**Trichomonas vaginalis** Lipophosphoglycan Mutants Have Reduced Adherence and Cytotoxicity to Human Ectocervical Cells

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The extracellular human pathogen **Trichomonas vaginalis** is covered by a dense glycocalyx thought to play a role in host-parasite interactions. The main component of the glycocalyx is lipophosphoglycan (LPG), a polysaccharide anchored in the plasma membrane by inositol phosphohexameride. To study the role of LPG in trichomondans, we produced **T. vaginalis** LPG mutants by chemical mutagenesis and lectin selection and characterized them using morphological, biochemical, and functional assays. Two independently selected LPG mutants, with growth rates comparable to that of the wild-type (parent) strain, lost the ability to bind the lectins *Ricinus communis* agglutinin I (RCA120) and wheat germ agglutinin, indicating alterations in surface galactose and glucosamine residues. LPG isolated from mutants migrated faster than parent strain LPG on sodium dodecyl sulfate-polyacrylamide gel electrophoresis, suggesting the mutants had shorter LPG molecules. Dionex high-performance anion exchange chromatography with pulsed amperometric detection analyses revealed galactosamine, glucosamine, galactose, glucose, mannose/xylene, and rhamnose as the main monosaccharides of **T. vaginalis** parent strain LPG. LPG from both mutants showed a reduction of galactose and glucosamine, corresponding with the reduced size of their LPG molecules and inability to bind the lectins RCA120 and wheat germ agglutinin. Mutant parasites were defective in attachment to plastic, a characteristic associated with avirulent strains of **T. vaginalis**. Moreover, the mutants were less adherent and less cytotoxic to human vaginal ectocervical cells in vitro than the parental strain. Finally, while parent strain LPG could inhibit the attachment of parent strain parasites to vaginal cells, LPG from either mutant could not inhibit attachment. These combined results demonstrate that **T. vaginalis** adherence to host cells is LPG mediated and that an altered LPG leads to reduced adherence and cytotoxicity of this parasite.
lial cells, indirectly suggesting that LPG is involved in adherence (27).

To study the role of T. vaginalis LPG in host-parasite interactions, we have constructed LPG mutants by chemical mutagenesis followed by selection with the galactose-specific lectin RCA120. Here we report the characterization of these T. vaginalis LPG mutants and demonstrate that mutants bearing truncated surface LPG are deficient in binding and lysing host cells.

MATERIALS AND METHODS

Parasites and media. Strain B7RC2 (ATCC 50167) was selected as it showed the highest adhesion and cytotoxicity to mammalian cells out of several strains tested in preliminary experiments (data not shown). TYM medium supplemented with 10% heat-inactivated horse serum (Invitrogen), 10 U penicillin (Invitrogen), 10 μg streptomycin (Invitrogen), 180 μM ferrous ammonium sulfate, and 28 μM sulfosalicylic acid was used to grow the parasites (12). Parasites were passaged daily (~10^6 parasites in 10 ml TYM). As an increased number of in vitro passages has been associated with reduced virulence in T. vaginalis (43), the parent and mutant parasites were passed a comparable number of times.

Chemical mutagenesis and lectin selection. Two independent mutagenesis experiments were carried out as previously described for Leischmania with slight modifications (35). Briefly, 3 × 10^7 parasites were grown overnight in complemented TYM and cultures were washed with TYM, resuspended to 3 × 10^7 cells/ml in medium, and incubated with 9 μg/ml ethyl methanesulfonate (Sigma) shaking (70 rpm) at 37°C for 4 h. No cell death was observed at this time. Mutagenized parasites were washed once with TYM. Approximately 50% cell death was observed for cells exposed to ethyl methanesulfonate after culturing overnight in TYM at 37°C. Cells were then grown for several generations before 10^7 parasites were selected using Rausin communis agglutinin I (RCA120).

For lectin selection, parasites were washed in TYM and incubated overnight at 37°C in TYM with 100 μg/ml RCA120 (Sigma). Agglutinated cells were removed by low-speed centrifugation. Lectin selection procedure was repeated three times before cloning of populations in soft agar (32). Parasite colonies that did not label with fluorescein isothiocyanate-RCA120 were cloned three more times by limiting dilution in 96-well tissue culture plates under anaerobic conditions in GasPak jars (Becton Dickinson and Co).

Fluorescent labeling of parasites. Overnight cultures of parasites were fixed in 5% Formalin at room temperature for 5 min. Fixed parasites were washed once with phosphate-buffered saline (PBS), and pellets were resuspended in a 1:500 dilution of fluorescein isothiocyanate-RCA120 or rhodamine-wheat germ agglutinin (Vector Laboratories). Stained parasites were examined using an Axioscope 2 epifluorescent microscope (Zeiss), and images were recorded with a Axiocam camera and processed with the Avisoision 3.2 program (Zeiss).

LPG isolation and staining. LPG was isolated by adapting the method for bacterial lipopolysaccharide isolation from Johnson and Perry (33) followed by solvent E extraction as used in Leischmania (35). Briefly, parasites were washed once with PBS and resuspended in 10 mM Tris, pH 8.0. Cells were sonicated on ice for 30 seconds and incubated at 37°C for 2 h with RNase A (2 μg/ml; Sigma) and DNase Q1 (2 μg/ml; Promega), followed by overnight incubation at 4°C with 2.5 μg/ml proteinase K (Roche). Polysaccharides were extracted by adding an equal volume of phenol (equilibrated to pH 8.0), and sonication on ice for 15 seconds. Phases were separated by centrifugation, and the aqueous phase was dialyzed overnight at 4°C and concentrated. LPG was then extracted with solvent E (H₂O, ethanol, diethylether, pyridine, NH₄OH, 15:15:5:10:0.02) before dialysis in PBS and further concentration. LPG was visualized by fractionation on sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) and subsequent staining with periodic acid silver.

Monosaccharide composition. The monosaccharide composition of purified LPG was determined by hydrolysis in 2 N trifluoroacetic acid at 100°C for 4 h, followed by Dionex high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) using a CarboPac PA1-column at the University of California–San Diego Glycotechnology Core Resource (http://glycotech.ucsd.edu/index.shtml). Preliminary sequence data was obtained from the Institute for Genomic Research through the website at http://www.tigr.org.

Adhesion to plastic. The parent and LPG mutant parasites were cultured overnight in TYM medium and washed once with warm (37°C) nonsupplemented TYM. Parasites were resuspended in warm TYM to 2.5 × 10^7 cells/ml, and 2 ml of the suspension were incubated in triplicate in 24-well plates for 4 h at 37°C. After incubation, wells were washed three times with nonsupplemented warm TYM to remove unattached parasites. Wells were then incubated with crystal violet (0.13% crystal violet/5:2 ethanol-formaldehyde) for 5 min and washed three times with PBS. Remaining dye was solubilized with 1% SDS in 50% ethanol, transferred to 96-well plates, and measured in a plate reader at 570 nm. The experiment was repeated three times.

Adherence to human vaginal ectocervical cells. The human ectocervical cell line Ect/EG6E7 (obtained from Ruina Fichorova) was grown as described (23) in keratinocyte-SFM ( Gibco) complemented with bovine pituitary extract (GIBCO), 100 ng/ml recombinant epidermal growth factor (GIBCO), 0.4 mM CaCl₂, 100 U/ml penicillin, and 100 μg/ml streptomycin (GIBCO). Ectocervical cells were seeded on 12-mm coverslips in 24-well plates at 5 × 10^5 cells/well in culture medium and grown to confluence at 37°C and 5% CO₂ for 2 days. Cells were washed once with PBS before the addition of parasites. Parasites were labeled with 10 μM CellTracker Blue CMAC as per the manufacturer’s instructions (Molecular Probes). Approximately 5 × 10^5 labeled parasites were washed and added to ectocervical cells in 1 ml PBS. Plates were spun at 500 × g for 5 min and incubated at 37°C and 5% CO₂ for 30 min. Coverslips were subsequently washed in PBS, fixed with 4% paraformaldehyde, and mounted on slides. Five × magnification fields were analyzed per coverslip. Fluorescent parasites adhered to host cells were counted using Scion Image for Windows, v. Beta 4.0.2 (Scion Corporation). The experiment was repeated at least three independent times.

Adherence inhibition by isolated LPG. LPG from parent and mutant parasites was quantified by densitometry on polysaccharide-stained SDS-PAGE gels relative to a known concentration of Salmonella lipopolysaccharide using the ImageJ program from the National Institutes of Health (http://rsb.info.nih.gov/j). Ectocervical cells were incubated with 150 μg of purified LPG in PBS at 37°C and 5% CO₂ for 1 h prior to the addition of fluorescently labeled parent parasites. The number of parasites attached to the monolayer was determined as described above.

Cytotoxicity to epithelial cells. Ectocervical cells were seeded in 48-well plates at 3 × 10^5 cells/well in culture medium and grown at 37°C and 5% CO₂ for 2 days. Cells were washed once with PBS before the addition of parasites. Approximately 1.5 × 10^6 parasites were washed and added to ectocervical cells in 1 ml PBS. Plates were spun at 500 × g for 5 min and incubated at 37°C and 5% CO₂ for 8 h. After incubation, 100 μl of parasite/cell culture supernatant was transferred to 96-well plates and allowed to equilibrate to room temperature for 30 min. Cytotoxicity was measured as a function of lactate dehydrogenase release from ectocervical cells using the CytoTox-ONE homogeneous membrane integrity assay as per the manufacturer’s instructions (Promega). Fluorescence was read using an FLx800 microplate fluorescence reader (Bio-Tek Instruments, Inc.). Paraste cytotoxicity was measured against 100% lysis of cells using lysis solution included in the CytoTox-ONE kit. Cytotoxicity values were calculated as follows: % cytotoxicity = 100 × [fluorescence reading of sample well] – [blank]/[fluorescence reading of 100% lysed cells] – [blank].

Statistical analyses. The adherence, LPG competition, and cytotoxicity data were analyzed using analysis of variance. Multiple comparisons of all pairs were performed using the method of Tukey-Kramer honestly significant difference (HSD) with an alpha level of 0.05 (Statistical software package JMPIN v3, SAS Institute).

RESULTS AND DISCUSSION

In this study, we have generated mutants of T. vaginalis that express truncated surface LPG with a modified sugar composition. These LPG mutants exhibited reduced adherence and cytotoxicity to human vaginal ectocervical cells. Mutant-derived LPG was incapable of inhibiting binding of the parasite to vaginal cells, in contrast to that observed using parent-derived LPG. These data demonstrate a critical role for LPG in the interaction of this extracellular parasite with host cells.

**T. vaginalis chemical mutants do not bind RCA120 or wheat germ agglutinin lectins.** Mutants of T. vaginalis strain B7RC2 (PA strain) were obtained by chemical mutagenesis with ethyl methanesulfonate and selection with the galactose-specific lectin RCA120. Two clonal populations (2E2 and 4-12), selected from independent mutagenesis experiments and used throughout the study, showed no surface reactivity.
when probed with fluorescein isothiocyanate-RCA120 or rhodamine-wheat germ agglutinin (Fig. 1). Mutant parasites also did not agglutinate in the presence of fluorescein isothiocyanate-RCA120 or rhodamine-wheat germ agglutinin, whereas strong agglutination of parent parasites was observed. The surface of all three parasite lines was found to be positively labeled with the mannose-specific lectin concanavalin A (data not shown).

The lack of reactivity with RCA120 and wheat germ agglutinin suggested that mutant parasites had lost surface terminal galactosyl and glucosamine residues, the main two monosaccharides in T. vaginalis LPG (47). Lectin-specific binding to the surface of the parent strain was demonstrated by showing that surface labeling with RCA120 and wheat germ agglutinin lectins can be blocked by competition with galactose and glucosamine, respectively (data not shown). Interestingly, the fluorescent lectins also labeled flagella of the parent strain with the same intensity as the cell membrane (inset, Fig. 1), indicating that T. vaginalis flagella may be covered by LPG, as reported for Leishmania flagella (46). This observation is noteworthy as previous studies have described adherence of trichomonads to host cells by flagella (13), raising the possibility that flagellum-mediated adherence may involve LPG.

A small percentage (1 to 4%) of lectin-positive cells were occasionally observed in the 2E2 and 4-12 mutants. Mutant parasites, however, remained negative for lectin binding after prolonged culture (>1 month of daily passage) and did not revert to the wild-type phenotype. Complementation studies to rescue lectin binding phenotype and uncover the mutated gene responsible for this phenotype were attempted without success. This is likely due to the combination of low transfection effi-

FIG. 1. Fluorescent lectin labeling of parent (PA) and LPG mutant parasites (2E2 and 4-12). Parasites were fixed with 5% formalin, washed with PBS, and labeled with the fluorescently labeled lectins fluorescein isothiocyanate-RCA120 (FITC-RCA120) or rhodamine-wheat germ agglutinin (Rho-WGA), as indicated.
ciency and the atypically large genome size (151 to 177 Mb) (36) of *T. vaginalis*, as the cosmid libraries used to attempt complementation were found to represent only half of the genome (data not shown).

*T. vaginalis* mutants do not have impaired growth in vitro. Chemical mutagenesis can simultaneously affect more than one gene, and altered parasite growth rates might influence adherence and cytotoxicity experiments. It was therefore important to demonstrate that the mutagenesis did not negatively affect the growth and morphological characteristics of the selected mutants. Light microscopic examination of the parasites did not reveal any morphological differences between the parent and the mutants. Analyses of growth rates over a 20-h period also showed no significant difference between the mutant and parent strains (Fig. 2). Based on growth rates, exponentially dividing cells were used to further characterize the adherent and cytotoxic properties of LPG mutants.

*T. vaginalis* chemical mutants have truncated LPG molecules. To determine if the lack of surface reactivity of mutant parasites for RCA120 and wheat germ agglutinin lectins was due to LPG loss, LPG was isolated from the parent and mutant parasites and subjected to SDS-PAGE analysis. Polysaccharide-specific silver staining of the gel revealed that the LPG of the two mutants migrated faster than that of parent cells, suggesting truncations in the polysaccharide chain (Fig. 3). Isolated LPG migrated as a broad band as previously described (48), with the parent LPG band migrating between 30 and 100 kDa, and the mutant LPG bands running between 25 and 50 kDa. Small shifts in mobility were observed between the two mutant lines, suggesting that they carry mutations in different LPG synthesis or assembly genes.

The polysaccharide nature of the isolated LPG was confirmed by staining LPG gels with the polysaccharide-specific periodic acid Schiff stain (data not shown). In contrast to the *Leishmania donovani* LPG chemical mutant R2D2 (35), which does not synthesize appreciable amounts of LPG, our *T. vaginalis* LPG mutants express LPG with apparent modification in polysaccharide content and/or structure. However, other *L. donovani* mutants that generate truncated LPG have also been produced (8).

LPG from mutant parasites has reduced levels of galactose and glucosamine. To further investigate the LPG from mutant parasites, the polysaccharide composition of isolated LPGs was determined by Dionex HPAEC-PAD (Table 1). The main monosaccharides resolved from the *T. vaginalis* parent strain LPG were galactosamine, glucosamine, galactose, glucose, mannose/xylose, and rhamnose (Table 1). The reduced relative levels of galactose and glucosamine in the mutant LPG are consistent with their lack of reactivity with RCA120 and wheat germ agglutinin (Fig. 1). Because the mutant parasite LPGs appear to be truncated (Fig. 3), the large reduction of galactose and glucosamine in the mutant LPGs (50%) indicates that the majority of these monosaccharides are located within the side chains and/or the terminal cap of the LPG. The reduction of these monosaccharides is accompanied by an in-

| Monosaccharide<sup>a</sup> | Parent | 2E2 | 4-12 |
|----------------------------|--------|-----|------|
| Rha                        | 23.4   | 23.9| 20.0 |
| GalNH2                     | 2.3    | 22.0| 10.9 |
| GlcNH2                     | 33.0   | 15.5| 9.9  |
| Gal                        | 18.9   | 5.1 | 3.2  |
| Glc                        | 6.0    | 20.4| 43.8 |
| Man/Xyl                    | 16.5   | 13.1| 12.1 |

<sup>a</sup> Rha, rhamnose; GalNH2, galactosamine; GlcNH2, glucosamine; Gal, galactose; Glc, glucose; Man, mannose; Xyl, xylose.
crease in galactosamine and glucose. Galactosamine, along with glucosamine is the main component of the glycan core of T. vaginalis LPG (49). The relative increased concentration of galactosamine in the LPG mutants is consistent with its presence primarily in the glycan core and further reinforces that the LPG of the mutants has sustained losses in the LPG side chains or cap.

Although glucose appears to be increased in the mutant LPGs, it is worth noting that glucose is a common environmental contaminant, and therefore this result should be interpreted with caution. An apparent difference in the relative concentration of monosaccharides also exists between the LPG mutants. Mutant 4-12, for example, has about half the amount of galactosamine of mutant 2E2. These differences between the mutants reflect the fact that the mutants were independently selected and suggest that these two strains have mutations in different steps of LPG biosynthesis.

The LPG monosaccharide composition of the parent strain shown in Table 1 differs in two aspects from that previously reported for another T. vaginalis strain (47, 48). One is the higher percentage of mannose/xylose, and second and most notable is the presence of rhamnose, which was not previously reported. Mannose and xylose frequently comigrate on Dionex HPAEC-PAD. Using different conditions to resolve the mannose/xylose peak, we determined that xylose is about 75% and 40% of the combined mannose/xylose peak for the parent and mutant LPGs, respectively. The difference between our results and those published previously for mannose can therefore be accounted for by the presence of xylose, another common contaminant in polysaccharide preparations. Whether the presence of rhamnose in T. vaginalis LPG reflects strain differences is unclear. The high percentage (>20%) of rhamnose strongly argues against its being a contaminant. Furthermore, rhamnose has recently been described in Tritrichomonas foetus LPG (50). Because rhamnose runs adjacent to galactosamine in Dionex HPAEC-PAD, we ran a rhamnose standard ramp and confirmed its presence and abundance in T. vaginalis parent and mutant strain LPGs (Fig. 4).

Rhamnose is a monosaccharide found in bacteria, some protists, and plants but is absent in mammals (1, 20, 28). It is notable that rhamnose is also not present in Leishmania LPG, the only well-characterized protist LPG (19, 38, 55). Analyses of the T. vaginalis genome database (www.tigr.org/tdb/tgi/tvgi) have revealed putative homologues for the genes encoding the core, raising the possibility that drug-mediated interference of rhamnose might disrupt the surface of the parent strain and the mutants suggests this monosaccharide is likely to be located in the main backbone or the glycan core, raising the possibility that drug-mediated interference of l-rhamnose synthesis or assembly might disrupt the surface of these parasites, rendering them avirulent (see results below).

In addition, disruption of this pathway in bacteria produces less-virulent organisms with truncated or rough lipopolysaccharide, the functional and structural analog to LPG in bacteria (11, 25).

**FIG. 4.** Dionex HPAEC-PAD rhamnose ramp. Chromatogram from HPLC analysis of monosaccharides using CarboPac PA-1 column. Monosaccharides from purified parent LPG (a) or standards (b) were resolved by Dionex HPAEC-PAD to confirm the presence of rhamnose in T. vaginalis LPG. The retention time for rhamnose in the standards coincides with the rhamnose peak in the LPG sample.

**LPG mutants show reduced adherence to vaginal epithelial cells in vitro.** To gain insight into the role of LPG in T. vaginalis pathogenesis, we focused on functional analyses of the LPG mutant parasites. Attachment and cytotoxicity to vaginal epithelial cells are critical for the virulence of this extracellular parasite (21, 43). Moreover, changes in LPG have also been associated with reduced virulence in Leishmania spp. (18, 55, 56). Because in vitro adhesion to plastic has been associated with in vivo virulence of T. vaginalis strains (29), we first tested if the parent and LPG mutants differed in attachment to plastic surfaces. The parasites showed significant differences in attachment to polystyrene surfaces (\( P < 0.0001 \); mean ± standard deviation: parent, 0.54 ± 0.118; 2E2, 0.08 ± 0.005; 4-12, 0.08 ± 0.003), with the adherence of the parent strain being >6x that of the mutants after four hours incubation (Fig. 5). Attachment differences between mutants were minor and not statistically significantly different.

We then tested our T. vaginalis LPG mutants for in vitro attachment to recently established human ectocervical epithelial cells Ect1/E6E7 (23). After 30 min incubation of fluoro-
cence labeled parasites on ectocervical cell monolayers, the parent strain showed a strikingly better attachment to ectocervical cells than the LPG mutants (Fig. 6a). The average number of parent parasites attached to the ectocervical cells per coverslip was seven fold higher than that of the LPG mutants (parent 2,841 ± 570.3 versus 2E2 391 ± 107.7 and 4-12 393 ± 138.5) (Fig. 6b). No significant differences in adhesion to ectocervical cells were observed between mutant parasites. These data indicate that the truncation in the LPG molecules of the mutant strains is responsible for reduced adherence to ectocervical cells, and provides the most direct evidence to date for a role for \textit{T. vaginalis} LPG in host cell adherence. The human ectocervical epithelial cell line used for these studies, although immortalized, has been shown to express epithelial differentiation proteins and immune mediators characteristic of primary cells from the human ectocervix (22, 23). These results therefore are likely to reflect properties of adherence in vivo (27). It is interesting to note the strong correlation between the ability to adhere to plastic and to ectocervical cells, as previously reported (29).

**LPG from the parent strain but not the mutant strains inhibits adherence to ectocervical cells.** To further demonstrate that LPG plays a critical role in \textit{T. vaginalis} adherence to host cells, and to determine whether the reduced adherence of the LPG mutant strains to host cells is directly attributable to their modified LPG, we performed competitive inhibition adherence assays. For these experiments, we analyzed the binding of parent parasites to ectocervical cells in the presence of LPG isolated from parent or mutant parasites. Preliminary experiments showed that inhibition of attachment with parent LPG was dose dependent and reached saturation at 150 µg/ml (data not shown). Parent LPG (150 µg/ml) inhibited parasite attachment to ectocervical cells significantly better than 2E2 or 4-12 LPG (Fig. 7). The number of attached parasites was reduced by 44% in the presence of parent LPG compared to the control untreated ectocervical cells. In contrast, 2E2 and 4-12 LPG only reduced parasite binding by 14% and 15%, respectively. These data show that \textit{T. vaginalis} adherence to ectocervical cells is mediated partially by LPG and that mutation of the LPG renders it incapable of mediating this host-parasite interaction.

The inability of parent LPG to completely abrogate adherence

**FIG. 5.** Adherence to plastic by parent and LPG mutant parasites. Parent (PA) and LPG mutant parasites were grown to mid-log phase, washed, and resuspended in TYM medium, and 5 × 10³ cells were distributed in triplicate in the wells of 24-well tissue culture plates. After 4 h incubation at 37°C and 5% CO₂, wells were washed three times and attached parasites were stained with crystal violet. Absorbance was then measured at 570 nm after dissolving the dye in SDS-ethanol. The graph depicts data from a representative experiment with standard deviations.

**FIG. 6.** Adherence of parent (PA) and LPG mutant parasites to human ectocervical cell monolayers. Mid-log-phase parasites were labeled with Cell Tracker Blue (Molecular Probes). Labeled parasites were then incubated for 30 min with ectocervical cell monolayers grown on coverslips in 24-well plates at 37°C and 5% CO₂. Coverslips were washed to remove nonadherent parasites, mounted, and visualized by fluorescence microscopy (a). Data are from three experiments showing the average number of parasites counted per coverslip with standard deviations (b).
of the parent strain to epithelial cells indicates that other factors in *T. vaginalis* are also involved in adherence to epithelial cells. This saturation of binding inhibition with LPG has previously been described for *Tritrichomonas foetus* (51), where a maximum 46% reduction in parasite binding to bovine vaginal epithelial cells was observed with 100 μg/ml of purified *Tritrichomonas foetus* LPG. Our binding inhibition data for *T. vaginalis* LPG closely mirrors the results obtained for *Tritrichomonas foetus*.

**LPG mutants are less cytotoxic to ectocervical cells.** To determine the relative cytotoxicity of the parent strain and LPG mutants to ectocervical cells, an in vitro assay that quantifies the release of lactate dehydrogenase as a measure of host cell intactness was used (Promega). As shown in Fig. 8, parent parasites were over 5 times more cytotoxic to ectocervical than mutant parasites. Differences in cytotoxicity between the LPG mutants were observed, with mutant 4-12 being twice as cytotoxic as mutant 2E2. This difference is statistically significant and was not expected since the adherence of the mutants to ectocervical cells was comparable (Fig. 6b). This difference in cytotoxicity may reflect limitations of the lactate dehydrogenase assay. The maximum incubation time recommended for this assay by the manufacturer is 9 h, whereas microscopic evaluation of ectocervical cell monolayers coinoculated with the parasites overnight show that parent strain parasites completely destroy and ingest the epithelial cell monolayer, while incubation with LPG mutant parasites resulted in almost fully intact monolayers (data not shown). Nonetheless, results from both assays illustrate that LPG plays a critical role in the interaction of *T. vaginalis* with host cells, as LPG mutants are significantly less cytotoxic to epithelial cells of the reproductive tract than the parent strain.

In summary, these studies comparing parental and mutant LPG on *T. vaginalis* demonstrate a direct role for LPG in the host cell adherence and cytotoxicity of this extracellular parasite in vitro. Future efforts towards determining the structures of parent and mutant LPGs should shed light on the critical properties of this parasite glycoconjugate required for host cell recognition and binding.

**ACKNOWLEDGMENTS**

This work was supported by a Burroughs Wellcome Scholar Award for Molecular Parasitology (to P.J.J.) and a Microbial Pathogenesis Training Grant (NIH-T32-AI07323) and NRSA grant (NIH-F31-AI066591) (to C.Y.O.).

We thank Julio-Cesar Carrero and Dara Chang for technical support, Bradley K. Hayes from the UCSD Glycotechnology Core Resource, San Diego, California, for advice on monosaccharide analyses, Raina Fichorova from Brigham and Women’s Hospital, Boston, Massachusetts, for the human ectocervical cells, and Kent Hill for use of the fluorescence microscope. Also, we thank our lab members for helpful discussions.

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