Membrane Insertion of Gap Junction Connexins: Polytopic Channel Forming Membrane Proteins

Matthias M. Falk, Nalin M. Kumar, and Norton B. Gilula

Department of Cell Biology, The Scripps Research Institute, La Jolla, California 92037

Abstract. Connexins, the proteins that form gap junction channels, are polytopic plasma membrane (PM) proteins that traverse the plasma membrane bilayer four times. The insertion of five different connexins into the membrane of the ER was studied by synthesizing connexins in translation-competent cell lysates supplemented with pancreatic ER-derived microsomes, and by expressing connexins in vivo in several eucaryotic cell types. In addition, the subcellular distribution of the connexins was determined. In vitro-synthesis in the presence of microsomes resulted in the signal recognition particle-dependent membrane insertion of the connexins. The membrane insertion of all connexins was accompanied by an efficient proteolytic processing that was dependent on the microsome concentration. Endogenous unprocessed connexins were detectable in the microsomes used, indicating that the pancreatic microsomes serve as a competent recipient in vivo for unprocessed full length connexins. Although oriented with their amino terminus in the cytoplasm, the analysis of the cleavage reaction indicated that an unprecedented processing by signal peptidase resulted in the removal of an amino-terminal portion of the connexins. Variable amounts of similar connexin cleavage products were also identified in the ER membranes of connexin overexpressing cells. The amount generated correlated with the level of protein expression. These results demonstrate that the connexins contain a cryptic signal peptidase cleavage site that can be processed by this enzyme in vitro and in vivo in association with their membrane insertion. Consequently, a specific factor or condition must be required to prevent this aberrant processing of connexins under normal conditions in the cell.

Channel-forming membrane proteins, similar to other plasma membrane (PM) proteins, are usually synthesized on ribosomes bound to the ER membrane and inserted cotranslationally into this membrane. Subsequently, the proteins are transported to the PM passing through the classical secretory pathway of the cell (Pfeffer and Rothman, 1987). For translocation into the ER membrane, the proteins need a signal specific for ER membrane targeting that is encoded in the amino acid sequence of the polypeptide chain, either as a cleavable signal sequence on the NH2 terminus and processed by signal peptidase, or as an internal signal anchor sequence (Blobel, 1980; Walter et al., 1984). After insertion into the ER membrane, the polypeptide domains fold into their final conformation, and they eventually oligomerize most likely with the help of different chaperones, folding enzymes, and metabolic energy (Rothman, 1989; Hendrick and Hartl, 1993). It can be concluded that connexins follow the secretory pathway based on subcellular connexin distribution studies presented in this paper and from studies reported by other investigators. Studies performed with wild-type connexin–expressing cells isolated from rat liver (this study; Rahman et al., 1993) and dog pancreas (acinar cells, this study), as well as tissue culture cell lines naturally expressing connexins (normal rat kidney fibroblasts [NRK 49F], Musil and Goodenough, 1991), or transfected with connexin cDNAs (BHK cells; this study) showed connexin proteins in the ER membranes, the Golgi membranes, and in the plasma membranes isolated from these cells.

Relatively little is known about the mechanisms that control the generation of the proper topology for membrane proteins. In particular, this is true for polytopic membrane proteins (class III membrane proteins) that traverse the membrane bilayer several times. Since naturally occurring membrane proteins generally adopt only one final orientation, the signals and mechanisms that direct a protein into its native membrane topology seem to be highly specific and extremely precise.

In this study, we analyzed the translocation and the transmembrane organization of the gap junction (GJ) proteins...
(connexins) in the membranes of the ER. Connexins, together with other aqueous channel-forming membrane proteins, represent specialized polytopic transmembrane proteins that are distinguished by the localization of charged amino acid residues within their transmembrane regions. At present, 12 different connexins have been cloned and sequenced from mammals. All represent structurally conserved, nonglycosylated members of a multigene family (Kumar and Gilula, 1992). Connexins traverse the PM bilayer four times oriented with their amino and carboxy termini located in the cytoplasm (NH2-COOH orientation). This membrane topology was suggested by hydropathy analyses of the primary amino acid sequences (Milks et al., 1988), and it is supported by various topological analyses of the connexin protein structure, such as site-directed antibody localizations combined with immunoelectron microscopy (Hertzberg et al., 1988; Milks et al., 1988; Laird and Revel, 1990; Rahman and Evans, 1991; El Aoumari et al., 1992), protease digestions of isolated GJ structures (Zimmer et al., 1987; Hertzberg et al., 1988), or the analysis of intramolecular disulfide bridges (Rahman and Evans, 1991). The NH2-terminal cytoplasmic domain of the connexins consists of a short stretch of 22–23 amino acid residues, and no cleavable signal sequence is present within the NH2-terminal region. Consequently, the first and/or one of the other transmembrane hydrophobic regions must function as an internal signal anchor sequence(s). After their membrane integration, connexins must oligomerize to form a functional GJ hemichannel (Makowski et al., 1977). At the PM, two hemichannels, one provided by each of two neighboring cells, associate to form a double-membrane hydrophilic channel that creates a cytoplasmic continuity.

In this study, we used standard coupled in vitro translation/membrane translocation assays supplemented with canine pancreatic microsomes, and the expression of connexin cDNAs in vivo in various eucaryotic cell types to analyze the processes that are involved in the membrane insertion of connexins. Five representatives of the connexin gene family have been studied. In addition, the distribution of the connexins in different subcellular membrane compartments was analyzed. The results indicate that several factors appear to influence the proper membrane integration and the transmembrane organization for this class of membrane proteins.

Materials and Methods

Special Materials

Rough microsomal membranes used in this study were either purchased from Promega Biotech (Madison, WI) or were prepared as described below. Salt-extracted microsomal membranes, purified signal recognition particle (SRP), and plasmid pBluescript encoding bovine preprolactin were kindly provided by Dr. D. Zopf and Dr. P. Walter (University of California, San Francisco, CA). Plasmid pCh2394-34 encoding the chicken ACHe α7 subunit was kindly provided by Dr. R. Schöpf (Center for Molecular Biology, Heidelberg, Germany). Antibodies α1-6, GAP 10, and anti bovine protein disulfide isomerase (PDI) were kindly provided by Dr. B. Riske (The Scripps Research Institute, La Jolla, CA), Dr. W. H. Evans (University of Wales, Heath Park, Cardiff, U.K.), and Dr. M. R. Jackson (The Scripps Research Institute, La Jolla, CA), respectively. A mouse hybridoma cell line producing a β1 GJ protein–specific monoclonal antibody (M12.13) was kindly provided by Dr. D. A. Goodenough (Harvard Medical School, Boston, MA).

cDNA Constructs

For the in vitro transcription of efficiently translated synthetic connexin RNAs, rat α1, human β1, rat β2, rat β3, and rat α3 GJ cDNAs were cloned into the BglII site of the transcription vector pSP64T (Krieg and Melton, 1984). The general method used for the construction involved mutation of a region ~90–30 bases upstream of the initiation codon to produce a BamHI site. Either the BamHI site in the polylinker or a BglII linker ligated to restriction enzyme–cut DNA was used to generate BamHI-BamHI or BamHI-BglII fragments of the GJ cDNAs. Specifically, the β1 GJ construct contains the nucleotides 51–996 of the human β1 GJ cDNA with a BglII linker ligated into the StuI site 82 bp downstream from the stop codon of the β1 GJ cDNA. The β1 GJ construct contains nucleotides 259 to ~2350 of the rat β2 GJ cDNA. The β2 GJ construct contains nucleotides 71–895 of the rat β2 sequence. The α3 GJ construct contains nucleotides 191–1608 of the α3 GJ cDNA with a BglII linker ligated into the AffIII site, after filling in the protruding ends with Klenow DNA polymerase. The α3 GJ construct contains nucleotides 201–1705 of the rat α3 GJ cDNA. Resulting constructs consisted of the connexin coding region and ~90–30 upstream bases and 100–300 downstream bases of the GJ cDNAs, flanked by the 5′ and 3′ noncoding regions of Xenopus β-globin cDNA.

Constructions of eucaryotic connexin expression vectors will be described in more detail in separate reports. Briefly, human β1 GJ protein was expressed in yeast cells (Saccharomyces cerevisiae) after the human β1 GJ cDNA was cloned into the BamHI site of the yeast expression vector YEP G2 containing a polyadenylation signal, a 2–μ plasmid origin of replication, and a galactose regulated promoter.

Rat α1 GJ protein was expressed in S9F cells after α1 cDNA was cloned into the baculovirus vector pAcC737, transfected into S9F insect cells, and followed by plaque purification as described in Stauffer et al. (1991) for β1 GJ cDNA.

Rat α1 and β1 GJ proteins were expressed in BHK cells after α1 and β1 cDNAs were stably transfected into these cells as described in Kumar and Gilula (1992). GJ cDNAs were cloned into the BamHI site of exon I of human growth hormone encoded in the eucaryotic expression vector pNUT, kindly provided by Dr. R. D. Palmer (University of Washington, Seattle, WA). With this strategy, the GJ cDNAs are under the control of an inducible metallothionin promoter that allows the expression of variable amounts of GJ protein. BHK cells were cotransfected with standard CaPO4 cotransfection procedures, and recombinants were selected by methotrexate treatment.

In Vitro Transcription, Translation, and Translocation Assays

All plasmids used for in vitro transcription were linearized, phenol chloroform extracted, and ethanol precipitated. Transcription reactions were performed with a Riboprobe transcription kit (Promega Biotech). In vitro translation reactions were performed in nucleate-free rabbit reticulocyte lysate or wheat germ extracts supplemented with [35S]methionine (Amersham Corp., Arlington Heights, IL). Reactions were generally performed for 30–60 min at 32°C and 28°C, respectively. Canine pancreas rough microsomal membranes or salt-extracted microsomal membranes were generally present at a final concentration of 1 Eq/10 μl of translation reaction. Purified SRP was used at a final concentration of 10 U/10 μl reaction volume. Canine pancreas rough microsomes were prepared as described by Walter and Blobel (1983). To test the influence of protease inhibitors on the proteolytic processing of connexins, all components of the translocation assays were mixed together except RNA. Protease inhibitors N-tosyl-L-phenylalanine chloromethyl ketone (TPCK), Nα-p-tosyl-lysine chloromethyl ketone (TLCK), diisopropyl fluorophosphate (DFP), E-64, and leupeptin were added to individual aliquots to a final concentration of 1 mM from 10× stock solutions and incubated for 10 min at 30°C before synthetic RNA was added. To modify reaction conditions from more reducing conditions (small amounts of DTT are present in the [35S]methionine and in the microsome preparations) to oxidizing conditions that allow posttranslational S-S bridge formation, oxidized glutathione was added in variable amounts from 0.1 to 5 mM final concentration to the translocation reactions, according to Marquardt et al. (1993).
lation, 15-μl aliquots were adjusted to 50 μl volume with buffer 1 consisting of 150 mM KOAc, 50 mM triethanolamine-acetic acid (pH 7.0), and 2.5 mM MgOAc. For alkaline extractions, aliquots were adjusted to pH 11.5 with 1 M NaOH, and adjusted to 50 μl volume. Samples were incubated on ice for 10 min, overlaid onto a 100-μl cushion of 0.5 M sucrose in buffer 1 or a 100-μl cushion of 0.2 M sucrose in 30 mM Hepes adjusted to pH 11.5, 150 mM KOAc, 2.5 mM MgOAc, and fractionated into a supernatant (S) and pellet (P) fraction. Proteins in the supernatant fractions were then precipitated with TCA and neutralized with saturated Tris base before processing for SDS-PAGE.

Protease Protection Assays. After translation, reactions were chilled on ice and 10-μl aliquots were diluted to 50 μl in buffer 1. Microsomes were stabilized by the addition of CaCl2 to a final concentration of 10 μM and incubated for 10 min. Either water or proteinase K (predigested for 30 min at 30°C) was added to final concentrations of 0.1 and 0.5 μg/ml, or trypsin (predigested in the same way) was added to final concentrations of 0.5 and 1 μg/ml from 10× stock solutions in water, respectively. In addition, where indicated, NP-40 was added to a final concentration of 1% from a 10% (wt/vol) stock in water. All digestes were incubated for 1 h on ice. Protease activity was blocked by the addition of 5 mM DFP, 1% SDS (final concentrations), and boiling for 5 min, following the method described by Chavez and Hall (1992). Samples were then diluted 10 times with a buffer (buffer 2) containing 150 mM NaCl, 1% NP-40, 0.5% deoxycholate, and 50 mM Tris, pH 7.5, chilled on ice, and processed for immunoprecipitation analysis, as described below. Immunoprecipitated polypeptide fragments were analyzed on special 20% SDS gels, allowing the resolution of small polypeptide fragments (Thomas and Kornberg, 1975).

Immunoprecipitations

Antipeptide antibodies raised against various regions of α1 and β1. GI protein β1, B, βE, βJ, and βS are described in Milks et al. (1988), α1J and α1S are described in Nishi et al. (1991), and GAP 10 is described in Rahman and Evans (1991). Antibody β1-6 was raised in rabbits against the first eight amino acids (MNWTGLYT) of β1 GJ protein (Risek, 1987). This antibody cross-reacts with the NH2-terminal domain of α1 GJ protein. Specificity of all antibodies was determined by dot-blot analysis as previously described (Milks et al., 1988). All immunoprecipitations were performed in 1-ml volumes in RIPA buffer consisting of 150 mM NaCl, 0.1% SDS, 1% NP-40, 0.5% deoxycholate, and 50 mM Tris, pH 7.5. 50 μl of α:10 sherry of protein A-Sepharose (prewashed for 1 h) plus 1–5 μl of antiserum were added to each sample and shaken for 1 h or overnight at 4°C, depending on the strength and specificity of the antibodies. Beads were sedimented by centrifugation and washed two times with RIPA buffer before the addition of SDS protein sample buffer.

Purification and Analysis of Connexin Isoforms from Tissues and Cell Culture

PMs were prepared from rat liver and dog pancreas using the method described by Stauffer et al. (1991) with the following modifications. One rat liver and ~20 g of a dog pancreas were used per preparation, respectively. 60,000 g pellets were resuspended in 5 ml buffer, and 10 ml of 67% (wt/wt) sucrose was added. Samples were placed in one tube each, overlayed with 25 ml of 30% (wt/wt) sucrose and centrifuged at 27,000 rpm for 105 min in a rotor (SW 28; Beckman Instruments, Inc.) PM were collected from the interface of the sucrose step gradient, containing either Golgi membranes, PM, or rough ER membranes, respectively, were harvested using a needle and syringe. Aliquots of the fractions were added directly to RIPA buffer and processed for immunoprecipitation, or processed further to prepare subcellular membranes. Subcellular membrane fractions were prepared by the method of Bole et al. (1986) with the following modifications. Cells were chilled on ice and scraped from the plate in 1 ml PBS containing 0.25 M sucrose, and then dispersed with 10-20 strokes in a tight-fitting dounce homogenizer. Opaque bands at the interfaces of the sucrose step gradient, containing either Golgi membranes, PM, or rough ER membranes, respectively, were harvested using a needle and syringe. Aliquots of the fractions were added directly to RIPA buffer and processed for immunoprecipitation, or aliquots were directly subjected to SDS-PAGE and transferred to nitrocellulose membranes for immunoblot analysis with connexin-specific anti-peptide antibodies. Bound antibodies were detected by coupled chemiluminescence. Immunoblots were stripped, reprobed with other connexin-specific antibodies, and stained again following the instructions of the manufacturer (Amerham Corp.).

Results

In Vitro Translation of Connexin Proteins

Complementary DNAs of five different members of the connexin gene family (α1 [Cx43], α3 [Cx46], β1 [Cx32], β2 [Cx26], β3 [Cx31], for connexin nomenclature see Kumar and Gilula [1992]) were cloned into the transcription vector pSP64T, and synthetic RNA was transcribed using SP6 polymerase (Krieg and Melton, 1984). This strategy allowed the synthesis of connexin-specific cRNAs that were translated much more efficiently than the natural connexin RNAs in competent cell lysates, such as rabbit reticulocyte lysates or wheat germ extracts. Resulting connexin translation products generated in the absence of microsomes corresponded to the molecular masses for these proteins as predicted from their amino acid sequences (Fig. 1, lanes 1, 2, 11, 12, 15, 16, 19, and 21). Identification of the translation products by specific antipeptide antibodies directed against different regions of the connexins (see below) provided additional confirmation for the synthesis of full-length connexin proteins.

Translation in the Presence of Microsomes Results in a Specific Processing of the Connexin Proteins

In previous studies, coupled in vitro translation/membrane
translocation assays supplemented with canine pancreatic microsomes (Walter and Blobel, 1983) have been found to integrate several type I, II, and III membrane proteins with a relevant membrane topology (e.g., Lipp and Dobberstein, 1986b; Spiess and Lodish, 1986; Zerial et al., 1986; Mayer et al., 1988). The microsomes, which are vesicles derived from the ER membranes of pancreatic acinar cells during isolation, are believed to contain all ER luminal and membrane proteins necessary for the successful translocation of secretory and membrane anchored proteins (Nicchitta and Blobel, 1993).

The translation of GJ proteins in the presence of canine pancreatic microsomes resulted in the generation of specific translation products that were ~2-2.5 kD smaller in size than the full-size proteins generated in the absence of microsomes, based on the electrophoretic mobility shift on the protein gels (marked as a' and b' GJ protein in the figures). The faster migrating translation products were generated with all connexin cRNAs translated in these assays (Fig. 1, lanes 3-10, 13, 14, 17, 18, 20, and 22). This finding was surprising because no general connexin modification has been described that could account for the observed increase in electrophoretic mobility. Endogenous b1 GJ proteins isolated from mammalian tissues correspond in their electrophoretic mobility to the full-size translation products (compare Fig. 3 A). Translocation reactions incubated for very short time periods always showed equivalent amounts of modified and unmodified translation products in the individual reactions, even at time periods when the translation of connexin polypeptides was still continuing (Fig. 1, lanes 3-8; shown for a1 GJ protein only). This result indicates that the processing occurred cotranslationally or, at least, in close relationship to the translocation reaction.

The Connexin Protein Processing Correlates with the Microsome Concentration

Performing translocation reactions with variable concentrations of microsomes showed that the amount of modified, faster migrating connexin protein was directly related to the concentration of microsomes in the translocation reactions, eventually reaching 100% completion (Fig. 2). In reactions containing low concentrations of microsomes (0.05-0.5 Eq/10 μl reaction volume, Fig. 2, lanes 2-6 and 12-16), only a small portion of the translation product was processed, while in reactions containing high microsome concentrations (1.5-2 Eq/10 μl reaction volume, Fig. 2, lanes 9 and 10, 19 and 20), the faster migrating connexin product was almost the only detectable product generated. Approximately equal amounts of processed and unprocessed connexins were generated when intermediate membrane concentrations (~0.75-1 Eq/10 μl reaction volume, Fig. 2, lanes 9 and 10, 19 and 20), the faster migrating connexin product was almost the only detectable product generated. Approximately equal amounts of processed and unprocessed connexins were generated when intermediate membrane concentrations (~0.75-1 Eq/10 μl reaction volume, Fig. 2, lanes 7 and 8, 17 and 18) were used in the experiments. Translocation reactions in all subsequent experiments (and in the experiments shown in Fig. 1) were carried out with intermediate membrane concentrations (~1 Eq/10 μl reaction volume), leading to the generation of complete and modified connexin proteins that appear as a double-band pattern on the fluorograms (α/α' and β/β' GJ protein).

Various control experiments exclude the possibility that the processing results from the cloning strategy (the same processing occurred with connexin cRNAs derived from...
Figure 2. Processing of connexins is correlated with the microsome concentration. Synthetic RNAs encoding α1 and β1 GJ proteins were translated in the reticulocyte lysate system for 1 h in the presence of increasing concentrations of microsomes (0.05–2 Eq/10 μl vol), resolved by SDS-PAGE, and visualized by fluorography.

Figure 3. Connexins isolated from natural sources correspond to the full-size connexins synthesized in vitro. (A) PM and RER membranes were isolated from rat liver and dog pancreas acinar cells and native β, GJ protein, endogenous to these membranes, was analyzed by immunoblot analysis using a connexin specific antibody as described in Materials and Methods. Complete cell lysates (lanes 1 and 4) and β, GJ protein synthesized in vitro in the absence (−) or presence (+) of microsomes (lanes 7 and 8) were analyzed in parallel. The RER membrane fraction isolated from dog pancreas (lane 6) is identical to the microsomal membranes used in the in vitro translocation assays. (B) Determination of alkaline PDE activity in subcellular fractions and cell lysates prepared from dog pancreas. The samples contained similar protein concentrations. Three independent preparations were analyzed. The activity of the PM marker enzyme was very low in the RER membranes containing fractions, indicating that the connexin detected in these fractions originated from the ER membranes of the pancreatic cells.

Figure 2. Processing of connexins is correlated with the microsome concentration. Synthetic RNAs encoding α1 and β1 GJ proteins were translated in the reticulocyte lysate system for 1 h in the presence of increasing concentrations of microsomes (0.05–2 Eq/10 μl vol), resolved by SDS-PAGE, and visualized by fluorography.

other in vitro transcription vectors), the translation system (reticulocyte lysates vs wheat germ extracts, Fig. 1 vs Fig. 5), the microsomes (five membrane batches prepared independently from individual canine pancreas were used), or the reaction conditions that were applied. Different control proteins (membrane anchored: acetylcholine receptor α7 subunit; secretory: prolactin, yeast α factor; cytoplasmic: β-globin; Fig. 4) translated in parallel to the connexin proteins showed the expected results described previously for these proteins. Reaction conditions were modified from more reducing conditions (presence of DTT) to oxidizing conditions (≤5 mM final concentration oxidized glutathione was added) to allow S-S bridge formation in the newly synthesized polypeptides (Marquardt et al., 1993). However, none of these conditions had any influence on the connexin processing (data not shown). Finally, the connexin processing could not have been caused by using microsomes pre-
pared from cells that normally do not synthesize gap junctions because pancreatic acinar cells normally express endogenous GJ proteins (Hertzberg and Skibbens, 1984). In fact, direct evidence was obtained for endogenous β1 GJ protein in the microsomal fractions prepared from rat liver and from the canine pancreas that was used for the in vitro translocation reactions (Fig. 3 A, lanes 3 and 6). The endogenous microsomal β1 GJ protein corresponded in size to unprocessed β1 GJ protein generated in the in vitro translation reactions in the absence of microsomes (Fig. 3 A, lanes 7 and 8). Subcellular membrane fractions were assayed for alkaline PDE activity to ensure that the detected GJ protein was derived from the RER membranes (Fig. 3 B).

**The Processed Connexins Are Integral Membrane Proteins**

To determine if the connexin translation products were cotranslationally translocated into the membranes, microsomes were pelleted at neutral and alkaline pHs. Supernatants and pellets were separated and analyzed by SDS-PAGE. At pH 7.0, as well as at pH 11.5, the faster migrating processed connexins (α1′/β1′) were abundant in the pellet fractions (Fig. 4, lanes 2 and 4, 14 and 16), while unprocessed connexins (α1 and β1) were located primarily in the supernatant fractions (Fig. 4, lanes 1 and 3, 13 and 15). This result indicated that the processed connexins were integrated into the microsomal membranes. However, it could not be determined unequivocally with this assay whether any of the unprocessed connexin proteins that were generated in parallel in the translocation reactions were also integrated into the microsomal membranes because polypeptides with distinct hydrophobic regions tend to associate with membrane surfaces even at alkaline pH (compare upper bands in Fig. 4, lanes 8 and 20, which represent unprocessed, non-membrane integrated prolactin precursors labeled as pPL). Therefore, full-length connexins that were present in the pellet fraction (Fig. 4, lanes 14 and 16) could represent either membrane-integrated or membrane-associated connexin polypeptides.

**Connexin Proteins are Proteolytically Cleaved**

**Similar to the Processing of a Cleavable ER Target Signal Sequence**

A possible proteolytic cleavage of the connexin proteins linked to their membrane translocation was studied by carrying out immunoprecipitations in the in vitro translation products that were generated in the presence of intermediate microsome concentrations. The products were immunoprecipitated with anti-peptide antibodies that corresponded to specific sequences of the α1 and β1 GJ proteins (Fig. 5 B). The specificity of the antibodies used was verified by dot blot analysis, as described previously (Milks et al., 1988). While antibodies directed against the extreme carboxy terminus (α1, β1, S; Fig. 5 A, lanes 5 and 11) or the intracellular loop region (α1, β1, J; Fig. 5 A, lanes 4 and 10) precipitated both faster migrating and full-length connexins, none of the three different NH2-terminal antibodies that were used (β1, B1, 1-6, Fig. 5 A, lanes 3, 8, and 9; GAP 10, Fig. 8 A, lane B; Fig. 8 B, lanes 1–4) precipitated the faster migrating products. Therefore, it was possible to conclude that the connexin polypeptides were proteolytically processed by a protease present in the microsomal preparation that specifically removed the NH2-terminal portion of the connexins.

**Additional information on the proteolytic processing of**

![Figure 4](https://jcb.rupress.org/)
Connexins are proteolytically processed when translated in vitro in the presence of microsomes, removing an NH2-terminal portion. (A) Synthetic RNAs encoding α1 and β1 GJ proteins were translated in the reticulocyte lysate system in the absence (−) or presence (+) of intermediate concentrations of microsomes. GJ protein-specific translation products were precipitated using anti-peptide antibodies specific for different topological domains of α1 and β1 GJ proteins. Immunoprecipitated polypeptides were resolved by SDS-PAGE and visualized by fluorography. While antibodies directed against the extreme carboxy-terminal domains of α1 and β1 GJ proteins (α1S, β1S, lanes 5 and 11) or an internal region of the proteins (α1J, β1J, lanes 4 and 10) precipitated both the full-size proteins (α1 and β1) and the faster migrating forms (α1' and β1'), neither of the two antibodies specific for the NH2-terminal sequence of the β1 GJ protein (β1 1-6, β1B, lanes 3, 8, and 9) recognized the faster migrating connexin products. (B) Schematic representation of the membrane topology for GJ proteins in vivo. The different topological domains of the proteins, located either extracellular (extracellular loops E1 and E2), in the cytoplasm (NH2-terminal domain, intracellular loop [I loop]), or within the membrane bilayer (transmembrane domains M1-M4) are indicated. The regions recognized by the different connexin specific anti-peptide antibodies (β1B, β1 1-6, GAP 10, β1E, β1J, α1J, β1S, and α1S) used in this study are schematically marked.

Figure 5. Connexins are proteolytically processed when translated in vitro in the presence of microsomes, removing an NH2-terminal portion. (A) Synthetic RNAs encoding α1 and β1 GJ proteins were translated in the reticulocyte lysate system in the absence (−) or presence (+) of intermediate concentrations of microsomes. GJ protein-specific translation products were precipitated using anti-peptide antibodies specific for different topological domains of α1 and β1 GJ proteins. Immunoprecipitated polypeptides were resolved by SDS-PAGE and visualized by fluorography. While antibodies directed against the extreme carboxy-terminal domains of α1 and β1 GJ proteins (α1S, β1S, lanes 5 and 11) or an internal region of the proteins (α1J, β1J, lanes 4 and 10) precipitated both the full-size proteins (α1 and β1) and the faster migrating forms (α1' and β1'), neither of the two antibodies specific for the NH2-terminal sequence of the β1 GJ protein (β1 1-6, β1B, lanes 3, 8, and 9) recognized the faster migrating connexin products. (B) Schematic representation of the membrane topology for GJ proteins in vivo. The different topological domains of the proteins, located either extracellular (extracellular loops E1 and E2), in the cytoplasm (NH2-terminal domain, intracellular loop [I loop]), or within the membrane bilayer (transmembrane domains M1-M4) are indicated. The regions recognized by the different connexin specific anti-peptide antibodies (β1B, β1 1-6, GAP 10, β1E, β1J, α1J, β1S, and α1S) used in this study are schematically marked.

Studies on the Transmembrane Organization of Connexins Generated in the Presence of Microsomes

The successful translocation of many secretory proteins and the integration of several transmembrane proteins have been studied using a protease protection assay. The assay is based on the analysis of polypeptides and polypeptide fragments that are protected from proteolytic degradation by the lipid bilayer of the microsomal vesicles. Since in polytopic membrane proteins, such as connexins, that traverse the membrane several times, several relatively small polypeptide fragments were expected to be protected, we combined the classical protease protection assay with the specific immunoprecipitation of protected protein fragments to obtain some information on the transmembrane organization of the processed connexin proteins. Assays were performed with β1 GJ protein since a complete set of anti-peptide antibodies recognizing all cytoplasmic and extracellular domains of the β1 GJ protein (NH2-terminal, intracellular loop, COOH terminal, and extracellular domains; Fig. 7) was available. Microsomes containing in vitro-translocated β1 GJ protein were incubated with different concentrations of trypsin or proteinase K (PK), respectively. Before the immunoprecipi-
Characterization of the connexin processing indicates a cleavage by signal peptidase. (A) The connexin cleavage was dependent on the cotranslational addition of microsomes. Synthetic RNAs encoding different GJ proteins were translated in the reticulocyte lysate system in the absence of microsomes for 20 min. After terminating the translation reactions, microsomes were added and incubated for an additional 30 min (lane 3). In parallel reactions, microsomes were pelleted, washed, and resuspended in fresh membrane buffer used for the preparation of microsomes. Microsomal supernatant (lane 4) or washed microsomes (lane 5) were then added cotranslationally. Standard translation reactions in the absence (lane 1) and presence (lane 2) of microsomes were done in parallel. All proteins in A, B, and C were separated by SDS-PAGE and visualized by fluorography. (B) The proteolytic processing was not inhibited by protease inhibitors. Translation reactions (reticulocyte lysate system) were preincubated with the protease inhibitors TPCK, TLCK, E-64, leupeptin, and DFP before the addition of synthetic connexin RNAs, as described in Materials and Methods. Only the results for β1 GJ protein are shown in A and B, representing similar results obtained with all connexins studied. (C) The connexin cleavage was dependent on SRP. RNAs encoding α1 and β1 GJ proteins, prolactin (pPL) and yeast α-factor were translated in the wheat germ system in the absence (−) or presence (+) of intermediate concentrations of microsomes. Either water (lanes 3, 6, 9, and 16), trypsin (0.5 mg/ml, lanes 4, 7, 10, and 17, and 1 mg/ml, lanes 11, 12, and 18), or proteinase K (PK, 0.1 mg/ml, lanes 5, 8, 13, and 19, and 0.5 mg/ml, lanes 14, 15, and 20), respectively, was added to aliquots of the membrane insertion reactions. Trypsin or PK, together with NP-40 (1% final concentration) was added to control aliquots (lanes 12 and 15). After blocking protease activity, GJ polypeptides and polypeptide fragments protected from proteolytic degradation were immunoprecipitated with anti-peptide antibodies specific for the NH2-terminal domain (β1B), the extracellular loops E1 and E2 (β1E), the intracellular loop region (β1J), and the COOH-terminal domain (β1S). Immunoprecipitations were analyzed on SDS-protein gels, allowing the resolution of small polypeptide fragments. Polypeptides were visualized by fluorography. A linear representation of the β1 GJ protein is shown under the fluorogram. Numbers for the amino acid residues for the different topological domains corresponding in their graphical pattern to Fig. 5 B, and the binding sites for the anti-peptide antibodies are given. The locations of radioactive labeled methionine residues are indicated by diamonds.
tably outside of the microsomes immunoprecipitated labeled β, polypeptide fragments of apparent sizes of ~9-10 and 15-17 kD that were protected from proteolysis by both proteases (Fig. 7, lanes 10, 11, 13, and 14). However, degradation of the intracellular loop was observed when detergent (1% NP-40, final concentration) was added before the addition of trypsin or PK (Fig. 7, lanes 12 and 15). No labeled protein fragments were immunoprecipitated under these conditions after trypsin treatment (Fig. 7, lane 12), and only a small polypeptide fragment with an apparent molecular mass of ~4 kD was resistant to PK digestion (Fig. 7, lane 15). β, GJ protein that was not exposed to proteases was precipitable with all the individual antibodies under the conditions used (Fig. 7, lanes 3, 6, 9, and 16).

Taken together, these results suggest that either the intracellular loop region of the NH2-terminally processed connexin protein was located inside the membrane vesicles, an orientation that differs from the protein in the native assembled junction channel, or that the intracellular loop was resistant to proteolytic degradation under the conditions applied. These possibilities will be considered in the Discussion.

NH2-terminally Processed Connexin Proteins Similar to the Ones Generated In Vitro Were Also Identified In Vivo

The translocation of connexins into the ER membrane was analyzed in vivo to determine if the observed signal peptidase processing is related to the normal membrane translocation process of connexins in vivo. Since the amount of connexin polypeptides present in the ER membranes is relatively low in normal connexin expressing cells (rat hepatocytes, dog pancreatic acinar cells, Fig. 3), making such an analysis relatively difficult, we expressed higher levels of the α and β, GJ proteins for this analysis by using heterologous eukaryotic protein expression systems (yeast, baculovirus, BHK cells, Fig. 8 A). Expressed connexins were analyzed either by immunoblot analyses or by the immunoprecipitation of radiolabeled proteins. Analysis of the expressed proteins showed that connexins with a processed NH2-terminal domain, similar to the products found in vitro, were detectable in all cell types analyzed (Fig. 8 A, left panel: β, GJ protein—transfected yeast cells, lanes 1, 3, 5; middle panel: insect cells infected with an α, GJ protein expressing baculovirus, lanes 10 and 11; right panel: BHK cells stably transfected with α or β, GJ protein, lanes 18, 19, 25, 27, 29, and 30). While yeast transfectants generated nearly equal amounts of processed and full-size connexins, the majority of the connexin protein generated in the infected insect cells, and especially in the transfected BHK cells, appeared to be the unprocessed full-size protein. Using the transfected BHK cells that allow the expression of variable amounts of protein (the connexin cDNAs are inserted behind a heavy metal inducible promoter), it was possible to demonstrate that the generation of the NH2-terminally processed connexin products correlated with the level of connexin expression (Fig. 8 A, right panel). Under moderate expression conditions (50 μM zinc), only intact α, or β, GJ protein was detectable (Fig. 8 A, lanes 14-16, and 22 and 23), while conditions of intermediate (100 μM zinc, Fig. 8 A, lanes 24 and 25) or high connexin protein expression (125-150 μM zinc, Fig. 8 A, lanes 17-19 and 26-30) resulted in the synthesis of increasing amounts of NH2-terminally cleaved (α' and β') Connexin proteins, as well as intact connexin proteins.

To determine the intracellular distribution of the different connexin products generated under these conditions in vivo, highly induced BHK cells expressing α, or β, GJ protein, respectively, were fractionated into subcellular membrane fractions consisting of PM, Golgi membranes, and rough ER membranes (microsomes). The cleaved connexin products were only detectable in the ER membrane containing fractions (Fig. 8 B, lanes 1, 5, and 9). This result indicates that the NH2-terminally cleaved connexin product generated in the transfected BHK cells was similar to the product generated in the in vitro translocation assays, and for example, was not related to the degradation of PM GJ structures. A small amount of processed connexin protein present in the PM fraction prepared from α, GJ protein expressing cells (Fig. 8 B, lane 6) was most likely related to a slight contamination of this subcellular fraction with some ER derived membranes. This was indicated by the presence of some PDI, an ER specific marker protein in the PM fraction (Fig. 8 C). Only the intact connexin product was detected in the Golgi fractions (Fig. 8 B, lanes 3, 7, and 11). Additional, slower migrating protein products present in Fig. 8 B, lanes 5-8; Fig. 8 C, lanes 4 and 5) and some other nonspecifically detected cellular proteins (especially in Fig. 8 B, lanes 14-4 and 9; Fig. 3 A, lane 3).

In summary, these results indicate that connexins can also be processed by signal peptidase during their translocation process into the ER membrane in vivo, and some mechanism prevents this processing under normal conditions.

Discussion

The results presented in this report provide direct evidence of the cotranslational membrane integration of connexin polypeptides into the ER membrane. The integration occurred in an SRP-dependent manner. The membrane integration was accompanied by an unprecedented and efficient proteolytic processing of the connexins when translocated in vitro into ER-derived microsomes, removing an amino-terminal portion of all connexins analyzed. Furthermore, the microsomes used in this analysis contained endogenous uncleaved full-length β, connexin, indicating that the pancreatic microsomes generated from the ER during isolation were competent to integrate connexins correctly when the cells were intact. An analysis of connexins generated in heterologous eukaryotic expression systems indicated that this proteolytic processing occurred also under in vivo conditions of elevated connexin expression when connexin polypeptides were translocated into the ER membrane. The expression of α, and β, GJ protein consistently resulted in the generation of variable amounts of processed connexin polypeptides (Fig. 8 A), independent of the expression system. Subcellular fractionation provided evidence that the ER membranes contained both the processed and full-length connexin polypeptides, while the Golgi membranes and the PM contained only full-length connexins (Fig. 8 B). BHK cells expressing different amounts of connexin protein also showed that the generation of the processed polypeptides correlated with the level of protein expression (Fig. 8 A).
Figure 8. Expression of connexin cDNAs in eucaryotic cells and analysis of the synthesized connexin polypeptides. (A) α1 and β1 GJ protein cDNAs were cloned into eucaryotic protein expression vectors and expressed in yeast, baculovirus, or stable transfected BHK cells, respectively. Yeast proteins were subjected to SDS-PAGE and transferred to nitrocellulose membranes for immunoblot analysis with the connexin-specific anti-peptide antibodies GAP 10, β1 J, and β1 S. Nontransfected yeast cell lysates were analyzed as controls. Connexins expressed in the presence of [35S]methionine in baculovirus-infected cells and BHK cells, respectively, were immunoprecipitated from complete cell lysates using the anti-peptide antibodies Bt 1-6, α1 J, α1 S (β1 GJ protein), or Bt B, Bt J and Bt S (β1 GJ protein), resolved by SDS-PAGE, and visualized by fluorography. Synthetic connexin RNAs translated in the absence (−) or presence (+) of microsomes were analyzed in parallel as controls. In all three systems, the NH2-terminally processed, faster migrating connexins (α1 and β1) were detected together with the unprocessed full-size connexins (α1 and β1). The BHK cell expression system allows the expression of variable amounts of connexin protein, since the connexin cDNA is inserted behind a heavy metal inducible promoter. Increasing the induction conditions (0.05, 0.1, 0.125, and 0.15 mM zinc) resulted in an equal increase in connexin expression for both full-size and NH2-terminally processed connexins (lanes 14–19 and 22–30). (B) Highly induced BHK cells expressing α1 or β1 GJ protein were separated into subcellular fractions containing either sER membranes, Golgi membranes, or PM, respectively, as described in Materials and Methods. Subcellular fractions were subjected to SDS-PAGE and transferred to nitrocellulose membranes for immunoblot analysis with GJ protein-specific anti-peptide antibodies GAP 10, β1 J, and β1 S. Complete cell lysates were analyzed in parallel as controls. Full-size (α1, β1) and NH2-terminally processed GJ proteins (α1J, β1J) were detected in the fractions as marked. (C) The presence of PDI using an antibody raised against bovine PDI was determined to verify the purity of the subcellular membrane preparations derived from β1 GJ protein–expressing BHK cells.

Processed connexin polypeptides that were possibly generated under normal expression conditions in wild-type cells most likely escaped detection because of their low abundance (Fig. 3).

Native connexin proteins isolated from PMs have been shown to be unprocessed by NH2-terminal amino acid microsequencing, or only the start-methionine was removed (Nicholson et al., 1981, 1987; Zimmer et al., 1987). This data was later confirmed by the cloning and sequencing of connexin cDNAs, showing that no cleavable ER target signal sequence is encoded at the NH2 terminus of the connexin proteins (for connexin sequences see Kumar and Gilula, 1992).

Analysis of the conditions that were associated with the proteolytic processing indicates that the connexins were not processed by one of the known proteolytic activities that function in the degradation of misfolded or improperly oligomerized proteins in the ER, a process generally referred to as "rapid ER degradation" (Amara et al., 1989; Klausner and Sitia, 1990; Bonifacino and Lippincott-Schwartz, 1991). First, these proteases normally do not process proteins in such a rapid reaction that is directly linked to membrane integration, such as observed in this study for the connexins (Fig. 1). Second, the proteolytic cleavage of the connexins could not be prevented by cysteine and serine protease inhibitors such as TPCK, TLCK, leupeptin, or E64 (Fig. 6 B), and these inhibitors have been reported to block the known enzymes related to this process (Wikström and Lodish, 1991; Urade and Kito, 1992; Yuk and Lodish, 1993).

However, a proteolytic processing of the connexin proteins...
by signal peptidase is indicated by the results in this study. All the following properties of the connexin processing are indicative of a cleavage by signal peptidase. First, the connexins were proteolytically processed in proximity to their NH2 terminus, removing the NH2-terminal portion of the proteins. This processing appears very similar to the cleavage of cleavable ER target signal sequences (Blobel, 1980; Walter et al., 1984). Second, the cleavage occurred concomitant with the insertion into the ER membrane, also typical for the processing of cleavable signal sequences (Fig. 1). Third, the proteolytic cleavage reaction was restricted to the lumen of the ER vesicles (Fig. 6 A). Signal peptidase is known to be present and active in the microsomes used in our studies (Evans et al., 1986). Finally, the cleavage was not affected by efficient serine protease inhibitors (Fig. 6 B, DFP, TPCK, and TLCK), a property described for signal peptidase and attributed to its unusual cleavage mechanism (Daly and von Heijne, 1992). Taken together, a potential proteolytic processing of connexin proteins by signal peptidase is indicated by the findings in this study, raising the question, what prevents this cleavage in vivo?

Some examples of an internal signal peptidase processing on "cryptic" sites were previously reported for type I (cytochrome P 450) and type II membrane proteins (invariant chain [I]], asialoglycoprotein receptor H1 subunit) after modifying the structure of the NH2-terminal domains of those proteins that altered their overall charge (Lipp and Dobberstein, 1986a; Schmid and Spiess, 1988; Szczesna-Skorupa et al., 1988). However, these examples clearly differ from our findings in that the connexins were translated as nonmutated, wild-type proteins.

To obtain some information on the transmembrane organization of the cleaved connexin polypeptides, protease protection assays (Morimoto et al., 1983; Scheele, 1983) were used in this study in combination with the immunoprecipitation of protein fragments protected from degradation. In previous studies, this protease protection assay has been used extensively to analyze the membrane insertion of proteins and to determine the membrane orientation of different types of membrane anchored proteins (Lipp and Dobberstein, 1986a; Spiess and Lodish, 1986; Zerial et al., 1986; Mayer et al., 1988; Chavez and Hall, 1992; Skach et al., 1994; Gafvelin and von Heijne, 1994). We found that the intracellular loop domain of the β1 connexin was protected from proteolytic degradation when trypsin or proteinase K were added exogenously to the microsomes, while it was degraded when the microsomes were permeabilized with detergent before the addition of protease (Fig. 7). This result indicates that either the intracellular loop domain was located inside the vesicle lumen, suggesting an inverted membrane topology for the NH2-terminally processed connexins (hypothesized in an earlier report, Falk et al., 1994), or that the intracellular loop was resistant to proteolytic degradation under the conditions applied. An inverted membrane topology would be consistent with the processing of a cryptic signal peptidase cleavage site motif (small, uncharged amino acid residues in positions -3 and -1; von Heijne, 1983) that exists in connexins in front of transmembrane region M1. Proteolytic processing at this site is suggested by the mobility shift of 2-2.5 kD observed on the protein gels.

However, an inversion of the transmembrane orientation of connexins is not simple to explain, and several observations argue against an inverted membrane topology. First, an inversion of the membrane topology of connexins would suggest that their COOH-terminal domain would also be located in the vesicle lumen, and thus is protected from proteolytic degradation. No such connexin fragments were precipitated in the protease protection assays using the COOH terminus-specific antipeptide antibodies BS (Fig. 7), indicating that the COOH terminus was correctly located outside the microsomal vesicles. Second, protease digestions of isolated PM-derived GJ structures indicate that the intracellular loop domain of β1 GJ protein is accessible to proteases, but it seems to be relatively resistant to a complete proteolysis (Milks et al., 1988). In digests performed with endoprotease Lys-C, the intracellular loop was only cleaved on its COOH-terminal side, while the NH2-terminal side resisted degradation. Therefore, it seems likely that the β1 GJ protein fragments that were precipitated in the protease protection assays with the β1J antibodies were generated by an incomplete degradation of the intracellular loop, rather than by its protection from proteolysis by the lipid bilayer of the membrane vesicles. Third, preliminary amino acid sequencing results obtained from radiolabeled NH2-terminally processed β1 connexin indicates that the potential signal peptidase cleavage site is located beyond the transmembrane region M1, suggesting that the extracellular loop E1 is correctly located in the lumen of the microsomes (unpublished results). Finally, in this context, it is also interesting to mention a study by Sipos and von Heijne (1993) that shows that the distribution of positively charged amino acid residues in the regions flanking the first transmembrane region of β1 connexin is consistent with its proposed transmembrane topology in the PM. Based on the distribution of positively and negatively charged amino acid residues flanking the transmembrane region, some models have been developed to predict the membrane orientation of bitopic membrane proteins (von Heijne and Gavel, 1988; Hartmann et al., 1989).

Recent data obtained in Escherichia coli suggests that these models could also apply for polytopic membrane proteins (Andersson and von Heijne, 1994; Gafvelin and von Heijne, 1994) to predict their transmembrane topology. Additional experiments and other approaches will be required to determine unequivocally the actual transmembrane orientation of the connexin proteins in the ER membrane.

The results of this study demonstrate that the connexins contain a "cryptic" signal peptidase cleavage site that can be processed by this enzyme in association with their membrane insertion. What then prevents this proteolytic cleavage under normal conditions in vivo? One possibility is that an additional factor exists that permits the connexin polypeptides to obtain a proper localization within the membrane bilayer, thus preventing their cleavage. A factor binding to cytoplasmic domains of the connexins is possibly indicated by the results obtained with the proteolytic degradation of connexins. Although potentially accessible to proteolytic degradation, the NH2-terminal domain and the intracellular loop region of connexins were found to be highly resistant to protease digestion (Zimmer et al., 1987; Milks et al., 1988; this study). Recently, Musil and Goodenough (1993) reported the oligomerization of α1 connexin after its exit from the ER in a cell culture line. Consequently, it seems
plausible to propose that a factor that may prevent the oligomerization process in the ER may also be involved in the integration of connexins into the ER membrane.

A recent report by Görlich and Rapoport (1993) suggests that only two membrane protein complexes (SRP receptor and Sec61p) and the TRAM protein are required for the successful integration of membrane proteins into synthetic liposomes. At least for connexin proteins, some other factor(s) appear to be involved in generating the proper organization of these proteins in the membrane bilayer. This issue will be addressed in future experimentation.

M. M. Falk is especially thankful to F.-U. Hartl and M. Wiedmann (Memorial Sloan-Kettering Cancer Center, New York), T. Rapoport (Max-Delbrück-Center for Molecular Medicine, Berlin-Buch, Germany), and C. Nicchitta (Duke University, Durham, NC) for valuable discussions. This work was supported by grants GM 37904 to N. B. Gilula, GM 37907 to N. B. Gilula and M. M. Falk, a grant from the Lucille P. Markey Charitable Trust, and a Deutsche Forschungsgemeinschaft grant Fa 264/1-1 to M. M. Falk.

Received for publication 14 April 1994 and in revised form 19 July 1994.

References

Amara, J. F., G. Leederkerken, and H. F. Lohid. 1989. Intracellular degradation of unassembled asialoglycoprotein receptor binding subunits: a pre-Golgi, non-lysosomal endoepitaxial cleavage. J. Cell Biol. 109:3315-3324.

Andersson, H., and G. von Heijne. 1994. Membrane protein topology: effects of the translocation of charged residues explain the 'positive inside' rule. EMBO (Eur. Mol. Biol. Organ.) J. 13:267-272.

Blobel, G. 1980. Intracellular protein topogenesis. Proc. Natl. Acad. Sci. USA. 77:1496-1500.

Bole, D. G., L. M. Hendershot, and J. E. Kearney. 1986. Posttranslational association of two connexin32 antigenic sites in intact and split rodent hepatocyte gap junctions. J. Cell. Biol. 107:1817-1824.

Gilmore, R., and G. Blobel. 1985. Translocation of secretory proteins across the membrane. Annu. Rev. Biochem. 54:349-384.
von Heijne, G., and Y. Gavel. 1988. Topogenic signals in integral membrane proteins. Eur. J. Biochem. 174:671–678.
Walter, P., and G. Blobel. 1983. Preparation of microsomal membranes for cotranslational protein translocation. Methods Enzymol. 96:84–93.
Walter, P., R. Gilmore, and G. Blobel. 1984. Protein translocation across the endoplasmic reticulum. Cell. 38:5–8.
Wikström, L., and H. F. Lodish. 1991. Nonlysosomal, pre-Golgi degradation of unassembled asialoglycoprotein receptor subunits: a TLCK- and TPCK-sensitive cleavage within the ER. J. Cell Biol. 113:997–1007.
Yuk, M. H., and H. F. Lodish. 1993. Two pathways for the degradation of the H2 subunit of the asialoglycoprotein receptor in the endoplasmic reticulum. J. Cell Biol. 123:1735–1749.
Zerial, M., P. Melancon, C. Schneider, and H. Garoff. 1986. The transmembrane segment of the human transferrin receptor functions as a signal peptide. EMBO (Eur. Mol. Biol. Organ.) J. 5:1543–1550.
Zimmer, D. B., C. R. Green, W. H. Evans, and N. B. Gilula. 1987. Topological analysis of the major protein in isolated intact rat liver gap junctions and gap junction-derived single membrane structures. J. Biol. Chem. 262:7751–7763.