Flavodiiron proteins (FDPs) constitute a recently established superfamily of soluble enzymes, thus far exclusively found in anaerobic and facultative aerobic organisms (2, 19, 54). Originally, the function ascribed to these proteins was the reduction of molecular oxygen to water as reported for *Desulfovibrio gigas* rubredoxin:oxygen oxidoreductase, the first thoroughly characterized protein of this type. This protein was found to utilize electrons derived from glycolysis for safe, four-electron reduction of dioxygen, thus protecting the anaerobic bacterium from the deleterious effects of oxidative stress (19). Later, some of these proteins were also shown to be involved in the reduction of nitric oxide in addition to their oxygen-reducing activity, thereby probably protecting the microbial organism against NO released during the immune response of the higher eukaryote host. The ratio of FDP activity toward oxygen and NO may differ substantially in various organisms; in some cases, FDP is almost exclusively reactive with oxygen, in others it is reactive with NO (20, 21, 43).

FDPs are modular proteins, with flavodoxin-like and metallo-β-lactamase-like domains as their core modules. This two-domain structure is found in the simplest and most common members of the family, named class A FDPs. These proteins are the terminal elements of a multicomponent electron transport chain that uses the reducing power of NAD(P)H to reduce and detoxify dioxygen and/or nitric oxide (41). Proximal electron donors to most class A FDPs are soluble electron transfer proteins. In the class A FDP rubredoxin:oxygen oxidoreductase from the sulfate-reducing bacterium *Desulfovibrio gigas*, the electron donor is a small protein, rubredoxin, that itself is reduced by an NADH:rubredoxin oxidoreductase (9, 10, 22). Besides rubredoxin, roles for other iron-sulfur flavoproteins in electron transport to FDPs have been suggested in several Archaea (41); coenzyme F420H2 is the electron donor for the FDP in the methanogenic archaeon *Methanothermobacter marburgensis* (44). The members of other FDP classes have additional domains fused to the C terminus that participate in electron transfer from the ultimate donor molecule [NAD(P)H] to the terminal electron acceptor (41).

While originally believed to be restricted solely to prokaryotes, recent progress in genome sequencing projects have revealed homologous protein sequences in the genomes of several “amitochondriate” anaerobic protists, mostly with parasitic lifestyles, such as *Trichomonas, Giardia, Entamoeba, Spirotrichonympha*, and a free-living Mastigamoeba (1, 2, 33, 42). *Giardia intestinalis* is the only eukaryotic organism to have had data on its FDP published recently. In line with what is known for the prokaryotic homologues, the giardial protein was shown to possess high oxygen (but not NO)-reducing activity and was therefore proposed to participate in protection against oxidative stress (13).

*Trichomonas vaginalis* is an anaerobic (or microaerophilic) protozoan parasite causing human trichomoniasis, the most common nonviral sexually transmitted infection (38), for which oxygen concentrations higher than those encountered in situ in the vagina (i.e., concentrations above ~60 μM) are toxic (17). The glucose metabolism of *T. vaginalis* is compartmentalized; while the reactions of classical glycolysis producing lactate, as well as the branch resulting in the formation of glyceral (8, 48) occur in the cytosol, a substantial portion of glycolytic carbon is diverted into the hydrogenosomes, a mitochondrion-related organelle where the reactions of extended glycolysis produce additional ATP by oxidative decarboxylation of pyruvate (47, 48). Typical in the trichomonad hydrogenosome is the presence of the iron-sulfur (FeS) cluster-containing enzymes pyruvate:ferrredoxin oxidoreductase (PFOR), hydrogenase, and the
electron carrier ferredoxin, which are involved in the generation of molecular hydrogen using electrons released from pyruvate (36). PFOR and hydrogenase are highly oxygen-sensitive enzymes (29, 32), and it is likely that the sensitivity of trichomonads to oxygen could at least in part be due to the inactivation of these key hydrogenosomal proteins.

*T. vaginalis* must cope with low oxygen concentrations in its natural environment and, accordingly, possesses defense mechanisms to combat oxidative damage caused by oxygen itself or by reactive oxygen species that arise either enzymatically or when the reduced prosthetic groups of enzymes such as flavins and FeS clusters come into contact with oxygen. Most eukaryotes utilize glutathione as a key redox buffer and antioxidant, but trichomonads lack this and similar thiols (17). Cysteine has been suggested as a major reducing buffer and antioxidant (17), and it is believed that the organism relies upon cytosolic NADH oxidase (reducing oxygen to water) and NADPH oxidase (reducing oxygen to hydrogen peroxide) to prevent the permeation of oxygen into the hydrogenosomes (31). Proteins of the peroxiredoxin cascade (11) are also important for cytosolic peroxide detoxification. The identified defense mechanisms of hydrogenosomes include superoxide dismutase activity (17, 30) and recently found putative peroxiredoxines that might provide protection against peroxides (39), but the protein that was suggested long ago to be responsible for oxygen uptake and detoxification has never been identified (6).

We describe here the properties of a class A FDP from *T. vaginalis* hydrogenosomes and suggest its role in the metabolism of oxygen and protection of the organelle.

**MATERIALS AND METHODS**

**Organism.** *T. vaginalis* strain T1 (J.-H. Tai, Institute of Biomedical Sciences, Taipei, Taiwan) was grown in Diamond’s TYM medium without agar as the carbon source in a shaking incubator at 37°C. The hydrogenosomes from 2 to 4 liters of culture were resuspended in a buffer containing 50 mM Tris-HCl (pH 7.6) at a flow rate of 1 ml/min using a BioLogic HR system (Bio-Rad). The purified recombinant enzyme was used for analysis. The isolated hydrogenosomes were subjected to Edman degradation (performed at the Protein/DNA Technology Center, Rockefeller University, New York, NY) to determine the amino-terminal sequences of the polypeptides. The heavier polypeptide was a homologue of the 51-kDa subunit (also called NuoX in bacteria) of the electron-input module of respiratory complex I (25), while the lighter polypeptide was found to be a homologue of bacterial FDPs and named TvFDP. Edman degradation also allowed determination of the processing site where the hydrogenosomal targeting peptide is cleaved from the preprotein by a proteinase processing peptidase.

**Expression and purification of TvFDP.** To express the His-tagged protein (His-FDP) in *E. coli* M15 cells, the bacteria were induced with 0.5 mM IPTG (isopropyl-β-D-thiogalactopyranoside) and grown for 7 h at 25°C in LB medium supplemented with 400 μM ammonium ferrous sulfate and 200 μM flavin mononucleotide (FMN). The harvested cells were homogenized by passage through a French press at 18,000 lb/in². The soluble fraction obtained by ultracentrifugation (250,000 × g, 1 h, 4°C) was applied to a Ni-NTA column (Qiagen) and eluted with a stepwise gradient of 20 mM imidazole (buffer A: 20 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol [pH 7.6]) and 400 mM imidazole (buffer B: 400 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol [pH 7.6]) at a flow rate of 1 ml/min using a BioLogic HR system (Bio-Rad). His-FDP-containing fractions were dialyzed overnight against 50 mM Tris-HCl (pH 7.6) and subjected to Edman degradation (performed at the Protein/DNA Technology Center, Rockefeller University, New York, NY) to determine the amino-terminal sequences of the polypeptides. The heavier polypeptide was a homologue of the 51-kDa subunit (also called NuoX in bacteria) of the electron-input module of respiratory complex I (25), while the lighter polypeptide was found to be a homologue of bacterial FDPs and named TvFDP. Edman degradation also allowed determination of the processing site where the hydrogenosomal targeting peptide is cleaved from the preprotein by a proteinase processing peptidase.

**Characterization of TvFDP.** The flavin cofactor was characterized by reversed-phase high-pressure liquid chromatography using a nucleosil 100-5 C18 column and a thin-layer chromatography (TLC) method (HPTLC-Alufolin; Merck). The TLC mobile phase consisted of n-butanol-acetic acid–water (6:2:8). The flavin was extracted from the protein with trichloroacetic acid at a final concentration of 10%, followed by centrifugation and supernatant neutralization with 1 M ammonium acetate (pH 7.0). The identity of the flavin was determined for recombinant His-FDP, as well as for TvFDPStrep purified from *T. vaginalis* hydrogenosomes according to the IBA Strep-tag protein purification protocol. FMN and FAD (Sigma) were used as standards. The amount of flavin in TvFDP and FDP-containing fractions was determined after protein precipitation with 80% trichloroacetic acid, using an extinction coefficient of 12.500 M⁻¹ cm⁻¹ at λ = 450 nm (21).

The iron content of TvFDP (His-FDP was used for analysis) was determined by using the 2,4,6-tripyridyl-1,3,5-triazine method (18).

**Isolation of hydrogenosomes and partial purification of TvFDP.** Hydrogenosomes were isolated from *T. vaginalis* homogenate by differential centrifugation (250,000 × g, 1 h, 4°C) at a flow rate of 1 ml/min using a BioLogic HR system (Bio-Rad). The supernatant was particularly enriched in two barely separable polypeptides with molecular masses of ~45 kDa. After transblotting onto a polyvinylidine difluoride membrane and Coomassie blue staining, the enriched bands were cut out and subjected to Edman degradation (performed at the Protein/DNA Technology Center, Rockefeller University, New York, NY) to determine the amino-terminal sequences of the polypeptides. The heavier polypeptide was a homologue of the 51-kDa subunit (also called NuoX in bacteria) of the electron-input module of respiratory complex I (25), while the lighter polypeptide was found to be a homologue of bacterial FDPs and named TvFDP. Edman degradation also allowed determination of the processing site where the hydrogenosomal targeting peptide is cleaved from the preprotein by a proteinase processing peptidase.

**Expression and purification of TvFDP.** To express the His-tagged protein (His-FDP) in *E. coli* M15 cells, the bacteria were induced with 0.5 mM IPTG (isopropyl-β-D-thiogalactopyranoside) and grown for 7 h at 25°C in LB medium supplemented with 400 μM ammonium ferrous sulfate and 200 μM flavin mononucleotide (FMN). The harvested cells were homogenized by passage through a French press at 18,000 lb/in². The soluble fraction obtained by ultracentrifugation (250,000 × g, 1 h, 4°C) was applied to a Ni-NTA column (Qiagen) and eluted with a stepwise gradient of 20 mM imidazole (buffer A: 20 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol [pH 7.6]) and 400 mM imidazole (buffer B: 400 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol [pH 7.6]) at a flow rate of 1 ml/min using a BioLogic HR system (Bio-Rad). His-FDP-containing fractions were dialyzed overnight against 50 mM Tris-HCl (pH 7.6) and subjected to Edman degradation (performed at the Protein/DNA Technology Center, Rockefeller University, New York, NY) to determine the amino-terminal sequences of the polypeptides. The heavier polypeptide was a homologue of the 51-kDa subunit (also called NuoX in bacteria) of the electron-input module of respiratory complex I (25), while the lighter polypeptide was found to be a homologue of bacterial FDPs and named TvFDP. Edman degradation also allowed determination of the processing site where the hydrogenosomal targeting peptide is cleaved from the preprotein by a proteinase processing peptidase.

**Characterization of TvFDP.** The flavin cofactor was characterized by reversed-phase high-pressure liquid chromatography using a nucleosil 100-5 C18 column and a thin-layer chromatography (TLC) method (HPTLC-Alufolin; Merck). The TLC mobile phase consisted of n-butanol-acetic acid–water (6:2:8). The flavin was extracted from the protein with trichloroacetic acid at a final concentration of 10%, followed by centrifugation and supernatant neutralization with 1 M ammonium acetate (pH 7.0). The identity of the flavin was determined for recombinant His-FDP, as well as for TvFDPStrep purified from *T. vaginalis* hydrogenosomes according to the IBA Strep-tag protein purification protocol. FMN and FAD (Sigma) were used as standards. The amount of flavin in TvFDP and FDP-containing fractions was determined after protein precipitation with 80% trichloroacetic acid, using an extinction coefficient of 12.500 M⁻¹ cm⁻¹ at λ = 450 nm (21).

The iron content of TvFDP (His-FDP was used for analysis) was determined by using the 2,4,6-tripyridyl-1,3,5-triazine method (18).

The native molecular mass of TvFDP was determined by gel filtration chromatography using a BioLogic HR system (Bio-Rad). The purified recombinant His-FDP, as well as the hydrogenosomal extract (prepared as described above) from FDP-overexpressing trichomonads (TvFDPWT), were run on a Superdex 75 (50 cm × 1.6 cm; GE Healthcare) column equilibrated with 400 mM imidazole–50 mM sodium phosphate buffer (pH 8.0) using a flow rate of 1 ml/min. The native molecular mass of the recombinant enzyme was calculated from the elution profile by Western blotting, followed by immunodetection of TvFDP using a specific rabbit polyclonal antiserum raised against His-FDP purified on an Ni-NTA column. The rabbit antiserum was prepared at the Institute of Parasitology, Academy of Sciences of the Czech Republic, České Budějovice, according to a published protocol (53).

Protein concentration was determined by the Lowry assay (34) using bovine serum albumin as a standard.

**Spectroscopic analysis of TvFDP.** The redox titrations were performed anaerobically at 25°C under an argon atmosphere by the stepwise addition of buffered sodium dithionite (250 mM Tris-HCl [pH 9.0]), in the presence of suitable redox mediators, in 50 mM Tris-HCl–18% glycerol (pH 7.5) as previously described (51). The samples were analyzed by visible spectroscopy (Shimadzu UV-1603 spectrophotometer) or by electron paramagnetic resonance (EPR) spectroscopy.
using an EMX Bruker spectrometer equipped with an Oxford Instruments ESR900 continuous flow cryostat. EPR spectra were recorded at 10 K, 9.39 GHz, 2.0 mW, and 1 mT of modulation amplitude. The electrodes (a silver/silver chloride combined electrode, or silver chloride and platinum electrodes) were calibrated against a saturated quinhydrone solution at pH 7.0, and the recorded potentials were normalized against that of the standard hydrogen electrode.

**Protein localization.** T. vaginalis cells overexpressing TvFDPHa were used for immunodetection of TvFDP. The cells were placed on glass slides coated with 3-aminopropyltriethoxysilane (Sigma), fixed with methanol (5 min), permeabilized with acetone (5 min) (both steps at −18°C), preincubated for 1 h in phosphate-buffered saline–0.25% bovine serum albumin–0.25% gelatin, and treated with antibodies as described previously (49). Anti-Ha tag monoclonal antibody (kindly provided by Patricia Johnson, University of California at Los Angeles) and hydrogenosomal malic enzyme polyclonal antiserum (15) were used as the primary antibodies. Anti-mouse immunoglobulin G (IgG) labeled with Alexa Fluor 488 (catalog no. A21202; Molecular Probes) and anti-rabbit IgG labeled with Alexa Fluor 549 (catalog no. A21207; Molecular Probes) were used as the secondary antibodies for fluorescent immunolocalization. Anti-TVFDPR rabbit polyclonal serum (see above) and anti-rabbit IgG antibody conjugated with alkaline phosphatase (ICN/Cappel) were used for Western blot analysis to visualize TVFDPR in T. vaginalis subcellular fractions obtained from untransformed, wild-type trichomonads and TVFDPW overexpressing trichomonads.

**Enzymatic reduction of TvFDP.** TVFDPR spectra were recorded on a Shimadzu UV-1601 spectrophotometer. The enzymatic reduction of TVFDPR (His-VDPR construct, 30 to 150 nmol) was monitored in stoppered cuvettes with a silicon septum in 860 µl of phosphate buffer (100 mM KH2PO4/KOH, 150 mM NaCl, 10% glycerol [pH 7.4]) using 44 mM pyruvate, 0.25 mM coenzyme A (CoA), 40 µM flavin adenine dinucleotide (FAD), 10 µM catalase (28 U), and glucose oxidase (28 U) to construct the calibration curve and as a positive control.

**Purification of PFOR.** In order to obtain a homologous enzymatic system capable of reducing TVFDPR, T. vaginalis PFOR was purified as follows: hydrogenosomes from 4 liters of culture were treated with 2% octylglucoside (MP Biomedicals) in 10 mM KH2PO4 (pH 6.8) on ice for 60 min and then centrifuged at 120,000 g for 30 min. The resulting membrane pellet containing PFOR was suspended in 50 mM ammonium ferrous sulfate, 100 mM potassium phosphate, 100 mM NaCl, 10% glycerol, and 1% deoxycholate (pH 7.4) for 60 min, followed by centrifugation at 120,000 g for 30 min. The supernatant fraction was mixed with 1 ml of Ni-NTA agarose (Qiagen), followed by incubation for 60 min at 4°C.

**RESULTS**

**Purification and properties of TvFDP.** Recombinant TvFDP with a His6 tag and without the hydrogenosomal targeting sequence (His-FDP) produced in M15 E. coli cells was purified by affinity chromatography on Ni-NTA agarose under native conditions. This procedure produced an almost homogeneous protein, as determined by SDS-PAGE (Fig. 1), with an approximate yield of 10 mg of His-FDP per liter of bacterial culture. The His-FDP polypeptide migrated on SDS-PAGE as a band with a molecular mass of ~45 kDa, a finding in agreement with the molecular mass calculated from the amino acid sequence without the hydrogenosomal targeting peptide.

To determine the native molecular mass of TvFDP, the hydrogenosomal extract from transformed T. vaginalis cells overexpressing native, nontagged TVFDPW, as well as the recombinant His-FDP isolated from E. coli, were analyzed by gel filtration chromatography. The native molecular mass of His-FDP was ~92 kDa. The TVFDPW in the elution profile was determined by SDS-PAGE and Western blot analysis with a specific antibody. TvFDPWT from the hydrogenosomal extract was eluted in the same elution volume as the purified recombinant protein (results not shown), suggesting that both the recombinant and the plasmid-encoded TVFDPW expressed in trichomonads exist as dimers in vivo.

The cofactor contained in the flavodoxin domain was identified by TLC and high-pressure liquid chromatography as noncovalently bound FMN (data not shown). This cofactor was established for both the recombinant protein and the protein containing the Strep tag (TVFDPW-Strep) isolated by affinity chromatography from T. vaginalis hydrogenosomes. Freshly isolated recombinant protein contained ~0.5 FMN and ~1.5 iron atoms per monomer. This stoichiometry indicates that a fully occupied protein contains one molecule of FMN and two

![FIG. 1. Spectroscopic analysis of recombinant T. vaginalis His-FDP. UV/Vis spectrum of pure His-FDP exhibits the typical features of flavoproteins. The inset shows the results of an SDS-PAGE analysis of purified recombinant His-FDP. Hom., bacterial lysate; FDP, purified His-FDP.](http://ec.asm.org/)
iron atoms per monomer; the incomplete occupancy is probably due to cofactor loss during purification. Pure, concentrated His-FDP was dark yellow and displayed a typical FDP UV/VIS spectrum (Fig. 1) dominated by the features of the flavin moiety, as the diiron center has a very low absorption.

**Sequence analysis.** TvFDP displayed homology with other described FDPs from both prokaryotic and eukaryotic sources. The residues implicated in binding the binuclear iron center (His94, His161, His240, Glu96, Asp98, and Asp180), as well as the conservative flavodoxin-like signature motif starting at position 271 (Val271) (41, 54), were all conserved in the T. vaginalis protein.

Soluble hydrogenosomal proteins are synthesized in the cytoplasm and typically contain a short amino acid presequence with a targeting function on their amino termini (3, 14). These targeting signals are cleaved by a specific hydrogenosomal processing peptidase (4) upon translocation of the protein into the organelle. TvFDP also possesses such a signal. The processing peptidase cleavage site was determined by amino acid sequencing via Edman degradation of mature TvFDP partially purified from hydrogenosomes; the cleavage site was found to be located between serine 11 and alanine 12 (Fig. 2). The conserved arginine residue typically located in the -2 position relative to the cleavage site in hydrogenosomal presequences was in an unusual -4 position in this case.

**Subcellular localization.** His-FDP was used to raise a specific antiserum that was subsequently utilized to determine the localization of native, nuclear-encoded TvFDP, as well as overexpressed TvFDPWT within trichomonad cells. Western blot analysis of T. vaginalis subcellular fractions showed that the TvFDP was expressed and specifically localized to the hydrogenosomal fraction (Fig. 3). To further verify the localization of TvFDP protein by immunofluorescence microscopy, T. vaginalis cells overexpressing TvFDPHa with an Ha tag at the C terminus were used. The protein was colocalized with the malic enzyme (hydrogenosomal marker enzyme) in the hydrogenosomal compartment (Fig. 4).

**Spectroscopic studies.** The visible spectrum of TvFDP was characteristic of a flavoprotein, with the main absorption bands at about 455 and 350 nm. Upon chemical reduction with sodium dithionite, there was no clear indication of the formation of a semiquinone radical, either of the blue or the red type, suggesting that the semiquinone form is not stabilized. Indeed, redox titration monitored by visible spectroscopy yielded a curve that could be fitted with two identical reduction potentials of approximately +25 mV for the FMN$_{red}$/FMN$_{ox}$ forms (Fig. 5). However, during the course of the enzymatic reduction by PFOR, a small band appeared at 398 nm during the early stages of the reduction (Fig. 6). Such a peak has not been observed in any of the other FDPs studied thus far, and its origin remains to be clarified.

Redox titration, followed by EPR spectroscopy, allowed us to both observe the signature of the diiron center and to determine its reduction potentials (Fig. 7 and Fig. 8). The diiron center had resonances at g = 1.955, 1.805, and 1.57, which is characteristic of the mixed valence [Fe(III)-Fe(II)]$^+$ state of a diiron site. By measuring the change in amplitude of the resonance at g = 1.8, the complete redox titration was constructed (Fig. 8) and analyzed with a Nernstian system of two consecutive one-electron transitions, yielding reduction potentials for the Fe(III)-Fe(III)/Fe(III)-Fe(II) and Fe(III)-Fe(II)/Fe(II)-Fe(II) transitions of 50 ± 20 and 190 ± 20 mV.

**Function.** In order to establish the physiological function of TvFDP, we studied the recombinant, His-tagged protein (His-FDP), which could be obtained in large quantities and displayed properties (i.e., FMN and iron cofactor, UV/VIS spectrum, and dimeric structure) similar to those of known FDPs. Studies of flavoproteins often utilize sodium dithionite as a reducing agent because it is effective at reducing iron to a lower oxidation state, allowing the iron to participate in the enzyme's catalytic cycle. The data obtained from these studies provide insights into the role of TvFDP in the hydrogenosomal compartment of T. vaginalis. Further investigations into the specific functions of this enzyme will be essential for understanding its biological importance.
reductant of the flavin moiety. This approach, however, is of little use when studying the physiological function of a protein whose presumed electron acceptor is oxygen. To reduce His-FDP enzymatically, we used a system consisting of pyruvate, CoA, PFOR purified from *T. vaginalis* hydrogenosomes, and purified recombinant *T. vaginalis* ferredoxin 1 (52). The primary electron source was pyruvate in a CoA-dependent PFOR reaction that oxidatively decarboxylates pyruvate, transfers electrons to ferredoxin, and releases acetyl-CoA. The reduced ferredoxin then served as a reductant for His-FDP in an anaerobic spectrophotometric assay. Catalytic amounts of PFOR and ferredoxin 1 fully reduced His-FDP in minutes. After complete reduction of His-FDP, the absorbance peak at 455 nm was totally bleached (Fig. 6). Upon introduction of air into the reaction mixture by opening the cuvette, His-FDP was immediately reoxidized (as documented by the regression of its spectral features to those of the native protein [Fig. 6]), indicating that an electron acceptor of TvFDP is indeed oxygen that is reduced with high affinity. When the cuvette was resealed again, the reduction of His-FDP resumed, indicating that the enzymatic system was not damaged by the short exposure to oxygen. To determine the product of the oxygen reduction, approximately 150 nmol of His-FDP was first fully enzymatically reduced as described above and then reoxidized with air. The mixture was subsequently analyzed by using the FOX method (55) for the presence of hydrogen peroxide. No hydrogen peroxide was detected, indicating that the product of oxygen reduction by TvFDP is water.

Since the ability of certain bacterial FDPs to reduce nitric
oxide has been well documented (21, 40, 45, 46), we tested the NO-reducing activity of His-FDP; NO was chemically generated by using DPTA-NONOate. The presence of NO in the reaction mixture was verified by a NO-specific electrode connected to an ISO-NO Mark II meter. Approximately 30 nmol of His-FDP was first enzymatically reduced and then combined with DPTA-NONOate to obtain 12 μM to 1 mM NO in an anaerobic assay mixture. No change in the spectrum of the reduced protein was observed with any concentration of NO tested over a period of 15 min (data not shown), indicating that TvFDP has negligible, if any, reactivity toward nitric oxide.

**DISCUSSION**

*T. vaginalis* hydrogenosomes harbor oxygen-sensitive proteins and are accordingly equipped with defensive enzymes to neutralize oxidative stress. These protective proteins include superoxide dismutase (17) and putative peroxidases such as thiol-dependent peroxidase and ruberythrin (39). However, the enzyme that is logically first in the line of defense, oxygen reductase, has remained elusive since its presence was first proposed in the 1970s (6). In the present study, we describe the properties of an FDP from *T. vaginalis* (TvFDP) that functions as a true oxygen reductase. This protein, first identified by proteomic methods (see Materials and Methods) during our search for other hydrogenosomal activities, localized to hydrogenosomes as demonstrated by immunoblotting with specific antibodies. The subcellular localization of TvFDP was further confirmed by immunofluorescence microscopy of transformed trichomonads expressing a tagged version of the enzyme. The organellar localization is consistent with the presence of a cleavable hydrogenosomal targeting peptide with a characteristic amino acid composition on the amino terminus of the conceptually translated protein (Fig. 2).

Analysis of cofactor content suggests that TvFDP binds two iron atoms and one FMN per monomer as in other homologues. Like most of its counterparts that have already been studied in this respect, native TvFDP is an ~92-kDa homodimer of identical 45-kDa subunits. This dimeric quaternary structure is essential for efficient electron transfer from the flavin of one monomer to the diiron center of the other monomer; these regions come into close contact due to the head-to-tail organization of each monomer, a feature common to all FDPs whose structures have been determined thus far. Within a single monomer, the redox centers are too far apart to allow electron transfer at a physiologically relevant rate (50). The reduction potentials determined here for the FMN and the diiron center are higher than those observed for the *E. coli* enzyme, the only FDP extensively characterized in this respect thus far (50, 51). Another remarkable characteristic was observed in the reduction potentials of the flavin transitions, which were almost identical; this has not been reported for any other FDP thus far. This precludes the formation of a stable flavin semiquinone form as observed in other FDPs and favors an almost two-electron oxidation of the flavin. Nevertheless, the measured reduction potentials are suitable for either NO or O2 reduction, and further studies in other FDPs are needed to determine whether these differences are enzymatically relevant. Due to the high reduction potentials of either substrate, the specificity toward these small molecules cannot be related to their redox parameters. The UV-visible and EPR spectroscopic data are similar to those of other FDPs characterized thus far. The data for the diiron center showed that the iron ions were antiferromagnetically coupled in the half-reduced state, yielding a total spin of S = 1/2. For the oxidized and reduced forms, it was not possible to obtain any EPR spectral signature; in the oxidized state the total spin state is most probably S = 0, while in the reduced form it could be either zero or S = 4 as observed for the *E. coli* enzyme.

In order to determine the activity and substrate specificity of TvFDP, we needed to reconstruct in vitro the electron transfer chain from an electron donor to the terminal acceptor, oxygen and/or nitric oxide. This was a nontrivial task, since *T. vaginalis* does not possess rubredoxin, the physiological reductant of some FDPs, and analysis of the *T. vaginalis* genome did not provide any clue about potential redox partners for TvFDP. Nevertheless, *T. vaginalis* encodes a large number of [2Fe-2S] ferredoxins, small electron carrier proteins linked to PFOR, hydrogenase, and a remnant of complex I (28). In addition, the genome also encodes several predicted hydrogenosomal iron-sulfur flavoproteins with unknown function but presumably involved in electron transfer (5). We have found that the enzymatic system consisting of PFOR purified from *T. vaginalis* hydrogenosomes and a homogeneous preparation of recombinant *T. vaginalis* ferredoxin 1 could effectively reduce TvFDP at the expense of electrons derived from pyruvate by the activity of PFOR. Strictly anaerobic conditions had to be used to follow the reduction of the flavin cofactor of TvFDP and also to prevent autooxidation of ferredoxin in the spectrophotometric assay. Alternatively, another redox system could be used to reduce TvFDP; this system consists of a purified hydrogenosomal remnant of complex I and recombinant *T. vaginalis* ferredoxin 1. In this case, the electron donor is NADH, but since its absorbance partially interfered with the absorbance of oxidized FMN of TvFDP in a spectrophotometric assay, the PFOR-based system was preferentially used. Under these experimental conditions, full reduction of TvFDP, monitored as disappearance of the absorbance of FMN, was achieved within
a few minutes. The reaction time depended on the concentration of PFOR and ferredoxin in the assay (Fig. 6). Upon introduction of air into the cuvette, TvFDP immediately reoxidized as documented by the regression of the spectral curve to the original, oxidized pattern (Fig. 6). The immediate reoxidation of the flavin cofactor indicates a fast reaction with a high affinity for oxygen; however, the kinetic parameters of the electron transfer could not be determined due to the autooxidative nature of the trichomonad ferredoxin, which precluded a continuous aerobic spectrophotometric assay.

To function as a protective, O₂ scavenging enzyme, TvFDP should reduce oxygen by four electrons to water. The product of oxygen reduction was determined by the FOX assay with enzymatically reduced His-TvFDP that was reoxidized by short exposure to air. No traces of hydrogen peroxide were detected, indicating that TvFDP indeed reduces dioxygen to water using four molecules of the one-electron carrier ferredoxin per cycle. It should be noted that FDPs do not reduce hydrogen peroxide (M. Teixeira, unpublished data). Experiments designed to verify the potential reactivity of enzymatically reduced TvFDP with nitric oxide, the known substrate of several members of the flavodiiron superfamily, did not show any such activity for the trichomonad protein, despite the broad range of NO concentrations and prolonged reaction times tested. This is similar to what was found for the only other eukaryotic FDP characterized, the protein from *G. intestinalis*, which is also virtually unreactive with NO (13).

Analysis of the primary structure of TvFDP shows that the enzyme belongs to the class A family of FDPs. Members of this family are the simplest representatives of the superfamily, consisting of only the core flavodiiron module, which is formed by a metallo-β-lactamase-like domain in the amino-terminal half of the protein and by a flavodoxin-like domain forming the rest of the polypeptide chain (41). The mature (without the hydrogenosomal targeting peptide) trichomonad protein is colinear over its entire length with its homologues, with conservation of all residues implicated in binding the binuclear iron center and a well-conserved flavodoxin signature motif at the start of the flavodoxin-like domain (Fig. 2).

Attempts have been made to attribute the selective specificity of some FDPs for oxygen (and not NO) to particular amino acid residues (44). This issue has been discussed in the light of data recently obtained for *Giardia* FDP (13), and we can only conclude that the results concerning both the *Giardia* (13) and the *Trichomonas* FDPs (the present study) invalidate the original hypothesis (44) and leave the question of the substrate specificity of FDPs unsettled. In addition, tryptophan 361 (W347 in reference 44), which is missing in the FDP from *M. marburgensis* but conserved in rubredoxin-specific FDPs, has been proposed to play a role in electron shuttling between rubredoxin and the FMN of rubredoxin-specific FDPs (44). However, neither *Trichomonas* nor *Giardia* FDPs, which have W361 conserved, have rubredoxins as electron donors, since there are no genes encoding these proteins (http://www.tichdb.org/trichdb/; http://www.giardidb.org/giardidb/). The absence of this tryptophan in *Methanothermobacter* FDP might be related to the nature of the proposed electron donor to FDPs from methanogens, enabling easy access to $E_{320} \text{H}_2$.

FDPs have already been subjected to a phylogenetic study, as particularly good candidates for genes that have been acquired by certain unicellular eukaryotes via lateral gene transfer from prokaryotic sources (1). Eukaryotic FDP homologues have thus far been invariably found in the genomes of anaerobic fermentative protists, most with parasitic lifestyles. This suggests that the role of FDPs in these eukaryotes is similar to that in prokaryotes, i.e., protection against oxidative/nitrosative stress. However, the only eukaryotic homologue studied to date is the FDP from *G. intestinalis* (13). *Giardia* (a member of Diplomonadida) is believed to share a common ancestor with trichomonads (23), and the *T. vaginalis* protein described in the present study is the closest relative to the giardial homologue (which, however, is natively a tetramer) (1), suggesting that this particular gene entered this eukaryotic lineage before the divergence of diplomonads and trichomonads. Notably, the sister group to diplomonad/trichomonad FDPs consists of the branch of archaeabacterial FDPs of methanogens (1), some of which have been shown to utilize only oxygen and not nitric oxide (44), like the homologues from both human parasites.

Class A FDP is not the only FDP present in *T. vaginalis*. In fact, four genes encoding FDP homologues could be identified in the genome of this parasite (5) (http://www.tigr.org/tdb/e2k1/tvg; locus numbers TVAG_036010, TVAG_129610, TVAG_049830, and TVAG_263800). One of these proteins is the subject of the present study; the remaining three are highly similar homologues that belong to a strongly separated clade and form a group with other eukaryotic sequences from *Entamoeba* and *Mastigamoeba* and with eubacterial sequences from *Clostridium* sp. (1). Unlike hydrogenosomal TvFDP, the genes of these three trichomonad homologues do not encode a clear amino-terminal extension with an organellar targeting function (and were not detected in the proteome of hydrogenosomes, unpublished data); thus, these putative proteins are likely localized to the cytoplasm. Because of their novel primary structure, these trichomonad homologues were proposed to constitute a fourth, D class of FDPs, together with clostridial proteins, that have an NADH:rubredoxin oxidoreductase and rubredoxin module fused to the flavodiiron core and could probably directly utilize NAD(P)H as an electron donor (50). In this respect, it should be mentioned that nitric oxide reducing activity has been detected in intact as well as lysed *T. vaginalis* cells; this activity has been ascribed to FDP partly because a cross-reactive band was visualized in a *T. vaginalis* cell lysate using a heterologous antibody against *E. coli* flavoredoxin (42). Since the molecular size of the denatured candidate protein (~60 kDa) was far from the predicted size of a class D FDP monomer (~95 kDa), its identity is doubtful. Nevertheless, if these class D FDP homologues are indeed expressed, the observed nitric oxide reducing activity (42) might be due to their presence in the *T. vaginalis* cytosol, where they could provide protection against the host immune response as well as against oxygen.

Finally, it should be stated that despite the evidence collected in vitro using a functional electron transport chain consisting of the *T. vaginalis* hydrogenosomal proteins PFOR, ferredoxin 1, and TvFDP (or complex I, ferredoxin 1, and TvFDP), the identity of the actual physiological components of the presumed pathway remains uncertain. This is largely due to the complexity of the predicted electron transporting pathways of hydrogenosomes. The core electron-generating reactions utilizing malate and pyruvate as the carbon and electron source, as coined by Müller (36), are still valid, but research...
over the last 15 years, as well as the recently completed annotation of the T. vaginalis genome (5), has shown that most already identified hydrogenosomal proteins involved in electron transfer and transport are coded by multiple and clearly distinct genes. These include seven genes encoding PFOR homologues, up to nine malic enzymes, seven ferredoxins, seven iron-sulfur flavoproteins, and up to nine putative hydrogenases, of which four are equipped with a predicted hydrogenosomal targeting peptide; many of these proteins are transcribed, also as indicated by complex EPR spectroscopic data (7, 16, 26–28; unpublished proteomic data). It is possible that certain PFOR homologues preferentially use particular ferredoxins, while others could function as electron acceptors of complex I or in iron-sulfur cluster assembly machinery (49).

The involvement of iron-sulfur proteins even in these known reactions also cannot be excluded, as indicated by preliminary experiments (unpublished data). Other possible sources of electrons besides pyruvate and NADH (generated by the malic enzyme), such as amino acids or α-glycerophosphate (6), could also play a role. Some combinations of substrate, electron-generating enzyme, and electron-transporting protein would likely be more efficient than others in shuttling electrons toward TvdFDP.

In summary, by fusing a hydrogenosomal targeting peptide to a class A FDP, most likely obtained from an anaerobic bacterium by lateral gene transfer, T. vaginalis equipped the hydrogenosomes with a new enzyme. Functional data suggest that this acquisition helped the parasite to cope with oxidative stress in these oxygen-sensitive organelles.

ACKNOWLEDGMENTS

This study was supported by the Czech Science Foundation grant 204/06/0944 to I.H. and by Ministry of Education, Youth, and Sports of the Czech Republic grant MSM021620588. Further support was provided by the Fundação para a Ciência e Tecnologia–Portugal, projects PTDC-BIA-PRO/67263/2006, and Structure, Dynamics and Functions of Proteins (REEQ/336/BIO/2005). V.L.G. is the recipient of a Ph.D. grant from Fundação para a Ciência e Tecnologia–Portugal (SFRH/BD/29428/2006).

We acknowledge the help of J. Vicente (Instituto de Tecnologia Química e Biológica) with the EPR-monitored redox titration. We thank Miša Marciničková (Charles University) for excellent technical assistance. Petr Jedelsky (Charles University) for mass spectrometry analyses, and Jan Mach (Charles University) for help with the figures.

REFERENCES


