

Science. Author manuscript; available in PMC 2007 November 17.

Published in final edited form as:

Science. 2007 January 12; 315(5809): 207-212.

Draft Genome Sequence of the Sexually Transmitted Pathogen Trichomonas vaginalis

Jane M. Carlton^{1,*,†}, Robert P. Hirt², Joana C. Silva¹, Arthur L. Delcher³, Michael Schatz³, Qi Zhao¹, Jennifer R. Wortman¹, Shelby L. Bidwell¹, U. Cecilia M. Alsmark², Sébastien Besteiro⁴, Thomas Sicheritz-Ponten⁵, Christophe J. Noel², Joel B. Dacks⁶, Peter G. Foster⁷, Cedric Simillion⁸, Yves Van de Peer⁸, Diego Miranda-Saavedra⁹, Geoffrey J. Barton⁹, Gareth D. Westrop⁴, Sylke Müller⁴, Daniele Dessi¹⁰, Pier Luigi Fiori¹⁰, Qinghu Ren¹, Ian Paulsen¹, Hanbang Zhang¹, Felix D. Bastida-Corcuera¹¹, Augusto Simoes-Barbosa¹¹, Mark T. Brown¹¹, Richard D. Hayes¹¹, Mandira Mukherjee¹¹, Cheryl Y. Okumura¹¹, Rachel Schneider¹¹, Alias J. Smith¹¹, Stepanka Vanacova¹¹, Maria Villalvazo¹¹, Brian J. Haas¹, Mihaela Pertea³, Tamara V. Feldblyum¹, Terry R. Utterback¹², Chung-Li Shu¹³, Kazutoyo Osoegawa¹³, Pieter J. de Jong¹³, Ivan Hrdy¹⁴, Lenka Horvathova¹⁴, Zuzana Zubacova¹⁴, Pavel Dolezal¹⁴, Shehre-Banoo Malik¹⁵, John M. Logsdon Jr.¹⁵, Katrin Henze¹⁶, Arti Gupta¹⁷, Ching C. Wang¹⁷, Rebecca L. Dunne¹⁸, Jacqueline A. Upcroft¹⁹, Peter Upcroft¹⁹, Owen White¹, Steven L. Salzberg³, Petrus Tang²⁰, Cheng-Hsun Chiu²¹, Ying-Shiung Lee²², T. Martin Embley², Graham H. Coombs²³, Jeremy C. Mottram⁴, Jan Tachezy¹⁴, Claire M. Fraser-Liggett¹, and Patricia J. Johnson¹¹

- 1 Institute for Genomic Research, 9712 Medical Research Drive, Rockville, MD 20850, USA
- 2 Division of Biology, Devonshire Building, Newcastle University, Newcastle upon Tyne NEI 7RU, UK
- 3 Center for Bioinformatics and Computational Biology, University of Maryland, College Park, MD 20742, USA
- 4 Wellcome Centre for Molecular Parasitology and Division of Infection and Immunity, Glasgow Biomedical Research Centre, University of Glasgow, Glasgow G12 8TA, UK
- 5 Center for Biological Sequence Analysis, BioCentrum-DTU, Technical University of Denmark, DK-2800 Lyngby, Denmark
- 6 Department of Biological Sciences, University of Calgary, 2500 University Drive NW, Calgary, Alberta T2N 1N4, Canada
- 7 Department of Zoology, Natural History Museum, Cromwell Road, London SW7 5BD, UK
- 8 Department of Plant Systems Biology, Flanders Interuniversity Institute for Biotechnology (VIB), Ghent University, Technologiepark 927, B-9052 Ghent, Belgium
- 9 School of Life Sciences Research, University of Dundee, Dow Street, Dundee DD1 5EH, UK

Supporting Online Material

www.sciencemag.org/cgi/content/full/315/5809/207/DC1

Materials and Methods

SOM Text Figs. S1 to S14 Tables S1 to S26 References

Data

^{*}To whom correspondence should be addressed. E-mail: jane.carlton@med.nyu.edu.

[†]Present address: Department of Medical Parasitology, New York University School of Medicine, New York, NY 10011, USA.

10 Department of Biomedical Sciences, Division of Experimental and Clinical Microbiology, University of Sassari, Viale San Pietro 43/b, 07100 Sassari, Italy

- 11 Department of Microbiology, Immunology, and Molecular Genetics, University of California, Los Angeles, CA 90095–1489, USA
- 12 J. Craig Venter Joint Technology Center, 5 Research Place, Rockville, MD 20850, USA
- 13 Children's Hospital Oakland Research Institute, 747 52nd Street, Oakland, CA 94609, USA
- 14 Department of Parasitology, Faculty of Science, Charles University, 128 44 Prague, Czech Republic
- 15 Department of Biological Sciences, Roy J. Carver Center for Comparative Genomics, University of Iowa, Iowa City, IA 52242–1324, USA
- 16 Institute of Botany, Heinrich Heine University, 40225 Düsseldorf, Germany
- 17 Department of Pharmaceutical Chemistry, University of California, San Francisco, CA 94143–2280, USA
- 18 Department of Microbiology and Parasitology, University of Queensland, Brisbane, Queensland 4072, Australia
- 19 Queensland Institute of Medical Research, Brisbane, Queensland 4029, Australia
- 20 Bioinformatics Center/Molecular Medicine Research Center, Chang Gung University, Taoyuan 333, Taiwan
- 21 Molecular Infectious Diseases Research Center, Chang Gung Memorial Hospital, Taoyuan 333, Taiwan
- 22 Medical Genomic Center, Chang Gung Memorial Hospital, Taoyuan 333, Taiwan
- 23 Strathclyde Institute of Pharmacy and Biomedical Sciences, John Arbuthnott Building, University of Strathclyde, Glasgow G4 ONR, UK

Abstract

We describe the genome sequence of the protist *Trichomonas vaginalis*, a sexually transmitted human pathogen. Repeats and transposable elements comprise about two-thirds of the ~160-megabase genome, reflecting a recent massive expansion of genetic material. This expansion, in conjunction with the shaping of metabolic pathways that likely transpired through lateral gene transfer from bacteria, and amplification of specific gene families implicated in pathogenesis and phagocytosis of host proteins may exemplify adaptations of the parasite during its transition to a urogenital environment. The genome sequence predicts previously unknown functions for the hydrogenosome, which support a common evolutionary origin of this unusual organelle with mitochondria.

Trichomonas vaginalis is a flagellated protist that causes trichomoniasis, a common but overlooked sexually transmitted human infection, with ~170 million cases occurring annually worldwide (1). The extracellular parasite resides in the urogenital tract of both sexes and can cause vaginitis in women and urethritis and prostatitis in men. Acute infections are associated with pelvic inflammatory disease, increased risk of human immunodeficiency virus 1 (HIV-1) infection, and adverse pregnancy outcomes. T. vaginalis is a member of the parabasilid lineage of microaerophilic eukaryotes that lack mitochondria and peroxisomes but contain unusual organelles called hydrogenosomes. Although previously considered to be one of the earliest branching eukaryotic lineages, recent analyses leave the evolutionary relationship of parabasalids to other major eukaryotic groups unresolved (2,3). In this article, we report the draft sequence of T. vaginalis, the first parabasalid genome to be described.

Genome structure, RNA processing, and lateral gene transfer

The *T. vaginalis* genome sequence, generated using whole-genome shotgun methodology, contains 1.4 million shotgun reads assembled into 17,290 scaffolds at ~7.2× coverage (4). At least 65% of the *T. vaginalis* genome is repetitive (table S1). Despite several procedures developed to improve the assembly (4), the superabundance of repeats resulted in a highly fragmented sequence, preventing investigation of *T. vaginalis* genome architecture. The repeat sequences also hampered measurement of genome size, but we estimate it to be ~160 Mb (4). A core set of ~60,000 protein-coding genes was identified (Table 1), endowing *T. vaginalis* with one of the highest coding capacities among eukaryotes (table S2). Introns were identified in 65 genes, including the ~20 previously documented (5). Transfer RNAs (tRNAs) for all 20 amino acids were found, and ~250 ribosomal DNA (rDNA) units were identified on small contigs and localized to one of the six *T. vaginalis* chromosomes (Fig. 1).

The Inr promoter element was found in ~75% of 5′ untranslated region (UTR) sequences (4), supporting its central role in gene expression (6). Intriguingly, the eukaryotic transcription machinery of *T. vaginalis* appears more metazoan than protistan (table S3 to S5). The presence of a *T. vaginalis* Dicer-like gene, two Argonaute genes, and 41 transcriptionally active DEAD-DEAH-box helicase genes suggests the existence of an RNA interference (RNAi) pathway (fig. S1). Identification of these components raises the possibility of using RNAi technology to manipulate *T. vaginalis* gene expression.

During genome annotation, we identified 152 cases of possible prokaryote-to-eukaryote lateral gene transfer (LGT) [tables S6 and S7 and Supporting Online Material (SOM) text], augmenting previous reports of conflicting phylogenetic relationships among several enzymes (7). The putative functions of these genes are diverse, affecting various metabolic pathways (fig. S2) and strongly influencing the evolution of the *T. vaginalis* metabolome. A majority (65%) of the 152 LGT genes encode metabolic enzymes, more than a third of which are involved in carbohydrate or amino acid metabolism (Fig. 2). Several LGT genes may have been acquired from Bacteroidetes-related bacteria, which are abundant among vertebrate intestinal flora (fig. S3).

Repeats, transposable elements, and genome expansion

The most common 59 repeat families identified in the assembly (4) constitute ~39 Mb of the genome and can be classified as (i) virus-like; (ii) transposon-like, including ~1000 copies of the first *mariner* element identified outside animals (8); (iii) retrotransposon-like; and (iv) unclassified (Table 2). Most of the 59 repeats are present in hundreds of copies (average copy number ~660) located on small (1- to 5-kb) contigs, and each repeat family is extraordinarily homogenous, with an average polymorphism of ~2.5%.

The lack of a strong correlation between copy number and average pairwise difference between copies (fig. S4) suggests that a sudden expansion of the repeat families had occurred. To estimate the time of expansion, we compared the degree of polymorphism among *T. vaginalis* repeats to the divergence between *T. vaginalis* and its sister taxon *T. tenax*, a trichomonad of the oral cavity (9), for several protein-coding loci (4). Our results indicate that repeat family amplification occurred after the two species split (table S8). Several families have also undergone multiple expansions, as implied by bi- or trimodal distribution of pairwise distances between copies (fig. S5). *T. vaginalis* repeat families appear to be absent in *T. tenax* but are present in geographically diverse *T. vaginalis* (4), consistent with the expansion having occurred after speciation but before diversification of *T. vaginalis*.

The large genome size, high repeat copy number, low repeat polymorphism, and evidence of repeat expansion after *T. vaginalis* and *T. tenax* diverged suggest that *T. vaginalis* has

undergone a very recent and substantial increase in genome size. To determine whether the genome underwent any large-scale duplication event(s), we analyzed age distributions of gene families with five or fewer members (4). A peak in the age distribution histogram of pairs of gene families was observed (fig. S6), indicating that the genome underwent a period of increased duplication, and possibly one or more large-scale genome duplication events.

Metabolism, oxidative stress, and transport

T. vaginalis uses carbohydrate as a main energy source via fermentative metabolism under aerobic and anaerobic conditions. We found the parasite to use a variety of amino acids as energy substrates (Fig. 2) (10), with arginine dihydrolase metabolism a major pathway for energy production (fig. S7) (11). We confirmed a central role for aminotransferases (Fig. 2 and table S9) and glutamate dehydrogenase as indicated previously (12,13); these pathways are likely catabolic but may be reversible to allow the parasite to synthesize glutamate, aspartate, alanine, glutamine, and glycine. Genes required for synthesis of proline from arginine (fig. S7) and for threonine metabolism (fig. S8) were identified. We also identified a de novo biosynthesis pathway for cysteine via cysteine synthase, an LGT candidate (fig. S8) (14), and genes encoding enzymes involved in methionine metabolism, including its possible regeneration (fig. S9).

Earlier studies indicated that de novo lipid biosynthesis in *T. vaginalis* is confined to the major phospholipid phosphatidylethanolamine (PE) (15), whereas other lipids, including cholesterol, are likely acquired from exogenous sources. We found an absence of several essential enzymeencoding genes in the synthesis and degradation pathways of nearly all lipids (4), in contrast to the PE synthetic pathway, which appears complete; however, experimental verification of these results is required.

T. vaginalis is microaerophilic with a primarily anaerobic life style and thus requires redox and antioxidant systems to counter the detrimental effects of oxygen. Genes encoding a range of defense molecules, such as superoxide dismutases, thioredoxin reductases, peroxiredoxins, and rubrerythrins, were identified (table S10).

T. vaginalis demonstrates a broad range of transport capabilities, facilitated by expansion of particular transporter families, such as those for sugar and amino acids (table S11). The parasite also possesses more members of the cation-chloride cotransporter (CCC) family than any other sequenced eukaryote, likely reflecting osmotic changes faced by the parasite in a mucosal environment.

None of the proteins required for glycosyl-phosphatidylinositol (GPI)-anchor synthesis were identified in the genome sequence, making *T. vaginalis* the first eukaryote known to lack an apparent GPI-anchor biosynthetic pathway. Whether *T. vaginalis* has evolved an unusual biosynthetic pathway for synthesis of its nonprotein lipid anchors, such as the inositol-phosphoceramide of surface lipophosphoglycans (16), remains to be determined.

Massively expanded gene families

Many gene families in the *T. vaginalis* genome have undergone expansion on a scale unprecedented in unicellular eukaryotes (Table 3). Such "conservative" gene family expansions are likely to improve an organism's adaptation to its environment (17). Notably, the selective expansion of subsets of the membrane trafficking machinery, critical for secretion of pathogenic proteins, endocytosis of host proteins, and phagocytosis of bacteria and host cells (table S12), correlates well with the parasite's active endocytic and phagocytic life-style.

Massively amplified gene families also occur in the parasite's kinome, which comprises ~880 genes (SOM text) encoding distinct eukaryotic protein kinases (ePKs) and ~40 atypical protein kinases, making it one of the largest eukaryotic kinomes known. The parasite has heterotrimeric guanine nucleotide—binding proteins and components of the mitogen-activated protein kinase (MAPK) pathway, suggesting yeast-like signal transduction mechanisms. Unusually, the *T. vaginalis* kinome contains 124 cytosolic tyrosine kinase—like (TKL) genes, yet completely lacks receptor serine/threonine ePKs of the TKL family. Inactive kinases were found to make up 17% of the *T. vaginalis* kinome (table S13); these may act as substrates and scaffolds for assembly of signaling complexes (18). ePK accessory domains are important for regulating signaling pathways, but just nine accessory domain types were identified in 8% (72/883) of the *T. vaginalis* ePKs (table S14), whereas ~50% of human ePKs contain at least 1 of 83 accessory domain types. This suggests that regulation of protein kinase function and cell signaling in *T. vaginalis* is less complex than that in higher eukaryotes, a possible explanation for the abundance of *T. vaginalis* ePKs.

T. vaginalis possesses several unusual cyto-skeletal structures: the axostyle, the pelta, and the costa (19). Most actin- and tubulin-related components of the cytoskeleton are present (table S15), with the exception of homologs of the actin motor myosin. In contrast, homologs of the microtubular motors kinesin and dynein are unusually abundant (Table 3). Thus, T. vaginalis intracellular transport mechanisms are mediated primarily by kinesin and cytoplasmic dynein, as described for Dictyo-stelium and filamentous fungi, raising the possibility that the loss of myosin-driven cytoplasmic transport is not uncommon in unicellular eukaryotes (20). Whether the structural remodeling of amoeboid T. vaginalis during host colonization (see below) is actin-based, as described for other eukaryotes, or driven by novel cytoskeletal rearrangements remains an open question.

We identified homologs of proteins involved in DNA damage response and repair, chromatin restructuring, and meiosis, the latter a process not thought to occur in the parasite (table S16). Of the 29 core meiotic genes found, several are general repair proteins required for meiotic progression in other organisms (21), and eight are meiosis-specific proteins. Thus, *T. vaginalis* contains either recent evolutionary relics of meiotic machinery or genes functional in meiotic recombination in an as-yet undescribed sexual cycle.

Molecular mechanisms of pathogenesis

T. vaginalis must adhere to host cells to establish and maintain an infection. A dense glycocalyx composed of lipophosphoglycan (LPG) (Fig. 3) and surface proteins has been implicated in adherence (22), but little is known about this critical pathogenic process. We identified genes encoding enzymes predicted to be required for LPG synthesis (table S17). Of particular interest are the genes required for synthesis of an unusual nucleotide sugar found in *T. vaginalis* LPG, the monosaccharide rhamnose, which is absent in the human host, making it a potential drug target. Genes (some of which are LGT candidates) were identified that are involved in sialic acid biosynthesis, consistent with the reported presence of this sugar on the parasite surface (23).

We identified eight families containing ~800 proteins (4) that represent candidate surface molecules (Fig. 3 and table S18), including ~650 highly diverse BspA-like proteins characterized by the *Treponema pallidum* leucine-rich repeat, TpLLR. BspA-like proteins are expressed on the surface of certain pathogenic bacteria and mediate cell adherence and aggregation (24). The only other eukaryote known to encode BspA-like proteins, the mucosal pathogen *Entamoeba histolytica* (25), contains 91 such proteins, one of which was recently localized to the parasite surface (26).

There are >75 *T. vaginalis* GP63-like proteins, homologs of the most abundant surface proteins of *Leishmania major*, the leishmanolysins, which contribute to virulence and pathogenicity through diverse functions in both the insect vector and the mammalian host (27). Most *T. vaginalis* GP63-like genes possess the domains predicted to be required for a catalytically active metallopeptidase, including a short HEXXH motif (28) (Fig. 3). Unlike trypanosomatid GP63 proteins, which are predicted to be GPI-anchored, most *T. vaginalis* GP63-like proteins have a predicted C-terminal transmembrane domain as a putative cell surface anchor, consistent with the apparent absence of GPI-anchor biosynthetic enzymes. Other *T. vaginalis* protein families share domains with *Chlamydia* polymorphic membrane proteins, *Giardia lamblia* variant surface proteins, and *E. histolytica* immunodominant variable surface antigens (Fig. 3 and table S18).

After cytoadherence, the parasite becomes amoeboid, increasing cell-to-cell surface contacts and forming cytoplasmic projections that interdigitate with target cells (19). We have identified genes encoding cytolytic effectors, which may be released upon host-parasite contact. *T. vaginalis* lyses host red blood cells, presumably as a means of acquiring lipids and iron and possibly explaining the exacerbation of symptoms observed during menstruation (29). This hemolysis is dependent on contact, temperature, pH, and Ca²⁺, suggesting the involvement of pore-forming proteins (30) that insert into the lipid bilayer of target cells, mediating osmotic lysis. Consistent with this, we have identified 12 genes (*TvSaplip1* to *TvSaplip12*) containing saposin-like (SAPLIP) pore-forming domains (fig. S10). These domains show a predicted six-cysteine pattern and abundant hydrophobic residues in conserved positions while displaying high sequence variability (fig. S11). The TvSaplips are similar to amoebapore proteins secreted by *E. histolytica* and are candidate trichopores that mediate a cytolytic effect.

The degradome

Peptidases perform many critical biological processes and are potential virulence factors, vaccine candidates, and drug targets (31). *T. vaginalis* contains an expanded degradome of more than 400 peptidases (SOM text), making it one of the most complex degradomes described (table S19). Of the three families of aspartic peptidases (table S20), *T. vaginalis* contains a single member of the HIV-1 retropepsin family that might serve as a putative candidate for anti-HIV peptidase inhibitors. Many studies have implicated papain family cysteine peptidases as virulence factors in trichomonads; we identified >40 of them, highlighting the diversity of this family. Cysteine peptidases that contribute to the 20S proteosome (ubiquitin C-terminal hydrolases) are abundant (117 members, ~25% of the degradome), emphasizing the importance of cytosolic protein degradation in the parasite. *T. vaginalis* has nine NlpC/P60-like members (table S20; several of which are LGT candidates), which play a role in bacterial cell wall degradation and the destruction of healthy vaginal microflora, making the vaginal mucosa more sensitive to other infections.

We also identified many subtilisin-like and several rhomboid-like serine peptidases, candidates for processing *T. vaginalis* surface proteins. In addition to the first asparaginase-type of threonine peptidase found in a protist, 13 families of metallopeptidases were also identified, as well as three cystatin-like proteins, natural peptidase inhibitors, which may regulate the activity of the abundant papain-like cysteine peptidases (table S20).

The hydrogenosome

Several microaerophilic protists and fungi, including trichomonads and ciliates, lack typical mitochondria and possess double-membrane hydrogenosomes, which produce adenosine triphophate (ATP) and molecular hydrogen through fermentation of metabolic intermediates produced in the cytosol. Although the origin of these organelles has been controversial, most evidence now supports a common origin with mitochondria (3). Few genes encoding homologs

of mitochondrial transporters, translocons, and soluble proteins were identified in the T. vaginalis genome (fig. S12), suggesting that its hydrogenosome has undergone reductive evolution comparable to other protists whose mitochondrial proteomes are reduced (e.g., Plasmodium). Because nuclear-encoded hydrogenosomal matrix proteins are targeted to the organelle by N-terminal presequences that are proteolytically cleaved upon import (32) similar to mitochondrial precursor proteins, we screened the genome for consensus 5-to 20-residue presequences containing $ML(S/T/A) X_{(1...15)}R(N/F/E/XF)$, $MSLX_{(1...15)}R(N/F/XF)$, or MLR(S/N)F(28) motifs. A total of 138 genes containing putative presequences were identified, 67% of which are similar to known proteins, primarily ones involved in energy metabolism and electron-transport pathways (fig. S13 and table S21).

The production of molecular hydrogen, the hallmark of the hydrogenosome, is catalyzed by an unusually diverse group of iron-only [Fe]-hydrogenases that possess, in addition to a conserved H cluster, four different sets of functional domains (fig. S14), indicating that hydrogen production may be more complex than originally proposed. The pathway that generates electrons for hydrogen (fig. S12) is composed of many proteins encoded by multiple genes (tables S22 to S24). Our analyses extend the evidence that *T. vaginalis* hydrogenosomes contain the complete machinery required for mitochondria-like intraorganellar FeS cluster formation (33) and also reveal the presence of two putative cytosolic auxiliary proteins, indicating that hydrogenosomes may be involved in biogenesis of cytosolic FeS proteins. Some components of FeS cluster assembly machinery have also been found in mitosomes (mitochondrial remnants) of *G. lamblia*, supporting a common evolutionary origin of mitochondria, hydrogenosomes, and mitosomes (3).

A new predicted function of hydrogenosomes revealed by the genome sequence is amino acid metabolism. We identified two components of the glycine-cleavage complex (GCV), L protein and H protein. Another component of this pathway is serine hydroxymethyl transferase (SHMT), which in eukaryotes exists as both cytosolic and mitochondrial isoforms. A single gene coding for SHMT of the mitochondrial type with a putative N-terminal hydrogenosomal presequence was identified. Because both GCV and SHMT require folate (fig. S12), which *T. vaginalis* apparently lacks, the functionality of these proteins remains unclear.

The 5-nitroimidazole drugs metronidazole (Mz) and tinidazole are the only approved drugs for treatment of trichomoniasis. These prodrugs are converted within the hydrogenosome to toxic nitroradicals via reduction by ferredoxin (Fdx) (fig. S12). Clinical resistance to Mz (Mz^R) is estimated at 2.5 to 5% of reported cases and rising (34) and is associated with decreases in or loss of Fdx (35). We identified seven Fdx genes with hydrogenosomal targeting signals (tables S21 and S25), the redundancy of which provides explanations for the low frequency of Mz^R and for why knockout of a single Fdx gene does not lead to Mz^R (35). Our analyses also provide clues to the potential mechanisms that clinically resistant parasites may use, such as the presence of nitroreductase (*NimA*-like), reduced nicotinamide adenine dinucleotide phosphate (NADPH)–nitroreductase, and NADH-flavin oxidoreductase genes (tables S23 and S26), which have been implicated in Mz^R in bacteria (36,37).

Summary and concluding remarks

Our investigation of the *T. vaginalis* genome sequence provides a new perspective for studying the biology of an organism that continues to be ignored as a public health issue despite the high number of trichomoniasis cases worldwide. The discovery of previously unknown metabolic pathways, the elucidation of pathogenic mechanisms, and the identification of candidate surface proteins likely involved in facilitating invasion of human mucosal surfaces provide potential leads for the development of new therapies and novel methods for diagnosis.

The analysis presented here of one of the most repetitive genomes known has undoubtedly been hampered by the sheer number of highly similar repeats and transposable elements. Why did this genome expand so dramatically in size? We hypothesize that the most recent common ancestor of T. vaginalis underwent a population bottleneck during its transition from an enteric environment (the habitat of most trichomonads) to the urogenital tract. During this time, the decreased effectiveness of selection resulted in repeat accumulation and differential gene family expansion. Genome size and cell volume are positively correlated (38,39); hence, the increased genome size of T. vaginalis achieved through rapid fixation of repeat copies could have ultimately resulted in a larger cell size. T. vaginalis cell volume is greater than that of T. tenax and related intestinal species Pentratrichomonas hominis (40) and T. gallinae (41), and it generally conforms to the relationship of genome size to cell volume reported for protists (41). T. vaginalis is also a highly predatory parasite that phagocytoses bacteria, vaginal epithelial cells, and host erythrocytes (42) and is itself ingested by macrophages. Given these interactions, it is tempting to speculate that an increase in cell size could have been selected for in order to augment the parasite's phagocytosis of bacteria, to reduce its own phagocytosis by host cells, and to increase the surface area for colonization of vaginal mucosa.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

References and Notes

- Global Prevalence and Incidence of Selected Curable Sexually Transmitted Infections. World Health Organization; Geneva: 2001. www.who.int/docstore/hiv/GRSTI/006.htm
- 2. Adl SM, et al. J Eukaryot Microbiol 2005;52:399. [PubMed: 16248873]
- 3. Embley TM, Martin W. Nature 2006;440:623. [PubMed: 16572163]
- 4. Materials and methods are available as supporting material on. Science. Online
- Vanacova S, Yan W, Carlton JM, Johnson PJ. Proc Natl Acad Sci USA 2005;102:4430. [PubMed: 15764705]
- 6. Schumacher MA, Lau AO, Johnson PJ. Cell 2003;115:413. [PubMed: 14622596]
- 7. Muller, M. Evolutionary Relationship Among Protozoa. Coombs, GH.; Vickerman, V.; Sleigh, MA.; Warren, A., editors. Chapman & Hall; London: 1998. p. 109-132.
- 8. Silva JC, Bastida F, Bidwell SL, Johnson PJ, Carlton JM. Mol Biol Evol 2005;22:126. [PubMed: 15371525]
- 9. Kutisova K, et al. Parasitology 2005;131:309. [PubMed: 16178352]
- 10. Zuo X, Lockwood BC, Coombs GH. Microbiology 1995;141:2637. [PubMed: 7582024]
- 11. Kleydman Y, Yarlett N, Gorrell TE. Microbiology 2004;150:1139. [PubMed: 15133073]
- 12. Lowe PN, Rowe AF. Mol Biochem Parasitol 1986;21:65. [PubMed: 3095639]
- 13. Turner AC, Lushbaugh WB. Exp Parasitol 1988;67:47. [PubMed: 2901980]
- 14. Westrop GD, Goodall G, Mottram JC, Coombs GH. J Biol Chem 2006;281:25062. [PubMed: 16735516]
- 15. Beach DH, Holz GG Jr, Singh BN, Lindmark DG. Mol Biochem Parasitol 1991;44:97. [PubMed: 2011157]
- Singh BN, Beach DH, Lindmark DG, Costello CE. Arch Biochem Biophys 1994;309:273. [PubMed: 8135538]
- 17. Vogel C, Chothia C. PLoS Comput Biol 2006;2:e48. [PubMed: 16733546]
- 18. Parsons M, Worthey EA, Ward PN, Mottram JC. BMC Genomics 2005;6:127. [PubMed: 16164760]
- 19. Benchimol M. Microsc Microanal 2004;10:528. [PubMed: 15525428]
- 20. Richards TA, Cavalier-Smith T. Nature 2005;436:1113. [PubMed: 16121172]
- 21. Ramesh MA, Malik SB, Logsdon JM Jr. Curr Biol 2005;15:185. [PubMed: 15668177]

22. Bastida-Corcuera FD, Okumura CY, Colocoussi A, Johnson PJ. Eukaryot Cell 2005;4:1951. [PubMed: 16278462]

- 23. Dias Filho BP, Andrade AF, de Souza W, Esteves MJ, Angluster J. Microbios 1992;71:55. [PubMed: 1406344]
- 24. Ikegami A, Honma K, Sharma A, Kuramitsu HK. Infect Immun 2004;72:4619. [PubMed: 15271922]
- 25. Loftus B, et al. Nature 2005;433:865. [PubMed: 15729342]
- 26. Davis PH, et al. Mol Biochem Parasitol 2006;145:111. [PubMed: 16199101]
- 27. Yao C, Donelson JE, Wilson ME. Mol Biochem Parasitol 2003;132:1. [PubMed: 14563532]
- 28. Single-letter abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; X, any amino acid; and Y, Tyr.
- 29. Lehker MW, Chang TH, Dailey DC, Alderete JF. J Exp Med 1990;171:2165. [PubMed: 2351937]
- 30. Fiori PL, Rappelli P, Rocchigiani AM, Cappuccinelli P. FEMS Microbiol Lett 1993;109:13. [PubMed: 8319880]
- 31. Klemba M, Goldberg DE. Annu Rev Biochem 2002;71:275. [PubMed: 12045098]
- 32. Bradley PJ, Lahti CJ, Plumper E, Johnson PJ. EMBO J 1997;16:3484. [PubMed: 9218791]
- 33. Sutak R, et al. Proc Natl Acad Sci USA 2004;101:10368. [PubMed: 15226492]
- 34. Schmid G, et al. J Reprod Med 2001;46:545. [PubMed: 11441678]
- 35. Land KM, et al. Mol Microbiol 2004;51:115. [PubMed: 14651615]
- 36. Kwon DH, et al. Antimicrob Agents Chemother 2001;45:2609. [PubMed: 11502537]
- 37. Leiros HK, et al. J Biol Chem 2004;279:55840. [PubMed: 15492014]
- 38. Cavalier-Smith T. Ann Bot (London) 2005;95:147. [PubMed: 15596464]
- 39. Gregory TR. Biol Rev Camb Philos Soc 2001;76:65. [PubMed: 11325054]
- 40. Honigberg, BM.; Brugerolle, G. Trichomonads Parasitic in Humans. Honigberg, BM., editor. Springer-Verlag; New York: 1990. p. 5-35.
- 41. Shuter BJ. Am Nat 1983;122:26.
- 42. Rendon-Maldonado JG, Espinosa-Cantellano M, Gonzalez-Robles A, Martinez-Palomo A. Exp Parasitol 1998;89:241. [PubMed: 9635448]
- 43. We are grateful to the *Trichomonas* research community and M. Gottlieb for support and guidance during the duration of this project. We thank M. Delgadillo-Correa, J. Shetty, S. van Aken, and S. Smith for technical support; T. Creasy, S. Angiuoli, H. Koo, J. Miller, J. Orvis, P. Amedeo, and E. Lee for engineering support; W. Majoros and G. Pertea for data manipulation; and S. Sullivan and E. F. Merino for editing. Funding for this project was provided by the National Institute of Allergy and Infectious Diseases (grants UO1 AI50913-01 and NIH-N01-AI-30071). The Burroughs Wellcome Fund and the Ellison Medical Foundation provided funds for *Trichomonas* community meetings during the genome project. The Chang Gung *T. vaginalis* Systems Biology Project was supported by grants (SMRPD33002 and SMRPG33115) from Chang Gung Memorial Hospital and Taiwan Biotech Company Limited. This whole genome shotgun project has been deposited at DNA Data Bank of Japan/European Molecular Biology Laboratory/GenBank under the project accession AAHC00000000. The version described here is the first version, AAHC01000000.

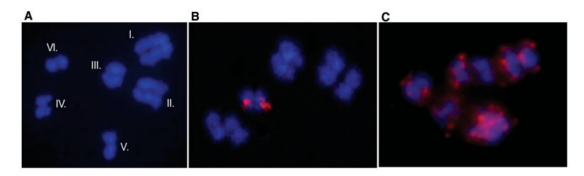


Fig. 1.Karyotype and fluorescent in situ hybridization (FISH) analysis of *T. vaginalis* chromosomes. (**A**) Metaphase chromosome squashes of *T. vaginalis* reveal six chromosomes (I to VI). (**B**) FISH analysis using an 18S rDNA probe shows that all ~250 rDNA units localize to a single chromosome. (**C**) In contrast, the *Tvmar1* transposable element (8) is dispersed throughout the genome.

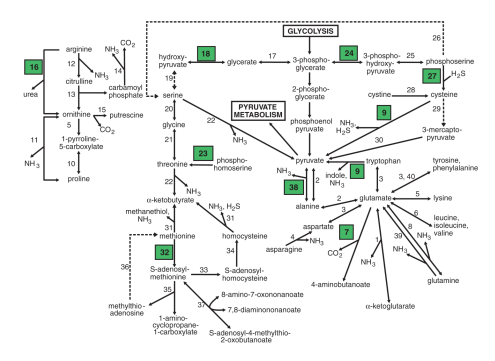


Fig. 2. Schematic of *T. vaginalis* amino acid metabolism. A complete description of enzymatic reactions (represented as numbers) is given in the SOM text. Broken lines represent enzymes for which no gene was identified in the genome sequence, although the activity would appear to be required. Green boxes indicate enzymes encoded by candidate LGT genes.

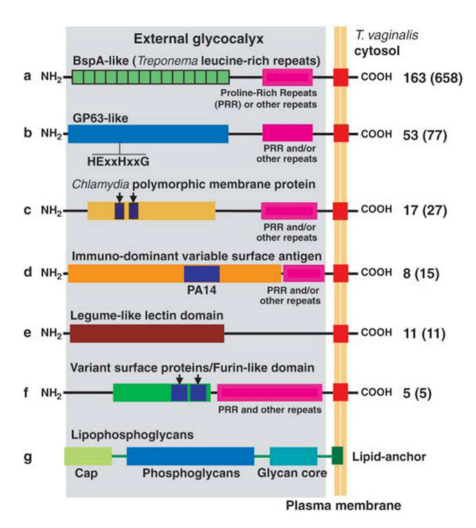


Fig. 3. Structural organization of putative *T. vaginalis* surface molecules involved in host cell adherence and cyto-toxicity (28). Candidate surface protein families (a to f) are depicted as part of the glycocalyx (gray shading), known to be composed of LPG (g). The number of proteins with an inferred transmembrane domain (red box) is indicated at right, with the size of the entire family shown in parentheses. Substantial length variation exists between and within families (proteins are not drawn to scale). Additional information can be found in table S18.

Table 1

Summary of the *T. vaginalis* genome sequence data. Assembly size (bp, base pairs) includes all contigs and differs from estimated genome size of ~160 Mb (4). The scaffold size is the minimum scaffold length, such that more than half the genome is contained in scaffolds of at least that length. The number of predicted genes may include low-complexity repeats or novel transposable elements rather than true *T. vaginalis* genes, but in the absence of decisive evidence these remain in the gene set. The number of evidence-supported genes includes those with either similarity to a known protein ($E < 1 \times 10^{-10}$, >25% length of protein) or similarity to an expressed sequence tag (>95% identity over >90% length of the gene). A total of 763 rDNA fragments (258 copies of 28S, 254 copies of 18S, and 251 copies of 5.8S) were identified.

Feature	Value
Size of assembly (bp)	176,441,227
G+C content (%)	32.7
No. of scaffolds	17,290
$N_{\rm so}$ scaffold size (bp)	68,338
Protein-coding ger	nes
No. of predicted genes	59,681
No. of evidence-supported genes	25,949
No. of genes with introns	65
Mean gene length (bp)	928.6
Gene G+C content (%)	35.5
Gene density (bp)	2956
Mean length of intergenic regions (bp)	1165.4
Intergenic G+C content (%)	28.8
Non-protein-coding	genes
Predicted tRNA genes	479
Predicted 5.8S, 18S, and 28S rDNA units	~250

NIH-PA Author Manuscript NIH-PA Author Manuscript

NIH-PA Author Manuscript

Table 2 Summary of highly repetitive sequences in the genome of *T. vaginalis*. The 31 unclassified repeat families have been collapsed into one group.

17.28h	Repeat name P	Putative identity	Copy no.	Length (bp)	Cumulative length (bp)	Average pairwise difference (%)
Proximary D5 protein 203 Vital Hyage tail fiber prot			12::21			
Properties Protein 203		u A		03.40	010	-
Phage tail fiber prot 2943		oxvirus D5 protein	203	2348	3/0,658	12
Hypothetical 2243		hage tail fiber prot	903	752	607,929	3.9
KilA-N terminal domain 867		lypothetical	2243	1037	1,879,762	3.0
Exercise total Exercise total Exercise total		ilA-N terminal domain	867	873	655,984	9.9
KilA-N terminal domain 831 KilA-N terminal domain 931 KilA-N terminal domain 931 KilA-N terminal domain 931 KilA-N terminal domain 931 Mariner transposase 982 Integrase 1842 Mutator-like profile 173 Mutator-like profile 173 Mutator-like profile 365 Integrase 54 Mutator-like profile 566 Endonuclease 49 Integrase 54 Integrase 56 Integrase 57 Integrase 68 Set (total) 19 Reverse transcriptase 66 Sasos Copia-like profile 6.5 Set (total) 33393 Copia-like profile 6.5 Set (total) 33393 Set (total) 31 unclassified familiess Set (total) 32 unclassified familiess Set (total)		oxvirus D5 protein	954	2637	1,721,187	4.6
KilA-N teminal domain 518		ilA-N terminal domain	831	663	503,959	3.6
Parine transposase Parametric profile		ilA-N terminal domain	518	699	298,778	3.1
Mariner transposase 982 Transposon Integrase 384 Mutator-like profile 282 Integrase 1842 1842 HNH andonuclease 173 173 Mutator-like profile 580 51 Integrase 51 56 Integrase 49 1148 Integrase 566 1148 Integrase 54 56 Integrase 19 56 Integrase 68 56 Integrase 68 68 Integrase 66 65 Integrase 66 7 Integrase 66 7 Integrase 66 65 Integrase 66 66 Integrase 66 66 Integrase 7 <t< td=""><td></td><td></td><td></td><td>1283</td><td>(6,038,257)</td><td>3.7</td></t<>				1283	(6,038,257)	3.7
Mariner transposase 982 Integrase 1842 1842 Integrase 1842 1842 Integrase 173 1842 Mutator-like profile 173 1842 Mutator-like profile 173 1842 Mutator-like profile 173 1842 Integrase 174 174 1844 Integrase 174 175 175 Integrase						
Integrase 384 Mutator-like profile 1842 Integrase 173 Mutator-like profile 173 Mutator-like profile 173 Mutator-like profile 260 Integrase 260 Integrase 275 Integrase 275 Integrase 276 Integrase 276 Integrase 276 Integrase 276 Integrase 276 Integrase 277		fariner transposase		1304	1,158,473	0.5
Mutator-like profile 282 Integrase 1842 HNH endonuclease 173 Mutator-like profile 365 Integrase 518 Integrase 518 Integrase 548 Integrase 549 Integrase 549 Integrase 549 Integrase 549 Integrase 549 Integrase 540 Integrase 54		ntegrase	384	2246	518,936	0.0
Integrase 1842 173 173 174 175		futator-like profile	282	2954	303,256	1.1
HNH endonuclease 200 Mutator-like profile 365 Integrase 51 Mutator-like profile 580 Integrase 51 Mutator-like profile 566 Integrase 54 Integrase 54 Integrase 54 Integrase 56 Integrase 68		ntegrase	1842	981	1,634,764	2.1
th Mutator-like profile 173 Mutator-like profile 365 Integrase 51 Integrase 75 Endonuclease 566 Integrase 1148 Integrase 54 Integrase 56 Integrase 56 Integrase 56 Integrase 56 Integrase 68 ge (total) Reverse transcriptase ge (total) 65 ge (total) 7 ge all 660		INH endonuclease	200	800	131,491	1.5
Mutator-like profile 365 Integrase 5180 Integrase 5180 Mutator-like profile 5580 Integrase 1148 Integrase 1148 Integrase 556 Integrase 556 Integrase 576 Integrase 576 Integrase 576 Integrase 576 Integrase 576 Integrase 576 Integrase 688 Integrase 698		futator-like profile	173	1129	137,526	0.7
Integrase 580 Integrase 51 Mutator-like profile 75 51 51 51 51 51 51 51		futator-like profile	365	2410	368,152	1.1
Integrase 75 Mutator-like profile 75 Endonuclease 566 Integrase 1148 Integrase 154 Integrase 148 Integrase 156 Integrase 150 Integrase		ntegrase	580	2433	636,901	1.5
Mutator-like profile 75		ntegrase	51	1323	51,185	1.6
Endonuclease 566 Integrase 1148 Integrase 54 Integrase 56 Integrase 56 Integrase 927 Integrase 68 Integrase 7 Integrase 7 Integrase 7 Integrase 7 Integrase 68 Integrase 68 Integrase 68 Integrase 7 Integrase 7 Integrase 68 Integrase 7 I		futator-like profile	75	2127	118,581	0.0
Integrase		ndonuclease	566	765	329,717	5.4
Integrase 1148 Integrase 54 Integrase 56 Integrase 7 Integrase 19 Integrase 19 Reverse transcriptase 6 Copia-like profile 6.5 J 31 unclassified families 10 660 1148 148 19 10 10 11 11 12 14 15 15 16 17 18 19 10 10 10 11 11 12 13 14 15 15 16 17 18 19 10 10 10 11 12 13 14 15 16 17 17 18 19 10 10 10 10 11 11 12 13 14 15 16 17 17 18 19 10 10 10 10 10 10 11 12 13 14 15 16 17 17 18 19 10 1		ntegrase	49	1151	45,564	2.5
Integrase 54 Integrase 56 Integrase 56 Integrase 56 Integrase 37 1ntegrase 927 10tegrase 19 19 Integrase 68 19 Integrase Copia-like profile 6 19 10tegrase 10tegrase 6 10tegrase 10tegrase 6 10tegrase 10tegrase 6 10tegrase 10t		ntegrase	1148	1141	1,175,921	2.6
Integrase 56 Integrase 37 Integrase 927 Integrase 19 Integrase 68 Integrase 68 Integrase 68 Copia-like profile 6.5 Integrase 6 Integrase 7 Integrase 6 Integrate 6 Integrase		ntegrase	54	1250	53,674	2.2
Integrase 37		ntegrase	56	1601	73,737	1.4
Integrase 927 19 68 19 68 19 68 19 68 19 69 19 69 69 69 69 69		ntegrase	37	606	27,708	1.4
Integrase 19 68 68 68 68 68 68 68 6		ntegrase	927	1184	807,711	2.5
Integrase 68 414		ntegrase	19	1124	15,254	1.3
However transcriptase 6 Retrotransposon Copia-like profile 6.5 31 unclassified families 793 (660)		ntegrase	89	1301	22,023	2.3
Reverse transcriptase 6 6 7 7 7 6.5 3.1 unclassified families 7 7 7 93 1 onclassified families 793 660	(1		414	1481	(7,610,574)	1.8
Reverse transcriptase 6 7 7 6.5 1) 6.5 31 unclassified families 1) 793			Retrotransposon			
Copia-like profile 7 I) 6.5 31 unclassified families 793 (660		everse transcriptase		2349	6,833	1.0
 4) 5) 31 unclassified families 1) 660 		opia-like profile	7	11001	16,504	7.5
31 unclassified families 793 660	erage (total)			6675	(23,337)	4.3
793						
099	erage (total)		793	1131	(24,617,704)	2.6
000	al all		099	1450	38,289,812	2 6
	stage an		000	00+1		C:-3

Table 3

Large gene families in *T. vaginalis*. Only gene families, or aggregates of gene families mediating a given process, for which there are more than 30 members and that have been assigned a putative function are listed. (See SOM text for subfamily organization.) The small guano-sine triphosphatases (GTPases) are the sum of Rab and ARF small GTPases only; all other GTPases are shown in table S12. ABC, ATP-binding cassette; MFS, major facilitator superfamily; MOP, multi-drug/oligosaccharidyl-lipid/polysaccharide; AAAP, amino acid/auxin permease.

Gene family (functional unit)	Members
Protein kinases	927
BspA-like gene family	658
Membrane trafficking: small GTPases	328
rDNA gene cluster	~250
Cysteine peptidase (clan CA, family C19)	117
Membrane trafficking: vesicle formation	113
ABC transporter superfamily	88
GP63-like (leishmanolysin)	77
MFS transporter family	57
MOP flippase transporter family	47
Cysteine peptidase (clan CA, family C1)	48
AAAP transporter family	40
Dynein heavy chain	35
P-ATPase transporter family	33
Serine peptidase (clan SB, family S8)	33
Membrane trafficking: vesicle fusion	31