**Trichomonas vaginalis** virulence factors: an integrative overview

Robert P Hirt

**ABSTRACT**

The elusive nature of *Trichomonas vaginalis*, the most common, non-viral, sexually transmitted pathogen has hampered our knowledge of its significance for human health for over 150 years. The combination of epidemiology, molecular cell biology, immunology and more recently genomics and other allied omics data, are all contributing at shedding new light onto what is increasingly recognised as a significant human pathogen leading to important health sequelae due to multifaceted interactions with its human host, the human microbiota, bacterial pathogens and viruses. The integrations of these various data are contributing in important ways to refining our understanding of the parasite pathobiology and virulent factors. Indeed, it is increasingly recognised that to rationalise the development of effective prophylactic and therapeutic treatments for human pathogens it is important to integrate the broadest possible spectrum of human-microbial-parasite-virus interactions in relation to qualitative and quantitative variations in the human innate and adaptive defence responses. This short review aims at providing an integrative overview of *T vaginalis* virulent factors by taking into account the importance of the human-microbiota-parasite-virus interplay in human health. It also highlights selected cellular characteristics of the parasite often overlooked in the biological and medical literature.

**INTRODUCTION**

Ever since its discovery in 1836 by the French medical doctor and microbiologist, Alfred F. Donné, the sexually transmitted obligate extracellular mucosal parasite *Trichomonas vaginalis* (TV) has progressively grown, rather sluggishly, from a perceived insignificant commensal to an important pathogen, inducing significant health sequelae in both men and women, and with adverse pregnancy outcomes.1 2 Some of the most recent developments in our understanding of TV pathobiology include its epidemiological association with HIV over and above high-risk members of the population in terms of exposure to sexually transmitted infections (STI),2 3 and a positive association with aggressive prostate cancers.4 The dramatic impact on human health of the HIV pandemic and resource-limiting conditions experienced by hundreds of millions of people worldwide have combined in highlighting the pathogenic significance of this common human parasite—one that is facilitating the spread of HIV and is associated with a number of reproductive complications including sterility, preterm birth, underweight newborn babies and cervical cancer.4

Significantly, TV infects a large fraction of individuals, in particular men, without overt symptoms, complicating its diagnosis and control.5 6 Furthermore, and particularly relevant in the context of a discussion on TV virulent factors, the parasite’s elusive nature also renders the characterisation of the molecular and cellular basis of its pathobiology less straightforward, as highlighted for similarly elusive bacterial pathogens.6 7 Indeed, there is growing recognition that the outcome of human-symbiont interactions (encompassing mutualists, commensals and parasite/pathogens) cannot be fully understood without considering the host immunological status and host-associated microbiota.7 There is a need to integrate the full range of human-microbe interactions in all its diversity to comprehend their different contributions and influence on each other, and how these impact both positively and negatively on human health.5 8 These considerations also underscore the limitations of the reductive approach implied by the original four Koch postulates and their molecular children, used to characterise pathogens and their virulence factors.6 8

Incorporating the host response to exposures to microbes is now considered to be an integral part of defining pathogens and their virulence factors.6 8 9 Indeed, these issues are increasingly recognised by the research community working on TV as illustrated in a number of recent reviews.4 9–12 Hence, an integrative overview of TV virulence factors is presented here, acknowledging the complex host-microbiota-parasite-virus interplay in influencing the outcomes of human-TV interactions.

**CELLULAR AND MOLECULAR BASIS OF TV VIRULENCE**

A number of recent reviews have discussed various aspects of well-characterised and more speculative virulence factors of TV10–13 Only a selection of TV virulence factors will be considered here and integrated with some of the most recent data on TV molecular cell biology and human innate and adaptive responses to TV. Some important gaps in our knowledge of TV pathobiology will also be highlighted. The evolutionary origin of the best-studied organelle of the parasite, the hydrogenosome, and the roles of hydrogenosomal enzymes in the parasite’s physiology and as a drug target were recently reviewed10 14 and are not covered here.

TV is a flagellated microbial eukaryote known to exist in several cellular forms (figure 1). The two best characterised forms are the trophozoite, a free-swimming, flagellated, pear-like cell, and an amoeboid form, with a pancake shape characterised by an important increase in surface contact; this is rapidly induced upon trophozoite contact in vivo with epithelial cells from the vagina, cervix, urethra, prostate and extracellular matrix (ECM) proteins...
Proteomics investigations. An initial study of the phospho-
investigate the molecular cell biology of sequence and its annotation represent an invaluable resource to relevance during the infection process. The draft genome aggregates will be an essential prerequisite to investigate their differentiations into these different forms. Identifying specific molecular markers for the different cellular forms and aggregates will be an essential prerequisite to investigate their relevance during the infection process. The draft genome sequence and its annotation represent an invaluable resource to investigate the molecular cell biology of TV by providing specific molecular leads and allowing comparative transcriptomics and proteomics investigations. An initial study of the phosphoproteome of TV trophozoites, amoeba and pseudocysts grown in vitro suggested differential protein phosphorylation profiles, consistent with specific signalling mechanisms occurring in the different cellular forms of the parasite that are induced upon specific environmental triggers.

Clinical isolates of TV have been accumulated over the years from many regions of the world. Comparisons of their capacity to bind and kill human (and other species) cell lines in vitro have demonstrated important variations between TV isolates. Accordingly, these capacities are considered important virulent traits for TV. One of the currently best-characterised adhesins mediating parasite binding to host tissue is TV lipoglycans (TvLG), the most abundant surface molecules of the parasite. Investigating TvLG also led to the identification of galecin-1, the only identified human receptor for TV so far. TvLGs are also known to modulate inflammatory responses of epithelial cells and macrophages. A proteomics survey of TV surface proteins identified a total of 411 proteins, confirming a number of in silico-predicted surface proteins, and also a host of novel and important candidate virulent factors. By contrasting strains with low versus high level of adhesion to vaginal epithelial cells (VECs) in vitro, 11 proteins were shown to be more abundant on the cell surface of the highly adhering strains. These include proteins annotated as hypotheticals—which have no sequence similarity to any other proteins in protein databases. A high level of expression of two of these hypothetical proteins increased the binding to VECs of a poorly binding TV strain. Other novel candidate cell surface virulence factors identified in this proteomics survey included three tetraspanins, which are membrane proteins involved in signalling modulating adhesion, motility and tissue invasion in other systems; all these are key processes underlying TV pathobiology.

The characterisation of one TV tetraspanin demonstrated that its expression and cellular localisation was modulated upon TV binding to VEC, and that it plays a role in regulating migration of the parasite through a surrogate ECM gel, strongly supporting tetraspanins as important TV virulence factors. These studies highlight how little we know about key aspects of TV biology, and how genomics and allied omics provide important tools to investigate host-parasite interactions. Related proteomics studies on TV are reviewed elsewhere.

During the infection process, TV actively phagocytoses human cells, bacteria and fungi to obtain nutrients. Similarly, receptor-mediated endocytosis by TV is also considered important in order to internalise nutrients and iron, and to neutralise host defence proteins. However, there are currently no functionally characterised TV genes encoding surface proteins mediating the specific binding process underlying phagocytosis or endocytosis. Bioinformatic analyses of candidate transmembrane proteins have identified a number of cytoplasmic tails possessing classic signals for endocytosis, some of which were supported by proteomics data (table 1). In silico identification of candidate genes regulating membrane trafficking have also identified a surprisingly large repertoire of proteins, with some gene families unexpectedly larger than those encoding human homologues (eg, ∼300 TrRab vs ∼70 HsRab GTases), further suggesting that phagocytosis and endocytosis are important processes for the parasite and represent promising targets for interfering with parasite virulence.

TV interactions with human-associated bacteria and viruses

The phagocytosis of human microbiota by TV, including bacteria and fungal cells, has the potential to induce dysbiosis
Table 1: Example of *Trichomonas vaginalis* (TV) candidate transmembrane proteins potentially mediating endocytosis*

<table>
<thead>
<tr>
<th>Protein family</th>
<th>RefSeq accession</th>
<th>Protein length</th>
<th>**[FY]**NPX[<strong>FY</strong>] motif</th>
<th>Acidic cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>BspA</td>
<td>XP_001308117</td>
<td>735</td>
<td>NENPIF</td>
<td>DDPFADFMDS</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001584393</td>
<td>1092</td>
<td>1FTNFLP</td>
<td>DDPFADDPDFDOP</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001321489</td>
<td>600</td>
<td>YDNPF</td>
<td>DDPFADQNFND</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001321493</td>
<td>582</td>
<td>YDNPF</td>
<td>DDPFADQNFND</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001321496</td>
<td>594</td>
<td>FDNPFL</td>
<td>DDPFADQNFNE</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001321499</td>
<td>447</td>
<td>FDNPFL</td>
<td>DDPFADQNFNE</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001321508</td>
<td>733</td>
<td>FDNPLY</td>
<td>DDPFADQFQNE</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001323867</td>
<td>405</td>
<td>LDNPFL</td>
<td>DDPFAEDFEEK</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001328761</td>
<td>549</td>
<td>FSNPFL</td>
<td>DDPFANDFEQG</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001328763</td>
<td>1142</td>
<td>YINPLF</td>
<td>DDPFADSFEDH</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001328767</td>
<td>834</td>
<td>FSNPFL</td>
<td>DDPFADSFEEH</td>
</tr>
<tr>
<td>BspA</td>
<td>XP_001312366</td>
<td>703</td>
<td>ADNPIF</td>
<td>DDPFADQNDND</td>
</tr>
<tr>
<td>Pmp</td>
<td>XP_001579443</td>
<td>593</td>
<td>NDNPLW</td>
<td>DDPFKDFEEEE</td>
</tr>
<tr>
<td>Pmp</td>
<td>XP_001331554</td>
<td>742</td>
<td>DENPLW</td>
<td>DDPFKDFEQEV</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001325298</td>
<td>624</td>
<td>NDNPFL</td>
<td>DDPFKDFEQEE</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001306144</td>
<td>569</td>
<td>IDNPFL</td>
<td>DPPRTEFEK</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001582856</td>
<td>712</td>
<td>NDNPFL</td>
<td>DDPFKDFEED</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001309734</td>
<td>812</td>
<td>NDNPFL</td>
<td>DDPFKDFEED</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001325816</td>
<td>605</td>
<td>YTNPLW</td>
<td>DDPVFDFEQE</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001581046</td>
<td>680</td>
<td>NDNPFL</td>
<td>DDPFKDFEED</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001325582</td>
<td>803</td>
<td>NDNPFL</td>
<td>DDPFKDNEI</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001325146</td>
<td>593</td>
<td>QENPLW</td>
<td>DDPFKDFOEQE</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001320330</td>
<td>526</td>
<td>NVNPLF</td>
<td>DDPFQDFEEQ</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001320335</td>
<td>1819</td>
<td>NDNPFL</td>
<td>DPPFPDNEED</td>
</tr>
<tr>
<td>Pmpf</td>
<td>XP_001299221</td>
<td>652</td>
<td>NENPLW</td>
<td>ENPNFLFDDEED</td>
</tr>
</tbody>
</table>

*Entries with at least one inferred transmembrane domain from two distinct TV protein families (BspA: Bacteroides surface protein A-like extracellular domain and Pmpf: Chlamydia polymorphic membrane protein-like extracellular domain) but with a shared cytoplasmic tails possessing two motifs ([FY]**NPX[**FY**] and acidic clusters) representing potentially signals for endocytosis—see ref. 11.*

Table 2: List of enzymes involved in glycan metabolism derived from bacterial lateral gene transfers (LGTs)*

<table>
<thead>
<tr>
<th>EC number</th>
<th>Enzyme name</th>
<th>RefSeq accession</th>
<th>Nearest neighbour in phylogeny</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.2.1.18</td>
<td>En-α-sialidase</td>
<td>XP_001319692</td>
<td>Bacteroidetes</td>
</tr>
<tr>
<td>3.2.1.23</td>
<td>β-galactosidase</td>
<td>XP_001581038</td>
<td>Bacteroidetes</td>
</tr>
<tr>
<td>3.2.1.52</td>
<td>β-N-acetylfucosaminidase</td>
<td>XP_001329989</td>
<td>Bacteroidetes</td>
</tr>
<tr>
<td>3.2.1.24</td>
<td>α-mannosidase</td>
<td>XP_001579222</td>
<td>Prokaryotes</td>
</tr>
<tr>
<td>3.2.1.25</td>
<td>β-mannosidase</td>
<td>XP_001322689</td>
<td>Proteobacteria</td>
</tr>
<tr>
<td>3.2.1.51</td>
<td>α-fucosidase</td>
<td>XP_001316088</td>
<td>Bacteria</td>
</tr>
<tr>
<td>4.1.3.3</td>
<td>N-acetylgalactosaminidase</td>
<td>XP_001323296</td>
<td>Pasteurellaceae (γ-proteobacteria)</td>
</tr>
<tr>
<td>5.1.3.8</td>
<td>Aroylglucosaminidase 2-epimerase</td>
<td>XP_001308218</td>
<td>Bacteroidetes</td>
</tr>
<tr>
<td>3.2.1.45</td>
<td>Glucosylceramidase</td>
<td>XP_001279673</td>
<td>Bacteroidetes</td>
</tr>
</tbody>
</table>

*For details about inference of LGT methodology and gene and enzyme characteristics see ref. 26.

Note that some of the listed enzymes could target O-glycans, N-glycans and/or gangliosides.*

*EC, Enzyme commission number.*

(imbbalanced microbial community) in the human microbiota, for example, as observed during bacterial vaginosis (BV). Consistent with this possibility, a correlation between the presence of TV and a vaginal microbiota characterised with low abundance of protective lactobacilli and higher proportions of *Mycoplasma, Prevotella* and other bacteria typically observed in BV, has been observed. Future work will be required to establish temporality and causality between BV and TV and their contribution (possibly synergistic) to increase host susceptibility to HIV. In addition, antibiotics can significantly disturb the host-microbiota-viruses balances, and are also increasingly recognised to contribute to various disease conditions. Hence, the potential negative impact of the treatment of TV with metronidazole/tinidazole should also be considered in this global framework of human–microbes interactions.

TV has been shown to form a symbiosis with *Mycoplasma hominis* in in vitro cultures, and *Mycoplasma* cells carried by TV could infect human cells, suggesting that TV could be a Trojan horse for the bacterium. Intriguingly, coinfections with TV-Mycoplasma were shown to consume larger amounts of free arginine than TV alone, which could contribute to reducing nitric oxide production by macrophages through depletion of free arginine in the vagina, thus potentially interfering with an important host defence mechanism. Another likely consequence of TV actively feeding on host microbiota, is the acquisition of a number of bacterial genes through lateral gene transfer (LGT). Of particular interest is an almost complete set of enzymes of bacterial origin capable of degrading glycans, and mostly acquired from members of the bacteroidetes, one of the most abundant bacterial lineage of mucosal microbiota and known to encode a rich set of glycan-targeting enzymes (Table 2). These TV enzymes of bacterial origins are thought to play important roles in degrading mucins and other human glycans, including those from the glycoalyx of epithelial cells, which represent the initial barrier the parasite needs to deal with to be able to colonise the host mucosal surfaces. Glycan degradation is also likely to provide an important source of energy for the parasite.

Tissue damage and inflammatory responses due to TV infections are thought to facilitate HIV entry and, in TV-HIV dually infected patients, to stimulate HIV production. Additionally, TV can also internalise human viruses through endocytosis. Viruses carried by TV could contribute to their transmission to a new host, as viruses internalised by TV were shown to be infectious for human cells for 2–6 days. The viruses are either released upon parasite death or secreted from TV-loaded viruses through the recycling route of the endocytic pathway. However, there is no evidence that any human virus can replicate in TV. Hence, as suggested for *Mycoplasma*, TV could be a Trojan horse for HIV and possibly other viruses.

These different considerations clearly illustrate the importance of studying a pathogen in the broader context of human–microbes interactions (figure 2).
parasite (TV1 and TV2) with distinct properties. JVI isolates are more sensitive to metronidazole, and are also significantly more likely to be infected with TVV than TV2, in a ratio ~3:1. In another study investigating the potential role of TVV in modulating the human innate immune system, it was established that TVV are sensed by the human toll-like receptor 3 (TLR3), and that this induces an innate response, including proinflammatory cascades, previously implicated in preterm birth and HIV-1 susceptibility. These new aspects of TV pathology should be taken into account in future research on TV pathology, and influence, for example, how clinical samples are processed for research and, possibly, diagnostics. For instance, to allow quantification of TVV, processing of both total DNA (to investigate TV; bacteria, etc...) and RNA will be required.

CONCLUSION

The aim of this short overview on TV virulence factors was to illustrate specific aspects of the intricate and complex interplays taking place between the parasite and its host. The complex interplay between the human urogenital tracts, TV and other human pathogens, human microbiota and TVV emphasise the importance of integrating the broadest possible spectrum of human–microbes interactions when studying human–TV interactions. The challenge will be to be able to take into consideration these dynamic host–microbes interactions to guide future research projects on TV molecular cell biology, immunology and epidemiology to eventually produce more refined and effective treatment protocols for STIs. The paper by Fiori et al., further illustrates the synergistic interactions between TV and Mycoplasma.

TVV dependent inflammation
Boosting of HIV transmission
TV induced dysbiosis
Mycoplasma transmission

TV dependent inflammation & tissue damage
Synergism with Mycoplasma

Figure 2 Interactions between Trichomonas vaginalis (TV), viruses, microbiota and the human host and their impact on human health. The diagram illustrates a TV-infected pregnant woman and her partner in the global context of human–microbe interactions. TV infections can affect the health status of both the adults and the embryo (eg, preterm birth and HIV transmission in utero). Rather than single linear human–pathogen interactions, it is the complex network of interactions between viruses-bacteria-eukaryotes and the human host that should be considered when investigating disease conditions.

Key messages

- The pathobiology of Trichomonas vaginalis (TV) is multifaceted involving direct and indirect interactions with host tissue, bacteria, human viruses and Trichomonas viruses.
- The genome sequence data of TV is being exploited to facilitate the study of the molecular basis of the parasite pathobiology.
- The immune response to TV is still poorly understood, and it will be essential to study its variations to rationalise the outcome of host–parasite interactions.

Acknowledgements

I thank Didier Ndeh for critical reading of an early draft of the manuscript. My apologies to the authors whose work could not be cited due to length restrictions.

Funding

Past support from the Wellcome Trust for my research on Trichomonas vaginalis is greatly acknowledged.

Competing interests

None.

Provenance and peer review

Commissioned; externally peer reviewed.

Open Access

This is an Open Access article distributed in accordance with the terms of the Creative Commons Attribution (CC BY 3.0) license, which permits others to distribute, remix, adapt and build upon this work, for commercial use, provided the original work is properly cited. See: http://creativecommons.org/licenses/by/3.0/

REFERENCES


Trichomonas vaginalis virulence factors: an integrative overview

Robert P Hirt

Sex Transm Infect 2013 89: 439-443 originally published online May 21, 2013
doi: 10.1136/sextrans-2013-051105

Updated information and services can be found at:
http://sti.bmj.com/content/89/6/439

These include:

References
This article cites 26 articles, 3 of which you can access for free at:
http://sti.bmj.com/content/89/6/439#BIBL

Open Access
This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 3.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/3.0/

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections

Open access (245)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/