNA(Fer).

For construction of the pCas9-gRNA construct targeting the mlf gene (TVAG_219770), a dual gRNA was constructed. For each of the gRNAs the 20 nt seed region of pMPAC:fkbp-cas9-gRNA(nlux) was replaced after the adenine reside start with 19 nucleotides of the mlf gene. The mlf sequences "g300" (5′-CAA AGA GTG CTG AGG ACT G-3′) and "g301" (5′-CCA AAG AGT GTG AGA GAC T-3′) were used to construct 19 nucleotides at the A in the seed region. Briefly, "g300" gRNA was constructed using PCR primers For-gRNA-F-300 and U6-KpnI-R. The resulting product was purified and used in a second PCR with primer U6_Sacl_F1. For "g301" gRNA was constructed using PCR primers For-gRNA-F-301 and U6_Sacl_R1. The resulting product was purified and used in a second PCR with primer U6-KpnI-F. Both PCR products were digested with KpnI, ligated together and gel purified. The resulting ligated product was then digested with SacI and ligated into pMPAC:fkbp-cas9-gRNA(nlux) producing the plasmid pMPAC:fkbp-cas9-gRNA(mlf).

Construction of the ferredoxin KO cassette utilized 1000 bp upstream of the ferredoxin-1 start codon and 1000 bp downstream of the stop codon. Briefly, the neo resistance gene was PCR amplified using primer For-Neo-Eco and Rev-Neo-Bam. The resulting product was cloned into pSC-A (Stratagene) to produce the plasmid pSC-A:neo. PCR amplification of the 5′ UTR of ferredoxin utilized For-Fer-Kpn and Rev-Fer-Eco. The product was then digested with KpnI and EcoRI and cloned into the pSC-A:neo plasmid resulting in plasmid pSC-A:5′UTR-Fd-neo. Amplification of the 3′ UTR of ferredoxin utilized For-Fer-Bam and Rev-Fer-Sac. The
resulting product was digested with BamHI and SacI and cloned into the pSC-A::5'UTR Fd-neo plasmid to produce plasmid pSC-A::5'UTR Fd-neo plasmid (term pKO-Fd).

Construction of the mif KO cassette utilized the same method used to generate the ferredoxin KO cassette except using mif gene-specific primers. Briefly, the 1000 bp upstream of the mif start codon and downstream of the stop codon were ligated to the Neo gene. PCR amplification of the 1000 bp 5' UTR upstream of the mif start codon utilized primers For-MIF-Kpn and Rev-MIF-Xho. The resulting product was digested with KpnI and XhoI and ligated to the pSC-A::neo plasmid resulting in plasmid pSC-A::5'UTR-mif-neo. PCR amplification of the 1000 bp 3' UTR downstream of the mif stop codon utilized primers For-MIF-Bam and Rev-MIF-Sac. The resulting products were digested with BamHI and SacI and ligated to pSC-A::5'UTR-mif-neo to produce plasmid pSC-A::5'UTR-mif-neo (term pKO-mif).

**Repair oligos/PCR.** Repair oligos were constructed to target mutations in the nhuc(stop) gene (Operon). Annealing of the 125 nt repair oligos utilized Repair-oligo-F and Repair-oligo-R. Annealing of the 50 nt repair oligos utilized Repair-oligo-F50 and Repair-oligo-R50. An equal concentration of each repair oligo (200 μg) was annealed in 1X NEB T4 ligase buffer to create a stock of anneal oligos. For PCR amplification of the 125 bp repair sequence, primers For-125bp and Rev-125bp were used on the annealed 125 bp oligos above. For full length PCR of nhuc, the digested (SacI/KpnI) and gel purified template of pMN:nhuc was used as a PCR template using primers For-pMN-5UTR and Rev-Nhuc-Kpn. The resulting product was treated with DpnI and then gel purified and amplified using the same primers. The final PCR product was precipitated, resuspended and quantified.

PCR amplification of the ferredoxin-1::neo knockout utilized the primers For-Fer-Kpn and Rev-Fer-Sac on a gel-purified product from a KpnI and SacI digestion of pKO-Fd. PCR amplification of the mif::neo knockout utilized the same method described above with mif-specific primers For-MIF-Kpn and Rev-MIF-Sac. The resulting PCR products were treated with DpnI and confirmed to be free of template plasmid. The final KO PCR product was precipitated, resuspended and quantified.

**Protein and Immunoblot analysis.** For Shield-1 (Clontech) titration, parasites were seeded at 1 x 10⁶ cells/ml and treated with the indicated concentrations for 24 hours. For all immunoblot analyses, total protein was extracted and analyzed by SDS-PAGE using equal amount of soluble protein (10 μg). Protein was then transferred to a membrane which was subsequently incubated with an anti-Cas9 antibody (Clontech). For TgVPDGH detection, the membrane was incubated with an anti-TgVPDGH antibody (Cocalico Biologicals). Images were captured and analyzed by Bio Rad Gel Doc and ImageLab software v. 5.1, Bio Rad). Resulting values were normalized to TgVPDGH signal and presented as the average ±/− the standard deviation from two independent analyses. For detection of nanoluminase-2xHA protein, total protein was extracted from parasites, resolved by SDS-PAGE, transferred to a PVDF membrane and incubated with an anti-HA antibody (Govance). For immunoblot analysis of the Ferredoxin-1 and MIF KO samples, total soluble protein from parasite samples was resolved by SDS-PAGE, transferred to a membrane and incubated with an anti-Ferredoxin-1β, anti-MIFβ and anti-TgVPDGH (Cocalico Biologicals) antibodies. Uncropped images for the corresponding immunoblots are included in Figure S5.

**Nanoluciferase activity assay.** A total of 1 x 10⁶ parasites were harvested per reaction (in duplicate), washed in ice-cold 1x PBS with 5% sucrose (w/v) and resuspended at a concentration of 1 x 10⁶ cells/100 μl in the same PBS/sucrose solution. Each 100 μl aliquot was mixed with 100 μl of completed NanoGlo luciferase assay reagent (Promega) and transferred to round bottomed white walled 96-well plates (Costar). Luciferase activity was analyzed by measurement with a Victor 3 model 1420 plate reader at 460 nm. Relative luminescence signal was measured in duplicate or triplicate for each assay and the average signal calculated ±/− standard deviation.

**Transfection of T. vaginalis using nucleofection and electroporation.** Parasites were collected by centrifugation and resuspended in 100 μl T-cell Parasite-1 or Parasite-2 buffers (Lonza) and the indicated amounts of DNA and parasites (1 x 10⁶ or 5 x 10⁶). Typically, 10 μg plasmid(s) and/or 100 μg of anneal oligos or PCR product was added. After five minutes of incubation at room temperature, parasites were nucleofected using an Amaxa nucleofector using the programs U-033, D-023 or X-100. For parasite electroporation, 5 x 10⁷ parasites were resuspended in 100 μl of completed TYM media and placed into electrocuvettes with the indicated DNA. Parasites were then electroporated using a single pulse of 350 V with 975 μF capacitance using a Bio Rad Gene Pulser IIβ. For both nucleofection and electroporation parasites were immediately resuspended into 50 ml of fresh completed TYM media. During recovery, parasites were treated with 1 μM Shield-1 where indicated.

Knockout of the ferredoxin-1 gene (TVAG_003900) in the MSA 1103 T. vaginalis strainβ utilized the nucleofection conditions (U-033 and T-cell buffer) with the substitution of pCas9-gRNA(Fd) or pMPAC_eEV and 50 μg of KO PCR product in duplicate. Parasites were recovered in completed TYM media for four hours and then selected for resistance to 50 μg/ml of G418. After an additional 16 hours, the parasites were harvested and resuspended in fresh completed TYM media and 100 μg/ml G418. After outgrowth from selection to 1 x 10⁶ parasites/ml, genomic DNA was extracted from ~1.5 x 10⁷ parasites samples and PCR screening was done for the presence of the KO allele at the ferredoxin-1 locus (see below). The cultures with signal positive for ferredoxin-1 disruption were diluted and re-screened using limiting dilution until clones were obtained. The clones were screened for the presence of the neo gene at the ferredoxin locus as well as for the presence of the wild type ferredoxin-1 gene. All PCR's were performed on 100 ng of purified genomic DNA and utilized NEB Phusion polymerase, per the manufacturers' instructions. Screening for the wild-type gene used primers For-003900-1 and Rev-003900-1 (PCR cycle: 95°C- 30 sec, 60°C- 30 sec, 72°C- 90 sec). Screening for the 5’ end of the ferredoxin-1::neo KO locus used primers For-003900-5UTR-1 and Rev-Neo-Int3 (PCR cycle: 95°C- 30 sec, 60°C- 30 sec, 72°C- 90 sec). Screening
for the 3’ end of the \textit{ferredoxin-1} \textit{neo} KO locus used primers For-Neo-Int2 and Rev-003900-3UTR-2 (PCR cycle: 95°C - 30 sec, 64°C - 30 sec, 72°C - 90 sec). All PCR products were confirmed by DNA sequencing.

Knockout of the \textit{mfj} gene (TVAG_219770) in \textit{T. vaginalis} strain BT2C2 utilized nucleofection conditions described for the knockout of \textit{ferredoxin-1} above, except the D-023 program and V-kit buffer were utilized, and pCas9-gRNA(\textit{mfj}) or water (for the negative control) and 100 μg of KO PCR product was used. BT2C2 parasites were recovered in completed TYM media for 24 hours and selected for resistance to 100 μg/ml of G418, and 30 μg/ml of puromycin. DNA was extracted from the parasites that received pCas9-gRNA(\textit{mfj}). Negative control parasites that did not receive pCas9-gRNA(\textit{mfj}) were only selected with 100 μg/ml of G418. After an additional 24 hours, the parasites that received pCas9-gRNA(\textit{mfj}) were re-dosed with the 100 μg/ml G418, and 30 μg/ml of puromycin and control parasites were re-dosed with just 100 μg/ml G418. Drug selected parasites were then sub-populated into 20 cells per well in a 96-well plate. When the parasites reached ~1 x 10^7 parasites/ml, genomic DNA was prepared for PCR screening. Initial screening was done utilizing the 3’ end of the \textit{mfj:neo} KO withprimers For-219770-5UTR-1 and Rev-Neo-Int4 (PCR cycle: 95°C - 30 sec, 55°C - 30 sec, 72°C - 90 sec). A subpopulation with the correct Neo replacement based on 5’ UTR PCR analyses was cloned and then screened for the presence of the NeoR gene in the \textit{mfj} locus using both the 5’ primer described above and the 3’ primer For-Neo-Int3 and Rev-219770-3UTR-1 (PCR cycle: 95°C - 30 sec, 55°C - 30 sec, 72°C - 90 sec). Screening for the wild-type \textit{mfj} gene used primers For-219770-1 and Rev-219770-1 (PCR cycle: 95°C - 30 sec, 55°C - 30 sec, 72°C - 30 sec). All PCR products were confirmed by DNA sequencing.

\textbf{RT-PCR analysis of gRNA expression.} Total RNA was isolated from ~1.5 x 10^7 parasites using the Direct-zol RNA Miniprep kit (Zymo) and then fractionated into small RNAs using the RNA Clean and Concentrator kit (Zymo). Small RNAs were further treated with the TURBO DNA-free kit (Ambion). Poly-A tails were then added using the Tailing Kit (Ambion). Small RNAs were quantified and cDNA created by utilizing an equivalent amount of RNA in the Supernscript III first strand synthesis system for RT-PCR (Invitrogen) using the provided oligo-dT\textsubscript{18} primer. All PCRs were performed on one microtiter of the resulting cDNA and utilized NEB Phusion polymerase, per the manufacturers’ instructions. For the gRNA, primers For-gRNA and Rev-gRNA were used to detect the gRNA and primers For-U6 and Rev-U6 were used to detect the U6 snRNA (for both, the PCR cycle: 98°C - 10 sec, 55°C - 20 sec, 72°C - 8 sec). All samples were resolved on agarose-TBE gels, stained with ethidium bromide and imaged by Bio-Rad Gel Doc, as above.

\textbf{Data Availability.} All data generated or analysed during this study are included in this published article (and its Supplementary Information files).

\textbf{References.}


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Author Contributions

B.D.J., Y.-P.C., A.S.-B. and P.J.J. conceived of and designed experiments. B.D.J., Y.-P.C., B.M.M., S.E.W. performed experiments and B.D.J., Y.-P.C., A.S.-B. and P.J.J. analyzed data. B.D.J. and P.J.J. wrote the manuscript. P.J.J. secured funding. All authors reviewed the manuscript.

Additional Information

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Chapter 5:

Examination of cytokine induction from human innate immune cells by *Trichomonas vaginalis* co-infected with *Mycoplasma hominis* and by the presence of *Tv* protein *TvMIF*
Abstract

Most infections caused by *Trichomonas vaginalis* (*Tv*) are asymptomatic but chronic and symptomatic infections do exist. To understand how the parasite transitions from acute to long-term infection, we characterized innate immune responses against *T. vaginalis* by setting up an *in vitro* co-culture system to detect cytokine responses elicited by *T. vaginalis* from human monocyte-derived dendritic cells and macrophages. By including *T. vaginalis* symbiont *Mycoplasma hominis* in the study, we found that majority of the cytokine responses are induced by *Tv* in the presence of *M. hominis* but not when *M. hominis* is absent. In addition, a laboratory adapted *Tv* strain triggers significantly stronger inflammatory responses than a clinical *Tv* strain, indicating strain variations could contribute to various infection outcomes. We also examined innate immune responses to *Tv* homologue (*TvMIF*) of human cytokine macrophage migration inhibitory factor (*HuMIF*). *TvMIF* does not induce significant amount of inflammatory cytokine responses from innate immune cells. This could be because the parasite has co-evolved with the host so the parasite homologue of the human cytokine preserves its anti-apoptotic effect on the parasite but does not trigger host immune response during infection. These results lay the groundwork for future studies on host immune responses to *T. vaginalis* infection.

Introduction

Immune responses induced by *T. vaginalis* in infected individuals are largely unknown but remain interesting given that majority of the infections are asymptomatic while chronic symptomatic cases also exist [1]. As a first step towards fighting the chronic *Tv* infection, it is critical to understand how human innate immune system such as monocytes, macrophages, and dendritic cells respond to the parasite. Human monocytes, derived from bone marrow, are known
to circulate and survey bloodstream [2]. When monocytes pass through capillary walls and enter tissues, they differentiate into macrophages which are 5-10 times larger in size and more phagocytic than monocytes [2]. The main defensive roles of macrophages involve phagocytosis, presenting foreign antigens, and initiating inflammatory responses by releasing small signaling molecules called cytokines [2]. Dendritic cells (DCs) develop from bone marrow-derived hematopoietic stem cells but the precise origin of dendritic cells is not well understood [3]. In vitro, DCs can be differentiated from bone marrow derived-monocytes by receiving granulocyte-macrophage colony stimulating factor (GM-CSF) and cytokine IL-4 [4]. DCs in tissues are activated by receiving signals such as cytokines from macrophages during an infection [5]. It is also known that during herpes simplex viral infection in mice, bone-marrow-derived precursors differentiate into vaginal epithelial DCs and become activated [6]. Understanding the type of responses DCs and macrophages mount against Tv will provide insights into how our innate immune system communicates with adaptive immune responses to result in diverse infection outcomes.

Cytokine responses induced by T. vaginalis and its symbionts or products have been examined sporadically. Lipoglycan on T. vaginalis surface [7], T. vaginalis infected with its endosymbiotic viruses (TVVs) [8], exosomes secreted by the parasite [9], live T. vaginalis, and its secretory proteins and lysates [10][11][12][13] induce cytokine production from epithelial host cells or host immune cells. Mycoplasma hominis is a natural T. vaginalis symbiont and the co-infection exists at a range from 5% to 90% of clinical cases depending on the regions of the isolation [14]. However, only recently, our laboratory systemically tested cytokine responses comparing both clinical and laboratory adapted T. vaginalis strains and both M. hominis-free and M. hominis-positive isogenic strains on primary human monocytes [15] and found that most of
the cytokine responses are triggered by *T. vaginalis* symbiont *M. hominis* instead of the parasite itself except for IL-8 [15], contradictory to the highly immunogenic properties of *T. vaginalis* found in previous reports [11][12][13]. As a result, it is worthwhile to revisit the cytokine responses induced by *T. vaginalis* in the presence and absence of *M. hominis* from other immune cells such as macrophages and dendritic cells.

*Trichomonas vaginalis* macrophage migration inhibitory factor (TvMIF) is homologue of human pleiotropic cytokine HuMIF [10]. Plenty of pro-inflammatory cytokine properties of mammalian MIFs were characterized in mouse models during infections [16][17][18]. For example, in MIF-depleted mice, pro-inflammatory cytokines from macrophages were reduced in response to *Mycobacterium tuberculosis* infection, or exposure to lipopolysaccharide (LPS) and gram-negative bacteria compared to the wild-type (WT) mice [16][17]. Like HuMIF, parasite MIF homologues modulate human immune responses. *Entamoeba histolytica* MIF [19], *Toxoplasma gondii* MIF [20], and *Plasmodium yoelii* MIF [21] and TvMIF [10] can induce pro-inflammatory cytokine production from host. On the other hand, helminth parasite *Brugia malayi* MIF promotes Th2 response that alleviates inflammatory response [22]. To gain a thorough understanding of immunomodulatory roles of TvMIF, we examined cytokine responses from human macrophages and dendritic cells that were previously unstudied.

We used human bone marrow-derived primary cells as our model because human is the natural host of *T. vaginalis*. Furthermore, current *T. vaginalis* mouse model requires immunosuppressive treatment that is not suitable for studying immune responses from the host [23]. Macrophages and dendritic cells are differentiated from primary human monocytes with a method that is standardly employed in the immunology field [4][24]. In this chapter, we will
present cytokine responses from human macrophages and dendritic cells to *T. vaginalis* and a secreted cytokine homologue TvMIF.

**Results**

**Differentiation of macrophages and dendritic cells**

By treating isolated primary human monocytes with granulocyte-macrophage colony stimulating factor (GM-CSF) only or GM-CSF + IL-4 for 4 days, we obtained monocyte-derived macrophages (MDM) or monocyte-derived dendritic cells (MDDC), respectively. We verified the differentiation by examining the surface markers such as CD11c and CD14 [25][26][27] and the cell sizes. MDM has higher CD11c and CD14 expression than monocytes on day 0 and it has a significant size increase shown by a shift of forward side scatter (FSC) to the right on a flow cytometry histogram (Fig. 5-1A). For MDDC, CD14 expression is reduced and the size increased (higher FSC) after the differentiation (Fig. 5-1B). After confirming that the differentiation was successful, we then co-cultured *T. vaginalis* with MDM or MDDC overnight and collected the supernatant for cytokine detection.

**Examination of cytokine production by live *T. vaginalis* in the presence or absence of *M. hominis***

We tested a laboratory adapted *Tv* strain G3 and a clinical isolate MSA1103, and *M. hominis*-negative and *M. hominis*-positive for both of these strains. Consistent with our previous examination using primary monocytes in our laboratory [15], most of the cytokine production from both MDM and MDDC are present in *M. hominis*-positive strain but not very much in *M. hominis*-negative strain (Fig. 5-2 & 5-3). In addition, we found that in *M. hominis*-positive strains, laboratory adapted G3 induced much more cytokine productions than clinical isolate
MSA1103 for almost all the inflammatory cytokines tested including IL-1β, IL-6, IL-8 and TNF-α from both MDDC and MDM (Fig. 5-2 & 5-3). IL-12 was only stimulated in MDDC but not MDM (Fig. 5-2E & 5-3E). The differences between G3- and MSA1103-stimulated cytokines were more dramatic in MDDC than MDM. To test if G3 contains more M. hominis than MSA1103, we performed qRT-PCR to measure M. hominis levels by using M. hominis-specific primers and normalized it to Tv housekeeping gene, β-tubulin. The amounts of M. hominis were similar in G3 and MSA1103 (Fig. 5-3G) so this suggests that G3 is more immunogenic than MSA1103 but only when M. hominis is also present.

Interestingly, anti-inflammatory cytokine IL-10 is also induced in M. hominis-positive strains and is significantly more in G3 than MSA1103 when produced by MDM (Fig. 5-3D). The result could suggest that T. vaginalis and M. hominis co-infection presents a mixture of antigens that stimulate both inflammatory and anti-inflammatory responses from the host immune cells.

**Examination of cytokine production in the presence of TvMIF**

After we confirmed the differentiation of MDM and MDDC as described in Fig. 5-1, we added 100 ng/ml, 1 µg/ml, 10 µg/ml of recombinant TvMIF (rTvMIF) or recombinant HuMIF (rHuMIF) or PBS to MDM or MDDC and the culture was incubated for 16h at 37ºC. We found that none of the rTvMIF or rHuMIF concentration tested induced significant pro-inflammatory cytokine production (IL-1β, IL-6, IL-8 and TNF-α) from MDDC (Fig. 5-4 & 5-5). 10 µg/ml of rHuMIF caused IL-6, IL-8 and TNF-α production from MDM but none of the rTvMIF induced the pro-inflammatory cytokines from MDM (Fig. 5-5).

**Discussion**
We examined a panel of cytokine responses induced by live *T. vaginalis* and rTvMIF in MDM and MDDC. By including *M. hominis*-positive, *M. hominis*-negative, laboratory adapted and clinical *Tv* strains, we observed that *Tv* symbiont *M. hominis* plays a significant role in the cytokine stimulation, consistent with our previous study that *T. vaginalis*-induced cytokines from human monocytes are greatly dependent on the presence of *M. hominis* [15]. In addition, all the cytokines tested in the *M. hominis*-positive samples were more abundant in the laboratory adapted strain than the clinical strain, suggesting that the immunogenic property of *M. hominis* is also dependent on *Tv* strains. The striking differences in the cytokine productions between strains and whether *M. hominis* is present could, at least partially, explain the diverse outcomes of *T. vaginalis* infection in humans.

We showed that *T. vaginalis* stimulates immunosuppressive cytokine IL-10 from MDDC and MDM, consistent with Song et al. observation that *T. vaginalis*-derived secretory products up-regulates anti-inflammatory IL-10 from MDDC [11]. On the other hand, the pro-inflammatory cytokines (TNF-α, IL-1β and IL-6) induced by *M. hominis*-positive *T. vaginalis* samples in MDM are consistent with previous observations with live *Tv*, *Tv* lysates and human plasma-opsinized live *Tv* [12]. Notably, we only saw abundant cytokines from MDDC and MDM using *M. hominis*-positive parasites whose presence was not examined in previous studies on cytokine stimulated by *T. vaginalis* or its products. These results emphasize the importance of taking *M. hominis* into account as majority of the pro-inflammatory responses produced by MDDC and MDM were only present in *M. hominis*-positive samples. In this study, we also observed strain-to-strain variations. As a result, it is critical to compare between different *Tv* strains in the presence or absence of *M. hominis* before drawing any conclusion whether a *T.*
vaginalis protein, secretory products, and Tv molecules have true immunogenic effects on the host.

Induction of either pro-inflammatory or immunosuppressive cytokines by Tv from immune cells have been reported in independent studies [1]. However, this is the first time that we found the presence both anti-inflammatory and immunosuppressive responses induced by Tv by the same cell types. The same observation was made in sepsis during pathogen infection and it was reasoned that the increase in pro-inflammatory cytokines is to fight off the infection and the immunosuppressive cytokines are made to prevent organ failure from excessive inflammation [28]. Moreover, it was also found that Mycoplasma-infected human tumor exosomes stimulated both pro-inflammatory IFN-γ and immunosuppressive IL-10 from B cells but these cytokines were absent in Mycoplasma-negative exosomes [29]. This further highlights that the immunogenic properties of Tv described previously may attribute to the presence of M. hominis.

Contrary to seen in human monocytes [15] and MDM, IL-12 that induces Th1 responses was elicited in MDDC by live Tv (Fig 5-2E & 5-3E). The different immune cell types used may contribute to the observed differences. Future analyses are required to determine whether our laboratory adapted strain that induces significantly more pro-inflammatory cytokines from both MDM and MDDC are more susceptible to attack by immune cells.

In this study, rHuMIF induced IL-6, IL-8 and TNF-α responses produced by MDM as HuMIF plays a pro-inflammatory role in human cells although only at high concentration (10 µg/ml). However, almost no cytokine tested was induced by rTvMIF or lower concentrations of rHuMIF. We suspect that this is because of the production method of the recombinant proteins we used. Both rTvMIF and rHuMIF have C-terminal His tags that can inhibit the oligomerization
of MIF proteins [30]. The trimeric MIF proteins are primarily responsible for binding to MIF receptor and induce the cytokine responses [31]. As a result, our rMIF with C-terminal His tags may not be able to induce the cytokine production. Another explanation for the difference is that most of the pro-inflammatory roles of mammalian MIFs were studied in MIF knockout mice during pathogenic infections [16][17][18][32]. Purified MIF proteins may not stimulate cytokine responses on isolated immune cells in the absence of an infection. We previously observed that rTvMIF was able to induce IL-8 signal from monocytes [10]. In several studies, macrophages and DC were found to produce less cytokines than their precursor monocytes when treated with the same agonists. Seow et al and Daigneault et al reported that macrophages produce less pro-inflammatory cytokines than monocytes in response to LPS and TLR1/2 agonist Pam3CSK4, respectively [33][34]. MDDC produce less pro-inflammatory cytokine than monocytes in response to LPS, *Staphylococcus epidermis* and TLR2 ligand [35]. The less abundant cytokine production from MDDC and MDM may explain the absence of rTvMIF-induced cytokine release.

TvMIF has been shown to bind HuMIF receptor CD74 [10] and both MIFs can inhibit the parasite death under nutrient starvation [36]. However, TvMIF may not be as immunogenic to the host as we hypothesized. This could be the parasite reaching a balance of mimicking host survival factor while avoiding the immune consequence of having a pro-inflammatory cytokine in the parasite. TvMIF may demonstrate an intricate example of co-evolution of host and parasite.

**Material and Methods**

*Trichomonas vaginalis* culture
*T. vaginalis* strains G3 and MSA1103 were cultured in complete Diamond’s media as previously described [37]. The cells were passaged for 2 weeks or less. *M. hominis*-negative *T. vaginalis* culture was treated with 50 µg/ml of chloramphenicol and 5 µg/ml of tetracycline daily for at least 3 days and *M. hominis*-positive *Tv* was cultured without the antibiotic treatment [15].

**Acquisition of human monocyte-derived macrophages and monocyte-derived dendritic cells**

Primary human monocytes were isolated by adherence to plastic by UCLA Virology Core. Monocyte-derived macrophages (MDM) and monocyte-derived dendritic cells (MDDC) were differentiated *in vitro* as previously described [4][15]. Briefly, 20 ng/ml of GM-CSF (Biolegend) was added to induce MDM differentiation. 20 ng/ml of GM-CSF and 20 ng/ml of IL-4 were added to induce MDDC differentiation. Both MDM and MDDC were differentiated for 4 days in RPMI 1640 media supplemented with 10% fetal bovine serum (Thermo Fisher Scientific), 100 U/ml penicillin + 100 µg/ml streptomycin (Thermo Fisher Scientific), GlutaMAX (Thermo Fisher Scientific) and MEM non-essential amino acids (Thermo Fisher Scientific) before co-culture with *Tv*. Successful differentiation was determined by increase in sizes for both cell types, increase in CD11c and CD14 for MDM, and decrease in CD14 for MDDC. The differentiated cells were stained with 20 µg/ml CD11c-FITC (BioLegend) and 50 µg/ml of CD14-PE (BioLegend) for 30 min at 4 °C in the dark. The stained cells were read on BD LSR Fortessa and the data were analyzed by FlowJo 7.6.

**Co-culture of MDM or MDDC with *Tv* and cytokine production analysis**
M. hominis-free and M. hominis-positive Tv were counted and resuspended in complete RPMI media to be incubated with MDM or MDDC at an multiplicity of infection (MOI) of 1 for 16 hours. 100 ng/ml lipopolysaccharide (LPS) (Sigma) alone or 100 ng/ml LPS + 1000 U/ml interferon gamma (IFN-γ) (BioLegend) was used as positive controls to stimulate cytokine production. After the incubation, the cells were centrifuged and the supernatants were collected for cytokine detection. Cytometric Bead Array (Becton-Dickenson) was used to measure IL-1β, IL-6, IL-8, IL-10, IL-12 and TNF-α cytokines. The results were read on BD LSR Fortessa and the data were analyzed by FlowJo 7.6.

**rTvMIF induction of cytokine**

Recombinant TvMIF protein was made according to the method described in [10]. In brief, TvMIF or HuMIF in pET200 expression vector grown in *Escherichia coli* BL21 Star™(DE3) cells (Thermo Fisher Scientific) was purified using His FastFlow Columns (GE healthcare) against the C-terminal His tags on both proteins. Endotoxin was removed from both proteins using Detoxi-Gel Endotoxin Removing Columns (Thermo Fisher Scientific). The protein concentrations were determined by Pierce BCA Protein Assay Kit (Thermo Fisher Scientific).

100 ng/ml, 1 µg/ml or 10 µg/ml of rTvMIF was added to MDM or MDDC to induce cytokine production. PBS and LPS (Sigma) were used as the negative and positive controls, respectively. The supernatants of MDM and MDDC were collected by spinning the cells and the cytokine detection was measured and analyzed by the same method described in *Tv*-induced cytokine analysis.

**Quantification of M. hominis in T. vaginalis by qRT-PCR**
*T. vaginalis* along with *M. hominis* total RNA was extracted with TRIzol (Life Technologies) and treated with TURBO DNase (Invitrogen). cDNA and qRT-PCR were performed using SuperScript III (Thermo Fisher Scientific) and Platinum SYBR Green qPCR SuperMix-UDG kits (Thermo Fisher Scientific). The *M. hominis*-specific primers were described in [38] and their Cq values were normalized to *T. vaginalis* β-tubulin gene Cq values [39].
Figure 5-1: MDM and MDDC differentiation were verified by the surface markers and their size changes. (A) After 4 days of differentiation, MDM gained CD11c and CD14 signals and had a size increase (FSC). (B) MDDC (day 4) had a lower CD14 and smaller size (FSC) compared to day 0.
Figure 5-2: Cytokines IL-1β, IL-6, IL-8, IL-10, IL-12 and TNF-α from MDDC are induced by *M. hominis*-positive live *T. vaginalis* and significantly more in the laboratory adapted strain G3 than the clinical strain MSA1103. (A-F) Unstimulated is PBS-treated. LPS or LPS + IFN-γ are the positive controls. Live G3 or MSA1103 parasites were co-cultured with MDDC at multiplicity of infection = 1 for 16 hours. * = p ≤ 0.05, ** = p ≤ 0.01, *** = p ≤ 0.001
Figure 5-3: Cytokines IL-1β, IL-6, IL-8, IL-10, IL-12 and TNF-α from MDM are induced by *M. hominis*-positive live *T. vaginalis* and significantly more in the laboratory adapted strain G3 than the clinical strain MSA1103. (A-F) Unstimulated is PBS-treated. LPS or LPS + IFN-γ are the positive controls. Live G3 or MSA1103 parasites were co-cultured with MDM at multiplicity of infection = 1 for 16 hours. (G) Comparison of *M. hominis* (Mh) quantity in each strain was quantified by qRT-PCR. The Mh rRNA-encoding gene was normalized to *T. vaginalis* housekeeping gene β-tubulin. The Mh levels are expressed as fold change (G3 = 1). ns = not significant *, ** = p ≤ 0.05, *** = p ≤ 0.01, **** = p ≤ 0.001
Figure 5-4: TvMIF does not induce significant amounts of pro-inflammatory cytokine responses from MDDC. (A-D) 100 ng/ml, 1 μg/ml, 10 μg/ml of rHuMIF or rTvMIF were added to the MDDC culture and the culture was then incubated for 16 hours. PBS-treated was the negative control and LPS-treated was the positive control.
Figure 5-5: HuMIF but not TvMIF induces significant amounts of pro-inflammatory cytokine responses from MDM. (A-D) 100 ng/ml, 1 μg/ml, 10 μg/ml of rHuMIF or rTvMIF were added to MDM culture and the culture was then incubated for 16 hours. PBS-treated was the negative control and LPS-treated was the positive control. * = p ≤ 0.05, ** = p ≤ 0.01
References


Chapter 6:

Summary and discussion
Summary

Trichomonas vaginalis is a human-infective protozoan that is highly prevalent yet commonly neglected [1]. With the development of molecular tools [2] and efforts from researchers in the field, understanding how the parasite establishes and maintains the infection has become possible. The goals of this dissertation are identification of survival factor of the parasite (Chapter 2), characterization of a Tv cadherin-like protein involved in host cell binding and killing (Chapter 3), efforts in adapting CRISPR-Cas9 to edit T. vaginalis genome (Chapter 4), and examination of immune responses to Tv and TvMIF from human macrophages and dendritic cells (Chapter 5). These studies reveal that the parasite depends on multiple factors to establish infection and the development of novel techniques is necessary to better understand the mechanisms underlying infection.

Insight into Tv survival factor and pathways

TvMIF is the first identified survival factor in T. vaginalis to our knowledge. The overexpression of TvMIF enhances the parasite survival more than 10-fold and knockout of the protein decreases their survival up to 16-fold under nutrient-poor conditions (Chapter 2). We show that TvMIF inhibits the apoptosis of the parasite by lowering reactive oxygen species in the parasite to enhance their survival (Chapter 2). Both TvMIF and huMIF are able to increase the parasite survival, suggesting the cross-talk between the parasite and its host (Chapter 2).

Identification of specific players involved in the survival pathway in T. vaginalis is laborious. Very few proteins that play a role in apoptosis in other eukaryotic systems are known in protozoan parasites. Metacaspases in yeast are believed to have the same function as mammalian caspases which are involved in apoptosis [3]. However, the specific roles of metacaspases in parasites are not entirely clear [4]. Other players in mammalian apoptotic
pathways such as pro-apoptotic factors/anti-apoptotic factors cannot be easily identified in the *T. vaginalis* genome based on primary sequence. As a result, to identify *T. vaginalis* survival pathways, transcriptomic and proteomic approaches that examine global differential expression are required. The lack of survival in TvMIF knockout (KO) cells suggests that comparison of wild-type (WT) cells and KO cells gene expression during serum starvation will provide information on unidentified factors involved in the survival pathway. In addition, *T. vaginalis* has conserved mitogen-activated protein kinases (MAPKs) and Akt that are known to be activated by phosphorylation downstream of huMIF activation. Phosphoproteome on WT versus TvMIF KO parasites under serum starvation may aid the process of identifying survival players.

HuMIF receptor CD74 is the key to huMIF downstream signaling including activating MAPK/Erk and Akt/PI3k pathways, and inhibiting apoptotic p53 [5][6][7]. We have made attempts to identify TvMIF receptor in *Tv* although unsuccessfully. Affinity chromatography using recombinant TvMIF and pulldown with tagged TvMIF expressed in *T. vaginalis* have not been able to reveal any promising candidates. For example, the adjusted p-values were not significant from 3 biological replicates or a candidate was not pulled down in independent experiments. Pulldown has always been challenging in *T. vaginalis* because commercially available tags used for pulldown lack specificity in the organism and the majority of the proteins in the genome are not annotated well. As a result, to identify a novel receptor in *T. vaginalis*, adaptation of new techniques is required. For example, proximity labeling which can label nearby proteins without strong interactions such as BioID used in *Toxoplasma gondii* [8][9] may assist in receptor identification. TurboID, an improved BioID that can label as quickly as 10 minutes and has stronger activity on cell surface may significantly speed up identification of surface receptors in *T. vaginalis* [10]. Alternatively, use of tandem affinity purification such as
PTP (ProtC-TEV-ProtA) tags employed in *Trypanosoma brucei* is likely to enhance the specificity of the identified products [11][12].

**Characterization of a *Tv* cadherin-like protein in host adherence and killing**

Previous surface proteome has allowed us to identify several adherent factors in *Tv* that mediate host binding [13]. As more and more *Tv* adherence factors such as the parasite exosomes and lipoglycan have been discovered, it became clear that host binding by *Tv* is mediated by a multifactorial process [13][14][15]. Cadherin-like protein (CLP) described in Chapter 3 was identified in the surface proteome comparing less adherent versus adherent parasite strains [13]. In addition, CLP was also found in the *Tv* exosomal proteome [16], indicating that the adherent factor on the parasite surface may mediate host binding by being delivered to another parasite or host via exosomal fusion. Using our CRISPR-Cas9 system in *Tv* to knock out the CLP gene to further characterize its role in host adherence and its role in exosome-mediated host adherence is currently underway and will provide insight to these questions.

A clumping phenotype was observed when CLP is overexpressed in the parasite. The behavior is reminiscent of *T. vaginalis* tetraspanin 8 (TvTSP8)-overexpressing parasites and highly adherent parasite strains [17]. The role of parasite aggregation in *T. vaginalis* infection is not entirely understood. Clumping increases the number of parasites in contact with the host and this behavior may enhance the chance of successful infection. In addition, when parasites aggregate, the outside parasites may be able to protect the parasites in the center from attack from host immune cells as a potential mechanism to increase parasite survival inside the host. These hypotheses are yet to be tested.

The interaction between human E-cadherin protein and tetraspanin Co-029 determines cell motility of colon cancer cell [18]. Several *T. vaginalis* TSP proteins have been found to be
localized to the parasite surface and re-localize during host contact [17][19]. The clumping phenotype of both TvTSP8-overexpressing and CLP-overexpressing parasites indicate that these two surface proteins may complement each other in host binding. Whether TvTSP8 or other parasite proteins act as CLP partners requires further investigation.

**Optimization of CRISPR-Cas9 techniques in T. vaginalis**

Adaptation of CRISPR-Cas9 system in *T. vaginalis* has significantly improved the ability to knock out (KO) genes in the parasite [20]. However, the need to use homology-directed repair CRISPR-Cas9 pathway in the parasite requires the researchers to build knockout (KO) cassette [20] and amplify the KO cassette by PCR which is extremely time-consuming and expensive. KO genes in a population also requires extensive PCR screening to obtain a parasite clone that has the gene replacement at the right locus. In addition, it is unfeasible to KO genes with multiple copies since that would require various selectable markers for multiple gene replacement and multiple rounds of screening. As a result, CRISPRi knockdown system using catalytically inactive Cas9 guided by guide RNAs to block transcription initiation or transcription elongation [21][22] is considered to be useful in *T. vaginalis*. For example, CRISPRi does not require cloning of KO cassettes and PCR screening for the right clone. Furthermore, targeting genes with multiple copies with CRISPRi is more feasible than CRISPR-Cas9 knockout as CRISPRi blocks the transcription of the same genes without the need to replace each of them with different drug selectable markers. For these reasons, repurposing CRISPR-Cas9 in *T. vaginalis* may provide a more efficient platform to control gene expression.

**More information is yet to be explored on immune responses to Tv**

From our data testing cytokine responses induced by live *T. vaginalis* in the presence or absence of *Mycoplasma hominis*, we found that *M. hominis* plays a significant role in inducing
cytokine releases from human macrophages and dendritic cells. In addition, the *M. hominis*-induced cytokine responses are remarkably higher in a laboratory adapted strain G3 than a clinical strain MSA1103 we tested. To draw a meaningful conclusion on whether clinical strains are less immunogenic than lab-adaptive strains, a significantly larger number of *Tv* strains will need to be tested.

An interesting follow-up question to our observation in Chapter 5 that a laboratory adapted strain G3 is significantly more immunogenic than a clinical strain MSA1103 is whether G3 is more susceptible to killing by host immune cells than MSA1103. For example, the induction of cytokines by G3 parasites may alert more immune cells to attack them than MSA1103 parasites. However, lacking a good animal model is likely to be an impediment to truly understanding host immune responses to *Tv*. As a result, our laboratory is now developing a male mouse model because female mice as *Tv* hosts have not been a successful model given that the infection often requires treatments of estrogens and/or immunosuppressants [23][24][25]. The existing data lack good quantification of the infection rates and evidence of long-term infection [23][24][25]. Treatment with immunosuppression resulted in disruption of host immune system that failed to reflect the immune responses during a natural infection. Once a mouse model is established, it will open doors to many hypotheses related to *T. vaginalis* infection such as how different each *Tv* strain is in terms of infectivity, how *Tv* regulates its gene expression during host contact, and how host immune cells respond to *Tv* infection in vivo. Efforts towards developing an animal model and other technologies for studying *T. vaginalis* are the keys to truly understanding how to prevent the infection.
References


