The Mammalian Calcium-binding Protein, Nucleobindin (CALNUC), Is a Golgi Resident Protein

Ping Lin, Helen Le-Niculescu, Robert Hofmeister, J. Michael McCaffery, Mingjie Jin, Hanjo Hennemann, Tammie McQuistan, Luc De Vries, and Marilyn Gist Farquhar

Division of Cellular and Molecular Medicine and Department of Pathology, University of California, San Diego, La Jolla, California 92093-0651

Abstract. We have identified CALNUC, an EF-hand, Ca\(^{2+}\)-binding protein, as a Golgi resident protein. CALNUC corresponds to a previously identified EF-hand/calcium-binding protein known as nucleobindin. CALNUC interacts with Goi3 subunits in the yeast two-hybrid system and in GST-CALNUC pull-down assays. Analysis of deletion mutants demonstrated that the EF-hand and intervening acidic regions are the site of CALNUC’s interaction with Goi3. CALNUC is found in both cytosolic and membrane fractions. The membrane pool is tightly associated with the luminal surface of Golgi membranes. CALNUC is widely expressed, as it is detected by immunofluorescence in the Golgi region of all tissues and cell lines examined. By immunoelectron microscopy, CALNUC is localized to cis-Golgi cisternae and the cis-Golgi network (CGN). CALNUC is the major Ca\(^{2+}\)-binding protein detected by \(^{45}\)Ca\(^{2+}\)-binding assay on Golgi fractions. The properties of CALNUC and its high homology to calreticulin suggest that it may play a key role in calcium homeostasis in the CGN and cis-Golgi cisternae.

The ER is well known to constitute a major storage compartment for intracellular Ca\(^{2+}\) that upon appropriate signaling is released into the cytosol and plays a fundamental role in cell physiology. In addition to its role in signaling, it has become evident that Ca\(^{2+}\) is also essential for many functions that take place within compartments of the exocytic pathway. For example, it is required for chaperone functions and quality control in the ER (Booth and Koch, 1989; Lodish and Kong, 1990; Sambrook, 1990; Lodish et al., 1992) and concentration and proteolytic processing of secretory proteins within regulated secretory granules (Chand and Hurtner, 1991; Carnell and Moore, 1994; Canaff and Brecher, 1996). The Golgi region has also been identified as a Ca\(^{2+}\)-enriched compartment based on ion microscopy (Chandra et al., 1991; Zha and Morrison, 1995). Recently, it has been demonstrated directly by electron energy loss spectroscopy (EELS),\(^1\) a new technique that allows high resolution calcium mapping of intracellular compartments (Grohovaz et al., 1996), that Golgi cisternae have a high Ca\(^{2+}\) level extending across the entire cis-trans axis of the Golgi (Pezzati et al., 1997). The question is how is this high Ca\(^{2+}\) concentration maintained in the Golgi? In the case of the ER, Ca\(^{2+}\) storage is believed to be maintained by multiple calcium-binding proteins including calnexin, calreticulin, GRP78 (BiP), GRP94, ERP72, protein disulfide isomerase, reticulocalbin, and ERC55 (Pozzan et al., 1994; Meldolesi and Pozzan, 1998) of which the most important appears to be calreticulin (Mery et al., 1996). Much less is known about the internal milieu of the Golgi. To date, only a single Golgi luminal Ca\(^{2+}\)-binding protein has been identified, Cab45, which, interestingly, has high homology to the ER Ca\(^{2+}\)-binding proteins reticulocalbin and ERC55 (Scherer et al., 1996).

\(^1\) Abbreviations used in this paper: \(\alpha\)-Man II, \(\alpha\)-Mannosidase II; \(\beta\)-gal, \(\beta\)-galactosidase; aa, amino acids; CGN, cis-Golgi network; CHX, cycloheximide; CRT, calreticulin; CV, carrier vesicle fraction; Cyt, cytosol; EELS, electron energy loss spectroscopy; ERGIC, ER–Golgi intermediate compartment; GAP, GTPase-activating protein; GH, Golgi heavy; GL, Golgi light; GDT, glutathione-S-transferase; NLS, nuclear localization signal; PFA, paraformaldehyde; PK, proteinase K; PNS, postnuclear supernatant; PVDF, polyvinylidene difluoride; RM, residual microsomes; TBS-T, TBS-0.05% Tween; TM, total membranes.
In this paper we have identified a second Golgi Ca\(^{2+}\)-binding protein which we call CALNUC with significant sequence homology to another ER Ca\(^{2+}\)-binding protein, calreticulin. We identified CALNUC in a yeast two-hybrid screen using the heterotrimeric G protein Go\(_{13}\) as bait. CALNUC corresponds to a known protein called nucleobindin (Miura et al., 1992; Wendel et al., 1995). Nucleobindin was thought to be a transcription factor based on its ability to bind DNA fragments in vitro, thus the name nucleobindin (Miura et al., 1992). Nucleobindin was first identified in culture supernatant of a B lymphocyte cell line established from mice prone to the autoimmune disorder, systemic lupus erythematosis (Kanai et al., 1986; Miura et al., 1994). The localization of nucleobindin has been problematic. It has been variously suggested to be a nuclear protein (Wang et al., 1994), a secreted protein (Miura et al., 1992; Wendel et al., 1995), and a resident ER protein, the latter based on its interaction with the cyclooxygenase isoenzymes 1 and 2 (Ballif et al., 1996).

Because of the intriguing diverse properties of this molecule, such as the EF-hand/calcium–binding domains, its homology to calreticulin, and its ability to interact with the Go\(_i\) subfamily of heterotrimeric G proteins (Mochizuki et al., 1995), we set out to characterize nucleobindin, hereafter referred to as CALNUC, and especially to define its localization in the hope of shedding light on its function. To our surprise, we found CALNUC both in cytosolic fractions and associated with Golgi membranes. The Golgi-associated form proved to be a Golgi resident protein concentrated in the cis-Golgi network (CGN) and cis-Golgi cisternae facing the Golgi lumen.

**Materials and Methods**

**Antibodies**

Rabbit antiserum raised against the rat homologue of nucleobindin (calvaria calcium-binding protein) was obtained from D. Heinegard (University of Lund, Sweden; Wendel et al., 1995). Rabbit antiserum to GM130 (Nakamura et al., 1995), α-adaptin, calnexin (Wada et al., 1991), and TGN38 were obtained, respectively, from Drs. Graham Warren (Imperial Cancer Research Foundation, London, UK), S. Schmid (Scripps Research Institute, La Jolla, CA), J.J.M. Bergeron (McGill University, Montréal, Canada), and G. Banting (University of Bristol, UK). Rabbit antiserum to α-Mannosidase II (α-Man II) was prepared as described (Velasco et al., 1993) and mouse mAb 53F3 against α-Man II was obtained from Dr. Brian Burke (Burke et al., 1982; University of Alberta, Canada). Affinity-purified anti-β-COP IgG, raised against the EAGE peptide (Duden et al., 1991) was characterized earlier (Hendrick et al., 1993). mAb G193 against ERGIC-53 was provided by Dr. H.-P. Hauri (Biozentrum, University of Basel, Switzerland).

**Construction of the Rat GC Cell Line and Yeast Two-Hybrid Screening**

Double stranded cDNA was generated from 7 μg poly(A)+ RNA isolated from rat GC pituitary cells (subcloned from GH\(_4\) cells) using a random primed cDNA synthesis kit (Stratagene, Inc., San Diego, CA). EcoRI and XhoI oligo synthesized adapter linkers were ligated onto the cDNA at the 5’ and 3’ end, respectively, and the library was inserted into the Gal4 activation domain pACT2 “prey” vector (CLONTECH Laboratories, Palo Alto, CA). The total library contains ~2 × 10\(^6\) independent clones.

The complete rat cDNA of Go\(_{13}\) was cloned into the Gal4 DNA-binding domain pGBT9 “bait” vector (CLONTECH Laboratories) as described (De Vries et al., 1995). The pGBT9Go\(_{13}\) “bait” vector was transformed into yeast strain HF7c (CLONTECH Laboratories). The transformed yeast colonies were selected on tryptophan (~Trp) selective plates, and after 6 d the plasmids from surviving colonies were analyzed for the presence of pGBT9Go\(_{13}\).

For interaction screening in the yeast two-hybrid system (Chien et al., 1991), 50 μg of the rat GC-cell cDNA library in the pACT2 vector was transformed into yeast HF7c(pGBT9Go\(_{13}\)) strain (Schiestl and Gietz, 1989). Approximately 10\(^6\) colonies were plated onto selective medium, and colonies that survived were scored for β-galactosidase (β-gal) activity by a colony lift assay (CLONTECH Laboratories). Plasmid DNA from the HIS\(^{−}/β-gal\) colonies was purified by transforming into Escherichia coli HB101 by electroporation. These plasmids were retransformed into the HFE7 strain alone or with various control plasmids, including the original pGBT9Go\(_{13}\) “bait” plasmid. Positive clones were grouped based on restriction analysis, and 24 fragments of different sizes were selected and sequenced from the 5’ or the 3’ end of the inserts by automated sequencing.

**In Vivo Interactions**

To construct a complete coding sequence of CALNUC, the insert of a clone containing the 5’-end was ligated to the insert of a 3’-end via an internal EcoRI site, and the reconstituted full-length insert was ligated into the pACT2 vector. Deletion mutants of CALNUC (CALNUCe1-272, CALNUCe1-320, and CALNUCe227-320) were generated by PCR using CALNUC-specific primers (sequences available upon request), subcloned into the pACT2 vector, and cotransformed with pGBT9Go\(_{13}\).

Full-length pACT2-CALNUC was cotransformed with the following Ga subunits in pGBT9 vectors: rat Go\(_{13}\) (De Vries et al., 1995), rat Go\(_{23}\), and mouse Go\(_{23}\) (obtained from P. Insel, University of California San Diego), rat Go\(_{i1}\) (from T. Kosasa, University of Texas, Southwestern Medical Center, Dallas, TX), rat Go\(_{i3}\) (from H. Bourne, University of California San Francisco), rat Go\(_{i3}\), from E. Neer (Brigham and Women’s Hospital, Boston, MA), and rat Go\(_{i3}\) (from E. Ross, University of Texas, Southwestern Medical Center). Interactions were analyzed qualitatively by a colony lift assay for β-gal using 5-bromo-4-chloro-3-indolyl-β-D-galactoside (Guarante, 1983).
Expression and Purification of Glutathione-S-Transferase–CALNUC Fusion Protein

For the production of recombinant glutathione-S-transferase (GST) fusion protein, CALNUC cDNA was subcloned into the pGEX-KG vector (Pharmacia Biotechnology, Inc., Piscataway, NJ), and transformed into E. coli (Y1090) cells. GST-CALNUC fusion protein was affinity purified from bacterial lysates on glutathione–Sepharose and used for immunization of rabbits, in vitro pull-down assays, and 45Ca\textsuperscript{2+} overlay.

Preparation, Affinity Purification, and Characterization of Anti-CALNUC IgG

Antibodies were raised in rabbits against purified full-length GST-CALNUC and used for protein analysis. To compare the primary structure of CALNUC, calreticulin (CRT), and Cab45 their amino acid (aa) sequences were aligned using GeneWorks 2.5.1 software (IntelliGenetics, San Francisco, CA).

Characterization of Anti-CALNUC IgG

Preparation, Affinity Purification, and Characterization of Anti-CALNUC IgG

Antibodies were raised in rabbits against purified full-length GST-CALNUC and used for protein analysis. To compare the primary structure of CALNUC, calreticulin (CRT), and Cab45 their amino acid (aa) sequences were aligned using GeneWorks 2.5.1 software (IntelliGenetics, San Francisco, CA).

Online BLAST searches were performed in the GenBank database (National Institutes of Health, Bethesda, MD) via the National Center for Biotechnology Information’s (NCBI) home page on the World Wide Web. PROSITE (Gowenw; IntelliGenetics, San Francisco, CA), BLASTP and Psort were used for protein analysis. To compare the primary structure of CALNUC, calreticulin (CRT), and Cab45 their amino acid (aa) sequences were aligned using GeneWorks 2.5.1 software (IntelliGenetics, San Francisco, CA).

In Vitro Interactions

Wild-type rat G\textsubscript{\alpha\textsubscript{3}} cDNA, activated rat G\textsubscript{\alpha\textsubscript{3}}(G204L), and inactivated G\textsubscript{\alpha\textsubscript{3}}(G203A) mutants, obtained from A. Spiegel (National Institutes of Health, Bethesda, MD), were subcloned into pBluescript SK+ (Stratagene, San Diego, CA). In vitro transcription/translation of the various forms of G\textsubscript{\alpha\textsubscript{3}} from the T7 promoter in pBluescript SK+ was performed using the TNT-coupled reticulocyte lysate system (Promega Corp., Madison, WI) in the presence of [\textsuperscript{35}S]methionine (in vivo cell labeling grade; Amersham Pharmacia Biotech Inc, Arlington Heights, IL) according to the manufacturer’s instructions. Purified GST-CALNUC fusion protein (6 μg) or GST control (6 μg) was immobilized on glutathione–Sepharose beads and incubated with 15,000 cpm [\textsuperscript{35}S]labeled, in vitro-translated G\textsubscript{\alpha\textsubscript{3}} from the T7 promoter in pBluescript SK+ was used for protein analysis. To compare the primary structure of CALNUC, calreticulin (CRT), and Cab45 their amino acid (aa) sequences were aligned using GeneWorks 2.5.1 software (IntelliGenetics, San Francisco, CA).

Database Searches

Online BLAST searches were performed in the GenBank database (National Institutes of Health, Bethesda, MD) via the National Center for Biotechnology Information’s (NCBI) home page on the World Wide Web. PROSITE (Glowenw; IntelliGenetics, San Francisco, CA), BLASTP and Psort were used for protein analysis. To compare the primary structure of CALNUC, calreticulin (CRT), and Cab45 their amino acid (aa) sequences were aligned using GeneWorks 2.5.1 software (IntelliGenetics, San Francisco, CA).

Figure 2. (A) Structure of CALNUC. Rat CALNUC is a 434-aa protein with an NH\textsubscript{2}-terminal signal sequence, a putative DNA-binding domain (BASIC REGION) with a nuclear localization signal (NLS), an acidic region flanked by two EF-hand motifs (EF-1, EF-2), and a leucine zipper motif. (B) aa 227–287 are required for interaction with Go\textsubscript{\alpha\textsubscript{3}}. Those deletion mutants spanning aa 227–287 containing two EF-hands and a central acidic region (CALNUC227–372, CALNUC320–372) bind to Go\textsubscript{\alpha\textsubscript{3}}, whereas, mutants that lack this region (CALNUC320–434, CALNUC320–372) do not. pGBT9Go\textsubscript{\alpha\textsubscript{3}} “bait” vector was cotransformed with various pACT2-CALNUC mutants into yeast strain HF7c. The transformed colonies were scored for β-gal activity by a colony lift assay. (C) Verification of the site of CALNUC’s interaction with Go\textsubscript{\alpha\textsubscript{3}} by an in vitro binding assay. [\textsuperscript{35}S]Go\textsubscript{\alpha\textsubscript{3}} (arrows) binds to GST-CALNUC1–16,434 (lane 6), GST-CALNUC227–372 (lane 2), or to control beads with GST alone (lane 1). GST-CALNUC fusion proteins bound to glutathione-agarose beads were incubated with [\textsuperscript{35}S]-labeled, in vitro translated Go\textsubscript{\alpha\textsubscript{3}} for 2 h, the bound proteins were separated by 10% SDS-PAGE and detected by autoradiography.
Table I. Interaction between CALNUC and Ga Subunits in Yeast Two Hybrid Filter Assay

| Bait        | β-gal Filter |
|-------------|--------------|
| Go13        | ++           |
| Go13 (Q204L)| ++           |
| Go13 (G203A)| ++           |
| Go14        | ++           |
| Go12        | +            |
| Go11        | +            |
| Go1         | ++           |
| Go4         | ++           |

The β-gal filter assay was performed on (Leu−, Trp+) plates, and the color intensity was scored after 8 h. −, no color; +, weak color; ++, intermediate color; ++++, strong color. Baits were constructed in pGBT9, and CALNUC prey vector was pACT2-CALNUC.

**Cell Culture**

AtT-20/D-16v pituitary cells, obtained from Dr. Richard Mains (Johns Hopkins University, Baltimore, MD), were maintained in DME medium (high glucose) supplemented with 10% (vol/vol) horse serum, 5% FCS. All culture media contained 100 U/ml penicillin G, 100 μg/ml streptomycin sulfate. Cells were used as confluent monolayers.

**Preparation and Treatments of Membrane Fractions from Cultured Cells**

Cells were washed twice with cold PBS and subsequently scraped into homogenization buffer containing protease inhibitors (0.5 mg/ml aprotinin, 0.5 mg/ml pepstatin A, 0.2 mg/ml leupeptin, 1 mM 4-(2-aminoethyl)-benzenesulfonyl-fluoride hydrochloride; ICN Biomedicals, Costa Mesa, CA), 0.5 mg/ml pepstatin A, 0.2 mg/ml leupeptin, 1 mM 4-(2-aminoethyl)-benzenesulfonyl-fluoride hydrochloride; ICN Biomedicals, Costa Mesa, CA), and 5 mM EDTA in TBS. They were then gently homogenized by 10 passages through a 28 1/2-G needle and nuclei and unbroken cells were removed by centrifugation (600 g for 3 min). Membrane pellets were prepared by centrifugation of the postnuclear supernatant (PNS) at 100,000 g for 1 h at 4°C in a Beckman TL-100 ultracentrifuge (Beckman Instruments, Inc., Fullerton, CA). The protein concentration of the membrane pellets was obtained by BCA assay (Pierce Chemical Co., Rockford, IL).

**Immunocytochemistry**

For immunofluorescence, cells were cultured on glass coverslips for 2–3 d, fixed for 1 h with 2% paraformaldehyde (PFA) in 0.1 M phosphate buffer, pH 7.4, and permeabilized with 0.1% Triton X-100 in PBS (10 min). They were then incubated for 1 h at room temperature with affinity-purified rabbit antibodies to CALNUC or mouse mAb to ERGIC-53 or Man II followed by cross-absorbed FITC- or Texas red-conjugated donkey anti-rabbit or anti-mouse F(ab′)2 (Jackson ImmunoResearch Laboratories, West Grove, PA). For double labeling, cells were incubated simultaneously with rabbit polyclonal anti-CALNUC and mouse mAb ERGIC-53 or Man II followed by incubation with appropriate anti-rabbit and anti-mouse conjugates. Cells were mounted in 25% PBS, 75% glycerol with 1 mg/ml p-phenylenediamine, and then examined with a Zeiss Axiophot equipped for epifluorescence (Carl Zeiss Inc., Thornwood, NY).

For immunogold labeling, cultured cells were fixed in 8% PFA, 100 mM phosphate buffer, pH 7.4 (15 min), followed by 4% PFA in phosphate buffer (1 h). Samples were then cryoprotected and frozen in liquid nitrogen as described earlier (McCaflery and Farquhar, 1995). Ultrathin cryosections were prepared and incubated with primary antibodies in 10% FCS/PBS for 2 h at room temperature followed by incubation in 5 or 10 nm gold, goat anti-rabbit or anti-mouse IgG conjugates (Amersham Pharmacia Biotech Inc.) for 1 h at room temperature. Sections were post-fixed (10 min) with 2% glutaraldehyde in 0.1 M phosphate buffer, pH 7.4, stained for 30 min in 2% uranyl acetate, adsorption stained for 10 min with 0.2% uranyl acetate, 0.2% methylcellulose, and 3.2% polyvinyl alcohol and then examined in a Philips CM10 or a JEOL 1200EX II electron microscope. For immunofluorescence, semithin cryosections (0.5–1.0 μm) were incubated as above except that incubation in primary antibodies was for 2 h at room temperature. For immunofluorescence and immunogold labeling of pituitary, liver, and kidney tissue, organs were perfusion-fixed with PLP fixative (2% paraformaldehyde, 0.75% lysine, 0.01 M sodium periodate in 0.1 M phosphate buffer, pH 6.2; McLean and Nakane, 1973) and processed as described for cultured cells.

**Preparation and Treatments of Membrane Fractions from Cultured Cells**

Cells were washed twice with cold PBS and subsequently scraped into homogenization buffer containing protease inhibitors (0.5 mg/ml aprotinin, 0.5 mg/ml pepstatin A, 0.2 mg/ml leupeptin, 1 mM 4-(2-aminoethyl)-benzenesulfonyl-fluoride hydrochloride; ICN Biomedicals, Costa Mesa, CA), and 5 mM EDTA in TBS. They were then gently homogenized by 10 passages through a 28 1/2-G needle and nuclei and unbroken cells were removed by centrifugation (600 g for 3 min). Membrane pellets were prepared by centrifugation of the postnuclear supernatant (PNS) at 100,000 g for 1 h at 4°C in a Beckman TL-100 ultracentrifuge (Beckman Instruments, Inc., Fullerton, CA). The protein concentration of the membrane pellets was obtained by BCA assay (Pierce Chemical Co., Rockford, IL).

**Preparation and Treatments of Membrane Fractions from Cultured Cells**

Cells were washed twice with cold PBS and subsequently scraped into homogenization buffer containing protease inhibitors (0.5 mg/ml aprotinin, 0.5 mg/ml pepstatin A, 0.2 mg/ml leupeptin, 1 mM 4-(2-aminoethyl)-benzenesulfonyl-fluoride hydrochloride; ICN Biomedicals, Costa Mesa, CA), and 5 mM EDTA in TBS. They were then gently homogenized by 10 passages through a 28 1/2-G needle and nuclei and unbroken cells were removed by centrifugation (600 g for 3 min). Membrane pellets were prepared by centrifugation of the postnuclear supernatant (PNS) at 100,000 g for 1 h at 4°C in a Beckman TL-100 ultracentrifuge (Beckman Instruments, Inc., Fullerton, CA). The protein concentration of the membrane pellets was obtained by BCA assay (Pierce Chemical Co., Rockford, IL).

**Preparation and Treatments of Membrane Fractions from Cultured Cells**

Cells were washed twice with cold PBS and subsequently scraped into homogenization buffer containing protease inhibitors (0.5 mg/ml aprotinin, 0.5 mg/ml pepstatin A, 0.2 mg/ml leupeptin, 1 mM 4-(2-aminoethyl)-benzenesulfonyl-fluoride hydrochloride; ICN Biomedicals, Costa Mesa, CA), and 5 mM EDTA in TBS. They were then gently homogenized by 10 passages through a 28 1/2-G needle and nuclei and unbroken cells were removed by centrifugation (600 g for 3 min). Membrane pellets were prepared by centrifugation of the postnuclear supernatant (PNS) at 100,000 g for 1 h at 4°C in a Beckman TL-100 ultracentrifuge (Beckman Instruments, Inc., Fullerton, CA). The protein concentration of the membrane pellets was obtained by BCA assay (Pierce Chemical Co., Rockford, IL).
For digestion with proteinase K (PK), membranes (100 μg) were suspended in 200 μl of reaction buffer containing 10 mM Tris-HCl (pH 7.8), 150 mM KCl, 2 mM MgCl₂, 2 mM CaCl₂, and 0.2 M sucrose followed by incubation with 10 or 20 μg of PK (Boehringer Mannheim GmbH, Mannheim, Germany) at room temperature for 30 min, after which 10 mM PMSF was added to stop the reaction. Proteins were separated by SDS-PAGE and immunoblotted with appropriate antibodies.

For alkaline extraction, membranes (100 μg) were suspended in 200 μl–0.2 M Na₂CO₃ (pH 11.5; Fujiki et al., 1982) containing protease inhibitors, incubated on ice for 30 min, and centrifuged at 100,000 g for 1 h at 4°C. Supernatants were kept on ice for later use, and pellets were resuspended in 200 μl-0.2 M Na₂CO₃ (pH 11.5), followed by SDS-PAGE and immunoblotting.

Triton X-114 extraction was carried out essentially as described earlier (Bordier, 1991) except that proteins in the aqueous and detergent phases were precipitated with 10% TCA for 30 min on ice followed by SDS-PAGE and immunoblotting.

**Preparation of Rat Liver Golgi Fractions**

Rat liver fractions were prepared by density gradient centrifugation and characterized as described previously (Saucan and Palade, 1994; Jin et al.,...
1996). In brief, total microsomes were adjusted to 1.24 M sucrose, loaded onto the bottom of a discontinuous sucrose gradient, and then centrifuged at 82,000 g. All procedures were done at 4°C in the presence of protease inhibitors. The resulting fractions were designated as Golgi light (GL) and Golgi heavy (GH), enriched in Golgi elements, carrier vesicle fractions 1 and 2 (CV1 and CV2), enriched in transport vesicles, and residual microsomes (RM). The protein concentration of each fraction was determined by BCA assay (Bio-Rad Laboratories, Hercules, CA). Fractions were then treated with PK, alkaline pH (11.5), or Triton X-114 essentially as described for membranes from cultured cells except that the Na2CO3-extracted samples were loaded on the top of a 6% sucrose cushion and centrifuged at 400,000 g for 25 min. The supernatants were collected and Golgi membrane pellets were resuspended in Tris-EDTA (10 mM Tris, pH 7.5, 1 mM EDTA) and processed for SDS-PAGE and immunoblotting.

For some experiments liver fractionation was carried out by the method of Leelavathi et al. (1970) as modified by Taylor et al. (1997). In brief, livers of Sprague-Dawley rats were perfused with 20 ml 6% (wt/vol) sucrose in 100 mM KH2PO4/KHPO4, pH 6.8, 5 mM MgCl2 buffer with protease inhibitors, homogenized in the perfusion buffer using a glass/teflon homogenizer and spun at 1,500 g for 10 min. The PNS was loaded onto a discontinuous sucrose gradient (1.3 M, 0.86 M sucrose, PNS, 0.25 M sucrose, 6 ml each) in a SW28 tube (Beckman Instruments, Inc.), centrifuged at 100,000 g for 90 min, and crude fractions designated SI, SII, and SIII were isolated with disposable syringes from the PNS/0.25 M sucrose, 0.86 M sucrose/PNS, and 1.3 M sucrose/0.86 M sucrose interphases, respectively. To prepare Golgi fractions depleted of cargo, rats were injected with 1.5 mg cycloheximide in 150 mM NaCl per kg body weight through the tail vein 90 min before liver excision (Taylor et al., 1997). Fractions were processed for 45Ca2+ overlay or immunoblotting.

**SDS-PAGE and Immunoblotting**

Proteins were separated on 10% SDS gels under reducing conditions, followed by transfer to PVDF membranes (Millipore Corp.). Membranes were blocked with 5% nonfat dry milk in TBS–0.05% Tween (TBS-T) and blotted with primary antibodies followed by incubation with HRP conjugated goat anti–rabbit IgG (1:3,000 in TBS-T), secondary antibody (Bio-Rad Laboratories), and detection by ECL (Pierce Chemical Co.). Quantitation of protein bands was done by densitometry using ScanAnalysis Software (Biosoft, Cambridge, UK).

**Results**

**Identification of CALNUC as a Ca2+-binding Protein that Interacts with Goi3**

To search for protein(s) that can interact with the heterotrimeric G-protein, Goi3, we used the yeast two-hybrid system to screen a rat GC cell pituitary library and isolated 24 inserts of different sizes that were sequenced and analyzed. 3 out of the 24 clones sequenced were identified as the rat homologue of human GAIP, a GTPase-activating protein (GAP) identified earlier (De Vries et al., 1995, 1996, 1998), 8 contained novel sequences, and 13 were identified as the rat calvaria calcium-binding protein (Wendel et al., 1995), the rat homologue of human nucleobindin (Miura et al., 1992). Using the BLAST program the sequence we obtained was found to match 100% the rat calvaria calcium-binding protein. Moreover, antibodies raised against the latter protein (Wendel et al., 1995) recognized GST-CALNUC, and GST-CALNUC was found to bind Ca2+ in a 45Ca2+-overlay (Fig. 3C). Important features of CALNUC are its signal sequence, two EF-hands with an intervening acidic region, leucine zipper, and basic region with a putative nuclear localization signal (NLS; Fig. 2A). CALNUC lacks a putative transmembrane domain, but examination of its hydrophobicity using the GeneWorks 2.5.1. program revealed a 15-aa hydrophobic region at its COOH terminus.

**Homology between CALNUC and Calreticulin**

Among the known calcium-binding proteins CALNUC was found to have 30% homology with rat CRT. The sequence similarity between rat, human, and bovine CRT and rat and mouse CALNUC was found throughout the two proteins, but it was highest in the P-domain (rat CRT198–297) and C-domain (rat CRT298–416) where it was 36 and 43%, respectively (Fig. 3), and lower (26%) in the N-domain (rat CRT1–197). Interestingly, the P- and C-domains of CRT are thought to contain, respectively, high affinity and low affinity Ca2+-binding sites. Three helix-loop-helix motifs (Type A repeats; Baksh and Michalak, 1996) that are similar to the EF-hand motif in CALNUC (Fig. 3) are found in the P-domain of CRT. The COOH-terminal region of CALNUC, like that of CRT, is also relatively acidic. Since the acidic segments of the C-domain of CRT were reported to represent low affinity, high capacity calcium-binding sites, the same may apply to CALNUC. Thus, the structure of CALNUC suggests that the COOH-terminal region may have low affinity, high-capacity binding sites at CALNUC343–352 and putative higher affinity-binding sites at the sites of the EF-hand motifs. There are also notable differences between the structure of CALNUC and CRT. For example, the three times repeated NPD/E motifs in the P-domain of CRT and the conserved KPEDWD motif found in both CRT and calnexin (Nash et al., 1994) are not found in CALNUC. The COOH-terminal KDELR ER retention signal found in CRT is also absent from CALNUC which is consistent with our finding...
Another striking feature of unknown significance shared by CALNUC and CRT are two short conserved aa sequences, AY(I/A)EE and QRLX(Q/E)E(I/E)E, found in the C-domain of CRT (rat CRT337–341 and rat CRT365–372). Further investigation is required to determine the functions of these domains.

When the sequences of CALNUC and Cab45 were compared, they also showed ~30% similarity throughout, but no striking homology was found except for a short aa sequence AANXE(E/D) identified in both Cab45 (aa 47–52 and CALNUC (aa 73–78).

**The Two EF-Hand Motifs of CALNUC and Intervening Acidic Region Are Required for Interaction with Goαi3**

Analysis of the CALNUC clones isolated from the yeast two-hybrid screen revealed that the smallest clone that interacted with Goαi3 was CALNUC227–434 (Fig. 2 B). The fact that the clone that coded for CALNUC227–434 interacted but CALNUC287–434 did not points to the importance of aa 227–287, the region containing the two EF-hands, for binding. To further analyze the Goαi3-binding domain several deletion mutants, i.e., CALNUC227–372, CALNUC320–372, and CALNUC227–320, were generated and their ability to interact with Goαi3 assessed in the yeast two-hybrid system. The smallest mutant capable of binding Goαi3 was CALNUC227–320. Once again, this deletion mutant includes both EF-hands and the intervening acidic region. The leucine zipper region, aa 320–372, does not appear to be required because CALNUC227–320 interacted with Goαi3, but CALNUC320–372 did not. From these data we conclude that the two EF-hands and intervening acidic region constitute the site of CALNUC’s interaction with Goαi3.

Similar results were obtained using an in vitro binding assay in which 35S-labeled, in vitro–translated Goαi3 was incubated with full-length or truncated GST-CALNUC fusion proteins bound to glutathione-agarose beads: Goαi3 bound to all GST-CALNUC fragments containing aa 227–287, but not to those lacking this region (Fig. 2 C). Thus, our results demonstrate by two different assays that the region that spans aa 227–287 containing the two EF-hand motifs and intervening acidic aa-rich region, is required for interaction with Goαi3.

**CALNUC Interacts with the Goγ and Goα Family of Ga**

We also used the two-hybrid system to test whether CALNUC interacts with other Go family members. As shown in Table I, CALNUC interacted with all members of the Goγ subfamily (Goα1, Goα2, Goα3, and Goα4) as well as Goα, but not with Goαq, Goα12, or Goα14. Based on the semi-quantitative β-gal filter assay, the strongest interaction was with Goα3 and Goα1 and interactions with Goγ and the other Goαi members were weaker. These results suggest that CAL-

---

**Figure 6.** Localization of CALNUC to Golgi cisternae in NRK cells. Gold particles are found largely or exclusively on 1–2 cisternae (Gc) on one side of the Golgi stack (arrowheads). Typically, >80% of the label is on the side facing the nucleus (n) assumed to be the cis side of the Golgi stack. Cells were fixed in 8% para-formaldehyde (PFA; 15 min) and 4% PFA (1 h). Ultrathin cryosections were prepared and incubated sequentially with affinity-purified anti-CALNUC IgG and 5- or 10-nm colloidal gold conjugated to goat anti-rabbit IgG. Sections were postfixed and stained as described in Materials and Methods. Bar, 0.1 μm.
NUC specifically interacts with members of the Goα and Gs subfamily of heterotrimeric G proteins.

We further investigated whether the interaction between Goα3 and CALNUC is nucleotide-dependent using the activated GTP-bound mutant, Goα3(Q204L), and inactivated, GDP-bound mutant Goα3(G203A), and found that both mutants bind to CALNUC as efficiently as wild-type Goα3 (Table I).

**CALNUC Is Localized to the Golgi Region by Immunofluorescence**

To determine the intracellular distribution of CALNUC we carried out immunofluorescence on a number of rat (GC pituitary cells, PC-12 cells, NRK cells, REF52 fibroblasts), mouse (AtT-20 pituitary cells), human (HeLa), and monkey (Vero) cell lines. To our surprise, we found that in all cells examined, CALNUC was localized to the Golgi region (Figs. 4 and 5 A). After double labeling CALNUC staining partially overlapped with both α-Man II, a Golgi marker, and ERGIC-53, a marker for the ER–Golgi intermediate compartment (ERGIC) and CGN (Fig. 4). We also checked CALNUC’s distribution by immunofluorescence in semithin cryosections of rat pituitary (Fig. 5, B and C), liver, and kidney. Again, in all cases CALNUC was localized to the Golgi region. Staining was very similar regardless of whether we used the antiserum against recombinant GST-CALNUC. In some cases (e.g., rat kidney) variable cytoplasmic staining was observed, but the predominant signal was in the Golgi region. CALNUC staining in the Golgi persisted in cells treated for 3 h with cycloheximide (data not shown).

**CALNUC Is Localized on Golgi Cisternae Concentrated on the cis Side of the Golgi Stack**

To determine whether CALNUC is associated with Golgi membranes or with other organelles located in the Golgi region we carried out immunogold labeling on ultrathin cryosections of AtT-20 and NRK cells (Fig. 6) and rat anterior pituitary (Fig. 7), kidney, and liver tissue. In all cases, labeling was concentrated on membranes of the stacked Golgi cisternae and associated vesicles and was more concentrated on cisternae and vesicles located on one side of the Golgi stack. The latter could be identified as the cis side based on the location of secretion granules (Fig. 7, A and B) and on double labeling with TGN38 (Fig. 7 C). TGN38 was clearly distributed on the opposite or trans side of the Golgi stack (Fig. 7 C). Counts of gold particles found on Golgi membranes after staining for CALNUC indicate that >80% of the gold particles were found on the one to two cis-most Golgi cisternae and associated vesicles that together constitute the cis-Golgi network (CGN; Farquhar and Hauri, 1997). Therefore, the immunoocytochemical findings indicate that CALNUC is associated with cis-Golgi cisternae and the CGN.

**CALNUC Is Found in Both Membrane and Cytosolic Fractions**

The results of immunofluorescence and immunogold labeling suggested that CALNUC is associated primarily with Golgi cisternae. To obtain further information on its localization, we assessed CALNUC’s distribution in membrane (100,000 g pellet) and cytosolic (100,000 g supernatant) fractions prepared from AtT-20, NRK, and REF-52 cells. In all cases, CALNUC was found in both membrane and cytosolic fractions, with the percent detected in the membrane fraction varying among different cell lines, i.e., 90% in REF-52 cells, 80% in AtT-20 cells, and 50% in NRK cells (Fig. 8). The CALNUC found in cytosolic fractions could represent a true cytosolic pool or a soluble pool released from the Golgi lumen during homogenization. However, it seems unlikely that all the CALNUC found in the cytosolic fraction could be released from the Golgi lumen because in NRK cells only 30% of the Cab45, a soluble luminal Golgi protein, was released into the cytosol under the same conditions (data not shown). Further study is required to clarify the relationship between the CALNUC associated with Golgi membranes and that found in cytosolic fractions.

**The Membrane Pool of CALNUC Is Tightly Associated with Golgi Membranes and Faces the Golgi Lumen**

The aa sequence of CALNUC contains a signal peptide and lacks putative membrane-spanning domains suggesting it is likely to be a soluble luminal protein or a peripheral membrane protein. To determine the topography of the membrane-associated pool of CALNUC we digested 100,000 g pellets prepared from AtT-20 cells with proteinase K (PK), β-COP, a peripheral coat protein (Duden et al., 1991) was completely digested by increasing amounts of PK (Fig. 9), whereas CALNUC and calnexin (Fig. 9), an integral membrane protein that faces the ER lumen (Wada et al., 1991), were resistant to PK digestion, suggesting that CALNUC is a luminal protein.

Next, AtT-20 cell membranes were subjected to alkaline extraction (pH 11.5) with Na2CO3. As shown in Fig. 10 A, most (90%) of the GM130, a Golgi-associated peripheral membrane protein (Nakamura et al., 1995), was removed but all of the calnexin, an ER integral membrane protein (Wada et al., 1991), and most of the CALNUC remained associated with the membrane pellet. When the time of al-
were resuspended in homogenization buffer to the same volume as supernatants. 50-μl membrane or cytosolic fractions were separated by 10% SDS-PAGE and immunoblotted with affinity-purified rabbit anti-CALNUC. CALNUC (63 kD) is detected in both membrane (M) and cytosolic (Cyt) fractions of all cell types tested. The percent of the total CALNUC associated with membrane fractions is 80% in AtT-20 cells (lane 2), 50% in NRK cells (lane 4), and 90% in REF-52 cells (lane 6).

Figure 8. CALNUC is found in both membrane and cytosolic fractions. Membrane (M) and cytosolic (Cyt) fractions were prepared by centrifugation of PNS at 100,000 g for 1 h. Membrane pellets were resuspended in homogenization buffer to the same volume as supernatants. 50-μl membrane or cytosolic fractions were separated by 10% SDS-PAGE and immunoblotted with affinity-purified rabbit anti-CALNUC. CALNUC (63 kD) is detected in both membrane (M) and cytosolic (Cyt) fractions of all cell types tested. The percent of the total CALNUC associated with membrane fractions is 80% in AtT-20 cells (lane 2), 50% in NRK cells (lane 4), and 90% in REF-52 cells (lane 6).

Figure 9. Membrane-associated CALNUC is a luminal protein. Membranes (100,000 g pellet) prepared from AtT-20 cells (see Fig. 8) were treated with the indicated concentrations of proteinase K (PK) at room temperature for 30 min in the presence and absence of detergent followed by immunoblotting with polyclonal antibodies to β-COP, calnexin, or CALNUC. β-COP, a peripheral coat protein, was completely digested by 50 (lane 2) or 100 (lane 3) μg/ml PK whereas CALNUC (lanes 2, 3, 6, and 7) and calnexin (lanes 6 and 7), a membrane protein that faces the ER lumen, were resistant to PK digestion. In the presence of detergent (1% Triton X-100) both CALNUC and calnexin were digested (lanes 4 and 8).

CALNUC Is Enriched in Golgi Fractions of Rat Liver

To obtain further biochemical evidence for the association of CALNUC with Golgi membranes, we carried out immunoblotting for CALNUC on Golgi fractions prepared from rat liver by a procedure designed to separate Golgi fractions from carrier vesicles and residual microsomes (Saucan and Palade, 1994; Jin et al., 1996). Unlike in the cultured cell lines, CALNUC was detected in membrane fractions but not in cytosolic fractions even when up to 400 μg of cytosolic protein was loaded on the gels (data not shown). As shown in Fig. 11A, CALNUC was most enriched in the two Golgi fractions (GL and GH). To determine the topology of CALNUC, the GH fraction was treated with PK in the presence or absence of detergent (Fig. 11B). CALNUC was resistant to PK digestion in the absence of detergent, but was completely digested in the presence of detergent as was Man II, a Golgi membrane protein that faces the Golgi lumen. Rab6, a membrane anchored protein that faces the cytosol, was digested under both conditions. When Golgi membranes were subjected to alkaline extraction (pH 11.5), ~80% of the CALNUC was released into the supernatant (Fig. 11C), but 20% remained associated with Golgi membranes even after treatment for up to 1 h. From these results we conclude that CALNUC is a peripheral membrane protein tightly associated with the luminal surface of Golgi membranes.

To establish whether CALNUC is a Golgi resident protein or a cargo protein in transit, we cleared rat hepatocytes of proteins in transit by treatment with cycloheximide for 90 min, using the procedure recently introduced to distinguish endogenous Golgi proteins from cargo (Taylor et al., 1997), followed by immunoblotting of Golgi fractions for CALNUC. We found that CALNUC was readily detected in Golgi fractions from cycloheximide (CHX) treated rats at levels indistinguishable from controls (Fig. 12), thus demonstrating that CALNUC is a bona fide, Golgi resident protein.

CALNUC Is a Major Calcium-binding Protein in Golgi Fractions

To directly demonstrate that endogenous CALNUC associated with Golgi membranes is capable of binding Ca$^{2+}$, we carried out $^{45}$Ca$^{2+}$ overlays on pooled Golgi fractions. We found that $^{45}$Ca$^{2+}$ bound to one major band at 63 kD (Fig. 13, lane 1) that corresponded in mobility to CALNUC (Fig. 13, lane 1). These results indicate that CALNUC is the major Ca$^{2+}$-binding protein in Golgi fractions as determined by $^{45}$Ca$^{2+}$ overlay.

Discussion

We have identified, characterized, and localized CALNUC, an EF-hand, calcium-binding protein, and shown that it is a resident protein of the CGN and cis-Golgi cisternae. CALNUC corresponds to the rat homologue of a previously identified mouse and human protein called nucleobindin, so named because it bound to DNA in vitro (Miura et al., 1992) and was thought to be located in the nucleus (Wang et al., 1994). We demonstrate here that CALNUC is localized to the Golgi region by immunofluorescence and that it is concentrated in cis cisternae and/or the CGN by immunoelectron microscopy in all tissues and cell lines examined. We further found that CALNUC is a bona fide resident Golgi protein as it is enriched in Golgi fractions prepared from rat livers depleted of cargo. Moreover, we demonstrated by $^{45}$Ca$^{2+}$ overlay that both recombinant CALNUC (see Fig. 1) and endogenous CALNUC (see Fig. 13) are capable of binding Ca$^{2+}$. In fact, the only band detected in Golgi fractions by $^{45}$Ca$^{2+}$ overlay was CALNUC.

Interestingly, CALNUC shows high homology to CRT and has several properties in common with CRT in that...
both are Ca\(^{2+}\)-binding proteins (Miura et al., 1994; Baksh and Michalak, 1996), both have been found to be secreted in autoimmune diseases (Kanai et al., 1993; Sontheimer et al., 1996), both have been found to be secreted in autoimmune diseases (Kanai et al., 1993; Sontheimer et al., 1996), both have been said to be localized to compartments of the exocytic pathway (Opas, 1996; Krause and Michalak, 1997). It is intriguing that CALNUC shows the highest homology to the P- and C-domains of CRT that both are Ca\(^{2+}\)-binding proteins (Miura et al., 1994; Baksh and Michalak, 1996), both have been found to be secreted in autoimmune diseases (Kanai et al., 1993; Sontheimer et al., 1996), both have been said to be localized to compartments of the exocytic pathway (Opas, 1996; Krause and Michalak, 1997). It is intriguing that CALNUC shows the highest homology to the P- and C-domains of CRT that are thought to represent, respectively, high affinity, low capacity and high capacity, low affinity binding sites for Ca\(^{2+}\) (Baksh and Michalak, 1996). Besides its role in binding Ca\(^{2+}\), CRT, as well as calnexin, has been shown to be a glucose lectin that functions as an ER chaperone involved in ER quality control by binding incompletely processed proteins with exposed glucose groups (Parlati et al., 1996; Helenius et al., 1997). However, neither the KPEDWD motif found in both CRT and calnexin that binds integrins (Krause and Michalak, 1997), nor the COOH-terminal KDEL ER retention/retrieval signal is present in CALNUC. We discovered that two motifs of unknown function, AY(I/A)EE and QRLX(Q/E)E(I/E)E, found in CALNUC are shared with CRT. These regions may reflect the common role of those two proteins in binding Ca\(^{2+}\) or another, as yet undisclosed function.

A problem that remains unanswered is the significance of CALNUC’s interaction with G\(_{\alpha}\) subunits. CALNUC has previously been shown to interact with G\(_{\alpha}\) subunits in the yeast two-hybrid system (Mochizuki et al., 1995), and we found here that it can interact with G\(_{\alpha}\) and other members of the G\(_{\alpha}\) subfamily as well as G\(_{\beta}\) but not other G\(_{\beta}\) subunits in the yeast two-hybrid system. We assume that cytosolic CALNUC interacts with G\(_{\beta}\) subunits since G\(_{\beta}\) subunits are mainly anchored through lipid anchors to the cytoplasmic surface of membranes (Mumby, 1997). It are thought to represent, respectively, high affinity, low capacity and high capacity, low affinity binding sites for Ca\(^{2+}\) (Baksh and Michalak, 1996). Besides its role in binding Ca\(^{2+}\), CRT, as well as calnexin, has been shown to be a glucose lectin that functions as an ER chaperone involved in ER quality control by binding incompletely processed proteins with exposed glucose groups (Parlati et al., 1996; Helenius et al., 1997). However, neither the KPEDWD motif found in both CRT and calnexin that binds integrins (Krause and Michalak, 1997), nor the COOH-terminal KDEL ER retention/retrieval signal is present in CALNUC. We discovered that two motifs of unknown function, AY(I/A)EE and QRLX(Q/E)E(I/E)E, found in CALNUC are shared with CRT. These regions may reflect the common role of those two proteins in binding Ca\(^{2+}\) or another, as yet undisclosed function.

A problem that remains unanswered is the significance of CALNUC’s interaction with G\(_{\beta}\) subunits. CALNUC has previously been shown to interact with G\(_{\alpha}\) subunits in the yeast two-hybrid system (Mochizuki et al., 1995), and we found here that it can interact with G\(_{\alpha}\) and other members of the G\(_{\alpha}\) subfamily as well as G\(_{\beta}\) but not other G\(_{\beta}\) subunits in the yeast two-hybrid system. We assume that cytosolic CALNUC interacts with G\(_{\beta}\) subunits since G\(_{\beta}\) subunits are mainly anchored through lipid anchors to the cytoplasmic surface of membranes (Mumby, 1997). It are thought to represent, respectively, high affinity, low capacity and high capacity, low affinity binding sites for Ca\(^{2+}\) (Baksh and Michalak, 1996). Besides its role in binding Ca\(^{2+}\), CRT, as well as calnexin, has been shown to be a glucose lectin that functions as an ER chaperone involved in ER quality control by binding incompletely processed proteins with exposed glucose groups (Parlati et al., 1996; Helenius et al., 1997). However, neither the KPEDWD motif found in both CRT and calnexin that binds integrins (Krause and Michalak, 1997), nor the COOH-terminal KDEL ER retention/retrieval signal is present in CALNUC. We discovered that two motifs of unknown function, AY(I/A)EE and QRLX(Q/E)E(I/E)E, found in CALNUC are shared with CRT. These regions may reflect the common role of those two proteins in binding Ca\(^{2+}\) or another, as yet undisclosed function.

A problem that remains unanswered is the significance of CALNUC’s interaction with G\(_{\beta}\) subunits. CALNUC has previously been shown to interact with G\(_{\alpha}\) subunits in the yeast two-hybrid system (Mochizuki et al., 1995), and we found here that it can interact with G\(_{\alpha}\) and other members of the G\(_{\alpha}\) subfamily as well as G\(_{\beta}\) but not other G\(_{\beta}\) subunits in the yeast two-hybrid system. We assume that cytosolic CALNUC interacts with G\(_{\beta}\) subunits since G\(_{\beta}\) subunits are mainly anchored through lipid anchors to the cytoplasmic surface of membranes (Mumby, 1997). It are thought to represent, respectively, high affinity, low capacity and high capacity, low affinity binding sites for Ca\(^{2+}\) (Baksh and Michalak, 1996). Besides its role in binding Ca\(^{2+}\), CRT, as well as calnexin, has been shown to be a glucose lectin that functions as an ER chaperone involved in ER quality control by binding incompletely processed proteins with exposed glucose groups (Parlati et al., 1996; Helenius et al., 1997). However, neither the KPEDWD motif found in both CRT and calnexin that binds integrins (Krause and Michalak, 1997), nor the COOH-terminal KDEL ER retention/retrieval signal is present in CALNUC. We discovered that two motifs of unknown function, AY(I/A)EE and QRLX(Q/E)E(I/E)E, found in CALNUC are shared with CRT. These regions may reflect the common role of those two proteins in binding Ca\(^{2+}\) or another, as yet undisclosed function.

A problem that remains unanswered is the significance of CALNUC’s interaction with G\(_{\beta}\) subunits. CALNUC has previously been shown to interact with G\(_{\alpha}\) subunits in the yeast two-hybrid system (Mochizuki et al., 1995), and we found here that it can interact with G\(_{\alpha}\) and other members of the G\(_{\alpha}\) subfamily as well as G\(_{\beta}\) but not other G\(_{\beta}\) subunits in the yeast two-hybrid system. We assume that cytosolic CALNUC interacts with G\(_{\beta}\) subunits since G\(_{\beta}\) subunits are mainly anchored through lipid anchors to the cytoplasmic surface of membranes (Mumby, 1997). It......
seems unlikely that the Golgi luminal pool would have the opportunity to bind Go subunits as CALNUC and Go subunits are on opposite sides of membranes. In this paper we focused on characterization of the luminal, Golgi membrane–associated pool of CALNUC, but the relationship between soluble and membrane-associated CALNUC and the interaction of CALNUC with Go subunits remain intriguing problems for the future.

Another unanswered problem is how the membrane pool of CALNUC is bound to the luminal surface of Golgi membranes. In cultured cells we found that CALNUC has a pH of 4.9 by isoelectric focusing (our unpublished observation), is tightly anchored to Golgi membranes and cannot be extracted by extended alkaline treatment. CALNUC lacks a putative transmembrane domain and is likely to represent a peripheral membrane protein which, like p62 (Jones et al., 1993), is so tightly bound to one or more Golgi proteins that it cannot be readily extracted by high pH treatment. However, we cannot rule out that CALNUC could be anchored by a lipid anchor such as palmitate or by a 15-aa hydrophobic region found at its COOH terminus. In the case of synaptobrevin and other SNARES it has been shown that 12 hydrophobic aas at the COOH terminus is sufficient for membrane anchoring (Kutay et al., 1993; Whitley et al., 1996).

CALNUC is only the second Golgi calcium-binding protein identified to date, with Calb45 being the first (Scherer et al., 1996). Calb45 differs from CALNUC in that it is a soluble Golgi resident protein found in the Golgi lumen and is completely extracted after high pH treatment. It also has a putative glycosylation site that is lacking in CALNUC.

What is the role of calcium-binding proteins in the Golgi? It has recently been shown that high level of Ca$^{2+}$ in the Golgi is comparable to that of the ER and is seen all across the stacked cisternae, cis to trans (Pezzati et al., 1997). Surprisingly, the concentration in the TGN is somewhat lower. We can speculate, based on studies on the ER, that Ca$^{2+}$-binding to CALNUC can serve one of two functions: (a) It may serve to maintain the high calcium levels required for Golgi functions such as sorting, lectin binding, and concentration of cargo into regulated secretory granules, or (b) binding of Ca$^{2+}$ may cause conformational changes in CALNUC that render it competent for specific, as yet unknown interactions. In addition, it is possible that CALNUC may bind other divalent cations. CRT has been shown to bind zinc and iron as well as calcium (Baksh and Michalak, 1996). Specific Golgi functions may have specific divalent cation requirements. For example, it has been shown that UDP-GalNAc-polypeptide N-acetylgalactosamine transferase, the enzyme that adds the initial galactose to mucin-type glycoproteins, requires Mn$^{2+}$ both in vitro (Sugiura et al., 1982) and in vivo (Kaufman et al., 1994).

This study opens a number of new and intriguing questions concerning the functions of CALNUC in Ca$^{2+}$ homeostasis in the Golgi, the requirements for Ca$^{2+}$ for Golgi functions, and whether heterotrimeric G proteins are involved in Ca$^{2+}$ homeostasis in the Golgi.

Helen Le-Niculescu is a graduate student in the Molecular Pathology Graduate Program at UCSD and is a Markey Foundation Fellow. This work was supported by National Institutes of Health grants DK 17780 and CA 58689 to Marilyn G. Farquhar.

Received for publication 20 March 1998 and in revised form 7 May 1998.

References

Baksh, S., and M. Michalak. 1996. Basic characteristics and ion binding to calreticulin. In Calreticulin. M. Michalak, editor. R.G. Landes Company, Georgetown. 11–26.

Ballif, B.A., N.V. Minnck, J.T. Barratt, M.L. Wilson, and D.L. Simmons. 1996. Interaction of cyclooxygenases with an apoptosis- and autoimmunity-associated protein. Proc. Natl. Acad. Sci. USA. 93:5544–5549.

Booth, C., and G.L.E. Koch. 1989. Perturbation of cellular calcium induces secretion of luminal ER protein. Cell. 59:729–737.

Bordier, C. 1991. Phase separation of integral membrane proteins in Triton X-114 solution. J. Biol. Chem. 256:1604–1607.

Burke, B., G. Griffiths, H. Reggio, D. Louisgard, and G. Warren. 1982. A monoclonal antibody against a 33-kD K Golgi membrane protein. EMBO (Eur. Mol. Biol. Organ.). J. 1:1621–1628.

Canaff, L., and V. Brecher. 1996. Secretory granule targeting of atrial natriuretic peptide correlates with its calcium-mediated aggregation. Proc. Natl. Acad. Sci. USA. 93:4883–4887.

Carnell, L., and H.P. Moore. 1994. Transport via the regulated secretory pathway in semi-intact PAC12 cells: role of intra-cisternal calcium and pH in the transport and sorting of secretogranin II. J. Cell Biol. 127:693–705.

Chanat, E., and W.B. Huttner. 1991. Mihelcucu-induced, selective aggregation of regulated secretory proteins in the trans-Golgi network. J. Cell Biol. 115:1505–1519.

Chandra, S., E.P.W. Kahle, G.H. Morrison, and W.W. Webb. 1991. Calcium sequestration in the Golgi apparatus of cultured mammalian cells revealed by laser scanning confocal microscopy and ion microscopy. J. Cell Sci. 100:747–752.

Chiun, C.T., P.L. Bartel, R. Sternglanz, and S. Fields. 1991. The two-hybrid system: a method to identify and clone genes for proteins that interact with a protein of interest. Proc. Natl. Acad. Sci. USA. 88:9578–9582.

De Vries, L., E. Elenko, L. Hubler, T.L. Jones, and M.G. Farquhar. 1996. GaIP is membrane-anchored by palmitoylation and interacts with the activated (GTP bound) form of Go subunits. Proc. Natl. Acad. Sci. USA. 93:15203–15208.

De Vries, L., E. Elenko, J.M. McCaffery, T. Fischer, L. Hubler, T. McQuistan, N. Watson, and M.G. Farquhar. 1998. RGS-GaIP, a GAP for Go, heterotrimeric G proteins, is located on clathrin coated vesicles. Mol. Biol. Cell 9:1224–1234.

De Vries, L., M. Mousli, A. Wurmser, and M.G. Farquhar. 1995. GaIP, a protein that specifically interacts with the trimeric G protein Go, is a member of a protein family with a highly conserved core domain. Proc. Natl. Acad. Sci. USA. 92:11916–11920.

Duden, R., G. Griffiths, R. Frank, P. Argos, and T.E. Kreis. 1991. β-COP, a 110 kD protein associated with non-clathrin-coated vesicles and the Golgi complex, shows homology to β-adaptin. Cell. 64:849–856.

Farquhar, M.G., and H.P. Hauri. 1997. Protein sorting and vesicular traffic in the Golgi apparatus. In The Golgi Apparatus. E.G. Berger and J. Roth, editors. Birkhauser, Basel. 63–129.

Fujiki, Y., A.L. Hubbard, S. Flower, and P.B. Lazarow. 1982. Isolation of intracellular membranes by means of sodium carbonate treatment: application to endoplasmic reticulum. J. Cell Biol. 93:97–102.

Grohovaz, F., M. Bossi, R. Pezzati, J. Meldolesi, and F.T. Tarelli. 1996. High calcium concentrations in the trans-Golgi network. EMBO (Eur. Mol. Biol. Organ.) J. 15:1124–1134.

Guarente, L. 1983. Yeast promoters and question of cloned genes in yeast. Methods Enzymol. 101:181–183.

Helenius, A., E.S. Trombetta, D.N. Hebert, and J.F. Simons. 1997. Calnexin, calreticulin and the folding of glycoproteins. Trends Cell Biol. 7:193–200.

Hendricks, L.C., M. McCaffery, G.E. Palade, and M.G. Farquhar. 1998. Disrup-
tion of endoplasmic reticulum to Golgi transport leads to the accumulation of large aggregates containing β-COP in pancreatic acinar cells. Mol. Biol. Cell. 4:413–424.

Jin, M., L. Saucan, M.G. Farquhar, and G.E. Palade. 1996. Rab1a and multiple other Rab proteins are associated with the transcytotic pathway in rat liver. J. Biol. Chem. 271:30105–30113.

Jones, S.M., J.R. Crosby, J. Salamero, and K.E. Howell. 1993. A cystolic complex of p62 and rab6 associates with TGN38/41 and is involved in budding of exocytic vesicles from the trans-Golgi network. J. Cell Biol. 122:775–788.

Kanai, Y., T. Katagiri, S. Mori, and T. Kubota. 1986. An established MRL/Mp-lpr/lpr cell line with null cell properties produces a B cell differentiation factor(s) that promotes anti-single-stranded DNA antibody production in MRL splen cell culture. Int. Arch. Allergy Appl. Immunol. 91:82–94.

Kanai, Y., K. Miura, T. Uehara, M. Amagai, O. Takeda, S. Tanuma, and Y. Kurosawa. 1993. Natural occurrence of Nuc in the sera of autoimmune-prone MRL/lpr mice. Biochem. Biophys. Res. Commun. 196:729–736.

Kaufman, R.J., M. Swooraop, and P. Murtha-Riel. 1994. Depletion of mananae within the secretory pathway inhibits O-linked glycosylation in mamalian cells. Biochemistry. 33:9813–9819.

Krause, K.-H., and M. Michalak. 1997. Calreticulin. Cell. 88:439–443.

Kutay, U., E. Hartmann, and T.A. Rapoport. 1993. A class of membrane proteins with a C-terminal anchor. Trends Cell Biol. 3:73–75.

Leeavathi, D.E., L.W. Estes, D.S. Feingold, and B. Lombardi. 1970. Isolation of a Golgi-rich fraction from rat liver. Biochim. Biophys. Acta. 211:124–138.

Lodish, H.F., and N. Kong. 1990. Perturbation of cellular calcium blocks exit of secretory protein from the rough endoplasmic reticulum. J. Biol. Chem. 265:10893–10899.

Lodish, H.F., N. Kong, and L. Wikstrom. 1992. Calcium is required for folding of newly made subunits of the asialoglycoprotein receptor within the endoplasmic reticulum. J. Biol. Chem. 267:12753–12760.

Maruyama, K., T. Mikawa, and S. Ebashi. 1984. Detection of calcium binding proteins by 45Ca autoradiography on nitrocellulose membrane after sodium dodecyl sulfate gel electrophoresis. J. Biochem. 95:511–519.

McCaffery, M., and M.G. Farquhar. 1995. Localization of GTPases (GTP-bind-Lodish, H.F., N. Kong, and L. Wikstrom. 1992. Calcium is required for folding proteins) by indirect immunofluorescence and immunoelectron microscopy. Methods Enzymol. 257:275–279.

McLean, W., and F.F. Nakane. 1973. A new fixative for immunoelectron microscopy, J. Histochem. Cytochem. 22:1077–1083.

Meldolesi, J., and T. Pozzan. 1998. The endoplasmic reticulum Ca2+ store: a view from the lumen. Trends Biochem. Sci. 23:10–14.

Mery, L., N. Masaeli, M. Michalak, M. Opas, D.P. Lew, and K.-H. Krause. 1993. A cystolic complex of p62 and rab6 associates with TGN38/41 and is involved in budding of exocytic vesicles from the trans-Golgi network. J. Cell Biol. 122:775–788.

Mochizuki, N., M. Hibi, Y. Kanai, and P.A. Insel. 1995. Interaction of the pro-