Architectural Organization of the Metabolic Regulatory Enzyme Ghrelin O-Acyltransferase*

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Background: Ghrelin O-acyltransferase (GOAT) is a membrane protein that is responsible for octanoylating the metabolism-regulating peptide hormone ghrelin.

Results: We have used a combination of approaches to determine the topology of GOAT.

Conclusion: We have shown that GOAT has 11 transmembrane-spanning domains and one reentrant loop.

Significance: These findings serve as a reference for other membrane-bound O-acyltransferase family members.

Ghrelin O-acyltransferase (GOAT) is a polytopic integral membrane protein required for activation of ghrelin, a secreted metabolism-regulating peptide hormone. Although GOAT is a potential therapeutic target for the treatment of obesity and diabetes and plays a key role in other physiologic processes, little is known about its structure or mechanism. GOAT is a member of the membrane-bound O-acyltransferase (MBOAT) family, a group of polytopic integral membrane proteins involved in lipid-biosynthetic and lipid-signaling reactions from prokaryotes to humans. Here we use phylogeny and a variety of bioinformatic tools to predict the topology of GOAT. Using selective permeabilization indirect immunofluorescence microscopy in combination with glycosylation shift immunoblotting, we demonstrate that GOAT contains 11 transmembrane helices and one reentrant loop. Development of the V5Glyc tag, a novel, small, and sensitive dual topology reporter, facilitated these experiments. The MBOAT family invariant residue His-338 is in the ER lumen, consistent with other family members, but conserved Asn-307 is cytosolic, making it unlikely that both are involved in catalysis. Photocross-linking of synthetic ghrelin analogs and inhibitors demonstrates binding to the C-terminal region of GOAT, consistent with a role of His-338 in the active site. This knowledge of GOAT architecture is important for a deeper understanding of the mechanism of GOAT and other MBOATs and could ultimately advance the discovery of selective inhibitors for these enzymes.

GOAT§ is a polytopic integral membrane protein required for the octanoylation of ghrelin, a weight- and glucose-modulating peptide hormone produced primarily in the stomach, upper gastrointestinal tract, and brains of vertebrates (1–3). Octanoylation is required for activation of the ghrelin receptor, GHSR-1a, and is the only known example of protein octanoylation in higher organisms (4). Because dietary fats and those from local fatty acid metabolism can be directly conjugated to ghrelin (5–7), it can be viewed as a “fat sensor” from the brain to the body or within the brain itself, modulating energy balance. Supporting this possibility, inhibition of GOAT in mice prevented weight gain on a high fat diet and improved glucose control (8) and may represent an attractive target for the treatment of obesity and diabetes. Additionally, ghrelin has been implicated in a host of other physiologic processes, including gastric motility (9, 10); learning, memory, and reward behavior (11–13); cardiovascular function (14–16); and survival in extreme starvation (17–19).

GOAT is a member of the membrane-bound O-acyltransferase (MBOAT) superfamily (20), which is composed of many multispanning membrane proteins from prokaryotes and eukaryotes, including 11 in humans. MBOAT members have

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6 The abbreviations used are: GOAT, ghrelin O-acyltransferase; MBOAT, membrane-bound O-acyltransferase; ER, endoplasmic reticulum; TM, transmembrane helix; IBV, infectious bronchitis virus; FC-16, Fos-Choline 16; DTR, dual topology reporter; TEV, tobacco etch virus.
been shown to be critical for lipid biosynthesis, sterol acylation, and acyl modification of secreted proteins, including ghrelin, Hedgehog (HHAT), Wnt proteins (PORC), and yeast GPI-anchored proteins (GUPI) (21–26). MBOAT proteins contain a highly conserved asparagine (Asn-307 in GOAT) and an invariant histidine (His-338 in GOAT) separated by 30 – 45 residues; these residues have been proposed to be catalytic. There are two additional regions of conservation in the family, at the N- and C-terminal boundaries of loop 5 on GOAT, where it adjoins surrounding hydrophobic regions. The topology of a few MBOATs has been analyzed, and the invariant histidine has been shown to be luminal in these cases (27, 28). Based on the modification of luminal, secreted proteins and on luminal location of the conserved histidine, the active site for the MBOAT family has been proposed to be conserved on the luminal face (2, 27).

For several MBOATs, including GOAT, mutation of this conserved histidine eliminates acyl transfer activity (2, 23, 28–31). Detailed studies of catalysis by these purified enzymes is generally hindered by the challenge of solubilization in active conformations (32–34); notable exceptions to this are ACAT1, also known as SOAT1, a robust enzyme that transfers long chain fats from coenzyme A (CoA) to cholesterol and has been extensively characterized (reviewed in Ref. 35), and HHAT, which N-palmitoylates hedgehog proteins (36). In contrast to other MBOAT members, for HHAT, some catalytic activity was retained upon alanine mutation of the conserved histidine (37). Thus, the assignment of this histidine as catalytic is not yet definitive.

GOAT octanoylates serine-3 of proghrelin in the ER lumen after signal peptide cleavage. A range of fatty acids can be processed by GOAT, and the most likely acyl donors are acyl-CoAs (reviewed in Ref. 34). Because fatty acyl-CoAs are known to be cytotoxic (e.g. those produced during fatty acid β-oxidation), the ER membrane represents a barrier that these acyl-CoAs must cross. In this regard, GOAT has been postulated to be a possible acyl-CoA transporter (2). Determination of the topology of GOAT is a critical step to defining its detailed structure and mechanisms. GOAT has been proposed to contain eight transmembrane helices (TMs) (2), but this has not been experimentally investigated.

In this study, we examine GOAT topology in detail, first computationally and then experimentally. Our data suggest that GOAT contains 11 TMs and segregate the invariant His-338 and conserved Asn-307 to opposite sides of the ER membrane. Photocross-linking binding studies map the binding of acyl ghrelin analogs to the C-terminal region of GOAT and demonstrate that purified GOAT can bind ligand as a monomer.

**EXPERIMENTAL PROCEDURES**

All reagents were purchased at the highest quality available from Sigma-Aldrich or Acros Organics unless otherwise indicated. Commercially available reagents were used without further purification.

Bioinformatics—GOAT sequences were identified using BLASTp. Annotation of chicken GOAT was corrected to include the first exon. Dog and green anole (Anolis carolinensis) GOAT sequences were identified by megaBLAST and tBLASTx searches from their draft genomes and annotated. Sequences were aligned using MUSCLE (38). Conserved domains and residues were identified using CDD (39), querying mouse GOAT against MBOAT pfam03062 (20). A battery of topology prediction servers were queried with GOAT sequences from various organisms, starting with mouse and human sequences. A multiple sequence alignment was submitted wherever possible. Older methods, such as TMHMM version 2.0 (40), TMpred (41), SOSUI (42), TopPred (43), and MEMSAT version 3 (44) gave very different results from species to species, even between the highly conserved human and mouse GOATs; the range was from 4 to 11 TMs. The consensus prediction servers TOPCONS (45) and ConPred II (46) both run a number of algorithms, including the above, in order to make better predictions. However, for GOAT, the results from both servers were similarly inconsistent between species and thus hard to interpret. HMMtop (47) and PolyPhobius (48) can use a multiple sequence alignment as input. Although this improved the results somewhat as compared with the aforementioned algorithms, these algorithms predict only seven and eight TMs, respectively. Both missed TMs predicted by MemBrain and MEMSAT-SVM that were subsequently validated experimentally. MemBrain and MEMSAT-SVM predictions were used as a basis for our study and are discussed under “Results.” For MemBrain, predictions for each organism were overlaid on the multiple-species alignment, and TM propensity plots were overlaid and aligned.

Cell Culture—HeLa cells were maintained in DMEM (Invitrogen) with 10% heat-inactivated FBS and penicillin/streptomycin. Transfections were done using FuGene 6 or FuGene HD (Promega) according to the manufacturer’s instructions, with 2 μg of GOAT DNA and 6 μl of reagent per 35-mm dish or well of a 6-well plate. SF9 cells were maintained in ISFM medium, an affordable serum-free synthetic medium, made as reported in (49), with minor modifications.7

Cloning—For human cell expression, GOAT-3xFLAG was cloned into the CAG vector as described previously (8) with a modified multiple-cloning site. N- and C-terminal tags were placed immediately before the first or after the last residue of GOAT without any extraneous sequence by incorporating tag sequences into oligonucleotide primers. Internally V5-tagged GOAT clones were prepared using fusion PCR (50). Individual point mutants were made using a modified QuickChange protocol (Stratagene). Asn-free and Lys-free GOAT were made in one round of polymerization using MISO mutagenesis, as described in Fig. 2a of a previous study (51), as were V5Glyc clones. N-terminal deletion ΔN-H1 starts with a Met at residue 32. ΔN-H1-2, ΔN-H1-3, and ΔN-H1-3 constructs start with Met-56 and ATG codons substituted at positions 81 and 109, respectively, to define the first residue of the constructs. ΔC-H10-11 and ΔC-H11 are truncated after residues 357 and 399, respectively. N-IVB-1-GOAT and N-IVB-2-GOAT are GOAT-3xFLAG with an N-terminal fusion of the first 20 and 10 residues, respectively, of the M glycoprotein from infectious bronchitis virus (IVB; an avian coronavirus). Asn-3 and Asn-6

7 T. E. Cleveland, 4th, M. D. Ward, M. S. Taylor, J. M. Kavran, and D. J. Leahy, manuscript in preparation.
of this sequence are glycosylated in IBV-M, previously called the E1 protein (52). All clones were fully sequence-verified. Baculovirus constructs were made using the Bac2Bac system (Invitrogen) according to the manufacturer’s instructions.

Selective Permeabilization Immunofluorescence—200,000 HeLa cells were plated on glass coverslips, transacted 8–16 h after plating, and incubated for 40 h. For complete permeabilization, cells were fixed in 3% paraformaldehyde for 10 min, and fixative was then quenched using PBS containing 10 mM glycine and 0.2% sodium azide (PBS/gly). Cells were incubated for 3 min in 0.5% Triton X-100 in PBS/gly and then washed twice with PBS/gly. For selective permeabilization, dishes were placed on ice, washed with ice-cold KHFM (20 mM HEPES, pH 7.4, 110 mM potassium acetate, 2 mM magnesium acetate), and kept on ice. The cells were permeabilized with 75 µg/ml digitonin (EMD Biosciences) in KHFM for 10 min. Cells were washed twice with ice-cold KHFM, moved to room temperature, and fixed as above.

Staining with primary and secondary antibodies was carried out for 20 min at room temperature by inverting coverslips onto Parafilm containing 45-µl drops of PBS/gly supplemented with 1% BSA and appropriate antibodies. Antibodies used were mouse anti-FLAG M2 (1:500; Sigma), rabbit anti-FLAG (1:1000; Sigma F7425), mouse anti-V5 (1:5000; Invitrogen), mouse anti-Myc 9E10 (1 µg/ml), rabbit anti-GFP (1:500; Molecular Probes), mouse anti-GFP (1:500; Roche Applied Science 11814460001), and rabbit anti-SSR1 (2 µg/ml; Sigma HPA011276; recognizes residues 230–286). Coverslips were washed and incubated analogously with secondary antibodies: Alexa Fluor® 488-conjugated anti-mouse IgG (1:1000) and Alexa Fluor 568-conjugated anti-rabbit IgG (1:1000). As controls, untransfected cells were stained identically, and primary antibodies were omitted. DNA was stained prior to imaging with Hoechst 33285 (0.1 µg/ml, 5 min), and coverslips were mounted using glycerol with 100 mM N-propyl gallate. Epifluorescent images were collected at room temperature using an Axioscop microscope with ×63 and ×40 objectives (Zeiss, Jena, Germany) equipped for epifluorescence using an ORCA-03G CCD camera (Hamamatsu, Japan) and iVision software (BioVision). For equal comparison of staining between permeabilizations, within each construct, for each antibody, an optimal exposure time was first determined, and then this fixed exposure time was used for both permeabilizations. Signal intensity was then assigned based on the optimal V5 exposure times as follows: ++ + +++, <30-ms exposure; + + +, 30–60-ms exposure; +++, 60–150-ms exposure; +++, 150–500-ms exposure; −, not clearly above background. Image layouts were done in Adobe Photoshop CS6 and Illustrator CS6.

For co-localization of GOAT with CFP-KDEL, coverslips were treated as above except that fixation was in methanol for 20 min at −20 °C. Confocal images were acquired with a Cascade QuantEM 512SC camera (Photometrics) attached to a Zeiss AxioImager (×63 objective) with a Yokogawa spinning disc confocal scanner and Slidebook software (Intelligent Imaging Innovations).

Synthesis—Peptide synthesis was performed using the Fmoc (N-(9-fluorenylmethoxycarbonyl) strategy. GO-CoA-Tat, GO-Tat, and GO-CoA-Tat F4BP were prepared as described previously (8). In brief, amide-linked ghrelin analogs were prepared in which Ser-3 in ghrelin was replaced with Alloc (allyloxy-carbonyl) protected 1,2-diaminopropionic acid, orthogonally deprotected using palladium catalysis, and then reacted with octanoic acid in the presence of HATU (1-(bis-(dimethyl-amino)methyl)imiumyl)-1H,1,2,3-triazolo(4,5-b)pyridine-3-oxide hexafluorophosphate), reacted with octanoic anhydride, or reacted with derivatives thereof. For ghrelin-28-Oct-diazirine, 7,7’-diazooctanoic acid was used at this step. This was prepared by using the method described in the literature (53), and its structure and purity were confirmed by 1H NMR. Synthesized peptides were purified using a reversed-phase C-18 column, and their MALDI and ES mass spectrometry data were consistent with the calculated values. The final concentrations of the compounds in aqueous solution for assay were determined by amino acid analysis. Ghrelin sequences synthesized correspond to human ghrelin (GSFSLSPHEQVRQSQKESKKPAKLPQPR).

GOAT Purification—Mouse GOAT with a C-terminal TEV-3xFLAG tag was expressed in SF9 cells at 3–6 ×10⁶/ml in 1–6-liter aerated spinner flasks using a 72-h baculoviral infection with a multiplicity of infection of ~10. Cells were harvested, and all subsequent steps carried out on ice or at 4 °C. Lysis was performed using a microfluidizer in 4 packed cell volumes of HBS + PI (50 mM HEPES, pH 7.0, 150 mM NaCl, 2 µg/ml aprotinin, 2.5 µg/ml leupeptin, 2 µg/ml pepstatin A, and 1 mM EDTA). Lysate was cleared for 10 min at 15,000 × g, and then microsomes were collected at 120,000 × g for 2–6 h, flash-frozen on liquid nitrogen, and stored at −80 °C. Microsomes were resuspended in 10 volumes of HBS + PI in a 40-ml Dounce homogenizer, solubilized for 1 h at 4 °C with 1% Fos-Choline 16 (FC-16; Anartrase), and cleared for 30 min at 100,000 × g. Supernatant was bound to pre-equilibrated FLAG-agarose in batch for 1–3 h and then transferred to a column, washed with HBS + PI + 0.05% FC-16 and with the same buffer containing 500 mM NaCl, and then washed with HBS without protease inhibitors and cleaved overnight with His6-TEV protease in 10 mM HEPES, pH 8.0, 100 mM NaCl, 0.005% FC-16, 2 mM DTT. Supernatant was collected, 20 mM imidazole was added, and then His6-TEV protease was removed using nickel affinity. The sample was then further purified by ion exchange chromatography (HiTrap SP FF, GE Healthcare), eluted in the same buffer containing 300 mM NaCl without imidazole, and concentrated to 2–10 mg/ml.

MALDI-TOF Mass Spectrometry—MALDI-TOF mass spectrometry of intact GOAT was performed by Tatiana Boronina in the Johns Hopkins Proteomics Core Facility using a Voyager DE-STR (Applied Biosystems). 3 µg of purified, tag-cleaved mouse GOAT was desalted using PLRP-S 300A/10–15-µm material (Polymer Labs/Agilent) packed in a pipette tip pre-equilibrated with 12% acetonitrile and 0.1% TFA; washed three times in the same buffer; and eluted with 90% acetonitrile, 0.1% TFA. 2 pmol of GOAT was spotted in a sinapinic acid matrix (Laser BioLabs) in either 50% ethanol, 0.1% TFA or 40% acetonitrile, 0.1% TFA. Calibration was with AB SCIEX C3 standards (5–17 kDa) in adjacent spots under identical conditions.

Analytical Ultracentrifugation—Sedimentation velocity analytical ultracentrifugation was done on a Beckman XL-I at 10 °C with an An-60Ti rotor at 40,000 rpm with a total run time of 12.5 h. Purified tag-cleaved GOAT samples in a buffer contain-
Topology of Ghrelin O-Acyltransferase

ing 10 mM HEPES, pH 7.0, 300 mM NaCl, and 0.0008% FC-16 were diluted to 0.8, 0.3, and 0.1 A280, corresponding to 6.8, 2.6, and 0.9 μM. Data shown (see below) are from the highest concentration; results from other cells were similar. Analysis was performed as described under “Results.”

**Competition Photocross-linking—** Each 200-μl reaction mixture in HBS + PI + 0.005% FC-16 contained 0.2–2.0 μM purified GOAT and 1 μM photocross-linking compound with or without 200 μM competitor. Competitors were preincubated for 1 h, and photocross-linkers were preincubated for 15 min at 4 °C. Reactions were cross-linked with a UV lamp with a 360-nm filter in a stirred quartz cuvette, water-cooled to 4 °C using a circulating bath. GO-CoA-Tat F4BP was exposed for 10 min, ghrelin-28-Oct-F4BP for 2 min, and ghrelin-28-Oct-diazirine for 15 min. Samples were then analyzed on SDS-PAGE, blotted for biotin (as above), and stained with colloidal Coomassie Brilliant Blue (Invitrogen) as a loading control.

**Microsomal GOAT Octanoyltransferase Assay—** SF9 microsomes containing GOAT-3xFLAG and control microsomes from empty vector SF9 cells were prepared as above except that lysis was performed using a Dounce homogenizer. Assays were carried out as described (2, 34) with modifications. 25 μg of SF9 microsomes (BCA) in 50 μl of HBS (pH 7.0) was incubated at 30 °C for 5 min with 10 μM C-terminally biotin-tagged human ghrelin (ghrelin-27-biotin), 60 μM palmitoyl-CoA, and 1 μM [3H]octanoyl-CoA (high specific activity; American Radiolabeled Chemicals). Reactions were quenched and solubilized by adding 1 ml of 2% SDS in TBS containing streptavidin beads and bound for >15 min. Beads were washed with 25 ml of TBS + 0.1% SDS on small columns and then scintillation-counted.

**Microsomal Photocross-linking and Partial Proteolysis—** Each 1200-μl cross-linking reaction contained 5 μM GO-CoA-Tat F4BP and 5 mg of total microsome protein from either GOAT-3xFLAG containing microsomes or control microsomes, which were made using empty virus-infected SF9 cells. Reactions were carried out as described (8), solubilized by diluting to 1.5 mg/ml with 1% FC-16 for 30 min at 4 °C, and then cleared for 30 min at 100,000 × g. 0.4–40 μg sequencing grade modified trypsin (Promega) or buffer was added to each sample and incubated for 15 min at 37 °C. Protease was quenched with 20 μg/ml aprotinin and 20 μg/ml leupeptin, and samples were bound to pre-equilibrated anti-FLAG-agarose (Sigma) overnight. Beads were washed three times with HBS + PI with 0.1% FC-16 and eluted with 1 mg/ml 3xFLAG peptide (Sigma) in the same buffer and then separated with SDS-PAGE. Biotin detection was used streptavidin (Pierce) followed by polyclonal anti-streptavidin (Abcam).

**RESULTS**

**Bioinformatic Predictions of GOAT Topology—** To predict the membrane topology of GOAT, we started with the assumption that topology should be conserved across species. 16 GOAT sequences were identified and, in the cases of dog, green anole, and chicken, annotated. A battery of Web-based topology prediction servers were then tested, but most individual algorithms and consensus prediction servers gave very different results from species to species, even between the highly conserved human and mouse GOATs; the range was from 4 to 11 TMs (see “Experimental Procedures”).

In contrast, two more recently developed programs, MemBrain (54) and MEMSAT-SVM (55), stood out in their consistency of results across species. MemBrain and MEMSAT-SVM incorporate newer machine-learning algorithms, phylogeny and PSI-BLAST strategies, and current knowledge about membrane protein structure, such as short and very long helices (<16 or >40 residues) and reentrant loops (also referred to as half-TMs or half-helices). Results from both programs were nearly identical, including predicted helical boundaries, except where described below. MemBrain-predicted TMs for each species are highlighted in Fig. 1, with a total of 12 candidate TMs identified in consensus. For clarity, we numbered TMs and loops in accordance with our final model. Corresponding TM propensities for each species are overlaid in Fig. 2A. Some species are predicted individually to be 11 or 13 TMs. Note that TM-1, -5, -8, and -9 were predicted to be either a TM or a reentrant loop, depending on species. TM-7 is predicted to be one long TM by MemBrain in 13 of 16 species, but in dogs, horses, and green anoles, it predicts this region as one TM and one reentrant loop.

TM-8 and TM-9 are predicted to be two separate TMs by MemBrain in 14 of 16 species, with the 2–3-residue turn between the TMs containing MBOAT invariant His-338 at the border of TM-9. Our data (see below) suggest that this prediction is correct. In contrast, MemBrain predicts one long TM in this region for rabbits and zebrafish. MEMSAT-SVM predicts this region as one long TM in all cases. No signal peptide is predicted for GOAT by MemBrain, MEMSAT-SVM, PolyPhobius, or SignalP (56). MemBrain does not predict orientation for helices, but MEMSAT-SVM predicts that the GOAT N terminus is luminal and that its C terminus is cytosolic.

We compared the new computational results with those obtained using a Kyte and Doolittle scanning hydrophy window (Fig. 2B). As reported previously (2), this method predicts eight TMs. In comparison, the consensus MemBrain results predict 12 TMs (Fig. 2, A and B (inset bar)).

**Topology Determination Using Selective Permeabilization—** To experimentally determine the topology of GOAT, we first verified its predicted localization to the ER by co-transfection with CFP-KDEL, an ER marker containing a strong, cleaved signal peptide and the KDEL retention sequence. Upon expression, GOAT-3xFLAG and CFP-KDEL colocalize (Fig. 3A).

To test the topology predictions, we designed GOAT constructs containing N-terminal and C-terminal tags. In addition, GOAT constructs were generated with internal epitope tags inserted in the loops between candidate TMs, aligning to gaps where possible (Fig. 1, red bars). Because the boundary between TM-10 and -11 was not clearly defined across species, we created two positions, 11a and 11b. Position 8b was added later due to masking of V5 epitopes installed at 8a (see below), and positions 8c and 8d were added with V5Glyc tags (below). We next screened FLAG, 3xFLAG, and V5 tags for compatibility with proper translation of the GOAT N terminus, which contains a non-cleaved internal signal sequence (validated below). The V5 tag was selected for testing because its net charge is neutral. Staining of an N-V5 tag was indistinguishable from C-terminal
### FIGURE 1. Bioinformatic analysis of GOAT sequences identifies 12 candidate TMs. Shown is GOAT multiple sequence alignment across 16 species with MemBrain-predicted candidate TMs highlighted in yellow; for clarity, TMs and loops are numbered in accordance with our final model. *, predicted to be either a full-length TM or a reentrant loop; **, TM-7 was predicted to be either one long TM or a reentrant loop plus one standard-length TM. c6-RL, candidate 6 reentrant loop; this region was found to be non-transmembrane, probably a reentrant loop. Highly conserved Asn-307 and invariant His-338 residues are highlighted in green; in every case, His-338 is predicted to be part of TM-9 (green / yellow highlight). Additional highly conserved MBOAT residues (NCBI conserved domain database pfam03062, 1.5 bits (39)) are highlighted in green in the top bar. Positions selected for epitope tag insertion are shown as red bars, and their localizations experimentally determined in this work are listed above. GOAT protein sequences were aligned using MUSCLE (38). Dog and green anole sequences were identified and annotated from their genomes; chicken annotation was corrected to include the first exon (which contains TM-1 and -2).

![Alignment Diagram]

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### Table: GOAT Multiple Sequence Alignment

| Species      | 1 | 2 | 3 | 4 |
|--------------|---|---|---|---|
| **Mouse**    |   |   |   |   |
| **Rat**      |   |   |   |   |
| **Rabbit**   |   |   |   |   |
| **Dog**      |   |   |   |   |
| **Wild-Boar**|   |   |   |   |
| **Human**    |   |   |   |   |
| **Chimpanzee**|   |   |   |   |
| **Rhesus-Mac.**|   |   |   |   |
| **Panda**    |   |   |   |   |
| **Marmoset** |   |   |   |   |
| **Horse**    |   |   |   |   |
| **Cow**      |   |   |   |   |
| **Green Anole**|   |   |   |   |
| **Chicken**  |   |   |   |   |
| **Sebrafish**|   |   |   |   |

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**N-terminus: Lumen**

**1: Cytosol**

**2: Lumen**

**3: Cytosol**

**4: Lumen**

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**5: Cytosol**

**6: Cytosol**

**7: Lumen**

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**8a: Cytosol**

**8b: Cytosol**

**8c: Cytosol**

**9: Lumen**

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**10: Cytosol**

**11a: Lumen**

**11b: Lumen**

**C-terminus: Cytosol**

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*TM-1* | *TM-2* | *TM-3* | *TM-4* | *TM-5* | *c6-RL* | *TM-6* | *TM-7* | *TM-8* | *TM-9* | *TM-10* | *TM-11* | --
3xFLAG-tagged GOAT, but N-FLAG and N-3xFLAG tags resulted in poor expression and aggregated protein (not shown).

Mouse GOAT clones were then made with various internal and terminal epitope tags. The V5 tag was installed at the N and C termini and at all internal positions except for 8c and 8d (Fig. 1, red bars). To ensure that the internal tag did not perturb overall topology, “flipping” the C terminus to the opposite side of the membrane, all clones contain a constant C-terminal 3xFLAG tag (except when C-terminally V5- or 3xMyc-tagged, as noted). We then investigated the topology of these clones using indirect immunofluorescence with selective permeabilization of the plasma membrane. Parallel wells were transfected with constructs of interest and either fully permeabilized with Triton X-100 or selectively permeabilized at the plasma membrane with digitonin (Fig. 3B). To verify the success of the permeabilization, in each experiment, we employed control proteins of known localization in parallel:

**FIGURE 2. Comparison of new versus prior GOAT topology models.** A, aligned overlay of MemBrain helical propensities by position for GOAT from 16 species. A darker color indicates a more consistent prediction. Candidate TMs are labeled c1–c12 at the top. †, candidate TM-6 was found to be non-TM. B, Kyte and Doolittle mean hydrophobicity profile for mouse GOAT, with an 18-residue scan window. *, TMs predicted by this method are shown above the hydrophobic peaks. Our proposed topology of GOAT is shown in the thick bar at the bottom of the graph, with TMs in black, luminal loops in green, cytosolic loops in pink, and a possible reentrant loop in yellow.
FIGURE 3. Mapping the topology of GOAT by selective permeabilization of the plasma membrane and indirect immunofluorescence. 

A, mouse GOAT bearing a C-terminal 3xFLAG tag is localized to the ER in HeLa cells, as seen by co-localization with co-transfected CFP-KDEL, an ER marker; images are a confocal projection. 

B, selective permeabilization experiments mapping the positions of internal V5 epitopes. Two dishes of HeLa cells were transfected with a GOAT cDNA bearing an internal V5 or Myc epitope tag; all constructs have a C-terminal 3xFLAG tag as an internal control. Selective permeabilization with digitonin revealed cytosolic but not lumenal epitopes, whereas full permeabilization with Triton X-100 (TX-100) revealed all accessible epitopes. Lumenal epitopes were thus visible with Triton X-100 only. 

As a control, two dishes were stained for proteins with known cytosolic and lumenal epitopes and permeabilized identically; in this case, we used endogenous SSR1 (cytosolic) and transfected CFP-KDEL (lumenal). Identical exposures and image normalization for both permeabilizations ensure fair side-by-side comparison. 

C, V5 signal strength in indirect immunofluorescence of V5-tagged constructs from Figs. 3B and 4D (see “Experimental Procedures”).
transfected lumenal CFP-KDEL and endogenous SSR1, an ER transmembrane protein with a cytosolic epitope. Both proteins were detected with Triton X-100 permeabilization, but only the cytosolic SSR1 epitope was detected with digitonin permeabilization.

The GOAT C terminus was consistently localized in the cytosol in all constructs, with the 3xFLAG tag readily detectable (Fig. 3B). The ability to detect the V5 tag varied from construct to construct (relative signal strength is shown in Fig. 3C), but overall, the technique was successful and allowed determination of topology at most positions. Notably, N and C termini were found to be on opposite sides of the membrane, requiring an odd number of TMs. Both positions 5 and 6 were found to be cytosolic, suggesting that the strongly predicted, hydrophobic c6 does not cross the membrane. We suggest that this segment represents a reentrant loop.

The C-terminal part of the protein was more challenging, with the V5 tag undetectable at position 8a, 8b, 9, or 11b. This was probably due to masking of the epitope rather than an expression problem, because the internally tagged proteins were expressed well, as revealed by C-terminal 3xFLAG staining (not shown). We therefore installed Myc tags in these positions as well as in positions 2, 4, 10, and 11a. The Myc tag was strongly detected at positions 8a and 8b, with clear cytosolic localization (Figs. 3B and 4D). Note that this localizes the conserved MBOAT fingerprint residue Asn-307 to the cytosol. Neither V5 nor Myc tag was visible at position 9 or 11b; the proteins were expressed, as seen by 3xFLAG staining, but these epitopes may have been masked in the context of the short loops. Myc tags at positions 2, 4, and 11a all gave results consistent with V5 tags at these positions but with a less robust signal/noise ratio. Myc signal from position 10 was not detectable.

To confirm the cytosolic localizations of positions 5 and 8, we made parallel constructs in human GOAT, repeated the selective permeabilizations (Fig. 4A), and expressed them in SF9 cells for activity assays (Figs. 4B and C). We also made two C-terminal deletion mutants, ΔC-H11 and ΔC-H10–11, predicted to lack the final TM and final two TMs, respectively; all
transf erase and a distance from the membrane of ~12–14 amino acids (30–40 Å), corresponding to the distance of the oligosaccharyltransferase active site from the membrane (57). This is typically achieved by fusing a complete, folded protein (or domain) to the membrane protein of interest, often with a C-terminal truncation. For example, in yeast, the SUC2-HIS4C DTR, installed as a C-terminal fusion with truncation, has been used extensively to map the topology of a number of proteins, including other MBOATs. This DTR is reliable when used under particular conditions and has provided much insight, but it can be problematic when positioned too close to the membrane (27, 58–60). Additionally, this DTR is a very large modification of ~130 kDa, much larger than the proteins of interest here, and it is inconvenient to clone in-frame without using yeast homologous recombination. Fusion of the SUC2-HIS4C to C termini (without truncation) has been shown to cause incorrect C-terminal localization in some cases (61, 62). Mapping internal sites by C-terminally truncating a membrane protein, as the tag is typically used, is likely to be of low reliability for a number of reasons. C-terminal truncations alone may result in major structural perturbations, resulting in the incorrect topology determination in the MBOATs ACAT1 and ACAT2 (35). Moreover, a very large protein insertion can further perturb the overall topology. In addition, the large size of the DTR combined with truncations of various lengths makes it challenging to verify that the fusion proteins are full-length.

We hypothesized that a much smaller, flexible hairpin-like DTR consisting of two V5 epitope tags flanking a strong consensus glycosylation site could provide a consistent context for oligosaccharyltransferase, increase sensitivity for immunostaining and immunoblotting, and minimize epitope masking while minimizing protein perturbation by retaining a short sequence. It could also be readily installed in-frame in the context of a full-length protein of interest.

To create a strong consensus glycosylation site, we queried the NetNGlyc server, a neural network-based N-glycosylation prediction tool,8 with candidate synthetic V5Glyc sequences in the centers of lumenal loops 2 and 4 on GOAT. We started with NXS and NXT motifs and found that NXS was at best weakly predicted to be glycosylated (38–53% confidence), but NXT was stronger, with NVT optimal but still only with 62% confidence (7 of 9 “jury” agreement). We iteratively tested a series of additional residues on either side, resulting in the sequence GYLNVTYY as a strong agreement (83 and 85% confidence when the sequence was installed in GOAT in positions 2 and 4, respectively, with 9 of 9 jury agreeing). The resulting V5Glyc tag is 36 amino acids long, and its complete sequence is GKPIPNPLLGLDST-GYLNVTYV-GKPIPNPLLGLDST (where underlining indicates the glycosylated Asn and hyphens separate the V5 sequences from the Glyc sequence).

We then installed the V5Glyc tag at all previously referenced positions in mouse GOAT and used it to probe topology by gel shift immunoblotting (Fig. 6, A–D) and selective permeabilization immunofluorescence microscopy (selected positions shown in Fig. 6E). The N terminus of GOAT was luminal with complete glycosyl-

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8 R. Gupta, E. Jung, and S. Brunak, manuscript in preparation.
TABLE 1
GOAT topology summary

| Position | V5/Myc Sel. Perm | Glyc | V5Glyc Sel. Perm | Other |
|----------|------------------|------|-----------------|-------|
| N-Term   | L+++             | L+++ | L+++            | All gly* |
| 1        | C+++             | C+++ | C+++            | Endogenous Asn** |
| 2        | L+               | L    | L+              |       |
| 3        | C+++             | C+++ | C+++            |       |
| 4        | L+               | L    | L+              |       |
| 5        | C+++             | C    | C+++            | K141 Ubq†, Mhc bGOAT, Catalytically active NXS/T Mutant*** |
| 6        | C+++             | C    | C+++            | NXS/T Mutant*** |
| 7        | L+++             | L+++ | L+++            |       |
| 8a       | C+++             | ND   | ND              | Mhc bGOAT, Three NXS/T Mutants in loop 8** |
| 8b       | C+++             | ND   | ND              |       |
| 8c       | ND               | ND   | C               |       |
| 8d       | ND               | ND   | C               |       |
| 9        | L                | L    | L+              |       |
| 10       | C++              | C    | C++             | Endogenous Asn** |
| 11a      | L                | L    | L+              |       |
| 11b      | −                | −    | L               |       |
| C-Term   | C+++             | C    | C+++            | C-3xFlag cyto in all constructs. C-3xMyc also cyto. |

*Experimental Procedures.*

*The banding pattern also did not change in response to hydroxyamine and hydrazine treatment of GOAT (Fig. 8F), suggesting that multiple banding did not arise from fatty acid esterification.*

*We were concerned that the two bands might represent a gel artifact because, like many other integral membrane proteins, GOAT aggregates when heated in SDS and therefore may not be fully denatured; it also runs at a smaller molecular weight than those in D. The anti-FLAG blot in D shows an additional band present (gray arrow) not seen in the anti-V5 blot for the N-terminal construct and position 1; this band is shown below to represent a protein starting with Met-56. Note that positions 8a and 8b are not shown; no full-length GOAT could be detected from these constructs. In C, two additional positions were tested in loop 8 (8c and 8d), and both expressed well and confirmed cytosolic location. E, selective permeabilization of key V5Glyc constructs reports lumenal location for challenging positions 9 and 11 and cytosolic location of loop 8; other previously suggested positions are also confirmed. As above, GOAT bearing the indicated internal V5Glyc tag was used, and all constructs have a constant C-terminal 3xFLAG tag. The N terminus and positions 5 and 7 are shown because they were the most robust in our initial survey. F, NXS or NXT codons (not V5Glyc tags) were installed in all loops longer than 24 amino acids: loops 5, 6, and 8. Loop 8 is longer, so three constructs were generated. None of these epitopes could be glycosylated, consistent with cytosolic locations.*

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*heLa cells. Mouse GOAT constructs with internal or terminal V5Glyc tags and a constant C-terminal 3xFLAG tag were lysed 20 h after transfection and treated with peptide-N-glycosidase F (PNGaseF) or mock-treated in identical buffer and then subjected to anti-V5 immunoblotting. Only lumenal positions can be glycosylated, and peptide-N-glycosidase F treatment cleaves off the oligosaccharide. Constructs in positions 8 and 8d were not detected from these constructs. In C, two additional positions were tested in loop 8 (8c and 8d), and both expressed well and confirmed cytosolic location. E, selective permeabilization of key V5Glyc constructs reports lumenal location for challenging positions 9 and 11 and cytosolic location of loop 8; other previously suggested positions are also confirmed. As above, GOAT bearing the indicated internal V5Glyc tag was used, and all constructs have a constant C-terminal 3xFLAG tag. The N terminus and positions 5 and 7 are shown because they were the most robust in our initial survey. F, NXS or NXT codons (not V5Glyc tags) were installed in all loops longer than 24 amino acids: loops 5, 6, and 8. Loop 8 is longer, so three constructs were generated. None of these epitopes could be glycosylated, consistent with cytosolic locations.*

**Topology of Ghrelin O-Acyltransferase**

FIGURE 6. V5Glyc tag allows mapping of GOAT topology using two different techniques. A–D, glycosylation gel shift assay for localization in transfected HeLa cells. Mouse GOAT constructs with internal or terminal V5Glyc tags and a constant C-terminal 3xFLAG tag were lysed 20 h after transfection and treated with peptide-N-glycosidase F (PNGaseF) or mock-treated in identical buffer and then subjected to anti-V5 immunoblotting. Only lumenal positions can be glycosylated, and peptide-N-glycosidase F treatment cleaves off the oligosaccharide. Constructs in positions 8 and 8d were not detected from these constructs. In C, two additional positions were tested in loop 8 (8c and 8d), and both expressed well and confirmed cytosolic location. E, selective permeabilization of key V5Glyc constructs reports lumenal location for challenging positions 9 and 11 and cytosolic location of loop 8; other previously suggested positions are also confirmed. As above, GOAT bearing the indicated internal V5Glyc tag was used, and all constructs have a constant C-terminal 3xFLAG tag. The N terminus and positions 5 and 7 are shown because they were the most robust in our initial survey. F, NXS or NXT codons (not V5Glyc tags) were installed in all loops longer than 24 amino acids: loops 5, 6, and 8. Loop 8 is longer, so three constructs were generated. None of these epitopes could be glycosylated, consistent with cytosolic locations.*

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using this technique, suggesting that they are distinctly covalent and non-interconverting structures.

We analyzed purified mouse GOAT in FC-16 detergent micelles by size exclusion chromatography. Under these conditions, the solution contains a mixture of detergent micelles containing GOAT and empty detergent micelles. The empty micelles are \( \sim 72.4 \text{ kDa} \) for FC-16, with an aggregation number of 178 detergent molecules/micelle (Anatrace). TEV-cleaved GOAT has a molecular mass of 50.6 kDa; as a monomer or dimer in micelles, the predicted mass would be \( \sim 123 \) or \( \sim 174 \) kDa, respectively. In size exclusion chromatography, GOAT ran as one major peak (Fig. 8C). As compared with standard (soluble) proteins, the size exclusion chromatography-calculated molecular mass for GOAT in its FC-16 micelle is 140.7 kDa. This size does not allow clear determination of whether purified GOAT is a monomer or dimer, although it is closer to a monomer position. Hydrophobic interactions between the GOAT protein and resin may result in non-ideal behavior. Note that the detergent does not result in detectable UV absorbance at these concentrations, as seen by the lack of a peak at \( \sim 70 \text{ kDa} \) corresponding to free detergent micelles.

To further investigate the quaternary structure of GOAT, we employed sedimentation velocity analytical ultracentrifugation. As shown in Fig. 8D, sedimentation velocity data using interference optical monitoring of GOAT were fit in SEDFIT to a C(s) model (65). Approximately 90% of the signal in the sample fit to a single peak at 1.9 S, which was normalized to 5.4 S when taking into account the temperature correction factor (20 °C, water). This sedimentation behavior corresponds to an approximate molecular mass of 110 kDa, consistent with monomeric GOAT in FC-16. Although the peak was slightly asymmetric, we were unable to resolve additional, smaller species. UV absorbance data (not shown) showed similar results. Fitting the data using the program dC/dT +, which employs alternative methods of fitting data to those used by SEDFIT, calculated a similar \( S \) value and identified one major species in the experiment (66).

We next analyzed intact, purified GOAT by MALDI-TOF mass spectrometry (Fig. 9A). In addition to the major GOAT peak at the expected size, a minor species with a mass of 44,428 Da was found, consistent with a protein initiated at Met-56. Further support that this 44,428 Da peak corresponds to the lower band in SDS-PAGE can be found in its altered glycosylation pattern. The addition of various glycosylation sequences to the N terminus of GOAT (Figs. 6D and 9B), caused a size increase in the upper but not the lower band, and only the top band shifted further with glycosylation, suggesting that the faster migrating species initiates downstream of the first ATG.

Because Met-56 is predicted to be the last (luminal) residue in TM-2, recovery of the lower band suggests that GOAT lacking its first two TMs might be stable. This may also be a physiologic product, and we note that Met-56 is the first ATG in the second exon of GOAT. We therefore made constructs for baculoviral expression from which the first, first two, first three, and first four helices had been removed (Fig. 9C). The size of the construct from which the first two helices have been deleted (AN-H1-2; start codon is Met-56) co-migrated with the lower band in the full-length construct, consistent with our hypothesis. However, none of the deletion mutants were active (Fig. 9E). We note that although human GOAT expressed at levels lower than mouse GOAT, it had similar activity. The two different human GOAT clones have Thr or Ala at position 46; we made both after noticing that the conserved Ala residue has also been reported as Thr (2). Indeed, a recorded human coding SNP specifies this polymorphism (rs7813902). Both clones performed identically in our assays.
To further confirm that Met-56 can act as an alternate start codon, we mutated this residue. Met-56 is conserved in all GOAT sequences evaluated except Green Anole, where it is Ile; we therefore made both M56A and M56I mutants in three different C-3xFLAG-tagged GOAT constructs and expressed them in HeLa cells (Fig. 9). The lower band was no longer detectable in any of these mutants. We note that the expression of the mutant proteins was lower than that of WT, and we loaded 5-fold more total protein in those lanes. The N-IBV1 construct is shown to calibrate the lower band position.

Additional Data Supporting the Model—Mouse GOAT is not N-glycosylated (Figs. 8E and 9B). However, two Asn residues are predicted to be candidates for glycosylation by the
NetNGlyc server; that neither can be glycosylated is consistent with their cytosolic localizations in our model. Asn-27 (79% and 9 of 9 jury, rating) is at the cytosolic border of TM-1, and Asn-366 (62% and 9 of 9 jury, rating) is in the middle of the cytosolic loop between TM-9 and -10. Conserved Asn-307 is in an NXXS motif but is both cytosolic and not predicted to be glycosylated (48%, 7 of 9 jury, rating).

Mouse GOAT has only three loops where central installation of NXXS or NXXT motifs by point mutations would also allow sufficient spacing from the membrane for potential glycosylation by oligosaccharyltransferase (12 residues on each side of the Asn, or 24 residues total). In loops 5 and 6, we installed sites on both sides of the reentrant helix (I149N and Q191N, NRS in both cases). These were predicted to be glycosylated by NetNGlyc (74 and 68%, respectively, both with 9 of 9 jury and + rating). Neither mutant GOAT could be glycosylated (Fig. 6F).

We installed three more potential sites in loop 8, with mutants I290N, L293N, and R298N making NWT, NET, and NIS sequences (prediction 51, 58, and 67%, respectively, jury 6 of 9, 5 of 9, and 8 of 9, respectively, + rating). These positions were not glycosylated (Fig. 6F).

Loop 10 (20 residues) already contains Asn-366, which is centrally located and in a non-standard but putative NXXC motif, so we made no changes here. In summary, the lack of glycosylation in any of these sites and mutants is consistent with our model.

Additionally, Lys-141 of GOAT made in SF9 cells was found to be ubiquitylated by electrospray mass spectrometry, with a trypsin site that was not cleaved and +114.04-Da shift corresponding to GG isopeptide linkage. This is consistent with...
cytosolic localization, because ubiquitylation can only occur on the cytosolic face of the ER membrane (67).

Characterization of the GOAT Active Site by Photocross-linking—We previously reported that both microsomal and solubilized, purified GOAT can be specifically photocross-linked by a biotinylated analog of the bisubstrate inhibitor of GO-CoA-Tat containing UV-activatable benzoylphenylalanine in place of Phe-4 (GO-CoA-Tat F4BP) (8). Note that solubilized GOAT is not active (32, 34); however, both with and without presolubilization, this cross-linking could be competed by the addition of excess, unlabeled GO-CoA-Tat, suggesting specificity of the reaction to the enzyme’s active site and the maintenance of a physiologically relevant fold. Therefore, to further map the active site of GOAT, we performed partial proteolysis of cross-linked microsomal GOAT (Fig. 10A). This was done by first cross-linking and then solubilizing in FC-16, digesting with trypsin, and then purifying C-terminal fragments using anti-FLAG. A 15-kDa fragment was detected in both anti-FLAG and biotin blotting, suggesting that at least some of the cross-linking occurs in approximately the C-terminal 15 kDa of GOAT (see “Discussion”). Attempts to characterize the exact locations of photocross-linking by mass spectrometry have so far been unsuccessful.

We next characterized the binding of various analogs and inhibitors to GOAT using photocross-linking compounds and competitors. Based on the fact that ghrelin product analogs with amide lipid linkage have high affinity for GOAT (2), we synthesized two C-terminally biotinylated ghrelin product analog photocross-linkers (ghrelin-28-Oct-diazirine and ghrelin-28-Oct-F4BP). Purified GOAT in FC-16 was then cross-linked with these three analogs alone or in the presence of an excess of the indicated unlabeled competitors. Separate subpanels represent separate experiments. Coomassie Brilliant Blue (CBB) staining is used as a loading control.

FIGURE 10. Characterization of GOAT active site by photocross-linking. A, partial proteolysis of covalently photocross-linked mouse GOAT reveals a 15 kDa cross-linked fragment. GO-CoA-Tat F4BP (biotin-labeled) was photocross-linked to microsomal mouse GOAT-3xFLAG or to control microsomes (Ctrl; from empty-virus infected cells). Microsomes were then solubilized, digested with trypsin, and isolated using anti-FLAG beads (see “Experimental Procedures”). Samples were then immunoblotted using streptavidin or anti-FLAG. Note that although the biotin cannot be removed from GO-CoA-Tat F4BP during trypsin digestion, the 3xFLAG tag contains multiple trypsin cleavage sites, and smaller tag-cleaved species may have been lost or may have fallen below the limit of detection. B, synthetic scheme to make ghrelin-28-Oct-diazirine, an acyl ghrelin product analog photocross-linker. As a complement to GO-CoA-Tat F4BP previously reported (8), we designed another set of photoaffinity labeling agents based on the fact that the amide analogs of the acyl ghrelin product have high affinity for GOAT (32). Photolabile diazirine was installed at the 7-position of the octanoic acid side chain, and biotinylated lysine was installed at the C-terminus of ghrelin-28. Ghrelin-28-Oct-F4BP was made similarly, using unlabeled octanoic anhydride and replacing Phe-4 with benzoylphenylalanine. C–E, photocross-linking cross-competition binding assay. Each of three biotinylated photocross-linking compounds (GO-CoA-Tat F4BP (C), ghrelin-28-Oct-diazirine (D), and ghrelin-28-Oct-F4BP (E)) was cross-linked to purified, solubilized mouse GOAT in the presence of an excess of the indicated unlabeled competitors. Separate subpanels represent separate experiments. Coomassie Brilliant Blue (CBB) staining is used as a loading control.
ing that their binding to GOAT overlaps where the bisubstrate binds. Both product cross-linker analogs, which presumably attach to GOAT in distinct ways because of the location of their photoactive groups, could also be competed by both GO-Tat and acyl ghrelin. These results suggest that the location of the photoactive groups does not significantly affect analog binding. Octanoyl-CoA and desacyl ghrelin, which may bind more weakly to GOAT or in a non-overlapping way, did not compete or only minimally competed against both cross-linker product analog compounds.

**DISCUSSION**

These studies establish the location of the 11 transmembrane-spanning domains and one reentrant loop of GOAT. Using a combination of phylogeny, computational prediction, epitope mapping, and induced glycosylation, we obtained strong convergent evidence for this 11-transmembrane domain model. This new model contains three more transmembrane-spanning domains than previously predicted and also suggests that one domain predicted by the recently developed programs is instead a reentrant loop. This model also indicates that the topology of GOAT is quite structurally complex. Although our evidence strongly supports this overall 11-transmembrane model, we recognize that polypolytopic membrane proteins are complex, and dynamic alterations of topology are formally possible. We also appreciate that the loop insertions employed, although small in size and unable to flip the C terminus into the lumen, may perturb local structure.

Three other MBOATs have been fully mapped topologically: human ACAT1 and ACAT2 (official symbols SOAT1 and SOAT2, reviewed in Ref. 35) and yeast Gup1p (68). GOAT, ACAT1, and Gup1p have somewhat similar topologies, but ACAT2 is quite different than the other three. All four MBOATs have in common that the invariant histidine (His-338) is luminal or buried in the ER membrane near the lumen; this was also shown for partially mapped yeast MBOATs Ale1p and Are1p (68). For GOAT, ACAT1, Gup1p, and Ale1p, the histidine is at the luminal tip of two TM5 separated by only a short 2–3-residue stretch. Are2p is similar, except the C-terminal TM is probably instead a reentrant loop. The conserved asparagine (Asn-307 in GOAT) was also found to be cytosolic in all six cases.

The topology models of GOAT and Gup1p are the most similar, with 12 distinct hydrophobic regions separated by relatively short hydrophilic loops: N terminus luminal, C terminus cytosolic, short terminal tails, a reentrant loop near the middle of the protein, and a relatively long cytosolic loop immediately N-terminal to the hydrophobic region containing the invariant His. However, for Gup1p, the N-terminal two regions were found to be reentrant loops, resulting in a model with nine TM5s.

ACAT1 topology has been studied rigorously using a variety of methods. Overall, its topology is relatively similar to GOAT, with nine TM5s, but ACAT1 has longer loops, two fewer helices, and opposite orientation of the N and C termini. In contrast, ACAT2 was shown to have only two transmembrane domains using selective permeabilization with two distinct epitope tags (29). These results are surprising, considering that ACAT2 has 10 regions predicted to be transmembrane or reentrant by MemBrain (data not shown), similar to the other MBOATs; some of the non-TM hydrophobic regions were modeled as reentrant loops. Also, with the exception of some subtle differences in predicted loop regions around the invariant histidine, the positions chosen should have efficiently interrogated these locations. It is possible, therefore, that ACAT2 has a distinct topology compared with the other MBOATs.

These studies have revealed that the GOAT C terminus can physically contact peptide and octanoyl-CoA substrates and that the bisubstrate analog encompasses both enzyme interaction surfaces. The luminal location of His-338 suggests that it could well be in the active site of the GOAT enzyme, consistent with the hypothesis of a conserved luminal active site for the MBOAT family. In contrast, the conserved extraluminal position of the conserved MBOAT residue Asn-307 indicates that it is unlikely to be directly involved in catalysis. However, its importance might be in substrate interactions, transport of substrates, or protein structural stability.

The finding of a FLAG-tagged ~15-kDa fragment with GO-CoA-Tat 4BP cross-linking maps the ghrelin binding site to the C-terminal region of GOAT. This could be consistent with cross-linking to the regions around TM-8 and TM-9, depending on whether the cross-linked compound causes a gel shift of this smaller GOAT fragment. In any event, these results suggest that the GOAT C-terminal region is directly involved in substrate binding and/or catalysis. Additionally, this region contains three lysophospholipid acyltransferase motifs essential for activity in this subset of MBOATs (69).

The quaternary structure of GOAT has not been investigated previously; the only MBOAT for which the quaternary structure is known is ACAT1, which is a homotetramer both in cell culture and in vitro (63). We show by analytical ultracentrifugation that purified GOAT in detergent micelles is a monomer (Fig. 8D). Taken together with the specific binding of ghrelin photocross-linking analogs to GOAT under nearly identical conditions (Fig. 10, C–E), this suggests that GOAT can bind ghrelin and analogs as a monomer.

Additionally, the V5Glyc glycosylation strategy developed here may be of more general utility for the analysis of membrane proteins. In particular, the flanked sequence seems to be generally well tolerated for loop insertion and efficiently processed by oligosaccharyltransferase. It gives a robust signal on both immunofluorescence and immunoblotting, and this dual topology reporter is far smaller than others previously employed, suggesting that it is likely to be better tolerated for membrane protein applications.

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