Structural Determination and Tryptophan Fluorescence of Heterokaryon Incompatibility C2 Protein (HET-C2), a Fungal Glycolipid Transfer Protein (GLTP), Provide Novel Insights into Glycolipid Specificity and Membrane Interaction by the GLTP Fold^{*S}

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HET-C2 is a fungal protein that transfers glycosphingolipids between membranes and has limited sequence homology with human glycolipid transfer protein (GLTP). The human GLTP fold is unique among lipid binding/transfer proteins, defining the GLTP superfamily. Herein, GLTP fold formation by HET-C2, its glycolipid transfer specificity, and the functional role(s) of its two Trp residues have been investigated. X-ray diffraction (1.9 Å) revealed a GLTP fold with all key sugar headgroup recognition residues (Asp⁶⁶, Asn⁷⁰, Lys⁷³, Trp¹⁰⁹, and His¹⁴⁷) conserved and properly oriented for glycolipid binding. Far-UV CD showed secondary structure dominated by α-helices and a cooperative thermal unfolding transition of 49 °C, features consistent with a GLTP fold. Environmentally induced optical activity of Trp/Tyr/Phe (2:4:12) detected by near-UV CD was unaffected by membranes containing glycolipid but was slightly altered by membranes lacking glycolipid. Trp fluorescence was maximal at ${\sim}355$ nm and accessible to aqueous quenchers, indicating free exposure to the aqueous milieu and consistent with surface localization of the two Trps. Interaction with membranes lacking glycolipid triggered significant decreases in Trp emission intensity but lesser than decreases induced by membranes containing glycolipid. Binding of glycolipid (confirmed by electrospray injection mass spectrometry) resulted in a blue-shifted emission wavelength maximum (~6 nm) permitting determination of binding affinities. The unique positioning of Trp²⁰⁸ at the HET-C2 C terminus revealed membrane-induced conformational changes that precede glycolipid uptake, whereas key differences in residues of the sugar headgroup recognition center accounted for altered glycolipid specificity and suggested evolutionary adaptation for the simpler glycosphingolipid compositions of filamentous fungi.

Self/nonself recognition is a universally important process, encompassing intercellular interactions ranging from vertebrate immune responses to somatic chimera formation in protists, filamentous fungi, sponges, ascidians, and tunicates. Self/ nonself discrimination becomes critical for filamentous fungi during hyphal fusion, which enables the exchange of cytoplasm and nuclei during the assimilative growth phase (1-4). Nonself recognition triggers a postfusion, programmed cell death process known as vegetative incompatibility, which leads to heterokaryon death. The benefits of vegetative incompatibility include prevention of transmission of deleterious cytoplasmic elements (*i.e.* viruses) as well as restriction of plundering by parasitic genotypes.

het genes are known to play a major role in vegetative incompatibility processes that occur in *Neurospora crassa* and *Podospora anserina. het* genes exhibit extensive polymorphism and generally encode proteins carrying a HET domain (1-4). Originally, this domain was linked to the *het-c2* gene of *P. anserina*, where it was shown to encode a protein similar in size and with limited sequence homology to mammalian glycolipid transfer proteins (GLTPs)⁵ (5, 6). HET-C2 was shown to stimulate *in vitro* inter-



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The on-line version of this article (available at http://www.jbc.org) contains supplemental Table S1 and Figs. S1–S7.

The atomic coordinates and structure factors (code 3KV0) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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⁵ The abbreviations used are: GLTP, glycolipid transfer protein; BisTris, 2-[bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol; POPC, 1-palmitoyl-2-oleoyl-L-phosphatidylcholine; Cer, ceramide; Lac, lactose; OPM, orientation of proteins in membranes; GM1, Gal β 1 \rightarrow 3GalNAc1 \rightarrow 4[Neu5Ac α 2 \rightarrow 3]Gal β 1 \rightarrow 4Glc β 1 \rightarrow 1Cer.

membrane transfer of galactosylceramide, but not sphingomyelin, ceramide, phosphatidylcholine, or cholesterol (7). Similar glycolipid specificity is displayed by mammalian GLTPs (8), which accomplish transfer using a distinct two-layer "sandwich motif," dominated by α -helices and without intramolecular disulfide bonds (9-11). A binding site for a single glycolipid molecule consists of a sugar headgroup recognition site located where the sugaramide portion of the glycolipid is tethered via multiple hydrogen bonds to the protein's surface. A hydrophobic pocket, lined with nonpolar amino acid residues, encapsulates a major portion of the nonpolar hydrocarbon chains of the glycolipid ceramide region. The novel architecture of GLTP and its lipid liganding pocket are distinguishing features compared with other lipid binding and transfer proteins, which use conformational folds dominated by the β -sheet (*i.e.* β -grooves/concave cups and β -barrels) or helical bundles stabilized by multiple disulfide-bridges (i.e. saposin folds (8-10)). Accordingly, the human GLTP fold serves as the prototype for the GLTP superfamily (12, 13). Nonetheless, other protein folds (e.g. saposin, nonspecific LTP, and phosphatidylinositol transfer protein) are able to bind/transfer glycolipid (14-21).

Currently, there is a lack of experimental data elucidating HET-C2 structure/conformation, leaving the issue of GLTP fold formation by HET-C2 unresolved. Also, HET-C2 transfer specificity for different glycolipids and the functional roles of the two Trp residues of HET-C2 in glycolipid binding and/or membrane interaction remain undefined. Herein, these issues are addressed. The different positioning of Trp residues in HET-C2 provides new insights into the membrane interaction region of the GLTP fold, whereas x-ray diffraction reveals the structural basis for the altered glycolipid specificity of the HET-C2 GLTP fold.

EXPERIMENTAL PROCEDURES

Expression and Purification of HET-C2-HET-C2 was expressed and purified as detailed for GLTP (22). The P. anse*rina* open reading frame encoding HET-C2 (NCBI GenBankTM number U05236) was subcloned (7) into pET-30 Xa/LIC (Novagen) by ligation-independent cloning. Transformed BL21 cells (Escherichia coli) were grown in Luria-Bertani medium at 37 °C overnight, induced with isopropyl 1-thio- β -Dgalactopyranoside (0.1 mM), and then grown for 16-20 h at 15 °C. Purification of rHET-C2 from soluble lysate protein was accomplished by Ni²⁺-NTA affinity chromatography. The N-terminal His-S tag was removed by factor Xa, yielding protein identical in sequence to native HET-C2. HET-C2 was repurified by FPLC SEC using a HiLoad 16/60 Superdex-75 preparation grade column (Amersham Biosciences). For crystallographic studies, HET-C2 was expressed using pET-SUMO vector (Invitrogen). The His-Sumo tag (N terminus) was cleaved using Ulp1 SUMO protease (Invitrogen). HET-C2 fractions obtained by SEC were pooled, centrifugally concentrated (10 kDa cut-off membrane), and verified for purity by SDS-PAGE.

Crystallization of HET-C2—Reductive methylation of surface lysines was carried out to aid crystallization as described previously (23). Methylated HET-C2 was purified by SEC, and the extent of methylation was assessed by matrix-assisted laser desorption ionization time-of-flight mass spectrometry. Methylated and unmethylated HET-C2 were concentrated to 30

mg/ml for crystallization screening. Diffraction quality crystals of methylated HET-C2 were obtained in 0.1 M BisTris (pH 5.5–6.5) and 25% polyethylene glycol 3350.

Structure Determination—For data collection at cryogenic temperatures, crystals were stabilized in mother liquor supplemented with 20% ethylene glycol. X-ray diffraction data for the methylated HET-C2 crystal was collected on beamline ID24-E at the Advanced Photon Source (Argonne National Laboratory). This data set was processed using the HKL2000 suite (24). Systematic absences indicated that the crystals belonged to space group P4₁2₁2 or P4₃2₁2. The volume of the crystal asymmetric unit was compatible with only one subunit and a protein volume per unit of molecular mass of 2.9 Å³ Da^{-1} with a calculated solvent content of 58.1% (v/v). The structure was solved by molecular replacement using PHASER (25) with human apo-GLTP (Protein Data Bank 1SWX) structure as a search model in space group P41212. Refinement was carried out using REFMAC (26) alternating with manual building in COOT (27). The final structure was validated using PROCHECK (28). Statistics for data collection, refinement, and final model geometry are summarized in Table 1. Secondary structural elements were assigned using DSSP (29).

Glycolipid Transfer Activity of HET-C2—Glycolipid intervesicular transfer activity of purified HET-C2 (or GLTP) was monitored by established assays involving glycolipid labeled with either radioactivity or fluorescent probes (30–33) (see supplemental material for details).

Circular Dichroism Spectroscopy-CD spectra were collected using a J-810 spectropolarimeter (JASCO) equipped with a CTC-345 temperature control system at 0.05 mM HET-C2 in 10 mM sodium phosphate (pH 7.4; 10 °C) while purging continuously with N₂. Spectral and temperature dependence measurements in the far-UV range (185-250 nm) were performed at a 2-nm bandwidth using a U-type quartz cell (0.148-mm path length) and in the near-UV range (250-320 nm) at 1 nm bandwidth with a rectangular cell (0.5-cm path length) in a custombuilt thermostated cell holder (34-37). Five accumulations were recorded, each at a 20-nm/min scan rate with a 2-s response time. The temperature dependence of ellipticity at 222 nm was measured using a 60 °C/h scan rate and 8-s response time. Solvent evaporation was prevented by placing a drop of oil (repeatedly boiled in water to remove soluble impurities) on the top of the sample cell. CD spectra in the far-UV range were smoothed using the JASCO noise reduction routine. The CDPro program suite (34), a modified version of three methods (SELCON3 (35), CONTIN/LL locally linearized approximation (38) of CONTIN (39), and CDSSTR (40)), was used to calculate HET-C2 secondary structure from far-UV CD spectra. Tertiary structure class was assessed using the CDPro CLUSTER program (37) (see supplemental material for details).

Measurement of Protein Concentration—HET-C2 concentration was measured at 280 nm using a DU 640 spectrophotometer (Beckman) at 1.8-nm bandwidth and a molar absorptivity of 17.12 mm^{-1} cm⁻¹, obtained by averaging results from four calculation methods (41–44). Spectra were corrected for turbidity by plotting the log dependence of the solution absorbance versus the log of the wavelength and extrapolating the linear dependence between these quantities in the 340–440 nm



range to the 240-300 nm absorption range, using the DU-640 scatter correction routine. Extrapolated absorbance values were subtracted from measured values, decreasing the apparent protein absorbance at ~ 280 nm by $\sim 15\%$.

Fluorescence Measurements—Trp fluorescence (295 nm excitation) was measured at 25 °C from 310 to 420 nm with a SPEX Fluoromax spectrofluorometer (Horiba Scientific) using excitation and emission bandpasses of 5 nm. Protein concentration was kept at $A_{295} < 0.1$ to avoid inner filter effects (45, 46). Fluorescence quenching experiments were performed by adding aliquots of acrylamide or KI (5 M) stocks in 10 mM sodium phosphate-saline buffer (pH 7.4) to HET-C2 (1 μ M; 2.5 ml) with constant stirring at 25 °C. With KI, 10 mM sodium thiosulfate was added to the stock solution to prevent the formation of I_3^- , which absorbs in the region of Trp fluorescence (45). Data were corrected for dilution effects and analyzed using the Stern-Volmer and modified Stern-Volmer equations as detailed in the supplemental material.

Membrane Interaction and Glycolipid Binding by HET-C2— To assess membrane interaction, vesicles consisting of POPC/ glycolipid (8:2) (1.5 mM) or POPC (1.5 mM) were added stepwise to HET-C2 (1 μ M) in small aliquots while monitoring Trp emission. Lipids also were presented to HET-C2 by microinjection after dissolving in ethanol as recently detailed by Zhai *et al.* (46). Measurements were performed under constant stirring by the addition of small aliquots (1 μ I) of GSL, dissolved in ethanol (0.1 mM), to protein (1 μ M; 2.5 mI). Fluorescence titration curves were analyzed according to Equation 1,

$$\epsilon - 1 = (\epsilon_b - 1) - K_d(\epsilon - 1)/mn$$
 (Eq. 1)

where K_d is the dissociation constant of lipid-protein complex, *m* is lipid concentration, and *n* is the number of lipid binding sites (47, 48). ϵ is the spectral parameter value (emission wavelength maximum (λ_{max} expressed as wavenumber) or relative intensity, *I*) accompanying lipid binding to HET-C2 at lipid concentration *m*. Thus, ϵ can represent either I/I_0 or (λ_{max})₀/ λ_{max} , where the subscript 0 indicates values in the absence of lipid. ϵ_b represents spectral properties of the protein-lipid complex. According to Equation 1, the slope of $\epsilon - 1$ versus ($\epsilon - 1$)/*m* yields K_d/n and represents the reciprocal of the protein/ lipid association constant.

Mass Spectrometry—HET-C2 and HET-C2-glycolipid complex (10 μ M protein) were analyzed using an Agilent MSD-TOF mass spectrometer (Santa Clara, CA) in 5 mM ammonium acetate by infusing directly into the electrospray source. Spectra were collected in positive mode over a 500–5000 *m*/*z* range using parameters optimized for complex stability (*e.g.* capillary, 3000 V; fragmentor, 300 V; skimmer, 60 V; Oct RF, 300 V; Oct DC, 32 V. Raw spectral data were transformed into relative molecular masses using the Agilent TOF Protein Confirmation software.

RESULTS

Structure and Glycolipid Specificity of HET-C2—Limited sequence homology between HET-C2 and porcine GLTP was first noted by Saupe *et al.* (5). Subsequently, glycolipid intermembrane transfer by HET-C2 was shown with radiolabeled and fluorescently labeled galactosylceramide (7). These find-

TABLE 1

X-ray data collection and refinement statistics for HET-C2 Values in parentheses are for the highest resolution shell.

Parameter	Value		
Data collection			
Wavelength (Å)	0.9792		
Space group	P41212		
Unit cell	1 1		
a, b, c	96.93, 96.93, 57.91		
α, β, γ	90, 90, 90		
Resolution (Å)	50.0-1.9 (1.97-1.9)		
Total reflections	379,083		
Unique reflections	22,551		
I/oI	36.8 (2.9)		
Completeness (%)	98.9 (93.9)		
Redundancy	8.7 (6.0)		
R _{merge}	7.2 (35.9)		
Refinement			
$R/R_{\rm free}$	21.6/24.8		
Average B-factor			
Protein	37.9		
Water	43.6		
No. of atoms			
Protein	1577		
Water	118		
RMSD ^a			
Bond lengths (Å)	0.016		
Bond angles (degrees)	1.512		
Ramachandran statistics (%)			
Mostly allowed regions	95.5		
Allowed regions	4.5		
Generously allowed regions	0.0		
Disallowed regions	0.0		

^{*a*} Root mean square deviation.

ings occurred prior to structural resolution of human GLTP and mapping of the glycolipid binding site in the GLTP fold (9-11). To define HET-C2 conformation and localize residues potentially involved in glycolipid binding, the HET-C2 structure was solved by x-ray crystallography. Formation of high quality crystals of apoHET-C2 was greatly facilitated by mild reductive methylation of surface Lys residues (23), enabling resolution to 1.9 Å by molecular replacement (human apoGLTP was the search model) (9, 10). The electron density throughout the polypeptide chain was of good quality except for residues 1–13 and 99, presumably due to disordering. The final crystallographic model was refined to R and R_{free} values of 21.6 and 24.8%, respectively (Table 1). The HET-C2 conformational architecture strongly resembles the human GLTP fold (9, 10), showing the distinctive two-layer "sandwich motif" dominated by α -helices (Fig. 1 and supplemental Fig. S1). Like GLTP, HET-C2 has eight α -helices with one layer consisting of $\alpha 1$, $\alpha 2$, α 6, and α 7, whereas α 3, α 4, α 5, and α 8 make up the other layer. Unlike GLTP, a small anti-parallel β -sheet is present near the N terminus of HET-C2. The two Cys residues (Cys¹¹⁸ and Cys¹⁶²) present in HET-C2 are too far apart for disulfide stabilization. The N-terminal region of HET-C2 is disordered, as in GLTP, but is more extended by ~ 16 amino acids (Fig. 1, B and C). Superpositioning of HET-C2 and human GLTP shows that most residues present in the sugar binding pocket and hydrophobic channel are conserved. In HET-C2, Asp⁶⁶, Asn⁷⁰, Lys⁷³, Trp¹⁰⁹, and His¹⁴⁷ are properly positioned to form a sugar headgroup binding site, selective for glycolipids (Fig. 2A). This putative sugar headgroup recognition center is located at the entrance to a hydrophobic pocket, where the nonpolar hydrocarbon chains of the glycolipid are expected to localize. The hydrophobic pocket is lined by 2 dozen nonpolar residues, most



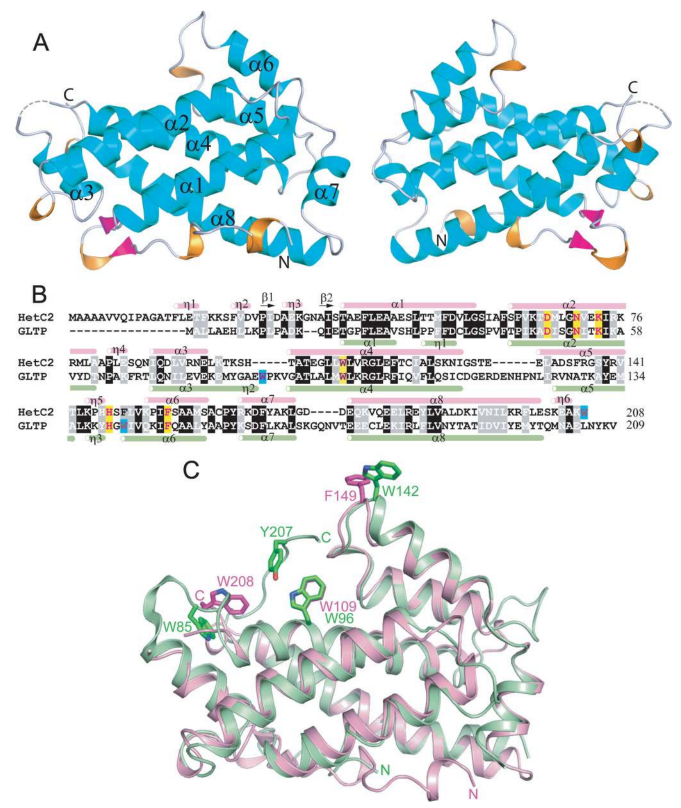


FIGURE 1. *A*, tertiary structure of HET-C2 in the apo-form. Front and back views of HET-C2 are shown with α -helices (*cyan*), 3₁₀ helices (*orange*), β -sheet (*magenta*), and loop segments (*light gray*) in a schematic diagram. *B*, structure-based sequence alignment between HET-C2 and human GLTP. Secondary structural elements of HET-C2 (*pink*) and GLTP (*light green*) are shown *above* and *below* the aligned sequences by *tubes* indicating 3₁₀ (η) and α -helices (α) and *arrows* indicating β -strands (β). Conserved residues that interact with the ceramide-linked sugar of glycolipids are *highlighted* in *yellow* in GLTP and HET-C2. Nonbinding site Trp residues are *highlighted* in *turquoise*. *C*, structural superposition of HET-C2 (*pink*) and human GLTP (*green*). Key aromatic residues are shown in *stick representation*. Trp¹⁴² of GLTP and Phe¹⁴⁹ of HET-C2 are localized on the surface and are completely accessible to the aqueous milieu. Human apoGLTP structure was solved previously (9, 10).



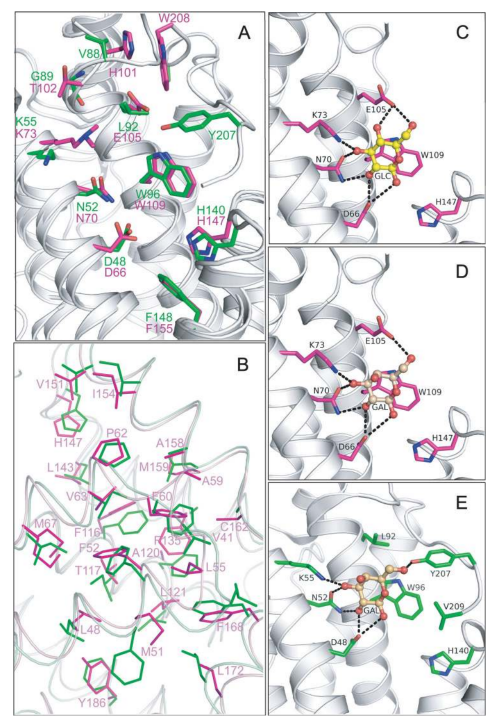


FIGURE 2. *A*, structural superposition of HET-C2 (*magenta*) and GLTP (*green*) showing conserved residues present in the pocket that interacts with the sugar-amide moiety of glycolipids. Lys⁷³ in HET-C2 is shown containing the exogenous dimethyl group introduced to facilitate crystallization. Human apoGLTP structure was solved previously (9, 10). *B*, residue homology and similarity between hydrophobic pockets of HET-C2 and human GLTP. For clarity, only HET-C2 residues have been labeled. The following positional correspondence is observed for HET-C2 and GLTP, respectively: Leu⁴⁸ = Leu³⁰, Met⁵¹ \approx Phe³³, Phe⁵² = Phe³⁴, Leu⁵⁵ = Leu³⁷, Ala⁵⁹ \approx Val⁴¹, Phe⁶⁰ = Phe⁴², Pro⁶² = Pro⁴⁴, Val⁶³ \approx Ile⁴⁵, Met⁶⁷ \approx Ile⁴⁹, Phe¹¹⁶ = Phe¹⁰³, Thr¹¹⁷ \approx Ile¹⁰⁴, Ala¹²⁰ \approx Pha¹⁰⁷, Leu¹²¹ = Leu¹⁰⁸, Phe¹³⁵ \approx Ala¹²⁶, Leu¹⁴³ = Leu¹³⁶, His¹⁴⁷ = His¹⁴⁰, Val¹⁵¹ = Val¹⁴⁴, Ile¹⁵⁴ = Ile¹⁴⁷, Ala¹⁵⁸ = Ala¹⁵¹, Met¹⁵⁹ \approx Leu¹⁵², Cys¹⁶² \approx Ala¹⁵⁵, Phe¹⁶⁸ = Phe¹⁶¹, Leu¹⁷² = Leu¹⁶⁵, Tyr¹⁸⁶ \approx Phe¹⁸³. C, docking of Glc onto the GSL headgroup binding site of HET-C2. Side chains of HET-C2 crystallization in wild type HET-C2, the exogenous dimethyl group added to Lys⁷³ to facilitate HET-C2 crystallization is not shown. *D*, docking of Gal onto the GSL headgroup binding site of HET-C2. The *color scheme* is the same as for *C* except that the galactose sugar ring is *colored yellow* more than of GLTP residues involved in Gal binding are shown in *scheme* is the same as for *D* except that the side chains of GLTP residues involved in Gal binding are shown in *green*. The galactose location is derived from Protein Data Bank entry 2EVL (10).

of them identical to those lining the hydrophobic pocket of GLTP (Fig. 2*B*).

Despite conservation of so many key residues in HET-C2, subtle structural differences raised the possibility of altered selectivity for glycolipids compared with GLTP, which can transfer a variety of glycolipids of differing structural complexity (8). The only glycolipid tested earlier with HET-C2 was GalCer (7). In fungi, glucosylceramide (GlcCer) is more widespread than GalCer, but both GalCer and GlcCer reportedly occur in filamentous fungi (49-51). We found that HET-C2 readily transfers GlcCer (Fig. 3), surpassing that of GLTP. LacCer is transferred by HET-C2 but not nearly as well as GlcCer or GalCer. Unlike GLTP, HET-C2 transferred negatively charged glycolipids (i.e. sulfatide or ganglioside GM1) very poorly.

To elucidate the structural basis for the glycolipid selectivity of HET-C2, comparative sugar docking was performed using high resolution GLTP-GalCer and GLTP-LacCer complexes (9, 10) as guides (Fig. 2, C-E). In HET-C2, Trp¹⁰⁹ serves as a stacking platform that facilitates the hydrogen bonding of Asp⁶⁶, Asn⁷⁰, and Lys⁷³ with Gal or Glc in the same way that Trp⁹⁶ facilitates the hydrogen bonding network of Asp⁴⁸, Asn⁵², and Lys⁵⁵ with Gal or Glc in human GLTP (9, 10). However, in HET-C2, residues corresponding to Tyr^{207} and Val^{209} of GLTP are absent, and Leu⁹² in GLTP is replaced by Glu¹⁰⁵ in HET-C2. Docking of Glc and Gal shows that Glu¹⁰⁵ of HET-C2 forms one hydrogen bond with Gal and two hydrogen bonds with Glc (Fig. 2, C and *D*), thus compensating for the absence of hydrogen bond formation by Gal (and Glc) with GLTP Tyr²⁰⁷ (Fig. 2*E*). The location of negatively charged Glu¹⁰⁵ seems to impede interactions with 3sulfo-GalCer (sulfatide).

Circular Dichroism of HET-C2— HET-C2 conformation in solution as well as in the presence of membrane vesicles containing or lacking



glycolipid was assessed by far-UV and near-UV CD. Fig. 4*A* shows far-UV CD spectra characteristic of high helical content in secondary structure. Calculations summarized in supplemental Table S1 indicate 56.6% helix, 8.8% β -structure, 12.1% β -turns, and 24.1% random. Incubation of HET-C2 with POPC vesicles either lacking or containing glycolipid resulted

in little change in the far-UV CD spectra and secondary structure calculations showing 54.5% α -helix, 9% β -structure, 12.8% β -turns, and 24% random (supplemental Table S1).

Near-UV CD spectra (Fig. 4*B*), which reflect tertiary structure, showed significant environmentally induced optical activity arising from the four Tyr and two Trp residues (270 –

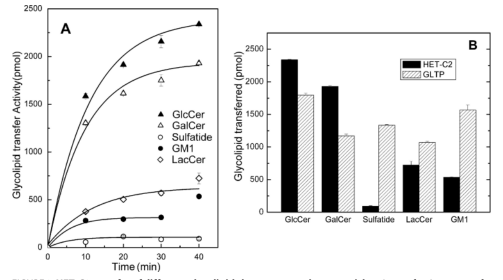


FIGURE 3. **HET-C2 transfer of different glycolipids between membrane vesicles.** *A*, transfer time course for different radiolabeled glycolipids by HET-C2 at 25 °C using a fixed protein amount (0.5 μ g). HET-C2 preferentially transfers glycolipids with headgroups consisting of a single uncharged sugar. *B*, comparison of HET-C2 and GLTP transfer of different glycolipids at 25 °C using fixed protein amounts (0.5 μ g) as detailed in the supplemental material. In contrast to HET-C2, GTLP readily transfers glycolipids with charged and complex sugar headgroups. The data points were fit using a first order exponential fitting routine. All data points represent duplicate assays (average values).

300 nm) of HET-C2. The negative signals at \sim 255, 262, and 268 nm probably belong to the 12 Phe residues; the prominent negative signal near ~277 nm probably originates from the four Tyr residues and/or two Trp residues; and the small negative peak near ~ 287 nm probably arises from the two Trp residues. When vesicles containing glycolipid were incubated with HET-C2, the near-UV CD signal was unaffected. Interestingly, incubation of HET-C2 with membranes lacking glycolipid resulted in slightly decreased near-UV CD signal from Tyr and/or Trp.

Thermally induced unfolding provided insights into HET-C2 stability at pH 7.4. The first derivative of the CD signal at 222 nm as a function of temperature (Fig. 4*C*) revealed a highly cooperative unfolding transition between 25 and

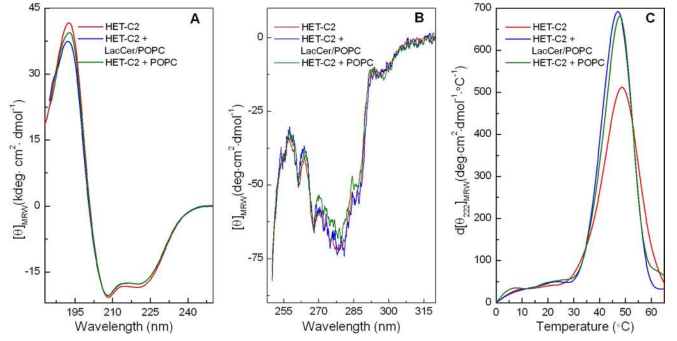


FIGURE 4. **Circular dichroism of HET-C2**. *A*, far-UV CD. The spectra indicate high helical content in HET-C2 with large negative $n-\pi^*$ transitions at 222 nm as well as $\pi-\pi^*$ transitions split into two transitions because of exciton coupling, resulting in negative bands at ~208 nm and positive bands at ~192 nm. Data are presented in units of molar ellipticity per residue. *B*, near-UV CD. Slightly altered spectral signals from Tyr/Trp are obtained by incubation of apoHET-C2 with membranes lacking glycolipid but not with membranes containing glycolipid, consistent with small conformational changes triggered by membrane interaction. *C*, derivative plots for temperature dependence of molar ellipticity at 222 nm. The relatively low midpoints of the unfolding transitions are consistent with a lack of conformational stabilization by intramolecular disulfides, as expected for a GLTP fold. ApoHET-C2 conformation is slightly destabilized by membrane interaction. Spectra were recorded at pH 7.4 as detailed under "Experimental Procedures."



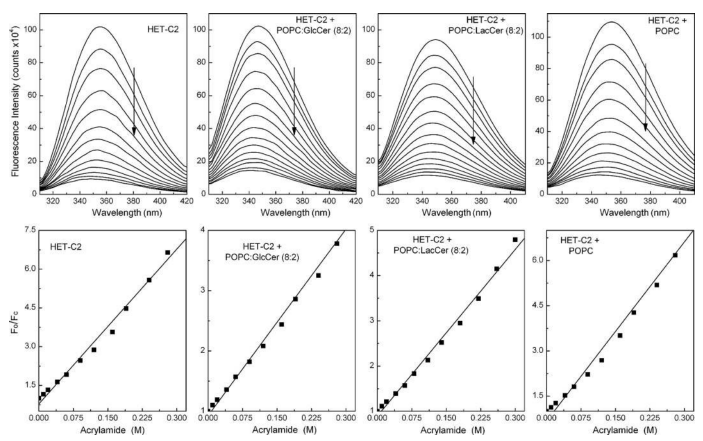


FIGURE 5. HET-C2 Trp emission quenching by acrylamide. Top, intrinsic Trp quenching in the presence of increasing concentrations of acrylamide (vertical arrows). Bottom, Stern-Volmer analyses of the acrylamide quenching response performed as described in the supplemental materials. The correlation coefficients are 0.999 except for LacCer/POPC vesicles (0.998). Values derived from the plots for quenching constants and Trp accessibility are summarized in Table 2.

55 °C, with a midpoint of ~49 °C. The relatively low unfolding temperature midpoint is consistent with a lack of stabilization by intramolecular disulfides. HET-C2 incubation with membranes lacking or containing glycolipid slightly decreased the unfolding transition temperature midpoint and decreased peak broadening, suggesting membrane-enhanced cooperativity of unfolding (Fig. 4*C*).

Intrinsic Fluorescence of HET-C2 Trp—Because Trp residues are important for glycolipid binding and membrane interaction of GLTP (8, 9, 45, 46, 52-55), we characterized the Trp emission of HET-C2. HET-C2 has two Trp and four Tyr residues in its 208-residue sequence. In contrast, GLTP has three Trp and 10 Tyr residues among its 209 residues. Sequence alignment shows that the number and location of intrinsic fluorescent residues differ greatly in HET-C2 and GLTP (Fig. 1B). One of the two Trp residues in HET-C2 aligns with GLTP Trp⁹⁶, a key residue for sugar headgroup interaction in the glycolipid binding site. The other HET-C2 Trp does not align with either of the remaining two GLTP Trp residues, one of which (Trp¹⁴²) seems to be directly involved in GLTP-membrane interactions. Superpositioning of the structures of HET-C2 and GLTP (Figs. 1C and 2A) shows that Trp¹⁰⁹ of HET-C2 is well located to play a similar role as GLTP Trp⁹⁶ within the glycolipid sugar headgroup recognition center. In contrast, Trp²⁰⁸ at the HET-C2 C terminus resides on the surface but in a location much different from that of GLTP Trp¹⁴². HET-C2 Trp was found to fluoresce maximally (λ_{max}) at ~355 nm (supplemental Fig. S3). Denaturation with 8 m urea caused the $\lambda_{\rm max}$ to red shift only ~ 2 nm. The red-shifted $\lambda_{\rm max}$ of wild-type HET-C2 suggests a relatively polar, average environment for the two Trp residues, consistent with surface localization rather than burial in the hydrophobic core (45, 56).

The accessibility and environment of the Trp residues were further evaluated using aqueous quenchers. Fig. 5 (top left) shows that acrylamide quenched >90% of the average Trp emission signal of HET-C2, consistent with exposure of Trp residues to the aqueous milieu. Quenching with KI resulted in a \sim 76% decrease in Trp emission signal (supplemental Fig. S6). The lower quenching efficiency by KI was consistent with charged residues residing near either or both Trp residues. With both quenchers, the quenching efficiency decreased slightly upon denaturation of HET-C2 with 8 M urea (supplemental Fig. S4). Stern-Volmer plots were linear under all conditions for HET-C2, consistent with dynamic quenching at physiological pH and ionic strength (Fig. 5, bottom). Modified Stern-Volmer analyses enabled quantitative determination of the fraction of accessible Trp residues (supplemental Figs. S5 and S6). In native HET-C2, Trp emission intensity was 100% accessible to both quenchers, whereas, in urea-denatured protein, Trp accessibility was reduced to 72% for KI (Table 2). Collectively, the data show that the two HET-C2 Trp residues are in a polar environment exposed to the aqueous milieu (45, 56), consistent with the surface localization shown by x-ray diffraction.



TABLE 2

Analysis of tryptophan fluorescence quenching

 K_{SV} and K_Q are Stern-Volmer and modified Stern-Volmer quenching constants, respectively, and f_a is fractional accessibility. The data represent the extent of quenching and accessibility achieved for both quenchers. The average values obtained for the quenching constants have S.D. values of 5% or less, whereas those of the f_a values are less than 1%.

Protein	Acrylamide			KI			Quenching	
Protein	$K_{\rm SV}$	K _Q	f_a	K _{SV}	K _Q	f_a	Acrylamide	KI
							%	%
HET-C2	19.97	15.4	1.0	6.9	7.75	0.97	90	76
HET-C2, denatured	15.48	8.8	1.0	4.48	8.6	0.72	87	70
HET-C2 + POPC/GlcCer (8:2) vesicles	10.33	10.7	0.94				84	
HET-C2 + POPC/LacCer (8:2) vesicles	12.88	11.5	0.96				88	
HET-C2 + POPC vesicles	19.45	12.6	1.0				88	

Incubation of HET-C2 with membrane vesicles in the presence of quenchers also provided insights into interactions with membranes and glycolipid. When the vesicles lacked glycolipid, the Trp signal was fully accessible to acrylamide quenching (Fig. 5, *top right*). In the presence of vesicles containing glycolipid, Trp emission was slightly protected from acrylamide quenching. The findings are consistent with membrane interaction of HET-C2 being transient and rapid during glycolipid acquisition/release rather than forming long lived protein-lipid complexes. It is also noteworthy (Table 2) that the Stern-Volmer (K_{SV}) and modified Stern-Volmer quenching constants (K_Q) were reduced in the presence of vesicles containing glycolipid, reflecting diminished quenching efficiency compared with HET-C2 Trp in the absence of vesicles.

Intrinsic Trp Fluorescence Changes Induced by Membranes Containing or Lacking Glycolipid-ApoGLTP exhibits a redshifted Trp λ_{max} (~348 nm). High resolution x-ray diffraction data and electrospray ionization mass spectrometry analyses show an absence of bound glycolipid in human GLTP expressed in *E*. coli (9–11, 46). The red-shifted Trp emission λ_{max} of HET-C2 (355 nm) suggested a lack of bound glycolipid, affording the opportunity to determine whether the Trp emission signal of HET-C2 becomes altered by uptake of glycolipid and/or by interaction with membranes containing or lacking glycolipid. Fig. 6 (top) shows that the stepwise addition of POPC vesicles containing glycolipid results in reduction of Trp emission intensity (25–29%) as well as a 6-nm λ_{max} blue shift (355 \rightarrow 349 nm) (Table 3). It is noteworthy that the λ_{max} blue shift ceases after a slight excess of glycolipid has been added, but the intensity continues to steadily decrease. In contrast, the stepwise addition of POPC vesicles lacking glycolipid (Fig. 6, top *right*) produces almost no λ_{max} blue shift (1 nm) but a substantial reduction in emission intensity (\sim 17%). The different Trp emission responses with membranes containing or lacking glycolipid suggested that both glycolipid binding and protein interaction with the membrane contribute to the response. The blue shift is indicative of glycolipid binding, whereas the intensity decrease reflects both glycolipid binding and membrane interaction. For this reason, the blue shift was used to assess the binding affinity of HET-C2 for different glycolipids. K_d values for the binding isotherms (Fig. 7, A and B) ranged from 0.15 to 0.2 μ M (Table 4). To obtain meaningful K_d values from the emission intensity changes induced by glycolipid binding (Fig. 7, C and D), corrections were made for the membrane partitioning contribution to the POPC vesicles over the glycolipid concentration range producing

blue shift (Table 4). With POPC vesicles lacking glycolipid, the intensity change yielded a K_d value of $\sim 5 \ \mu\text{M}$ for the membrane partitioning of HET-C2.

Presentation of Lipids to HET-C2 via Ethanol Microinjection— Recently, Zhai et al. (46) showed that microinjection involving small lipid aliquots dissolved in ethanol is useful for loading the glycolipid binding site of GLTP while minimizing the accumulation of excess membrane interface in solution, thus providing a means to distinguish emission changes induced by glycolipid binding from changes produced by nonspecific partitioning to membrane interface. Fig. 6 (bottom) shows the Trp emission response of HET-C2, titrated with lipid using the EtOH microinjection approach. With each successive injection of glycolipid, the HET-C2 Trp emission λ_{max} became progressively more blue-shifted, whereas fluorescence intensity systematically decreased (25-30%). The emission responses observed for GlcCer, GalCer, and LacCer followed similar patterns. K_d values were similar to those obtained by vesicle addition.

Formation of a wild type HET-C2-glycolipid complex by EtOH microinjection of glycolipid was verified by electrospray ionization mass spectrometry. Fig. 8 shows spectra obtained by direct infusion of glycolipid-free HET-C2 plus HET-C2-GlcCer complex under nondenaturing conditions. Although ions corresponding to monomeric, glycolipid-free rHET-C2 (28,147 Da) dominate, there is clear evidence for formation of monomeric complex (rHET-C2 + GlcCer (28,735 Da)).

In the case of microinjection of ethanol-solubilized POPC, a non-glycolipid that is not transferred by HET-C2, no λ_{max} blue shift was observed. Nonetheless, substantially decreased Trp intensity (\sim 17%) was evident (Fig. 6, *bottom right*). The latter finding was unexpected because ethanol-injected POPC has little effect on GLTP Trp emission intensity (46). The data suggested that interaction of HET-C2 with membrane alters the local environment of Trp²⁰⁸ either directly or by triggering conformational changes that precede glycolipid uptake. To evaluate further, membrane docking by HET-C2 was analyzed using the orientation of proteins in membranes (OPM) computational approach (57) and by assessing HET-C2 surface hydrophobicity (58, 59). In OPM analyses, optimal rotational and translational positioning of the protein with respect to the lipid bilayer is achieved by minimization of protein transfer energy from water to the membrane hydrocarbon core, approximated as decadiene nonpolar solvent, and to interfacial regions characterized by water permeation profiles. In the approximation, protein binding to the membrane is driven by hydrophobic



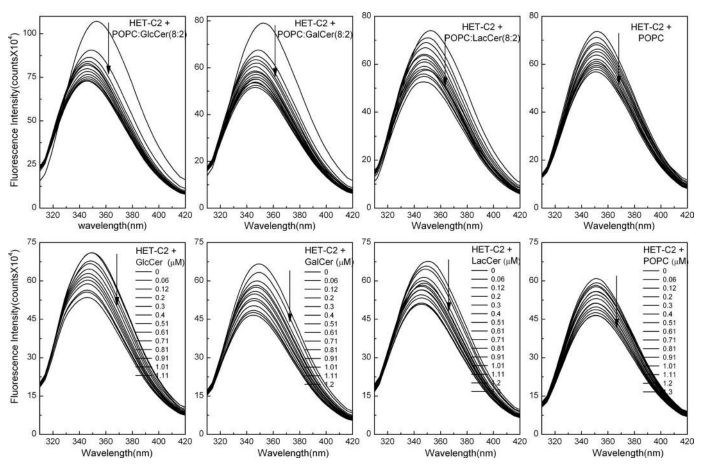


FIGURE 6. **Changes in HET-C2 Trp emission induced by lipids.** *Top*, POPC vesicles containing or lacking glycolipid were introduced in stepwise fashion with 5-min incubation times between injections. The *vertical arrows* indicate increasing vesicle concentration. The first few injections of membrane vesicles containing GalCer or GlcCer (\sim 1.2 μ M total available glycolipid by the third injection) induce a strong intensity decrease and λ_{max} blue shift. Additional injections result in Trp emission intensity decreases of similar magnitude to those observed with POPC vesicles lacking glycolipid. The λ_{max} blue shift occurred only when the vesicles contained glycolipid. *Bottom*, stepwise microinjection of lipids, dissolved in ethanol, was used to mix lipids with HET-C2 with 5 min incubation times between injections. The *vertical arrows* indicate increasing lipid concentration. The starting glycolipid injection and each incremental addition corresponded to \sim 10-fold lower levels than with the vesicle additions (*top*). Fluorescence measurements were performed as described under "Experimental Procedures." Control injections enabled correction for dilution effects.

TABLE 3

Trp fluorescence changes in HET-C2 induced by interaction with membranes containing/lacking glycolipid and by ethanol injection of glycolipid or phospholipid

Each glycolipid derivative contained octanoyl acyl chains. ν , wavenumber.

Glycolipid	Small unilamellar vesicles			Ethanol injection			
	Quenching	Blue shift	$\Delta \nu imes 10^4$	Quenching	Blue shift	$\Delta \nu imes 10^4$	
	%	nm	cm^{-1}	%	nm	cm^{-1}	
GalCer	29	6	0.49	30	5	0.41	
GlcCer	26	7	0.57	25	4	0.33	
LacCer	25	6	0.49	25	6	0.49	
POPC	17	1	0.08	23	0	0	

interactions and opposed by desolvation of polar and charged groups. The three-dimensional structure of apoHET-C2 aided the identification of potential membrane-binding sites, protein tilt angles, and membrane penetration depths. OPM analyses (Fig. 9A) suggest that membrane docking involves Phe¹⁴⁹, Leu¹⁵⁰, Pro¹⁵³, Ile¹⁵⁴, and Ala¹⁵⁷ of helix 6 and Ile⁵⁸ of the adjacent α 1–2 loop, which together form a hydrophobic patch (Fig. 9*B* and supplemental Fig. S7). In contrast, Trp²⁰⁸ and nearby residues do not seem to directly participate in the initial docking of HET-C2 to membranes.

DISCUSSION

HET-C2 Forms a GLTP Fold—Our x-ray diffraction data (1.9 Å) establish that HET-C2 utilizes a GLTP fold to accomplish glycolipid transfer, even though other protein folds (e.g. saposins, nonspecific LTP, and phosphatidylinositol transfer protein) also are capable of binding/transferring glycolipids (14-21). Absolute conservation and homologous positioning are observed for Asp⁶⁶, Asn⁷⁰, Lys⁷³, Trp¹⁰⁹, and His¹⁴⁷ in HET-C2 compared with Asp⁴⁸, Asn⁵², Lys⁵⁵, Trp⁹⁶, and His¹⁴⁰ in GLTP, residues known to form a sugar headgroup recognition center and confer selectivity for glycolipid binding (9, 10) (Figs. 1*B* and 2*A* and supplemental Fig. S2). In the hydrophobic pocket that encapsulates the lipid nonpolar chains, 14 of 24 residues are absolutely conserved, and another nine residues have similar physicochemical features (Fig. 2B). Far-UV CD analyses support GLTP fold formation by HET-C2, indicating a preponderance of helical secondary structure as well as a relatively low unfolding transition temperature midpoint (\sim 49 °C). Similar features are exhibited by porcine and bovine GLTPs (54, 60, 61) as well as by human GLTP.⁶ Our x-ray data



⁶ R. K. Kamlekar, R. Kenoth, S. Y. Venyaminov, and R. E. Brown, unpublished observations.

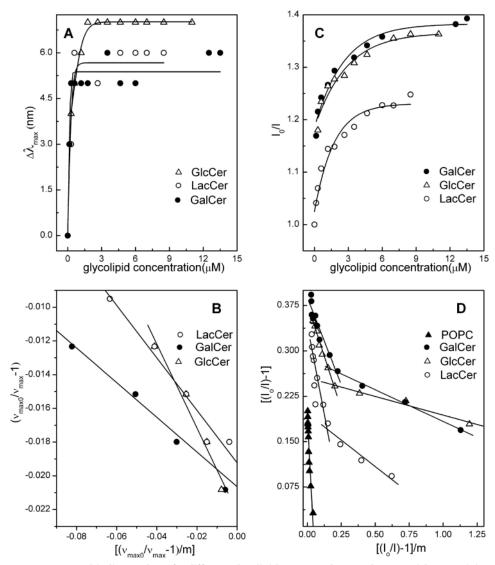


FIGURE 7. **HET-C2 binding analyses for different glycolipids presented as membrane vesicles containing or lacking glycolipid.** *A*, blue shift in Trp emission wavelength maximum ($\Delta \lambda_{max}$) of HET-C2 induced by POPC vesicles containing different glycolipids. *B*, analyses of $\Delta \lambda_{max}$ blue shift data expressed as wavenumber (ν). *C*, intensity (*I*) change in Trp emission of HET-C2 induced by POPC vesicles containing different glycolipids. *D*, analyses of intensity change data. *Subscript 0* indicates values in the absence of added vesicles. Calculations utilized the approach of Bashford *et al.* (47) and were performed as described under "Experimental Procedures."

TABLE 4

HET-C2 binding constants for glycolipids

 K_d values were calculated from changes in emission intensity and from the λ_{\max} blue shift. K_d I is calculated from the emission intensity changes resulting from the addition of vesicles containing glycolipid after correction against vesicles lacking glycolipid. Saturation of glycolipid binding is indicated by cessation of the λ_{\max} blue shift. K_d II represents values calculated from emission intensity changes occurring even after saturation of the λ_{\max} blue shift and corresponding to both glycolipid binding and HET-C2 partitioning to the membrane interface. Note that all of the calculations were performed assuming that only half of the glycolipid in the vesicles is available for binding.

	K_d				
Vesicle composition	Wavelength	Intensity change			
	shift/ $\Delta \nu$	Ι	II		
GlcCer/POPC GalCer/POPC LacCer/POPC POPC	$\begin{array}{c} 0.177 \pm 0.10 \\ 0.108 \pm 0.01 \\ 0.13 \pm 0.04 \end{array}$	$\begin{array}{c} 0.13 \pm 0.10 \\ 0.11 \pm 0.01 \\ 0.41 \pm 0.27 \end{array}$	$\begin{array}{c} 0.92 \pm 0.32 \\ 0.60 \pm 0.03 \\ 1.49 \pm 0.43 \\ 5.26 \pm 1.39 \end{array}$		



rectify discrepancies with previous three-dimensional modeling of HET-C2 regarding identification of key residues of the sugar head group recognition center (11). In our three-dimensional jigsaw model (data not shown), the HET-C2 core region precisely superimposed with the crystal-lographic structure, but a different location emerged for C-terminal residues (Trp²⁰⁸), and no structure was obtained for the first 38 amino acids of the N terminus.

HET-C2 Membrane Interaction and Glycolipid Uptake-Interaction of HET-C2 with PC vesicles containing or lacking glycolipid produces no significant conformational changes in secondary structure as detected by far-UV CD. However, mixing with lipid vesicles does affect the thermal dependence of HET-C2 unfolding, including a slight lowering of the transition temperature midpoint $(1-2 \degree C)$ and increased cooperativity of the unfolding transition (Fig. 4C). The near-UV CD signal, originating from environmentally induced optical activity of Phe, Tyr, and Trp, is largely unaffected by incubation with PC vesicles containing glycolipid, consistent with diffraction data showing only subtle changes in tertiary folding as a consequence of glycolipid binding by GLTP (9, 10, 46). Nonetheless, when the vesicles lack glycolipid, a hint of tertiary change is observed in the near-UV CD signal associated with Tyr and/or Trp, prompting the further study of intrinsic Trp fluorescence in HET-C2.

The involvement of Trp in GLTP

functionality is well established (8, 9, 45, 46, 51-55). In HET-C2, one (Trp¹⁰⁹) of the two Trp residues aligns with GLTP Trp⁹⁶, a key residue for sugar headgroup interaction in the glycolipid binding site. However, the other HET-C2 Trp residue (Trp²⁰⁸) does not align with either of the other two GLTP Trp residues. Fluorescence data show an emission wavelength maximum (λ_{max}) at \sim 355 nm and accessibility to aqueous quenchers, consistent with a relatively polar, average environment for the two Trp residues and localization on the protein surface. The crystallographic data (Figs. 1, 2, and 9 and supplemental Fig. S2) indicate that Trp¹⁰⁹ is well positioned to play a similar role as Trp⁹⁶ within the glycolipid sugar headgroup recognition center of GLTP. Trp²⁰⁸ at the HET-C2 C terminus is surface-accessible but at a different location than GLTP Trp¹⁴², a residue often linked to the GLTP-membrane interaction domain. Trp^{208} remains close to the $\alpha 3$ - $\alpha 4$ loop via a stacking interaction with His¹⁰¹ (Fig. 2A).

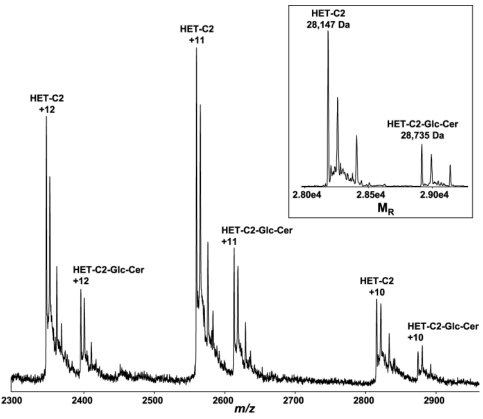


FIGURE 8. **Electrospray ionization mass spectrometry analysis of HET-C2-glycolipid complex.** Three main charge states are observed from the direct infusion of the HET-C2:*N*-octanoyl GlcCer complex (2:1 molar ratio) under nondenaturing conditions. The transformed spectra (see *inset*) result in molecular masses of 28,147 Da for rHET-C2 and 28,735 Da for the rHET-C2-*N*-octanoyl GlcCer complex. The N-terminal His₆-S tag accounted for the molecular mass being higher than wild type HET-C2 (23,179 Da).

With HET-C2, incubation with membrane vesicles containing glycolipid induces substantial decreases (25-30%) in Trp emission intensity and blue shifts in emission λ_{max} (6–7 nm). With human GLTP, glycolipid binding also induces changes in Trp emission fluorescence (i.e. ~40% intensity decrease and ${\sim}12\text{-nm}$ blue shift in $\lambda_{max})$ attributable almost entirely to stacking of the ceramide-linked sugar over Trp⁹⁶ of the sugar headgroup recognition site (46). However, with HET-C2, POPC vesicles lacking glycolipid also elicit significant intensity decreases (\sim 17%), complicating determinations of glycolipid binding affinity based on intensity changes. Fortunately, the blue shifts in emission λ_{max} depend on glycolipids being present in the vesicles. When the sugar headgroup is changed from glucose to galactose or lactose, the dramatic blue shift persists, thus providing a means to calculate glycolipid binding affinity. The resulting K_d values for various glycolipids indicate moderate binding strength by HET-C2, consistent with a role in glycolipid transfer. The K_d values also agree reasonably well with previous values for human GLTP calculated from the Trp intensity decrease. The K_d value for HET-C2 partitioning to POPC membranes is estimated to be \sim 5 μ M, which seems reasonable, considering values reported for other membrane interaction motifs (e.g. pleckstrin domain, range from 1.6 to 2.9 μ M) (62).

A fundamental difference in the HET-C2 Trp response compared with human GLTP is the significant (\sim 17%) decrease in emission intensity induced by mixing with POPC via ethanol microinjection. Similar treatment leaves the GLTP Trp emission intensity nearly unaltered (46), despite the apparent involvement of Trp¹⁴² in the membrane interaction of GLTP. It is noteworthy that Trp¹⁴² of GLTP shows spatial homology with Phe¹⁴⁹ of HET-C2, whereas Trp²⁰⁸ of HET-C2 localizes more similarly to Trp⁸⁵ (Fig. 1C). OPM and surface hydrophobicity analyses support direct involvement of Phe¹⁴⁹ in the initial docking of HET-C2 to the membrane interface (Fig. 9), but this is not the case for Trp²⁰⁸, despite seemingly favorable surface accessibility and positioning near the glycolipid binding site. Collectively, the findings point to the $\alpha 1-2$ loop and helix 6 (including Phe¹⁴⁹) as defining the initial docking site with membranes, whereas the membrane-induced changes in Trp²⁰⁸ fluorescence seem to reflect conformational changes in HET-C2 immediately preceding glycolipid uptake.

Glycolipid Selectivity of HET-C2— The preference of HET-C2 for the two neutral monoglycosylceramides, GlcCer and GalCer, reflects a more narrowly focused selectivity for certain glycolipid types. Inclusion of a second sugar markedly

slows transfer activity, whereas the addition of negatively charged sulfate to GalCer (*i.e.* sulfatide) nearly completely abolishes transfer. Low transfer rates also are observed with ganglioside GM1. It is noteworthy that the glycolipids most preferred by HET-C2 are naturally occurring in filamentous fungi, whereas the others are not (49–51). The more focused glycolipid selectivity of HET-C2 is shared by a plant glycolipid transfer protein recently discovered in Arabidopsis and designated AtGLTP1. West et al. (63) found that AtGLTP1 transfers GlcCer significantly faster than GalCer (~8-fold) and much faster than LacCer (~32.5-fold), whereas SM was not transferred at all. The strong preference of AtGLTP1 for GlcCer is not shared by GLTP or HET-C2, which transfer GlcCer and GalCer at roughly similar rates but faster than AtGLTP1. Modeling of AtGLTP1 indicates a GLTP fold and absolute conservation of residues most critical for glycolipid anchoring and selectivity (Asp⁵², Asn⁵⁶, Arg⁵⁹, Trp⁹⁹, and His¹³⁸ for AtGLTP1 *versus* Asp⁴⁸, Asn⁵², Lys⁵⁵, Trp⁹⁶, and His¹⁴⁰ for GLTP). Thus, to account for the selectivity of GlcCer over GalCer by AtGLTP1, West et al. (63) proposed that the OH4 hydroxyl in glucose, but not in galactose, is sufficiently close to interact with the amine group of Asn⁹⁵ of AtGLTP1. It is noteworthy that negatively charged glutamic acid (Glu¹⁰⁵) in HET-C2 occupies the site of Asn⁹⁵ in AtGLTP1 and of Leu⁹² in human GLTP but that HET-C2, like GLTP, transfers GlcCer as fast or faster than GalCer. It is also noteworthy that all hydrogen bond classes observed in previous protein-carbohydrate interactions (64, 65)

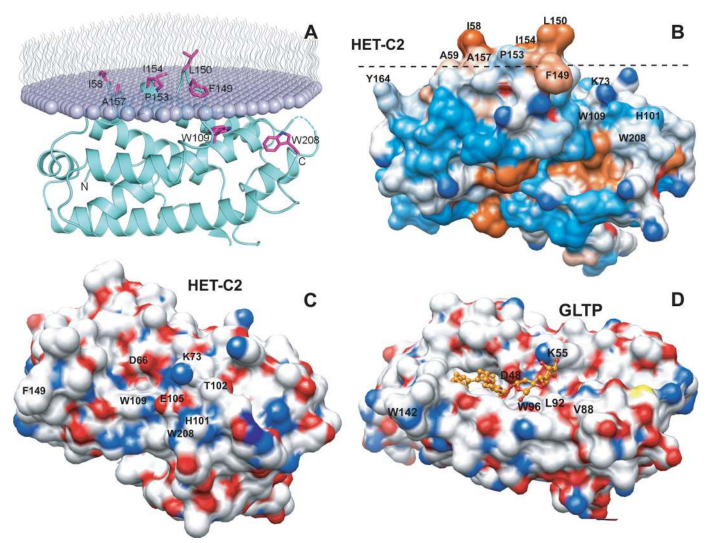


FIGURE 9. **Surface topography of predicted membrane docking site of HET-C2 and sugar headgroup recognition site of HET-C2 and GLTP.** *A*, HET-C2 orientation and positioning during membrane docking. The OPM computational approach (57) was used to identify residues involved in the initial docking of HET-C2 with the membrane interface. The lipid molecules comprising half of the membrane are shown as *lavendar-colored* with *wavy lines* representing the lipid hydrocarbon chains. HET-C2 helices are *cyan-colored*, and important side chain residues are shown in *magenta*. *B*, surface hydrophobicity of HET-C2 (Protein Data Bank code 3KV0). Mapping was performed using Chimera (58, 59), which relies on the Kyte-Doolittle scale to rank amino acid hydrophobicity, with *blue* indicating most hydrophilic, *white* equaling 0.0, and *orange-red* being most hydrophobic. The *dotted black line* corresponds to the membrane interface oriented as shown in *A*. *C*, surface topology of the GSL sugar headgroup binding site of HET-C2, the exogenous dimethyl group added to Lys⁷³ to facilitate HET-C2 crystallization is not shown. Glu¹⁰⁵, Thr¹⁰², and His¹⁰¹ obstruct the surface adjacent to Trp¹⁰⁹. *D*, surface topology of the GSL sugar headgroup binding site of human GLTP. Surface residue *color* reflects charge status (*red*, negative; *blue*, positive; *white*, neutral) for the GLTP-18:1 LacCer complex (Protein Data Bank code 1SX6) (9). 18:1 LacCer is depicted in *gold* with the hydrocarbon chains of ceramide (*left*) disappearing into the hydrophobic tunnel. The open, unobstructed surface adjacent to Trp⁹⁶ allows for broader selectivity for binding of various glycolipids.

are seen when HET-C2 interacts with the GSL sugar headgroup. Among the classes are bidendate hydrogen bonding involving Asn⁷⁰, bifurcated hydrogen bonds involving Asp⁶⁶ (and Glu¹⁰⁵ in the case of glucose), and cooperative hydrogen bond formation involving the multiple sugar hydroxyl groups and amino acid side chains.

Compared with GLTP, the surface region adjacent the sugar recognition center of HET-C2 displays considerably different topology (Fig. 9, *C* and *D*). In GLTP, this region has the appearance of a wide groove that flows unobstructed and openly away from Trp⁹⁶, enabling accommodation of sugar headgroups of varying length and complexity, whereas in HET-C2, residues adjacent to Trp¹⁰⁹ form impediments that constrict this region, effectively generating a surface pit best able to accommodate

glycolipids with single, neutral sugar headgroups. Two structural differences in HET-C2 accounting for the altered surface topology are 1) a shorter C terminus ending with Trp²⁰⁸ and 2) a $\alpha 3 - \alpha 4$ loop shorter by 5 residues (amino acids 96–101) than the $\alpha 3 - \alpha 4$ loop (amino acids 78–88) of GLTP with compositional differences, such as negatively charged Glu¹⁰⁵, His¹⁰¹, and Thr¹⁰² at the same respective locations as Leu⁹², Val⁸⁸, and Gly⁸⁹ in GLTP. Steric hindrance and charge repulsion effects by Glu¹⁰⁵ are likely to impede sulfatide binding. Glu¹⁰⁵ also forms a water-bridged hydrogen bond with His¹⁰¹, helping orient its imidazole ring for stacking (~3.4 Å) against the Trp²⁰⁸ indole ring. Nearby Thr¹⁰² is bulkier than Gly⁸⁹ of GLTP and is $C\beta$ -branched, restricting possible conformations and placing bulkiness near the protein backbone. Thus, the net effect of the



shorter $\alpha 3 - \alpha 4$ and C-terminal loops is formation of the bulky $His^{101} - Trp^{208}$ stack next to Glu^{105} and Thr^{102} , leading to the pitlike morphology of the HET-C2 sugar headgroup recognition center that favors a neutral monoglycosyl headgroup. It is tempting to speculate that the focused selectivity of the HET-C2 GLTP fold for specific glycolipid types reflects evolutionary divergence and adaptation for the simpler glycolipid compositions of filamentous fungi compared with mammals.

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