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Eukaryotic UDP-Galactopyranose Mutase (*GLF* Gene) in Microbial and Metazoal Pathogens†

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Galactofuranose (Gal_f) is a novel sugar absent in mammals but present in a variety of pathogenic microbes, often within glycoconjugates that play critical roles in cell surface formation and the infectious cycle. In prokaryotes, Gal_f is synthesized as the nucleotide sugar UDP-Gal_f by UDP-galactopyranose mutase (UGM) (gene *GLF*). Here we used a combinatorial bioinformatics screen to identify a family of candidate eukaryotic *GLFs* that had previously escaped detection. *GLFs* from three pathogens, two protozoa (*Leishmania major* and *Trypanosoma cruzi*) and one fungus (*Cryptococcus neoformans*), had UGM activity when expressed in *Escherichia coli* and assayed in vivo and/or in vitro. Eukaryotic *GLFs* are closely related to each other but distantly related to prokaryotic *GLFs*, showing limited conservation of core residues around the substrate-binding site and flavin adenine dinucleotide binding domain. Several eukaryotes not previously investigated for Gal_f synthesis also showed strong *GLF* homologs with conservation of key residues. These included other fungi, the alga *Chlamydomonas* and the algal phleovirus *Feldmannia irregularis*, parasitic nematodes (*Brugia*, *Onchocerca*, and *Strongyloides*) and *Caenorhabditis elegans*, and the urochordates *Halocynthia* and *Cionia*. The *C. elegans* open reading frame was shown to encode UGM activity. The *GLF* phylogenetic distribution suggests that Gal_f synthesis may occur more broadly in eukaryotes than previously supposed. Overall, *GLF*/Gal_f synthesis in eukaryotes appears to occur with a disjunct distribution and often in pathogenic species, similar to what is seen in prokaryotes. Thus, UGM inhibition may provide an attractive drug target in those eukaryotes where Gal_f plays critical roles in cellular viability and virulence.

In many microbes, the sugar galactofuranose (Gal_f) is an important constituent of glycoconjugates comprising major portions of the cell surface (17). In prokaryotes, Gal_f constitutes a key part of the mycobacterial cell wall and occurs in lipopolysaccharide (LPS) O-antigen domains, extracellular capsules, and polysaccharides (17, 19). In fungi such as *Aspergillus*, Gal_f is a major component within the cell wall and structural glycoproteins (13, 17). In pathogenic protozoa, Gal_f residues are key components of abundant surface glycosylphosphoinositol-anchored glycoconjugates, such as lipophosphoglycan (LPG) and glycoinositolphospholipids in *Leishmania*, and of mucins, glycosylphosphoinositol-anchored proteins, and lipids in *Trypanosoma cruzi* (6, 18). In contrast, Gal_f residues have not been found in humans or other metazoans (17), suggesting that inhibition of Gal_f synthesis could be an attractive target for chemotherapy in pathogens when its role(s) is critical for survival or virulence (17, 32).

The Gal_f synthetic pathway has been best studied for prokaryotes. Genetic and biochemical studies have shown that Gal_f arises through the action of UDP-galactopyranose mutase (UGM) (EC 5.4.99.9), which catalyzes the rearrangement of UDP-galactopyranose (Gal_p) to UDP-Gal_f, the substrate of cellular UDP-Gal_f transferases which participate in path-

ways such as LPS or cell wall biosynthesis (15, 16). The gene encoding UGM, *GLF*, was first located and hypothesized as such by Reeves and colleagues as part of genetic and structural studies of *Escherichia coli* K12 O antigen (27). It was then definitively identified and studied in *E. coli*, *Klebsiella*, and *Mycobacteria* (10, 15, 16). UGM is a flavin-dependent enzyme, and the *E. coli* enzyme structure has been solved (21). A detailed picture of the enzymatic mechanism involving a novel form of flavin-dependent catalysis has been developed (24). High-throughput assays for inhibitor screens have been developed, and a number of compounds showing activity against UGM activity and/or bacteria have been identified (22, 25, 28).

Less is known about the Gal_f synthetic pathway in eukaryotes. Previous efforts had not yielded the eukaryotic enzyme responsible for synthesis of their UDP-Gal_f substrate, although as in prokaryotes this was thought to arise by conversion of UDP-Gal_p to UDP-Gal_f (29). In the parasitic protozoan *Leishmania*, several genes encoding putative UDP-Gal_f transferases have been identified, including *LPG1*, which has been implicated in the synthesis of the core of the abundant surface glycoconjugate LPG (7, 20). Notably, the *Leishmania* genome encodes at least 6 candidate UDP-Gal_f transferases (34), and there are more than 20 related genes present in *Trypanosoma cruzi* (34) (unpublished data). No candidate UDP-Gal_f transferases have been reported in fungi, although they must exist given the number of Gal_f-containing glycoconjugates known in these organisms. In this report we have used a bioinformatics approach to identify the eukaryotic *GLF* gene family. We ex-

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† Supplemental material for this article may be found at <http://ec.asm.org/>.

pressed four diverse members of this family in *E. coli* and demonstrated their activity in vivo and/or in vitro.

MATERIALS AND METHODS

PCR and construction of pET3 derivatives. For *Leishmania major*, PCR oligonucleotides SMB2179 (5'-GCATGCCATATGAGCGCTGACAAGGTGGT CATAATC) and SMB2180 (5'-GCATGCGGATCTACGAGGCCGTCGAC GACCATGTGCA) (underlined bases represent sites added for subsequent cloning) were based upon the open reading frame (ORF) Lm18.0200 annotated in release 4.0 of the *Leishmania major* genome (www.genedb.org/genedb/leish). The PCR template was genomic DNA of *L. major* Friedlin (MHOM/IL/81/Friedlin) clone V1, and amplification was performed with 35 cycles of 30 s at 93°, 45 s at 50°, and 150 s at 68°C. For *Trypanosoma cruzi*, PCR oligonucleotides SMB2182 (5'-GCATGGCCATATGGCAGAATTATTGACACCGAAA ATTG) and SMB2230 (5'-GCGATAGGATCTCACATATCCTTCTGCAGT AGT) were based upon the second of two ORFs (Tc00.1047053511277.600 and Tc00.1047053507993.160) annotated in release 3.0 of the *Trypanosoma cruzi* genome project (www.genedb.org/genedb/tcruzi). The PCR template was DNA from the CL Brener strain of *T. cruzi* used for genome sequencing (kindly provided by D. Barthomoleu, The Institute for Genomic Research), and amplification was performed with 35 cycles of 30 s at 93°C, 45 s at 49°C, and 150 s at 68°C. For *Cryptococcus neoformans*, PCR primers Galf-S (5'-GGAATTCATA TGCCGTCAGACTTGATTTT) and Galf-AS (5'-CGGGATCCCTACTTCA AGCGCCTCTCGA) were based upon ORF177.m02862 (The Institute for Genome Research; www.tigr.org/tdb/e2k1/cna1/index.shtml). Total RNA was isolated from *C. neoformans* serotype D strain JEC43 α and converted to cDNA using the SuperScript First-Strand synthesis system as recommended by the manufacturer (Invitrogen, Carlsbad, CA). PCR amplification using HIFI *Taq* polymerase (Invitrogen, Carlsbad, CA) was performed with 30 cycles of 60 s at 94°C, 60 s at 55°C, and 120 s at 72°C. For *C. elegans*, gene predictions for the *GLF* ORF H04M03.4 were obtained from WormBase (<http://www.wormbase.org>). One internal region had not been determined experimentally, and we found by sequence analysis of several cDNAs (expressed sequence tags [ESTs] yk1442e06, yk1480e04, and yk1626a12 from Yuji Kohara, National Institute of Genetics, Mishima, Japan; GenBank accession no. DN856302-6; and ORF clone 10013@E10, obtained from Open Biosystems, Inc., Huntsville, AL) that the current gene model had missed a short intron, resulting in a predicted 9-amino-acid insertion (SVYCFLEEV) not encoded by any of the three cDNAs sequenced. The *C. elegans* H04M03.4 ORF contained an internal NdeI site, requiring a series of PCR and cloning steps starting with the 10013@E10 cDNA template prior to insertion into the pET3A NdeI-BamHI sites.

All PCRs yielded products with the expected sizes, which as necessary were digested with NdeI and BamHI, ligated to pET-3A vector DNA previously digested with the same enzymes, and transformed into *E. coli* DH10B or DH5 α , yielding pET3a-LmGLF, pET3a-CnGLF, pET3a-TcGLF, or pET3a-CeGLF (lab strains B5234, B5300, B5330, and B5425, respectively). The authenticity of candidate recombinants was confirmed by DNA sequencing (GenBank accession numbers AY900624, AY900625, AY900626, and BK005688, respectively). For *T. cruzi* *GLF*, our sequence corresponded well with the provisional Tc00.1047053511277.600 ORF, with three differences (GTG \rightarrow GCA/Val \rightarrow Ala in the second codon and a silent A \rightarrow G transition at nucleotide 1413). The first two differences may represent "cross priming" of the 5' oligonucleotide used in PCR.

For rescue of Gal_f-dependent LPS biosynthesis, the pET-GLF constructs above were introduced into *E. coli* strain CWG288+pWQ70 (10), yielding strains B5364, B5363, B5365, and B5431, respectively. As described in more detail below, CWG288 contains a deletion of the *E. coli* *rfb* locus, while plasmid pWQ70 contains the *rfb* locus of *Klebsiella pneumoniae* bearing a deletion of the *GLF* gene (see reference 10 for the complete genotype and characterization of these). For enzymatic assays, two of the pET-GLF constructs were introduced into the *E. coli* host strain BL21(DE3) (B5304), yielding pET-LmGLF/BL21 (DE3) and pET-CnGLF/BL21(DE3) (B5235 and B5303, respectively). Comparative phylogenetic analyses were conducted using MEGA version 3.0 (11).

LPS Western blotting. For immunoblotting, cells were pelleted from 1 ml of an overnight *E. coli* culture, suspended in 50 μ l of sample buffer (62.5 mM Tris, 2% sodium dodecyl sulfate, 10% glycerol, 2.5% β -mercaptoethanol, and 0.05% bromophenol blue), and boiled for 10 min. Samples were diluted fivefold further with sample buffer, and 5 μ l was loaded on a discontinuous gel consisting of a 4% stacking gel above a 12.5% Tris-glycine separating gel (12). Fresh preparations gave the best results. Gels were run at 200 V for 40 min and electroblotted to nitrocellulose (Hybond-ECL; Amersham Biosciences) overnight at 54 mA. Western blotting was performed with rabbit anti-*Klebsiella* galactan I (generously

provided by Whitfield and Clarke, U. Guelph) at a titer of 1:10,000 for 2 h at room temperature in 5% milk in TBST (10 mM Tris, 150 mM NaCl, 0.05% Tween 20 [vol/vol]). Binding was visualized with peroxidase-conjugated goat anti-rabbit immunoglobulin G (Jackson ImmunoResearch, catalog no. 111-035-003) at a titer of 1:20,000 in TBST for 1 h at room temperature followed by detection by chemiluminescence (Western Lightning, Perkin-Elmer Life Sciences, catalog no. NEL100).

UGM enzyme assays. One-liter cultures of *E. coli* BL21(DE3)/pET3a-CnGLF expressing *Cryptococcus GLF* (B5303), *E. coli* BL21(DE3)/pET3a-LmjGLF expressing *Leishmania GLF* (B5235), or *E. coli* BL21(DE3) without plasmid were grown overnight at 37°C in LB broth. Strains bearing pET3a-GLF constructs were grown in the presence of 100 μ g/ml ampicillin. All cultures were induced by addition of 0.5 mM isopropyl- β -D-thiogalactopyranoside, followed by incubation at 22°C with shaking at 90 rpm for 3 h. Cultures were harvested by centrifugation, and cell pellets were resuspended in 20 ml 100 mM potassium phosphate buffer (pH 7.0) containing 0.15 M NaCl, 15% glycerol, 10 mM flavin adenine dinucleotide (FAD), 0.1 mM phenylmethylsulfonyl fluoride, 1 μ M pepstatin A, and 1 μ M leupeptin. The cells were broken by three passages through a French press and centrifuged at 20,000 \times g for 40 min at 4°C. The supernatant fractions were frozen at -80°C until use.

UDP-Gal_f was synthesized enzymatically from UDP-Gal_p and purified as described previously (14). Activity assays were performed in 96-well PCR plates. Each well contained 50 μ M UDP-Gal_f, 100 mM phosphate buffer at pH 7, 1 mM MgCl₂, 15% glycerol, protein (80 to 10,000 ng *E. coli* extract expressing *Cryptococcus GLF*, 35 to 4,400 ng *E. coli* extract expressing *Leishmania GLF*, or 27,400 ng control *E. coli* BL21(DE3)), and 20 mM sodium dithionite (to reduce the FAD to FADH₂) in a 20- μ l volume. FAD was added in various amounts depending on how much protein was added (final concentrations of 1.6 to 1,000 μ M). Reactions were incubated at 30°C for 6 min and stopped by the addition of 40 μ l 95% ethanol. Samples were analyzed by high-performance liquid chromatography (HPLC) as described previously (14), except that the solvent system was modified by elution of the sugar nucleotides for 10 min with 200 mM KH₂PO₄ followed by a 600 mM KH₂PO₄ wash. Yields of UDP-Gal_f were calculated from the relative fraction of UDP-Gal_f and UDP-Gal_p in the samples and the knowledge that 1 nmol of UDP-Gal_f was originally added to each well.

RESULTS

Identification of candidate GLFs. Although the first prokaryotic *GLF* was identified in 1996, previous efforts to detect eukaryotic *GLFs* had not been successful. Recently the genomes of several eukaryotic pathogens known to synthesize Gal_f progressed towards completion, and we applied a combinatorial bioinformatics screen to these genomes in an effort to identify candidate *GLFs*. Briefly, these involved a combination of (i) BLAST searches with prokaryotic *GLFs*, (ii) searches for flavin binding domains, and (iii) appropriate phylogenetic distribution, specifically occurrence in fungi and protozoa known to synthesize Gal_f but absence from taxa lacking Gal_f. Since the presence of Gal_f has not been systematically addressed for many microbial and metazoan species, we applied the latter criterion conservatively. We focused these studies initially on the *Leishmania major* genome, using ORF predictions developed by the *Leishmania* genome project (release 4.0, August 2004; see www.genedb.org/genedb/leish).

While many ORFs satisfied each of these criteria individually (albeit often quite weakly), a single candidate *L. major* ORF was identified in the combined screen. The Lm18.0200 *GLF* sequence predicted a 491-amino-acid protein, which showed a weak relationship to the Pfam family of flavin-containing amine oxidoreductases (PB010804; $e = 6 \times 10^{-4}$), including a potential FAD binding motif at amino acids 6 to 19 (Fig. 1, underlined by parallel lines). The three-dimensional structure of *E. coli* UGM has shown that the FAD binding domain actually encompasses a much larger portion of the protein (21), and limited sequence conservation homology was

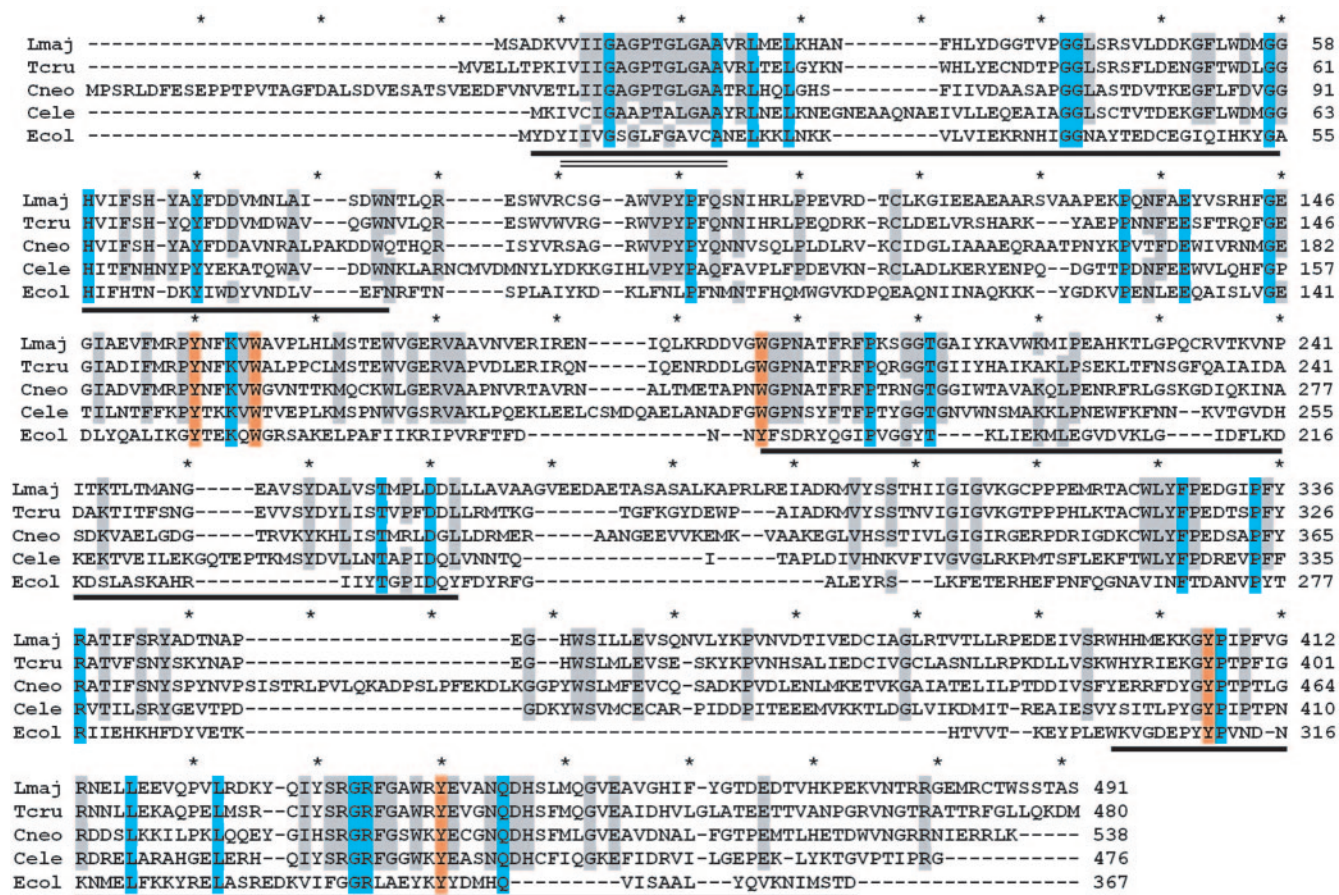


FIG. 1. Alignment of key eukaryotic and prokaryotic *GLF* proteins. The predicted proteins encoded by the *L. major*, *T. cruzi*, *C. neoformans*, and *C. elegans* *GLF* ORFs were aligned with that of *E. coli* using methods implemented in the Clustal program (4); the unedited alignment is shown. The amino acid positions are shown on the right, and asterisks mark 10-amino-acid intervals in the alignment. Residues identical in 5/5 sequences are shaded in blue, and residues identical in 4/5 sequences are shaded gray. Residues proposed as part of UDP-Gal_p binding site are shaded orange (all but one of these are 5/5 identities). The dark underlined regions correspond to the flavin-binding region of *E. coli* UGM (21); a smaller region showing homology to the Pfam family flavin-containing amine oxidoreductases (PB010804; residues 6 to 19 of the *L. major* UGM) is underlined by two parallel lines.

also evident throughout this region (Fig. 1, underlined by dark lines). BLAST searches of a variety of databases, including unfinished microbial genomes, revealed a number of strong BLAST hits ($P < 10^{-40}$), most of which were annotated as hypothetical proteins although several were annotated as amine oxidases (Table 1; Fig. 2). Notably, in every case where sequence from organisms capable of Gal_f synthesis was analyzed, strong BLAST hits to Lm18.0200 were present. These included the protozoans *Leishmania infantum* and *T. cruzi* and the fungi *Aspergillus*, *Cryptococcus*, and *Neurospora* (Fig. 2; Table 1). Correspondingly, homologs of the candidate *L. major* *GLF* were not detected in vertebrates, nor in *Trypanosoma brucei*, *Saccharomyces cerevisiae* or *Schizosaccharomyces pombe*, species where Gal_f may be absent. Prokaryotic *GLF*/UGMs yielded weak BLAST scores (>0.2) across only the N-terminal portion of the *Leishmania* UGM, which undoubtedly contributed to the eukaryotic genes being overlooked previously.

Comparison of the predicted proteins encoded by the *L. major*, *T. cruzi*, and *Cryptococcus neoformans* *GLFs* to that of *E. coli* UGM/*GLF* (Fig. 1; Table 1) shows that the overall similarity to prokaryote *GLFs* is relatively low. Most regions of

sequence conservation fall within the FAD binding domains determined previously from the structure of *E. coli* UGM (Fig. 1, underlined regions) (21). Significantly, genetic and structural analyses have identified a number of residues possibly involved in UDP-Gal_p binding, and with few exceptions these residues were identical in prokaryotic and eukaryotic *GLF* proteins (Fig. 1, residues shaded in orange). These residues were also conserved within the strong database hits noted elsewhere in this section (data not shown). The candidate eukaryotic *GLF* proteins showed a high degree of identity to each other (Fig. 1 and 2; Table 1).

While the correlation of *GLF* candidates with species known to carry out Gal_f synthesis was encouraging, strong BLAST hits were also obtained to sequences from a number of organisms not known to synthesize Gal_p, including the nematodes *C. elegans* and *Caenorhabditis briggsae*, the urochordate *Halocynthia*, and the algal *Feldmannia irregularis* virus (Table 1). A number of ESTs encoding partial *GLFs* were found in parasitic nematodes, including *Brugia malayi*, *Onchocerca volvulus*, *Strongyloides stercoralis* and *Strongyloides ratti*, *Heterodura glycine*, and *Meloidogyne hapla* and *Meloidogyne arenaria* (see Table S1 in

TABLE 1. Genes showing strong similarity to eukaryotic *GLFs*^a

Species	Group	Species contains Gal? ^b	Gene	Annotation ^c	BlastP <i>P</i> value	% Identity
<i>Leishmania major</i>	Protists	YES	Lmj18.200	UGM	(query)	100
<i>Trypanosoma cruzi</i>	Protists	YES	Tc00.1047053511277.600 Tc00.1047053507993.160	UGM	10 ⁻¹⁶⁰	58
<i>Aspergillus nidulans</i>	Fungi	YES	EAA63683	Hyp.	10 ⁻¹¹⁷	45
<i>Neurospora crassa</i>	Fungi	YES	EAA27372	Hyp.	10 ⁻¹¹⁷	44
<i>Magnaporthe grisea</i>	Fungi	UNK	EAA55038	Hyp.	10 ⁻¹¹⁵	44
<i>Gibberella zeae</i>	Fungi	UNK	EAA75642	Hyp.	10 ⁻¹¹³	44
<i>Cryptococcus neoformans</i>	Fungi	YES	EAL19520	UGM	10 ⁻¹¹¹	42
<i>Chlamydomonas reinhardtii</i>	Green algae	UNK	Table S2 ^d		10 ⁻¹¹⁰	42
<i>Ustilago maydis</i>	Fungi	UNK	UM03094	Hyp.	10 ⁻¹⁰⁴	40
<i>Geobacter sulfurreducens</i>	Eubacteria	UNK	AAR34886	Flavin amine oxidase	10 ⁻¹⁰³	44
<i>Desulfovibrio vulgaris</i>	Eubacteria	UNK	AAS94778	Flavin amine oxidase	10 ⁻⁹¹	40
<i>Desulfovibrio desulfuricans</i>	Eubacteria	UNK	ZP00346806	Protoporphyrinogen oxidase	10 ⁻⁸⁵	37
<i>Feldmannia irregularis</i> virus	Algal virus	UNK	AAR26880	Hyp.	10 ⁻⁸⁴	38
<i>Caenorhabditis briggsae</i>	Nematodes	UNK	CAE72630	Hyp.	10 ⁻⁶⁸	34
<i>Caenorhabditis elegans</i>	Nematodes	UNK	AAD12787	UGM	10 ⁻⁶⁸	34
<i>Ciona intestinalis</i>	Nematodes	UNK	Table S2		10 ⁻⁶²	31
<i>Halocynthia roretzi</i>	Urochordates	UNK	BAB20903	HrTLC1	10 ⁻⁴⁶	26
<i>Pyrococcus horikoshii</i>	Archea	UNK	B71153	Hyp.	10 ⁻²⁷	27
<i>Pyrobaculum aerophilum</i>	Archea	UNK	AAL62855	Hyp.	10 ⁻²⁶	27
<i>Yersinia pseudotuberculosis</i>	Eubacteria	PROB	CAB63294	O-antigen <i>WhyH</i>	10 ⁻¹⁸	22
<i>Methanococcus maripaludis</i>	Archea	UNK	CAF30324	Hyp.	10 ⁻¹⁴	21
<i>Klebsiella pneumoniae</i> (prokaryotic GLF)	Eubacteria	YES	Q48485	UGM	0.2	<10% (only N-term) ^e

^a The genes are ordered based on their BlastP *P* value, based on searches with the *L. major* *GLF* query sequence.

^b YES, evidence showing occurrence of Gal_f in one or more glycoconjugates; PROB, Gal_f suspected to be present; UNK, not investigated or established.

^c Hyp., hypothetical protein. Annotations of *L. major*, *T. cruzi*, *C. neoformans*, and *C. elegans* *GLFs* reflect GenBank depositions arising from this work.

^d See the supplemental material.

^e N-term, N-terminal.

the supplemental material), and we were able to assemble complete ORFs for the urochordate *Ciona intestinalis* and the alga *Chlamydomonas reinhardtii* (Table 1; see Tables S1 and S2 in the supplemental material). Several strong hits ($P < 10^{-85}$) were obtained to eubacterial sequences annotated as “amine oxidases,” although enzymatic data supporting these assignments were lacking. Strong hits ($P < 10^{-28}$ to 10^{-15}) were also obtained with sequences from three archaea (*Pyrococcus horikoshii*, *Pyrobaculum aerophilum*, and *Methanococcus maripaludis*) (Table 1). A phylogenetic tree depicting relationships among the eukaryotic candidates is shown in Fig. 2, and an alignment of the *C. elegans* ORF H04M03.4 is included in Fig. 1. As observed for the fungal and protozoal homologs, candidate flavin-binding regions and UDP-Gal_f binding residues were conserved in these predicted proteins (Fig. 2; also data not shown). Genes or proteins showing significant relationship were not observed in BLAST searches of the genomes of *Giardia lamblia*, *Trichomonas vaginalis*, *Entamoeba*, apicomplexans including *Toxoplasma* and *Plasmodium*, *Tetrahymena*, or several plant genomes including *Arabidopsis*.

The imperfect concordance between the occurrence of eukaryotic *GLF* sequences and Gal_f synthesis could arise from incomplete knowledge of Gal_f synthesis and/or an incorrect assignment of this gene family (or individual members therein) as encoding active UGMs. Thus, we sought confirmation that representative candidate *GLF* ORFs encoded UGM activity.

Rescue of Gal_f-dependent LPS synthesis in *E. coli*. We developed an in vivo complementation assay for UDP-Gal_f synthesis in *E. coli*, adapted from work by Whitfield and colleagues (10). These authors showed that expression of the *Klebsiella rfb*_{KPO1} locus in an *E. coli* strain deleted for the endogenous LPS *rfb* locus led to the synthesis of LPS O1 antigen bearing the repeating unit [→3)-β-D-Gal_f(1→3)-α-D-

Gal_f-(1→). Notably, inactivation of the *Klebsiella GLF* gene *rfbD*_{KPO1} abrogated LPS biosynthesis, which could be restored by episomal expression of Kp*GLF*. We surmised that this would provide a rapid and convenient test for the activity of

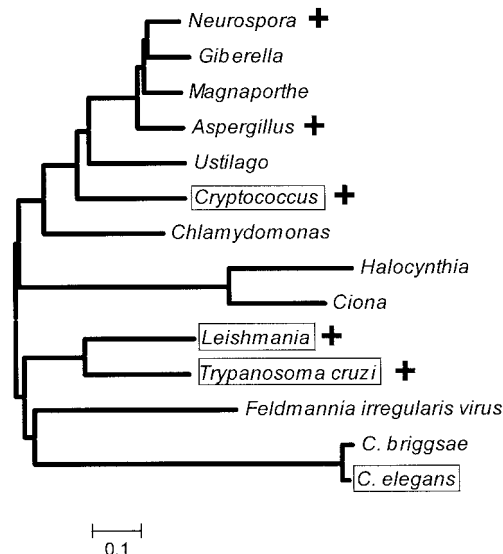


FIG. 2. Phylogenetic tree of the eukaryotic *GLF* family ORFs. A minimum evolution tree for the predicted protein sequences of candidate eukaryotic *GLFs* was constructed using algorithms implicated in the MEGA3 software package (11). A “+” marks species where Gal_f has been detected; Gal_f has not been examined in the other taxa shown. Species where the *GLF*-encoded UGM activity was confirmed in this work are boxed. For information on the specific sequences analyzed, see Table 1 and Table S2 in the supplemental material. The scale at the bottom corresponds to the fraction of amino acid sequence difference.

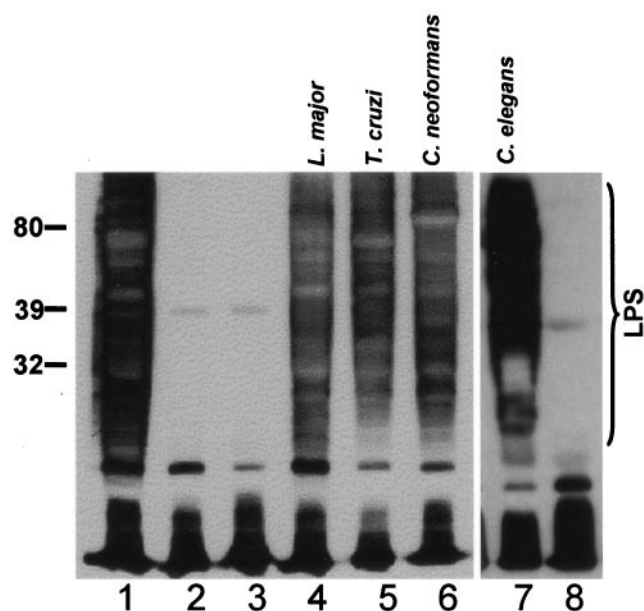


FIG. 3. In vivo rescue of Gal_f-dependent LPS synthesis in *E. coli*. Bacterial lysates were prepared and subjected to Western blot analysis with rabbit anti-*Klebsiella* galactan I antisera as described in Materials and Methods. The migration of molecular mass markers (in kDa) is shown on the left, and the region corresponding to migration of LPS is shown by the bracket on the right; the antiserum also reacts with low-molecular-weight material (perhaps lipid A and LPS core), but the identification of O antigen is unequivocal. All strains derive from the *E. coli rfb* deletion strain CWG288. Plasmids introduced were as follows: lane 1, pWQ71 (intact *Klebsiella rfb* locus; positive control); lane 2, pWQ70 (pWQ71 with deletion inactivating *GLF*); lane 3, pWQ70+pET3a (vector control); lane 4, pWQ70+pET3a-LmjGLF; lane 5, pWQ70+pET3a-TcGLF; lane 6, pWQ70+pET3a-CnGLF; lane 7, pWQ70+pET3a-CeGLF; and lane 8, pWQ70 + inactive mutant pET3a-CeGLF (clone 1.1, strain B3597, which contains a 1-nt deletion at position 129 introducing a premature stop amongst other PCR-generated mutations). The experiment shown in lanes 7 to 8 was performed separately from those shown in lanes 1 to 6.

potential heterologous *GLFs*. Accordingly, we obtained the *L. major*, *T. cruzi*, *C. neoformans*, and *C. elegans* *GLF* ORFs by PCR and introduced them into the bacterial expression vector pET3a. These were then introduced into *E. coli* strain CWG288 (deleted for the endogenous *rfb* locus) containing plasmid pWQ70, which bears the *Klebsiella rfb*_{KPO1} locus with an inactivated *GLF* (*rfbD*) gene (10). LPS expression was monitored by Western blotting with rabbit anti-galactan I antibody, which recognizes a [Gal_f-Gal_p] repeating unit (5).

We confirmed that the intact *Klebsiella rfb* locus restored Gal_f-containing O1 LPS biosynthesis in *E. coli* strain CWG288 transformed with pWQ71, which bears an intact *rfb* locus, while plasmid pWQ70 bearing the inactivated *GLF* failed to do so (Fig. 3, lane 1 versus lane 2). Transformation of strain CWG288/pWQ70 with any of the four eukaryotic species' pET3a-*GLF* constructs (*L. major*, *T. cruzi*, *C. neoformans*, or *C. elegans*) resulted in synthesis of LPS reactive with anti-*Klebsiella* galactan I antisera, at levels comparable to those of the controls (Fig. 3, lanes 4 to 7). As expected, neither pET3a nor a frameshift mutant *C. elegans* *GLF* rescued LPS expression (Fig. 3, lanes 3 and 8, respectively). These data

suggested that the eukaryotic candidate *GLFs* mediated Gal_f synthesis when expressed heterologously in *E. coli*.

While our data suggest that similar levels of LPS restoration were seen in the experimental versus control samples, some restraint is required in interpreting these data. As shown below, the eukaryotic UGMs are largely insoluble when expressed in *E. coli*. Thus, differences in the degree of eukaryotic *GLF* rescue of LPS expression may reflect either intrinsic differences in catalytic activity of the enzymes or "noise" arising from differences in expression and/or folding. Nonetheless, these data suggest that the eukaryotic UGMs are able to provide sufficient UDP-Gal_f under these conditions for LPS synthesis comparable to that of authentic bacterial UGMs.

UGM activity of *L. major* and *C. neoformans* *GLFs*. To confirm directly that candidate *GLFs* encoded UGM activity, we first expressed high levels of protein in *E. coli* using the pET3a system. While abundant expression was evident in sodium dodecyl sulfate-PAGE analysis, the majority of protein appeared in insoluble inclusion bodies (data not shown). Nonetheless, soluble protein extracts were prepared, and UGM activity was assayed in the reverse direction, following the conversion of UDP-Gal_f to UDP-Gal_p by HPLC (14). Since the *E. coli* strain BL21(DE3) lacks *GLF*, there was no endogenous background UGM activity in crude preparations (15) (Fig. 4E and F). In contrast, with extracts from cells expressing *L. major* or *C. neoformans* *GLFs*, the UDP-Gal_f substrate was converted to UDP-Gal_p (Fig. 4C and D) in a protein-dependent manner (Fig. 4A and B). These data establish that these eukaryotic *GLFs* encode proteins with UGM activity. In the future, studies of purified soluble and active eukaryotic UGMs will address the catalytic properties of these enzymes relative to those of prokaryotes in more detail.

DISCUSSION

By using a combinatorial bioinformatics approach incorporating criteria including sequence homology, protein motifs, and phylogenetic associations with Gal_f synthesis, we identified a candidate *Leishmania* *GLF* gene and from this a large family of potential eukaryotic *GLFs*. We used a rapid in vivo assay of *GLFs* expressed in *E. coli* as a first screen for UGM activity of several candidate eukaryotic proteins (Fig. 3), which were confirmed subsequently in enzymatic assays (Fig. 4). While the eukaryotic UGM/*GLF* proteins show some relationship to the previously described prokaryotic enzymes, the amino acid sequence divergence was extensive, and the correlation of potential *GLF* homologs with organisms known to synthesize Gal_f was imperfect. Perhaps for these reasons, eukaryotic *GLFs* had remained elusive despite efforts undertaken by many researchers.

Despite the extensive divergence, eukaryotic UGMs share key properties with those of prokaryotes. The eukaryotic proteins contain clear flavin binding motifs, and most importantly, a number of residues implicated by functional or structural criterion in substrate binding were conserved (Fig. 1). Preliminary analysis suggests that the eukaryotic UGMs can be readily modeled to the *E. coli* UGM structure determined previously, with a number of sequence insertions potentially mapping to external loops (data not shown). As in prokaryotes, eukaryotic UGMs lack obvious secretory signals and are likely to be

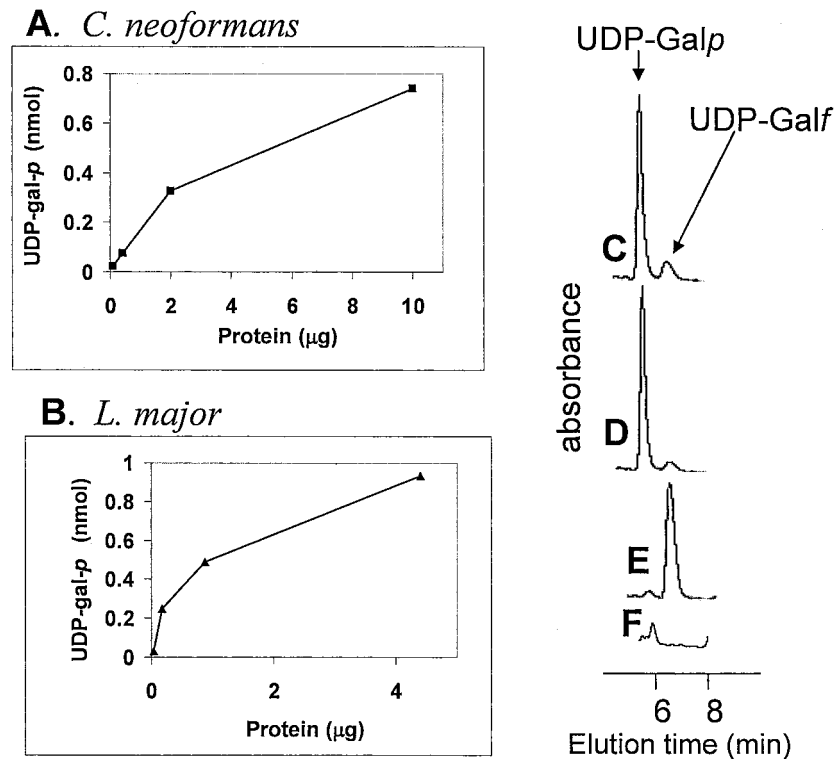


FIG. 4. UGM assays of eukaryotic *GLF*s expressed in *E. coli*. Protein-containing supernatants were prepared, and UGM activity was assayed by monitoring conversion of UDP-Gal_f to UDP-Gal_p by HPLC (15) as described in Methods. Panels A and B show the production of UDP-Gal_p as a function of added protein extract. (A) Extract from *E. coli* expressing *C. neoformans* *GLF* [BL21(DE3)/pET3a-Cn*GLF*]; (B) Extract from *E. coli* expressing *Leishmania* *GLF* [BL2(DE3)/pET3a-Lmj*GLF*]. Panels C-F show representative HPLC traces and controls used to generate the data in panels A and B. (C) The 10- μg protein point from (A); (D) the 4.4- μg protein point from (B); (E) 28 μg of protein incubated from *E. coli* BL21(DE3) incubated under standard assay conditions. In the standard assay, the total amount of UDP-Gal_f added was 1 nmol. Panel F shows 28 μg of protein incubated from *E. coli* BL21(DE3) incubated without UDP-Gal_f, showing that the small peak in (E) comes from endogenous UDP-Gal_p or UDP-Glc from *E. coli*.

cytoplasmic. Since the bulk of eukaryotic glycoconjugate synthesis typically occurs within the secretory pathway, this suggests that one or more nucleotide sugar transporters must recognize and transport UDP-Gal_f to luminal compartments.

We found a number of genes with strong homology to *GLF*s encoding eukaryotic UGMs in species not known to synthesize Gal_f (Fig. 1 and 2; Table 1; see Tables S1 and S2 in the supplemental material). These predicted proteins also showed conservation of key UGM substrate-binding residues (Fig. 1, data not shown). Since the phylogenetic tree presented in Fig. 2 showed that members of both major branches of the eukaryotic *GLF* family had UGM activity, by parsimony we believe that most or all of these candidate *GLF*s encode proteins with UGM activity as well. Given the paucity of information on Gal_f in eukaryotes outside of fungi and protozoa, these new eukaryotic *GLF*s raise the possibility that Gal_f synthesis may occur more widely in species than previously supposed. For fungi such as *Ustilago*, *Magnaporthe*, and *Giberella*, this was unsurprising, but the occurrence of eukaryotic *GLF*s in the nematodes *C. elegans*, the urochordates *Ciona* and *Halocynthia*, and the algae *Chlamydomonas* and algal virus *Feldmannia* (Fig. 1; Table 1; see Tables S1 and S2 in the supplemental material) was unanticipated. Notably, our data show that *C. elegans* *GLF* encodes a protein with UGM activity. Additionally, we found strong *GLF* homologs in a variety of unfinished genome and

EST surveys, including *Histoplasma capsulatum*, which contains Gal_f within sphingolipids (2), and nematode parasites of humans (*Brugia malayi* and *Onchocerca volvulus*), animals (*Strongyloides*), and plants (*Heterodura* and *Meloidogyne*) (see Table S1 in the supplemental material) (3, 33).

Assuming that the presence of eukaryotic *GLF* within a species is predictive of Gal_f synthesis, our analyses suggest that the phylogenetic distribution of the *GLF*/Gal_f synthetic pathway shows some features reminiscent of those seen in prokaryotes. First, the *GLF*/Gal_f pathway occurs only sporadically, in widely divergent species, and second, it is often found in pathogenic/parasitic species from both the microbial and metazoan worlds. Why pathogens show an affinity for inclusion of Gal_f in their metabolomic repertoire is unknown; potential explanations might include the strong immunogenicity of Gal_f and/or its ability to adopt novel structural conformations (17, 31), both of which could contribute to microbial virulence and survival. Notably, the *GLF* relationships depicted in Fig. 2 or evident in Table 1 often do not closely follow those of the species involved. Potentially, the predilection of pathogens for Gal_f could be a contributing factor to the sporadic distribution of Gal_f synthesis among species, perhaps by lateral gene transfer mechanisms as seen in bacterial LPS O antigens (19). The presence of a *GLF* within the algal virus *Feldmannia* (Fig. 2) is

interesting in this light, since viruses often can be transmitted laterally among species.

Gal_f-containing glycoconjugates are often dominant features of the surface of many protozoans and fungi, making Gal_f synthesis a potential target for chemotherapy. However, the role of Gal_f conjugates in the survival and virulence of eukaryotic microbes appear to vary greatly among species. For example, in fungi, Gal_f comprises only a small portion of the *Cryptococcus* capsule (30) but a major fraction of the abundant galactomannans in *Aspergillus* (13). Interestingly, in several global surveys of gene function using RNA interference approaches, inhibition of the *C. elegans* *GLF* encoded by gene H04M03.4 showed a variety of deleterious effects (1, 8, 23); future studies will be required to establish whether this involves Gal_f-containing glycoconjugates. Preliminary tests of a panel of prospective prokaryotic UGM inhibitors (22) suggest that they fail to inhibit the eukaryotic UGMs studied here (unpublished work), in keeping with the extensive amino acid sequence divergence (Fig. 1). Thus, it will be necessary to identify eukaryote-specific UGM inhibitors in the future. Interestingly, it has recently been suggested that several of the inhibitors of the *Mycobacterium tuberculosis* UGM may also act against bacteria independently of UGM (28).

For *Leishmania*, genetic studies suggest that parasites lacking Gal_f glycoconjugates, such as LPG and glycoinositolphospholipids, retain virulence in mammalian infections, although their ability to be transmitted by the insect vector sand fly is greatly reduced (26, 34, 35). Whether a similar conclusion pertains to *Trypanosoma cruzi*, where Gal_f-containing molecules are also highly abundant in infective stages, awaits confirmation (18). Given that the *Leishmania* and *T. cruzi* genomes encode numerous potential UDP-Gal_f transferases (34), genetic inactivation of their *GLFs* (which occur in one or two copies, respectively; Table 1) may offer an easier definitive test (9). Interestingly, *T. brucei*, which is thought to be evolutionarily more closely related to *T. cruzi* than *Leishmania*, lacks *GLF* and Gal_f-containing conjugates. While this could arise from evolutionary loss of *GLF* in this lineage, the complex disjunct distribution and relationships of *GLF* genes in other eukaryotic taxa suggest that other explanations cannot yet be ruled out.

In summary, we have identified a large eukaryotic *GLF* family and provided evidence that members of the two major groups within this family possess UGM activity. This in turn has permitted predictions about the occurrence of Gal_f synthesis among species and its frequent appearance within both microbial and metazoan pathogens. Future work will focus on exploitation of UGM-targeted inhibitors in chemotherapy and determination of the structure and role of Gal_f in those lineages for which information is currently lacking.

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