Structure of LEP100, a Glycoprotein That Shuttles between Lysosomes and the Plasma Membrane, Deduced from the Nucleotide Sequence of the Encoding cDNA

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Abstract. LEP100, a membrane glycoprotein that has the unique property of shuttling from lysosomes to endosomes to plasma membrane and back, was purified from chicken brain. Its NH₂-terminal amino acid sequence was determined, and an oligonucleotide encoding part of this sequence was used to clone the encoding cDNA. The deduced amino acid sequence consists of 414 residues of which the NH₂-terminal 18 constitute a signal peptide. The sequence includes 17 sites for N-glycosylation in the NH₂-terminal 75% of the polypeptide chain followed by a region lacking N-linked oligosaccharides, a single possible membrane-spanning segment, and a cytoplasmic domain of 11 residues, including three potential phosphorylation sites. Eight cysteine residues are spaced in a regular pattern through the luminal (extracellular) domain, while a 32-residue sequence rich in proline, serine, and threonine occurs at its midpoint. Expression of the cDNA in mouse L cells resulted in targeting of LEP100 primarily to the mouse lysosomes.

Several lysosomal membrane glycoproteins have been identified by immunological methods, isolated, and their overall biochemical characteristics described (1, 6, 14, 20, 21, 27). All of these proteins bear large numbers of N-linked oligosaccharides and have apparent molecular weights (determined by SDS-PAGE) in the range 80,000 to 120,000, whereas apparent molecular weights of the core polypeptides are in the range 40,000 to 60,000. The high level of glycosylation should contribute substantially to a region of fixed negative charge on the luminal surface of the membrane and may be related to protection of lysosomal membrane proteins from hydrolytic destruction or may have some as yet unknown biological significance. The functions of these lysosomal membrane glycoproteins are not known, but the functions may include mediation of ion permeation, export of digestion products, and facilitation of the vesicular transport that operates to deliver materials to the lysosome for digestion.

LEPI00 resembles the other lysosomal membrane glycoproteins in some general physicochemical properties but appears to be unique in that it shuttles between lysosomes, endosomes, and the plasma membrane (21, 22). The distribution of LEPI00 among these membrane systems is profoundly altered by agents that affect lysosomal pH (22). We have suggested that LEPI00 may be involved in the shuttling process, and we have reported a remarkable congruence between patches of this protein on the plasma membrane of chloroquine-treated cells and the distribution of membrane-associated clathrin (22). Thus, we expect that some features of LEPI00 reported here will prove to be general for lysosomal membrane glycoproteins, whereas other features will relate to functions unique to LEPI00.

Materials and Methods

Materials

Endoglycosidase H and 1-deoxymannojirimycin were purchased from Genzyme Corp., Boston, MA. Tetremethylrhodamine-labeled goat anti-rat Ig was purchased from CooperBiomedical Inc., Malvern, PA. [³⁵S]Methionine was purchased as “Trans³⁵S-label” from ICN Radiochemicals, Irvine, CA, and contained some other ³⁵S-labeled compounds in addition to methionine. mAb to LEPI00 (LEPI00-IgG) was purified and derivatized as previously described (21). It was used as iodinated antibody or after coupling with fluorescein isothiocyanate (FITC-LEPI00-IgG) or coupling to Sepharose 4B beads.

Cloning of the Encoding DNA for LEPI00

LEPI00 was purified from a detergent extract of chicken brain microsomes by immunoprecipitation with LEPI00-IgG (21) coupled to Sepharose 4B beads. The eluted protein was subjected to SDS-PAGE, electrophoretically from the gel (16), and subjected to automated Edman degradation and analysis to determine the NH₂-terminal amino acid sequence (15). The amino acid sequence was FDVDRSTGKVCIANLTAFYESVEYKXGQK, where X represents uncertain residues (see Results). A part of this sequence was used in designing an oligonucleotide that might hybridize with LEPI00 mRNA and cDNA. A set of 26mer oligonucleotides containing deoxyxynosine at six positions of codon third-base ambiguity (26) were designed to bind to all possible encoding sequences for the amino acid sequence TGKVCIAN: the sequences were 5'-TTIGCIAATAT(A/G)CAIAAC(C/T)TTICCGT-3'. This set of four oligonucleotides (³⁵P-labeled with polynucleotide kinase) was used to identify several clones (2) by screening about 300,000 plaques.
from unamplified chick brain cDNA libraries (30) in λgt10 (17). Two of these clones were later determined to encode the entire LEP100 protein. The longer of these clones is described in this report.

DNA Sequencing
The cDNA inserts of positive clones were subcloned into the Eco RI site in the plasmids pEMBL8 and pEMBL18 (7). DNA sequencing by the dideoxy method (28), with [35S]dideoxyadenosine 5'triphosphate, established that the cDNAs exactly encoded the NH2-terminal amino acid sequence of LEP100. Further DNA sequencing was done on subcloned fragments on and fragments truncated by timed digestions with exonuclease III (8). We were unsuccessful in many attempts to clone the larger Eco RI fragment into the sequencing vectors pEMBL or Bluescript in an orientation suitable for exonuclease III deletions and sequencing from the 3' to 5' end of the encoding DNA. However, we did succeed in subcloning the full-length cDNA into pEMBL8 and 18 in both orientations after partial Eco RI digestion of the original clone in lambda. Five synthetic oligonucleotides were used as sequencing primers to obtain the nucleotide sequence in the 3' to 5' direction. Generally two or three sequencing runs were analyzed for each segment. Most of the cDNA, including the entire coding sequence, was determined by sequencing in both directions (Fig. 1).

Hydrophobicity and Helicity Analyses
For analysis of hydrophobicity and alpha-helicity, MacGene Plus software (Applied Genetic Technology, Fairview Park, OH) was used. These programs employ algorithms developed by Kyte and Doolittle (19) and by Garnier et al. (10), respectively.

Determination of the Number of N-linked Oligosaccharide Chains
Chicken fibroblasts were grown to confluence and then transferred to medium containing 60 µg/ml 1-deoxymannojirimycin (3, 9) and lacking methionine. After 2 h [35S]methionine (150 µCi/ml) was added for 6 h. The biosynthetic intermediate of LEP100 with high-mannose N-linked oligosaccharides was isolated by immune precipitation with LEP100-IgG covalently linked to Sepharose 4B beads (21) and subjected to timed digestion with endoglycosidase H (31, 32). The digested samples were analyzed by SDS-PAGE and fluorography, and the number of N-linked oligosaccharide chains was estimated by counting the number of stepwise reductions in apparent molecular weight.

Expression of LEP100 cDNA in Mouse L Cells
An expression plasmid, pSV2CAT (12) with the restriction site 5' to the SV40 early promoter/enhancer region changed to a Sal I site, was modified as follows. The small Eco RI to Bam HI fragment was removed with destruction of the restriction sites. Then the small Hind III to Eco RI fragment, containing the 5' end of the CAT gene, was removed, and the polylinker from M13mp8 (25) was ligated into the Hind III–Eco RI sites. The resulting shuttle vector (30) was cut with Eco RI, and the large Eco RI fragment of the cDNA encoding LEP100 was cloned into the Eco RI site. The orientation of the insert cDNA was deduced from electrophoretic analysis of Pst I restriction enzyme digestion fragments. The initial expression clone used in the experiments reported here contained two copies of LEP100 cDNA in tandem.

Mouse Ltk- cells (18) were grown in DME-MEM containing 10% FCS. Cells were cotransfected by the calcium phosphate precipitate method (13) as described by Small and Scangos (29), with 0.1 µg of pH5tks5A0.2 DNA (24) and ~1 µg of LEP100 expression plasmid DNA per 100-mm-diameter tissue culture plate. Cultures were switched to HAT medium (23) after 24 h and maintained in this medium. Clones were picked ~2 wk later and maintained in HAT medium.

The number of LEP100 molecules expressed in the transfected cells was estimated by determining the number of binding sites for 125I-labeled LEP100-IgG on fixed cells made permeable by treatment with saponin as previously described (21). No specific binding was found unless the cells had been cultured in medium containing 10 mM butyrate (11) to enhance expression of the SV40 early promoter. (LEP100-IgG does not bind to any antigen in nontransfected mouse L cells with or without butyrate treatment.)

The distribution of LEP100 molecules in expressing cells was visualized by fluorescence microscopy after labeling fixed, permeabilized cells with FITC-LEP100-IgG. In some experiments cells were double-labeled with FITC-LEP100-IgG and mAb to LAMP1 or LAMP2, mouse lysosomal membrane proteins (6). For double immunofluorescence labeling, conditioned medium containing the anti-LAMP antibody was used first, followed by tetramethylrhodamine-labeled anti-rat IgG and FITC-LEP100-IgG.

Results and Discussion
Identification of the cDNA Encoding LEP100
LEP100 was purified from a detergent extract of chick brain microsomes by immune precipitation followed by SDS-PAGE (see Materials and Methods). The NH2-terminal amino acid sequence of LEP100, determined by sequential automated Edman degradation and product analysis, was PheSerValArgAspSerThrGlyLysValCysIleLeuAlaXLeuThrValAlaAsnLeuThrArgGluTyrLysXXXGlyGlnLys. Position 15 was deduced to be Asn on the basis of failure to identify any residue with or without reduction and alkylation of the protein before Edman degradation, and positions 26 and 27 were suspected to be Cys or Ser. (These predictions were confirmed by the nucleotide sequence.) With Asn assigned to position 15, the sequence includes a potential glycosylation signal (Asn-Leu-Thr), and the lack of a signal for Asn would suggest that it might be completely in the N-glycosylated form. If this were the case, the NH2-terminal end of the molecule would be on the lumenal side of the lysosomal membrane.

A set of 26-mer oligonucleotides based on a part of the NH2-terminal amino acid sequence and containing deoxyinosine at six positions of codon ambiguity was used to screen an unamplified chick brain cDNA library in λgt10. Two positive clones containing cDNA inserts of ~2.0 and 2.2 kb and each containing an internal Eco RI site were found; the longer clone appeared to include all of the sequence of the shorter clone. The longer cDNA was subcloned into the plasmids pEMBL8 and pEMBL18 (7) as full-length insert in both orientations and as Eco RI fragments. Initial nucleotide sequencing by the dideoxy method revealed that the larger Eco RI fragment contained the nucleotide sequence that encoded precisely the amino acid sequence determined for the NH2-terminal end of LEP100.

Nucleotide Sequence of the cDNA and Deduced Amino Acid Sequence of LEP100
The nucleotide sequence of the cDNA and the deduced amino acid sequence of LEP100 are shown in Fig. 2. The amino acid sequence begins with an 18-residue signal peptide and is followed by 396 residues, the first 30 of which exactly match the NH2-terminal amino acid sequence obtained for the mature protein, confirming the identification
Figure 2. Nucleotide sequence and deduced amino acid sequence of a chick brain cDNA encoding LEPI00. Nucleotide residues are numbered in the 5'-to-3' direction: the first residue of the translation initiation triplet ATG is given the number 1, and nucleotides 5'-3' to this residue are indicated by negative numbers. The sequence of only a portion of the 3' untranslated region of the cDNA is reported; the untranslated region is less than 100 bases long and does not contain a poly-A tail. (Genomic cloning reveals, in fact, that the Eco RI site at the 5' end of the cDNA is a natural site in the genomic sequence rather than due to Eco RI linker addition in construction of cDNA libraries [unpublished observations].) The 17 glycosylation sites are marked by triangles, cysteine residues are indicated by ovals, and potential phosphorylation sites by rectangles. The signal peptide is boxed, and the putative membrane-spanning region is indicated by M.

Pattern of N-Glycosylations

As shown in Fig. 2, there are 17 potential N-glycosylation sites in the LEPI00 molecule. An estimate of the actual number of N-linked oligosaccharide chains on the LEPI00 molecule was made from analysis of the products of partial deglycosylation by endoglycosidase H. For this determination, a high-mannose intermediate form of LEPI00 labeled with [35S]methionine was isolated from chick fibroblasts and subjected to timed digestion by endoglycosidase H (see Materials and Methods). The digested samples were analyzed by SDS-PAGE and fluorography, and the number of high-mannose N-linked oligosaccharide chains was estimated by counting the number of stepwise reductions in apparent molecular weight of the LEPI00 molecule as individual oligosaccharide chains were removed (Fig. 3). The entire experiment was done three times. Although at the high molecular weight end of this digestion series the digestion products were never well resolved, presumably due to heterogeneity in the size of the high-mannose N-linked oligosaccharides, nevertheless, it appears that there are approximately 17 steps in the series. In the absence of endoglycosidase H digestion, the high-mannose intermediate form of LEPI00 always appeared predominantly as a single band on fluorographs of SDS-PAGE, suggesting little heterogeneity in total glycosylations per molecule. These results are consistent with the deduced amino acid sequence and with the hypothesis that each of the 17 possible sites for N-glycosylation is indeed usually glycosylated.

Other Features of the Primary Structure of LEPI00

In membrane and secretory proteins with multiple cysteine residues there are usually intrachain disulfide bridges. Eight cysteine residues occur in the LEPI00 sequence, distributed in a remarkably regular pattern in which pairs of cysteines of Asn at position 15 and Ser at positions 26 and 27. Note that whereas nucleotides are numbered from the translation start codon, the aminoacyl residues are numbered from the phenylalanine that is the NH2-terminal residue of the mature protein. A search for homologous DNA and protein sequences in the Los Alamos National Laboratory GenBank failed to uncover any significant homologies.

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Figure 3. Estimate of number of N-linked oligosaccharides. A high-mannose form of LEP100 labeled with [35S]methionine was isolated from cultured chick fibroblasts and subjected to timed digestion with endoglycosidase H. Digestion conditions were 34 mU/ml endoglycosidase H in 0.1% SDS, 2% 2-mercaptoethanol, 60 mM sodium acetate, pH 5.8, at 37°C. At intervals up to 8 h, aliquots were removed and incubated for 2 min in SDS sample buffer at 100°C to stop the enzymatic reaction. Enzyme-treated samples were analyzed by SDS-PAGE and fluorography. In the graph, the position of each discernible band in the digestion series relative to the position of the undigested form is plotted against band number. The resulting curve is extrapolated to the position of the undigested band, yielding an estimate of N-linked oligosaccharides. The inset photograph shows a fluorograph of one digestion series, with positions of countable bands and extrapolated positions marked. In this example, there is some heterogeneity in apparent mobility of the undigested LEP100.

have intrapair spacings of 38, 36, 38, and 37 residues, respectively, whereas interpair spacings are 75, 50, and 69. We suggest that pairs of adjacent cysteine residues may form disulfide bridges that participate in stabilization of a set of four hydrophilic domains. In this hypothetical model of higher-order structure the first three such domains are glycosylated whereas the fourth, closest to the membrane-terminus, is unique in lacking N-linked oligosaccharides.

In the center of the amino acid sequence there is a 32-residue segment rich in Ser and Thr and containing seven proline residues (positions 172, 175, 183, 187, 189, 193, and 196). In the helicity plot (10) shown in Fig. 4, this central region shows a markedly low probability of containing alpha-helical structure. This central region of the LEP100 polypeptide separates pairs of the hypothetical domains mentioned above and might be a region of flexibility in the protein.

A hydrophobicity plot (19), based upon calculations that ignore the N-linked oligosaccharides, is shown in Fig. 4. It seems probable that the single very hydrophobic region near the COOH-terminus is a membrane-spanning region. Previously we reported evidence that LEP100 is an integral membrane protein (21). The evidence included (a) demonstration that LEP100 is a glycoprotein that could only be isolated from membrane fractions by detergent extraction, and (b) demonstration that LEP100 partitions into the detergent phase of a Triton X-114 extract. Because all of the N-glycosylated sites surely occur on the luminal side of the lysosomal membrane, any transmembrane segment within the glycosylated NH2-terminal 75% of the molecule would have to pass twice through the membrane. No such segment exists. The only candidate for a membrane-spanning segment is the 25-amino acid stretch from positions 361 to 385. Just past this putative membrane-spanning segment there is a set of three positively charged amino acids that presumably constitute a "stop-transfer" sequence (4) that may serve during biosynthesis to prevent extrusion of the nascent polypeptide into the lumen of the endoplasmic reticulum and thenceforth to stabilize LEP100 in the lipid bilayer. Given that LEP100 is an integral membrane protein and that hydrophobic segment 361-385 is the only candidate for a membrane-spanning segment, we conclude that this segment is a bona fide membrane-spanning one.

There is a very short amino acid sequence from the membrane-spanning segment to the COOH-terminus, constituting the cytoplasmic domain. We had expected a larger cytoplasmic domain for the LEP100 molecule because we postulate that such a domain might be involved in membrane–cytosol interactions and/or membrane–membrane recognition events. The codistribution of membrane-associated clathrin and LEP100 in chloroquine-treated cells (22) suggested the possibility that LEP100 might interact directly with clathrin or with some cytosolic component that links LEP100 and clathrin. This possibility is not excluded by our findings, although we feel that the lack of a bulkier protein mass on the cytosolic face diminishes the likelihood of this possibility being correct. The cytosolic domain of LEP100 contains three potential phosphorylation sites: one serine, one threonine, and one tyrosine. We reported previously (21) that we could not detect any phosphorylation of LEP100 isolated from tissue cultured fibroblasts. In those cells ~98% of LEP100 molecules occur in lysosomal and endosomal membranes. It may be that the state of phosphorylation is related to the distribution of LEP100 in the cells. If this were
the case, then LEP100 might be detectably phosphorylated in chloroquine-treated cells where ~30% of LEP100 molecules are in the plasma membrane subtended by patches of clathrin (22).

Fig. 5 is a cartoon that incorporates many features of LEP100 and also some of the speculations about its higher-order structure mentioned previously.

Expression of LEP100 in Mouse L Cells
The large Eco RI fragment of LEP100 cDNA was cloned into an expression vector derived from pSV2CAT (12, 30) and cotransfected into mouse Ltk- cells together with a plasmid containing the herpes virus tk gene (24). No expression was detected transiently, but many tk+ clones were isolated in HAT medium. Expression of LEP100 could be activated in some of these clones by maintaining them in medium containing 10 mM butyrate (11). Under these conditions expression approached maximal levels after ~48 h, at which time expression was comparable with that in chick fibroblasts, as judged by the binding of 125I-labeled LEP100-IgG to fixed, permeabilized cells (data not shown).

When expressed in the mouse L cells, LEP100 is correctly targeted to lysosomes (Fig. 6). This was confirmed by double-label immunocytochemistry. Fixed, permeabilized

Figure 5. Cartoon depicting some aspects of LEP100 structure. Each circle represents an amino acid in the mature LEP100 molecule. NH2- and COOH-terminal ends are indicated, and the membrane-spanning segment is represented as a projected helix. The proline-rich central segment is kinked to indicate its probable nonhelical, non-beta-sheet structure. Symbols representing N-linked oligosaccharide chains are attached to the asparagine residues of the glycosylation signals.
cells of the transfected mouse L cell lines were labeled with mAb against the lysosomal membrane proteins LAMP1 and LAMP2 (6) followed by tetramethylrhodamine-labeled goat anti-rat IgG and FITC-LAMP1-IgG. Codistribution of intracellular fluorescence from the anti-LAMP antibodies and LEP100-IgG was found. Furthermore, in the presence of 100 μM chloroquine, LEP100 redistributed until about half of the molecules were in the mouse L cell plasma membrane (unpublished observations of Paul Mathews in our laboratory). Thus, the expression experiments verify the identification of the LEP100 cDNA (based initially upon nucleotide sequence match-up with the NH₂-terminal amino acid sequence of LEP100) and provide a basis for study of relationships between LEP100 structure and its targeting to lysosomes and shuttling between membrane systems.

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