Abstract. The 17-juxtamembrane cytoplasmic residues of the polymeric immunoglobulin receptor contain an autonomous basolateral targeting signal that does not mediate rapid endocytosis (Casanova, J. E., G. Apodaca, and K. E. Mostov. Cell. 66:65-75). Alanine-scanning mutagenesis identifies three residues in this region, His656, Arg657, and Val660, that are most essential for basolateral sorting and two residues, Arg655 and Tyr668, that play a lesser role in this process. Progressive truncations suggested that Ser664 and Ile665 might also play a role in basolateral sorting. However, mutation of these residues to Ala or internal deletions of these residues did not affect basolateral sorting, indicating that these residues are probably not required for basolateral sorting. Two-dimensional NMR spectroscopy of a peptide corresponding to the 17-mer signal indicates that the sequence Arg658-Asn-Val-Asp661 has a propensity to adopt a B-turn in solution. Residues COOH-terminal to the B-turn (Arg662 to Arg669) seem to take up a nascent helix structure in solution. Substitution of Val660 with Ala destabilizes the turn, while mutation of Arg657 to Ala does not appear to affect the turn structure. Neither mutation detectably altered the stability of the nascent helix in the COOH-terminal portion of the peptide.

Polarized epithelial cells have two plasma membrane domains: the apical domain facing the external environment, and the basolateral domain facing the internal milieu. These domains display striking differences in protein and lipid composition (Caplan and Matlin, 1989; Simons and Wandering-Ness, 1990; Rodriguez-Boulan and Nelson, 1989; Gumbiner, 1990; Nelson, 1992; Mostov et al., 1992; Rodriguez-Boulan and Powell, 1992). In MDCK cells, most newly made plasma membrane proteins are directly targeted from the TGN to either the apical or basolateral surface (Rodriguez-Boulan and Nelson, 1989; Simons and Wandering-Ness, 1990; Le Bivic et al., 1990; Lisanti and Rodriguez-Boulan, 1990). Until recently, it had been presumed that targeting to the apical cell surface is a signal-mediated process, whereas targeting to the basolateral surface occurs by "default" (Simons and Wandering-Ness, 1990; Bomsel and Mostov, 1991). However, studies using wild-type and mutant forms of plasma membrane proteins (examples: low-density lipoprotein receptor [LDLR}], lysosomal protein Igpl20, Fc receptor [Fc-R-II], lysosomal acid phosphatase [LAP], and polymeric immunoglobulin receptor [plgR]) expressed in MDCK cells showed that the cytoplasmic domain of these proteins contain active sorting signals mediating targeting from the TGN to the basolateral surface (Hunziker et al., 1991; Casanova et al., 1991; Mostov et al., 1992). When mutations in the cytoplasmic tail inhibited basolateral targeting, the proteins were instead delivered to the apical plasma membrane.

The basolateral sorting signals so far analyzed can be placed into two categories. Most of the signals appear to be colinear with Tyr-containing signals for endocytosis via clathrin-coated pits (Hunziker et al., 1991; Brewer and Roth, 1991; Le Bivic et al., 1991; reviewed in Mostov et al., 1992). In several other cases (e.g., the transferrin receptor and vesicular stomatitis virus G protein) basolateral sorting signals appear not to overlap with endocytosis signals (Hunziker et al., 1991; Dargemont et al., 1993; Thomas et al., 1993). The cytoplasmic domain of the LDLR contains two basolateral signals, one of each type. The membrane proximal basolateral signal is colinear with the signal for endocytosis, while the membrane distal signal does not overlap with an endocytosis signal (Matter et al., 1993).

For those basolateral signals that are colinear with coated-pit internalization motifs, it has been suggested that both signals may share common structural features (Mostov et al., 1992). The three-dimensional structure of the tyrosine-containing endocytosis signal of the transferrin receptor was predicted by computer modeling to be a type I B-turn (Col-
sis signals, a \( \alpha \)-turn is an important part of both signals. How-
pere to cause sorting from the Golgi apparatus to the basolateral surface. This promoter-based expression vector, pCB6 (Brewer and Roth, 1991), using standard techniques (Maniatis et al., 1982).

Materials and Methods

Generation of Cell Lines Expressing Mutant plgRs

All mutants were generated by oligonucleotide site-directed mutagenesis according to the method of Kunkel (Kunkel, 1985). The cDNA encoding the plgR was subcloned into the vector M13mp8 as described previously (Mostov et al., 1986). Unless otherwise mentioned, mutations were constructed using a mutant DNA generated by insertion of a stop codon instead of 1760 as a starting template. In frame deletion of amino acids 656–658 was performed using a mutagenic oligonucleotide that was complementary to nucleotides 2131–2142 '5'-CCG CAC AGG AGG GAC AAC CGA GTA GTT 3' to the deleted segment. In frame deletions of Ser664 or Ile665 were performed in the context of the full-length plgR. For deleting Ser664 we used an oligonucleotide complementary to nucleotides 2155–2166 and 2170-2182 of the plgR cDNA. An oligonucleotide complementary to nucleotides 2158–2169 and 2173–2185 was used to delete Ile665. The mutagenic oligo-
nucleotide (5'-CCG CCC AGG AGG GAC AAC CGA GTA GCT TCC ATC GCA-GA 3') was used to scramble the order of the sequence NVDR (659–661) to DNVR (the nucleotides encoding the DNVR residues are underlined). The sequences of primers for truncation of residues (i.e., mutating the relevant codon to a stop codon) and alanine substitutions are available on request. The mutagenized sequences were confirmed by dideoxy sequencing using the Sequenase kit (United States Biochemical Corp., Cleveland, OH). For the expression of mutants, the replicative form phage DNA was isolated and the cDNA fragment was prepared by en-
donuclease BglIII digestion and agarose gel electrophoresis. The isolated cDNA fragment was then ligated into the BglIII sites of the cytomegalovirus promoter-based expression vector, pCB6 (Brewer and Roth, 1991), using standard techniques (Maniatis et al., 1982).

MDCK strain II cells were transfected by calcium phosphate precipita-
tion, followed by selection with G418 sulfate (GIBCO-BRL, Gaithersburg, MD) (0.7 mg/ml in growth medium for 14–18 h). Individual clones were isolated with glass cylinders, grown to confluence in plastic dishes, and tested for plgR expression by metabolic labeling and immunoprecipitation as described previously (Breitfeld et al., 1989b). The polarity of all clones was determined by methionine uptake and secretion of an endogenous gly-
coprotein complex (Urban et al., 1987; Casanova et al., 1991; Okamoto et al., 1992). All cell lines were assessed to be similarly polarized. Addition-
ally, to obtain a population of pooled clones, \( \sim 50-200 \) clones on a single 10-cm dish were grown to confluence. All experiments were performed using cell lines of passage number 2 through 15 after cloning.

Cell Culture

MDCK (strain II) cells were grown in MEM (CellGro; Mediatech, Washington, D.C.) supplemented with 5% FBS (HyClone, Logan, UT), 100 U/ml penicillin, 100 U/ml streptomycin, and 0.25 \( \mu \)g/ml Fungizone in 5% CO\(_2\), 95% air. In all cases, cells (>2 x 10\(^5\) cells/ml) were cultured on Transwell polycarbonate filter units (0.4-μm pore size; 24-mm diameter; Costar Corp., Cambridge, MA) for 4–5 days before the experiments (Breitfeld et al. (1989b). Fresh medium was added every other day after plating onto filters.

Vectorial Delivery of the plgR

Normally the plgR is delivered directly from the TGN to the basolateral surface. The plgR is then endocytosed and sorted into transcytotic vesicles that carry it to the apical surface (Breitfeld et al., 1989a). After reaching the apical surface an endogenous protease cleaves the extracellular, ligand

The plgR basolateral sorting signal is one of the best defined examples of a basolateral signal that is independent of endocytosis. As little is known about the sequence require-
ments and structural features of such signals, we performed a detailed mutational analysis of the 17-residue signal in order to identify individual amino acids that might play an important role in basolateral targeting of the plgR. We used several strategies, including progressive truncations from the COOH terminus, alanine scanning mutagenesis, and internal deletions. By using high-resolution two-dimensional \( \text{H} \) NMR spectroscopy we also assessed the nature of potential structural elements in a synthetic 17-mer peptide corresponding to this signal.

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binding domain and releases this domain into the apical medium. This cleaved fragment is called secretory component (SC). With the wild-type plgR +15% of the molecules leaving the TGN do not reach the basolateral surface. SC isoforms are targeted directly to the apical surface and then cleaved to SC. The basic principles of the protease sensitivity assay for vectorial delivery of newly synthesized plgR to the basolateral surface were described earlier (Casanova et al., 1991). We have modified the assay to more accurately and quantitatively data (Apodaca et al., 1993). A pair of filters containing confluent monolayer of MDCK cells were used for each determination. Cells were starved in MEM lacking cysteine (obtained from UCSP Cell Culture Facility; supplemented with 5% dialyzed FBS) for 15 min at 37°C. Then, the cells were pulse-labeled for 15 min at 37°C in starvation medium supplemented with [35S]cysteine (1,130 Ci/mmol, 10.5 mCi/ml; New England Nuclear, Boston, MA), to a final concentration of 1.75 μCi/μl, and washed three times quickly with warm MEM-BSA (MEM containing Hank's balanced salts, 20 mM Hepes, pH 7.3, antibiotics, and 0.6% BSA). Both filters were chased for 90 min at 37°C, to allow the cohort of labeled receptors to reach the cell surface. The first filter, which served as control, was chased in the presence of MEM-BSA (1 ml in the apical and basolateral compartments). The second filter was chased in the continuous presence of MEM-BSA supplemented with V8 protease (25 μg/ml; Boehringer-Mannheim Biochemicals, Indianapolis, IN) at the hasolateral medium and MEM-BSA without added protease in the apical compartment. The administration of V8 protease yielded more consistent and accurate results than the previously used TPCK-trypsin. At the end of the chase period, both the apical and basal media of the first (control) filter and the apical medium of the protease-treated filter were harvested and SC was immunoprecipitated as described below. Filters were washed three times with ice-cold MEM-BSA containing 0.5% FBS, excised, and subjected immmediately to immunoprecipitation as described below. For all mutants described here we performed 10-13 analyses, each of which consisted of a control filter and a V8-treated filter.

**Immunoprecipitation**

Cells on filters were boiled for 5 min in 1 ml of 0.5% SDS lysis buffer (0.5% SDS, 150 mM NaCl, 5 mM EDTA, 100 U/ml Trasylol, 20 mM Tris[hydroxymethyl]aminomethane-HCl, pH 8.1), and vortexed vigorously for 15 min. Basal and apical media were transferred to a microfuge tube containing 30 μl of 10% SDS, and boiled for 5 min. Triton dilution buffer (0.5% SDS, 0.1% Triton X-100, 100 mM NaCl, 5 mM EDTA, 100 U/ml Trasylol, 0.1% Na2MoO4, 2 mM PMSF, 50 mM N-trimethylaminoacetic acid-HCl, pH 8.6) was then added to each sample. Samples were pre-cleared twice with Sepharose CL-2B (Bietrifeld et al., 1989b), and plgR molecules were immunoprecipitated by overnight incubation (4°C; continuous end-over-end rotation) with 30 μl of protein G-Sepharose (15% slurry; Sigma Immunochemicals, St. Louis, MO) to which polyclonal sheep anti-SC IgG had been covalently coupled using 20 μg/ml N-hydroxysuccinimide ester (Gentest Laboratories, Woburn, MA). Beads were washed three times with "mixed micelle buffer" (1% Triton X-100, 0.2% SDS, 150 mM NaCl, 5 mM EDTA, 8% sucrose, 0.1% Na3MoO4, 10 U/ml Trasylol, and 20 mM triethanolamine-HCl, pH 8.6) followed by one wash with "final wash" buffer (i.e., mixed micelle buffer lacking detergent) and prepared for SDS-PAGE as described (Bietrifeld et al., 1989b; Casanova et al., 1991).

**Quantitative Analysis of Vectorial Delivery**

Fig. 2 depicts a typical delivery assay performed on the T870-stop mutant receptor. The radioactivity levels of bands corresponding to plgR and SC immunoprecipitated from cells and media, respectively, were determined by a PhosphorImager (Molecular Dynamics, Sunnyvale, CA). The total amount of radiolabeled plgR immunoprecipitated from V8 non-treated cells (T(plgR<sup>V8</sup>)<sup>-</sup>) over the 90-min chase at 37°C was determined. T(plgR<sup>V8</sup>)<sup>-</sup> = plgR<sup>V8</sup> + A[SC]<sup>V8</sup> + [BSC]<sup>V8</sup>, where plgR<sup>V8</sup> is the radioactivity level measured for plgR immunoprecipitated from V8 non-treated cells, and [A[SC]<sup>V8</sup> + [BSC]<sup>V8</sup>] are radioactivity levels measured for SC immunoprecipitated from the apical and basolateral media of the same cells, respectively. The amount of radiolabeled plgR in cells was detected under equivalent experimental conditions but in the continuous presence of V8 protease at the basolateral medium is reduced (compare label intensities of bands sized receptors delivered to the basolateral surface was determined as follows:

**Basolateral Delivery (%)** = \(\left(1 - \frac{T_{plgR^{V8}}}{T_{plgR^{V8} - plgR^{V8}}} \right) \times 100\)

SC comprises the major portion of the extracellular ligand-binding domain of the receptor and it contains all but one of the cysteine residues labeled with 35S. Once plgR molecules arrive at the apical surface they are efficiently cleaved by an endogenous protease to SC which is released into the apical medium (Casanova et al., 1991). This is in contrast to the basolateral endogenous protease which acts inefficiently, releasing small amounts (<5% of the total label; see Fig. 2, lane 5) of SC to the basal medium. The high efficiency of plgR cleavage at the apical surface was confirmed in control experiments in which V8 protease was included in the apical medium during the entire chase period. In all cases, the apically added V8 had no effect on the amount of plgR immunoprecipitated from cells (compare the intensities of plgR bands in lanes 3 and 5 in Fig. 2), indicating that there was not a substantial pool of plgR that had reached the apical surface and had not been cleaved to SC. Apcal SC can in principle have two sources: plgR delivered directly from the TGN to the apical surface and plgR that was first delivered to the basolateral surface and then transported to the apical surface via endocytosis. However, direct apical targeting was assessed by measuring the amount of apically released SC in the presence of basolateral V8 (band [A[SC]<sup>V8</sup> in lane 6) which effectively eliminated the population of transcytosing plgR. Therefore, the fraction of plgR delivery from the TGN to the apical plasma membrane was calculated as follows:

**Apcal Delivery (%)** = \(\left(\frac{[A[SC]^{V8}]}{T_{plgR^{V8} - plgR^{V8}}} \right) \times 100\)

In experiments performed on mutant receptors in the context of the truncated T870-stop receptor, the average amount (of many experiments) of SC immunoprecipitated from the apical medium of V8 non-treated cells was roughly comparable to that released into the apical medium of V8-treated cells. This can be directly attributed to the fact that all plgR mutants were severely blocked in endocytosis and subsequent transcytosis. The fact that V8 treatment largely did not affect direct apical delivery of plgR also suggests that V8 had no general effects on receptor traffic from the Golgi apparatus to the plasma membrane, and that V8 does not cross the tight cell monolayer from the basolateral to the apical compartment. The latter notion is also favored by the following two observations. First, the trans-monolayer electrical resistance of monolayers treated with basolateral protease (lane V8) was not significantly different from non-protease-treated monolayers (data not shown), implying that the tight junctions are intact and maintain their barrier activity in the presence of the protease. Second, if either basolateral or apical medium that contained protease is subjected to immunoprecipitation (at the end of the 90-min chase and after boiling with SDS to inactivate the protease) with anti-SC antibodies, a ladder of bands with molecular weights smaller than that of SC is observed. These bands presumably represent degradation products of receptors that arrived at the cell surface during the chase and were cleaved by the V8 protease. In experiments where V8 was added to the basolateral medium, if the protease had leaked into the apical medium and cleaved plgR at the apical surface (or cleaved SC in the apical medium) we would have expected to find this ladder in the immunoprecipitates from the apical medium. However, in all experiments where SC was immunoprecipitated from the apical medium of monolayers that were treated with basolateral protease, neither this ladder nor any other bands except for SC were ever observed, suggesting that such leakage did not occur.

The advantage of this protease-based delivery assay over surface biotinylation-based assays is that it directly measures the integrated amount of plgR inserted into the basolateral and apical plasma membranes over the entire chase period. Conceiviably, rapid endocytosis and transcytosis may enable receptor to avoid the exogenous protease, a phenomenon which might lead to inaccurate determination of basolateral delivery. However, as indicated above, all mutants were severely blocked in endocytosis (but not basalateral delivery), causing receptor accumulation on the basolateral cell surface and its continual exposure to the V8 protease.

**Peptide Synthesis**

Peptides representing the wild-type 17-mer basolateral targeting signal of plgR (residues 653-669; see Fig. 1 a), and the two mutant signals were synthesized on a Milligen/Bioresearch 9050 automated peptide synthesizer, using FMOC synthetic chemistry (Stanford University Medical Center, Protein and Nucleic Acid Facility, Beckman Center). All peptides synthesized were purified by preparative reverse-phase high pressure liquid chromatography (preparative column was a Millipore 25 mm × 10 cm C18,

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2D NMR

Proton NMR spectra were acquired using a spectrometer (GN-500; General Electric Co., Wilmington, MA). Experiments were run in the phase-sensitive mode with quadrature detection in both dimensions. Spectral widths were 6,024 Hz. All data processing and analysis were performed using software developed for a Sun work station by M. Day and D. Kneller in the laboratory of I. D. Kuntz (University of California, San Francisco, CA). Base-line correction has been described (Basus, 1984). Sample concentrations were approximately 7-9 mM in 10% D2O/90% H2O and included 1 mM sodium (trimethylsilyl)propionate (TSP) as an internal reference. Sample pH was adjusted to 4.6, without correction for the deuterium isotope effect, by addition of small amounts of NaOH and HCl.

For homonuclear Hartmann-Hahn (HOHAHA) spectra with MLEV-17 (Bax et al., 1982) the spectrometer was modified by inclusion of a 6-W amplifier as the transmitter with a γH of 8 kHz as described (Basus et al., 1988). A 75-ms mixing time was used. The H2O signal was suppressed with the DANTE sequence (Morris and Freeman, 1978). The period (4 s) incorporated the recycle time. The number of t1 increments was 512, each with four acquisitions and 4,096 data points in t2. One dummy scan was performed prior to each t1 increment. The data, in both dimensions, were modified by a Gaussian function with a line broadening component of −1.5 Hz and a multiplier of 0.1. The spectra were zero-filled once in each domain.

The temperature coefficients of the amide chemical shifts for P-WT and P-660A were determined from HOHAHA spectra acquired at 2, 5, 10, 15, 20, 25, 30, and 35 or 40°C.

Nuclear Overhauser effect (NOESY) spectra were acquired at 5°C using the procedure of States et al. (States et al., 1982). A 200-ms mixing time was used. The H2O resonance was selectively removed by use of a jump-and-return read pulse, plus a short (3 ms) pulsed field gradient at the start of the mixing period (Basus, 1984; Plateau and Guéron, 1982). A recycle time of 2.5 s was used. Spectra were acquired with 1,024 t1 increments, each with 16 acquisitions and 4,096 data points in t2. One dummy scan was performed before each increment. The data were multiplied with a squared sine-bell function shifted 50° in t2 and 80° in t1. The spectra were zero-filled once in the t1 domain and twice in the t2 domain.

Results

Identification of Specific Residues That Comprise the Basolateral Sorting Signal of the plgR

To identify specific residues required for basolateral sorting, we systematically mutated the 17-residue basolateral sorting signal in the cytoplasmic domain of the plgR. The mutagenesis strategies and designation of each mutant receptor are summarized in Fig. 1. Mutant cDNAs were transfected into MDCK cells, and cell lines stably expressing mutant receptors were generated. All amino acid substitutions and most other mutations were constructed in the background of the mutant receptor T670-stop (i.e., the codon for T670 is replaced by a stop codon), which truncates the cytoplasmic domain at residue 669. Since the 17-residue cytoplasmic segment contains the information required for efficient basolateral targeting, we used the T670-stop template as a simplifying model to dissect the basolateral sorting signal. Moreover, we employed this strategy to eliminate possible contributions from signals that might exist COOH-terminal to the 17-mer segment, between residues 670 and 755.

Vectorial delivery of each mutant receptor from the TGN to the basolateral surface was quantitatively determined by combining a pulse-chase procedure with protease treatment (Apodaca et al., 1993) as detailed in Materials and Methods and Fig. 2. MDCK cells expressing mutant plgRs were grown on polycarbonate Transwell filters. Receptors were metabolically labeled for 15 min with [35S]Cys, and delivery of these newly synthesized (radiolabeled) receptors to the cell surface was followed over a 90-min chase period. The radiolabeled receptors were either chased in plain medium (control cells) or chased in the presence of V8 protease in the medium bathing the basolateral surface. In the latter case, plgR molecules delivered from the TGN to the basolateral surface were proteolyzed and could not be immunoprecipitated as full length plgR. Consequently, a reduction in radiolabeled receptor was observed (Fig. 2), reflecting the integrated amount of receptors that arrived at the basolateral surface over the chase period. In contrast, receptors delivered directly to the apical plasma membrane were not proteolyzed by an exogenous protease (which is absent from the apical medium), but instead were efficiently cleaved by an endogenous apical plasma membrane protease to SC that was released into the apical medium. As SC comprises the major part of the radiolabeled plgR, SC immunoprecipitated from the apical medium reflects the fraction of plgR directly targeted to the apical surface. In all cases, 80–85% of the newly synthesized plgR was chased to the cell surface (both apical and basolateral) at the end of the 90-min incubation. Thus, whenever a mutation caused a decrease in basolateral targeting, a proportional increase in apical delivery was detected. In addition, although expression levels varied from clone to clone over a two- to fourfold range, we found that the results described below did not vary significantly with expression levels.

Successive Truncations

Figure 1. Mutagenesis of the 17-residue basolateral sorting signal. The cytoplasmic amino acids comprising the 17-residue basolateral sorting signal of the plgR are depicted using the single letter code. Oligonucleotide-mediated mutagenesis was used to construct the mutant receptors. Single alanine substitutions (e.g., substitution of arginine at position 655 to alanine) were all constructed in the context of the truncated T670-stop receptor. Truncation of residues was performed by mutating the codon that encodes for each of the indicated residues (e.g., threonine at position 670) to a stop codon (termed T670-stop).

Alanine Substitutions

of the segment 656-HRR-658 (HRR656-658-Δ), and the mutant in which the sequence 659-NVDR-662 was shuffled to DNRV (NVDR659-662-DNRV) were all constructed in the context of the truncated T670-stop receptor. Truncation of residues was performed by mutating the codon that encodes for each of the indicated residues (e.g., threonine at position 670) to a stop codon (termed T670-stop).
Figure 2. Determination of biosynthetic targeting to the basolateral cell surface by protease-based delivery assay. Filter-grown MDCK cells expressing the T670-stop mutant receptor were pulse labeled with [35S]cys and chased in MEM/BSA in the absence of label and in the absence (lanes 1, 4, and 5) or in the continuous presence of V8 protease in either the basal (lanes 2 and 6) or the apical chamber (lane 3). The plgR was immunoprecipitated from cell lysates and soluble SC was immunoprecipitated from the apical (A) and basal (B) media. Immunoprecipitates were analyzed by SDS-PAGE and fluorography. The radioactivity levels of the relevant bands were determined by a PhosphorImager and designated as: plgR(VS-); [A]SC(VS-); [B]SC(VS-); [A]SC(VS+). Basolateral delivery percent is the fraction of radiolabeled plgR proteolized by the basolaterally administered V8 protease. Apical delivery percent is the fraction of plgR cleaved to SC at the apical surface and in the presence of basolateral V8 as indicated under Materials and Methods. In this case, basolateral delivery and apical delivery are 75 and 7%, respectively.

Initially, we resolved individual residues involved in basolateral targeting, by systematically truncating the 17-mer segment from the COOH terminus (Fig. 3). Truncation at R669 (R669-stop) had no effect on basolateral targeting compared to the mutation T670-stop. Truncation at Y668 (Y668-stop) yielded a small but significant decrease in basolateral targeting (from 75 to ~60%) which was accompanied by a corresponding increase (from 6 to 25%) in apical cleavage of plgR to SC. The greater sensitivity of apically released SC is due to the low baseline level of apical targeting (~6%) exhibited by the unimpaired mutant receptors. Truncations at residues S667 and G666 had no further effect compared to Y668-stop. Note that while almost all of the residues comprising the 17-mer signal are highly conserved across species, glycine at position 666 is not conserved; serine and arginine residues occupy the glycine position in rat and human plgRs, respectively (Banting et al., 1989; Krajci et al., 1992). In addition, mutation of S667 to an alanine in the full-length plgR had no effect on any of the receptor's phenotypes, including basolateral targeting (Casanova et al., 1990). Taken together, these data suggest that S667 and G666 are not essential for basolateral targeting of the plgR. A further decrease in basolateral delivery was observed upon truncation at residues I665 or S664. Residues Y668, I665, and S664 are thus potentially implicated in basolateral sorting of the plgR.

The functions of residues Y668, I665, and S664 were further examined by substituting each amino acid with an alanine in the context of the truncated T670-stop mutant. Alanine was chosen as it is less likely than other residues to impose electrostatic or steric effects, or to alter the main chain conformation (Cunningham and Wells, 1989). In contrast to the results obtained by the truncation studies, the results summarized in Fig. 4 show that the relatively conservative mutation I665-A or the less conservative change of S664-A had virtually no effect on basolateral delivery of the plgR. In contrast, substituting Y668 for an alanine resulted in a relatively minor reduction in basolateral delivery, with values similar to those obtained for the truncated Y668-stop mutant. Therefore, a tyrosine at position 668 appears to be important for basolateral sorting, while the role of S664 and I665 remains unclear, as the Ala mutations and truncations yielded contradictory results.

To resolve this discrepancy we made individual internal deletions of S664 or I665. These were made in the context...
of a pIgR with a full-length cytoplasmic tail, so as to minimize possible secondary effects due to conformational instability of shortened peptides or proximity of the negative charge contributed by the COOH terminus (see Discussion). Both of these mutants revealed efficient basolateral delivery with values comparable to the wild-type receptor (i.e., 80 ± 4% [mean ± S.E., n = 10] of the newly synthesized receptor is sorted directly from the TGN to the basolateral surface and 16 ± 1.5% is targeted to the apical surface). These data suggest that S664 and I665 are not essential for basolateral sorting.

We next constructed mutations that affected relatively large portions of the cytoplasmic domain in the region between R653 and V660. The positively charged cluster HRR656-658 was deleted in the context of the T670-stop template. The basolateral delivery of this mutant (designated HRR656-658-Δ) was significantly inhibited. About 45% of the mutant receptor was directed to the basolateral surface and 40% to the apical plasma membrane (data not shown). Similar results were obtained upon scrambling the sequence NVDR659-662 to DNRV659-662 (NVDR659-662-DNRV). This indicates that the order of residues between N659 and R662 is critical for basolateral targeting.

To define specific residues that mediate basolateral targeting in the region of R655-V663, we systematically replaced individual amino acids with an alanine. Residue R653, which abuts the transmembrane region, was not mutated as its positively charged side chain probably locks the transmembrane domain of the receptor in the lipid bilayer. R653 is followed by residue A654 in rabbit and human pIgRs (Banting et al., 1989; Krajci et al., 1992). The rat pIgR, however, has a valine residue at 654 (Banting et al., 1989), suggesting that this residue may not be essential for basolateral delivery. The results of the alanine-scanning mutagenesis (Fig. 4) demonstrate that substitutions of H656, R657, or V660 yield mutant receptors with non-polarized expression. Approximately equal amounts (40–45%) of the radiolabeled mutant receptors reached the apical and the basolateral surfaces. Similar data were obtained when mutations H656-A, R657-A, and V660-A were constructed in the context of the full-length wild-type tail (Aroeti, B., and K.E. Mostov, unpublished results). These results further support the involvement of those residues in basolateral targeting of the pIgR, and suggest that their sorting activity is not affected by the remainder of the cytoplasmic tail. They also imply that the effects that the indicated Ala substitutions had on the targeting of the T670-stop receptor are probably not due to conformational instability of the shorter tail.

Similar to Y668, substitution of R655 with an alanine yielded a smaller but significant effect on basolateral delivery, with only ~60% of the receptor targeted to the basal surface and 15–20% delivered apically (Fig. 4). In summary, our results suggest that H656, R657, and V660 are major determinants for basolateral targeting of the pIgR, whereas R655 and Y668 play a lesser, but yet significant role in the sorting process.

### 2D-NMR Secondary Structure Analysis of Synthetic Peptides Corresponding to Wild-type and Mutant 17-mer Basolateral Sorting Signals

We acquired two-dimensional proton NMR spectra of three synthetic 17 residue peptides in an attempt to characterize conformational features that might be relevant to pIgR basolateral targeting. One of these peptides (P-WT) corresponds to residues 653-669 of the pIgR, while the other two contain point mutations that were found to inhibit significantly basolateral targeting. For one of these peptides (P-R657A), alanine replaced arginine at the position corresponding to residue 657 of the receptor and for the other peptide (P-V660A), alanine replaced valine at the position corresponding to residue 660. The spectra were obtained at 5°C to stabilize as much as possible any ordered conformations and at pH 4.6 to minimize amide proton exchange with water. The relatively low pH introduces the possibility that the ionization state of H656 is not the same as found in vivo.

Proton resonance assignments for spin systems characterized by intra-residue through-bond connectivities were obtained from HOHAHA spectra (Bax et al., 1982). NOESY spectra (Wagner et al. 1981) were acquired to obtain sequential inter-residue through-space connectivities, that is C=H-
NH\textsubscript{i+1}, C\textsubscript{Hi}-NH\textsubscript{i+1}, and NH\textsubscript{i}-NH\textsubscript{i+1} NOEs. Here, i and i+1 signify residues adjacent in sequence. Although arginine accounts for about one-third of the amino acid composition in each peptide, the resonances of the arginines were not completely degenerate and each arginine spin system was uniquely assigned. For example, in the P-WT NOESY spectrum, unique C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs were found for the NH\textsubscript{i} and COOH-terminal arginines and two C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs were readily identified for all tri-residue sequences containing an arginine as the middle residue, with the exceptions of H656-R657-R658 and R657-R658-N659. For these two arginines, the C\textsubscript{Hi} resonances are nearly degenerate so that the intra-residue C\textsubscript{Hi}-NH cross peak of R658 overlaps the C\textsubscript{Hi}-NH\textsubscript{i+1} NOE of R657 and R658. However, unique C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs were found for residues H656 and R657 and R658 and N659 in the NOESY spectrum of P-WT (and of P-V660A). Along with unambiguous C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs found for sequential residue pairs, the assignments were confirmed in many cases by identification of C\textsubscript{Hi}-NH\textsubscript{i+1} and NH\textsubscript{i}-NH\textsubscript{i+1} NOESY cross peaks. Fig. 5 shows the fingerprint and the NH-NH regions of the P-WT NOESY spectrum. Additional NOEs belonging to C\textsubscript{Hi}-NH\textsubscript{i+2} connectivities are also seen in the fingerprint region. The chemical shift assignments for P-V660A and P-R657A are entirely consistent with each other and with those of P-WT. Assignment diagrams and summaries of all inter-residue NOEs for the three peptides are shown in Fig. 6.

While the strong C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs found in the P-WT spectrum are typical of a peptide without ordered structure, the presence of NH\textsubscript{i}-NH\textsubscript{i+1} NOEs and medium range cross peaks (Fig. 5) indicates that P-WT also has a propensity to adopt ordered conformations that interconvert with disordered conformations. This interconversion must occur rapidly as only one complete set of resonances are found. For the following discussion, it is convenient to consider separately three segments of P-WT: residues R653-R658, residues R658-D661, and residues D661-R669.

There is considerable overlap of the C\textsubscript{Hi} and NH resonances for residues R655-N659 of P-WT (Fig. 5 a). This overlap precludes identification of NOEs indicative of ordered structure. With the exception of a relatively weak NH\textsubscript{i}-NH\textsubscript{i+1} NOE between H656 and R657 (Fig. 5 b), no other backbone-backbone cross peaks, associated with ordered structure, were identified. Thus, no correlation can be made between ordered conformations involving residues H656 and R657 of P-WT and the mutational studies that showed these two residues are important for proper basolateral targeting.

Beginning with the residue pair, R658/N659, NH\textsubscript{i}-NH\textsubscript{i+1} NOEs were identified for all subsequent pairs of residues except in one case where chemical shift degeneracy precluded identification (Fig. 6 a). Of these NOEs, the cross peak between residues V660/D661 is approximately twice as intense as all others, indicating closer average approach of those amide groups to each other. The identification of this NOE along with the weaker NH\textsubscript{i}-NH\textsubscript{i+1} NOE between N659/V660 and the presence of a C\textsubscript{Hi}-NH\textsubscript{i+2} NOE between N659 and D661 suggests that residues 658 to 661 have a significant tendency to adopt a β turn (Wüthrich et al., 1984; Dyson et al., 1988b). This turn is tentatively assigned as type I based on the relative intensities of the two NH\textsubscript{i}-NH\textsubscript{i+1} NOEs. The residue V660, shown by mutational studies to be important in basolateral targeting, would reside in position i+2 of such a turn.

A series of NH\textsubscript{i}-NH\textsubscript{i+1} NOEs, such as that found for residues 658 to 669, is characteristic of the so-called nascent helix (Dyson et al., 1988b). This type of structure was originally described for a peptide (derived from a helical region of myohemerythrin) as a conformational ensemble of turn-like structures and which upon addition of trifluoroethanol was stabilized as a helix with long-range order. Consecutive NH\textsubscript{i}-NH\textsubscript{i+1} NOEs are expected for helical structures, but neither C\textsubscript{Hi}-NH\textsubscript{i+2}, C\textsubscript{Hi}-NH\textsubscript{i+1}, nor C\textsubscript{Hi}-NH\textsubscript{i+3} cross peaks diagnostic of a regular α- or 3-10-helix with long-range order were identified in the spectrum of P-WT. Further, the circular dichroism spectrum of P-WT, recorded in the presence of 33% (vol/vol) trifluoroethanol at pH 4.6 and 5°C, was not characteristic of helix, showing only one minimum at 195 nm (spectrum not shown). Taken together, these observations suggest either that residues 658 to 669 are not part of a stable helix in the context of full-length plgR or that other medium and long-range interactions, which are not adequately mimicked by trifluoroethanol, are needed to stabilize helical structure between residues 658 to 669 of the plgR. On the other hand, the presence of multiple C\textsubscript{Hi}-NH\textsubscript{i+1} NOEs between residues 658-669 in conjunction with the NH\textsubscript{i}-NH\textsubscript{i+1} NOEs is clear evidence for transitory turn-like structures in P-WT.

As mentioned above, besides the NOESY spectrum of P-WT, the spectra of two peptides derived from the sequences of mutant plgR's were recorded. Comparison of NOESY spectra of peptides containing the endocytosis signal and mutated sequences for the LDLR, LAP, and the insulin receptor proved useful in correlating structure with function (Bansal and Gierasch, 1991; Eberle et al., 1991; Backer et al., 1992; Lehmann et al., 1992). As noted in the introduction, those studies correlated loss of a β-turn-like structure and function upon mutation. Replacement of R657 with alanine in P-R657A results in very little change in the NOESY spectrum (summarized in Fig. 6 b). Two additional NH\textsubscript{i}-NH\textsubscript{i+1} NOEs and an additional C\textsubscript{Hi}-NH\textsubscript{i+2} NOE between residues S667 and R669 were identified. But the NH\textsubscript{i}-NH\textsubscript{i+1} NOEs could not be found in the spectrum of P-WT due to resonance overlap. The weak C\textsubscript{Hi}-NH\textsubscript{i+2} NOE is also found in the spectrum of P-V660A (Fig. 6 c), indicating that it is not unique to the spectrum of P-R657A. Further, after normalizing the intensities of the NH\textsubscript{i}-NH\textsubscript{i+1} NOEs of P-WT and P-R657A to the N659 amide side chain exchange peak of each respective spectrum (data not shown), it was clear that no perturbation of identifiable conformations of P-WT occurs on substitution of an alanine at position 657. The normalized NH\textsubscript{i}-NH\textsubscript{i+1} NOE intensities are virtually the same in both spectra. Thus, alanine at position 657 does not disrupt structure distal to the substitution.

The NOESY spectrum of P-V660A is somewhat more informative than that of P-R657A. After normalizing the intensities of the NH\textsubscript{i}-NH\textsubscript{i+1} NOEs of P-WT and P-V660A in the manner described above, it was apparent that the relative intensity of the V660/D661 NOE decreases by approximately 50%, suggesting a destabilization in the putative β-turn-like conformation involving residues 658-661. Because of resonance overlap, it was impossible to determine if the C\textsubscript{Hi}-NH\textsubscript{i+1} cross peak of residues 659 and 661 is similarly affected. However, new cross peaks between C\textsubscript{Hi} of A654...
Figure 5. Portions of the 500 MHz spectrum of P-WT (pH 4.6, 5°C, 200 ms mixing time). (a) The fingerprint region. The connectivities between R653 to R657 and from R658 to R669 are traced out. Sequential C=H – NH,1 cross peaks are labeled with the appropriate residue numbers corresponding to those of the plgR. Intra-residue C=H – NH connecting sequential cross peaks are unlabeled. Also labeled are C=H – NH,2 cross peaks. The peaks labeled with an asterisk (*) could not be assigned. These cross peaks were not eliminated when the peptide was purified by cationic (Mono S) FPLC chromatography, followed by reverse phase chromatography suggesting that there is a second slowly interconverting conformer of P-WT present. The cross peak labeled with two asterisks (**) could not be unambiguously assigned and may be either an NOE between C=H of R657 and NH of V660 or one between C=H of R658 and NH of V660. (b) The NH – NH region. The connectivities between residues N659 to R662 and V663 to R669 are traced out. The labeling scheme is as above. This portion of the P-WT NOESY spectrum was contoured at a level twofold greater than that for the fingerprint region.
for P-WT and P-V660A (Fig. 7). A decrease in the temperature coefficient (−Δδ/ΔT) of an amide proton resonance is usually taken as a measure of protection of that proton from exchange with solvent due to (intramolecular) hydrogen bonding (Dyson et al., 1988a). For P-WT, the amide of D661 appears to be most protected from solvent. This observation is consistent with others, outlined above, that D661 would occupy the i + 3 position of a β-turn, and its amide proton would therefore be expected to hydrogen bond with R658 in position i of the turn. However, contrary to expectations, the temperature coefficient of the amide proton of D661 in P-V660A, did not increase but rather decreased further. Also, the temperature coefficient of R662 decreased significantly. The observed decrease in the temperature coefficient of the amide proton of D661 may be due to an increased propensity of the side chain carboxyl to hydrogen bond with the backbone amide. Nonetheless, the significant changes in the temperature coefficients of D661 and R662 indicate local alterations in structure and are therefore consistent with a decrease in stability of the proposed turn involving residues R658-D661.

Discussion

Sequence Requirements for the Basolateral Sorting Signal

We have systematically examined the sequence requirements for the pIgR basolateral targeting signal and its possible secondary structure in solution using several mutational strategies and two-dimensional NMR spectroscopy. The results of these experiments are summarized in Fig. 8.

Progressive truncations from the COOH terminus of a pIgR containing only the 17-residue basolateral signal suggested that Y668, I665, and S664 may be important for basolateral sorting. Substitution of Y668 with Ala confirmed
that this residue plays a role in basolateral sorting. Previously we found that basolateral sorting was not affected upon mutating Y668 to Cys in the full-length plgR (Okamoto et al., 1992). In contrast, as mentioned above alanine substitution of Y668 partially diminished basolateral targeting. It is possible that the Ala substitution at position 668 disrupted the basolateral sorting signal, whereas the Cys substitution did not. In addition, unlike the quantitative basolateral delivery assay applied in this study, the assay used to determine basolateral targeting of the mutant Y668 to Cys (Okamoto et al., 1992) was less quantitative and most likely not as sensitive in detecting small changes in basolateral targeting.

In contrast to the truncation results (Fig. 3), substitutions of I665 and S664 and Ala did not alter basolateral sorting, so the role of these residues was unclear. One possibility is that they are important, but the basolateral signal could tolerate an Ala at these positions. However, even though truncation of these residues had an effect on targeting, one cannot exclude the possibility that factors other than the truncated residues contributed to the altered phenotype. For example, at neutral pH the COOH-terminal carboxyl group bears a negative charge. Progressive truncation of residues from the COOH terminus changes the proximity of the charged moiety to possible sorting determinants residing in the non-truncated part of the tail. Consequently, the structure of the nontruncated region (and the sorting signal) might be disturbed. The negative charge may also interfere with interactions between the basolateral sorting machinery and a sorting signal possibly located in the non-truncated region. In both cases, a mutant receptor with impaired basolateral sorting activity might be observed even though the truncated residues are not part of the actual sorting signal.

To resolve the apparent contradiction between the Ala mutations and truncations at residues S664 and I665, we individually deleted these residues. We made these in the context of the full-length plgR to avoid possible complications from moving the charged C-terminus too close to the putative signal. Also we reasoned that the structure of the signal in the context of the full-length tail might be more resistant to a non-specific destabilization due to removal of a residue. In both cases these deletions did not disrupt basolateral sorting. Unlike Ala mutations, internal deletions might be expected to have a high propensity to disrupt structure, even if the deleted residues are not truly part of the signal. The finding that the deletions of S664 or I665 did not alter basolateral sorting therefore can be taken as particularly strong evidence that these residues are not part of the basolateral signal. In conclusion, the preponderance of data (from Ala mutations and internal deletions) suggests that these residues are not part of the signal. The effects of truncations at these residues are most likely due to secondary effects of the truncations.

We then analyzed the region from 656 to 663, primarily by alanine scanning mutagenesis. The alanine mutations that yielded the largest in basolateral sorting were at H656, R657, and V660 (see Fig. 4), and we conclude that these residues are particularly important for basolateral sorting. Alanine mutations at R655 and Y668 had smaller effects on basolateral sorting (Fig. 4), suggesting that these residues play a lesser role in this process. Obviously, one should keep in mind that alanine scanning only examines the ability of these positions to tolerate this particular substitution. Nevertheless, alanine scanning is widely accepted as a useful approach to identify functionally important residues.

Basolateral sorting seems to be dependent on the positively charged segment RHR655-657. The importance of polar and charged residues has also been suggested in previous studies on internalization and basolateral targeting signals (Kistakis et al., 1990; Trowbridge, 1991; Le Bivic et al., 1991; Matter et al., 1992). Interestingly, this positively charged cluster is located only two residues away from the transmembrane region of the molecule. In a previous study, the transferrin receptor internalization signal YXRF was shown to retain its activity only when separated by at least seven residues from the transmembrane region (Collawn et al., 1990). Our study suggests that the cytoplasmic residues RHR655-657 can be quite close to the plasma membrane and still mediate basolateral sorting of the plgR.

Negatively charged phosphatidylserine molecules are concentrated in the cytoplasmic leaflet of the plasma membrane (Alberts et al., 1989), and presumably also in the Golgi. The juxtamembrane positively charged cytoplasmic residues may serve to shield the sorting determinant from this negatively charged surface and thereby facilitate sorting. Alternatively, the positively charged residues may directly interact with components of the basolateral sorting machinery. Recent observations proposed that heterotrimeric G proteins play a role in targeting from the TGN to the basolateral surface (Stow et al., 1991; Bomsel and Mostov, 1992; Pimplikar and Simons, 1993). This class of G proteins is known to be activated by positively charged peptides, such as the wasp venom mastoparan. An intriguing possibility is that the juxtamembrane basic residues of the basolateral sorting signal of the plgR interact with a G protein.

Are there homologues of the plgR basolateral targeting signal in other proteins? In a recent study, alignment of a 13-amino acid sequence implicated in basolateral sorting of the LDLR with the 17-mer sequence in the plgR revealed five consensus residues: H/R819, S/N820, D/S822, T/S825, S828 in the LDLR aligned with residues R658, N659, D661, S664, S667 in the plgR (Yokode et al., 1992). Residue V660, which functions in basolateral sorting of the plgR, aligns with Q821 in the LDLR, pointing to a lack of residue conservation at the indicated position for these two receptors. However, to test the possibility that glutamine may still substitute for valine in the sorting determinant of the plgR, we mutated V660 to glutamine in the context of the truncated T670-stop mutant. The mutant receptor V660-Q showed impaired
basolateral delivery with values similar to those observed for the mutant V660-A (basolateral delivery = 35 ± 5% and apical delivery = 48 ± 5% [mean ± S.E., n = 10]). Moreover, mutagenesis studies on the basolateral sorting signal of the LDLR demonstrated that three of the consensus residues are not required for basolateral targeting (Matter et al., 1992). Tyrosine 824, which is part of the distal basolateral sorting determinant of the LDLR, is not conserved; it is V663 in the plgR. Taken together, these observations argue against the partial homology suggested between the distal basolateral signal of the LDLR and the basolateral signal of the plgR.

In the case of the transferrin receptor the single cytoplasmic Tyr is important for endocytosis, but is not necessary for basolateral targeting (Hunziker et al., 1991; Dargemont et al., 1993). The cytoplasmic domain of the transferrin receptor also contains the sequence RQVD27-30, which resembles the tight-turn forming sequence RNVD658-661 in the basolateral signal of the plgR. Interestingly, when a 36-cytoplasmic residue segment encompassing the RQVD sequence (and the Tyr-internalization signal) was deleted, a 20% reduction in the basolateral distribution of the transferrin receptor in MDCK cells was observed (Dargemont et al., 1993). It is therefore plausible that RQVD is part of a basolateral signal in the transferrin receptor.

Possible Structure of the Basolateral Sorting Signal

To obtain structural insights into the basolateral sorting signal, we studied the solution structures of P-WT, a peptide corresponding to the 17-residue basolateral sorting signal. Two-dimensional NMR spectroscopy suggested that a plausible ordered conformation of the peptide contains two structural motifs. The sequence RNVD658-661 tends to adopt a β-turn possibly of the type I conformation, whereas residues COOH-terminal to that turn contain a nascent helix structure which appear to be less stable than that of residues 658-661. Of the three crucial residues identified by the alanine scan, one residue (V660) is in position i+2 of the turn. The other two residues (H656 and R657) immediately precede the turn and are in a region for which ordered structure could not be identified. Tyrosine 668, which plays a lesser role in sorting, is located in the nascent helix. Arginine 655 precedes the turn, and as for H656 and R657, the structure in this region could not be identified.

As far as we know, this is the first structural data for a basolateral targeting signal that does not overlap with an endocytosis signal. Intriguingly we found that this signal, like endocytosis signals, contains an apparent β-turn. Our results raise the possibility that β-turns are a common feature of basolateral signals, including both those that do and do not overlap with endocytosis signals. The ability of a particular signal to function as a basolateral signal and/or endocytosis signal might then be determined by the presence and location of specific side-chains. In particular, endocytosis signals contain a Tyr (or other aromatic residue) at position i or i+3, whereas the β-turn in the basolateral signal of the plgR lacks aromatic residues, but contains a bulky hydrophobic Val at position i+2.

The ordered conformations proposed here are low energy conformers for P-WT in solution and are thus the best available starting points for considering the in vivo structure of the sorting signal. However, several caveats must be considered. First, the peptide may assume a different conformation when anchored to the membrane via the membrane spanning segment. Indeed, Bansal and Gierasch reported that a peptide corresponding to the transmembrane domain and part of the cytoplasmic tail of the LDLR exhibited an enhanced stability of the nascent helix in the region of the tight-turn internalization signal (Bansal and Gierasch, 1991). Second, the 17-residue signal is ordinarily part of the 103-amino acid cytoplasmic domain of the plgR and may have a different conformation in the context of the complete domain. However, the observation that this 17-residue segment can be functionally transplanted to a heterologous protein and can act as an autonomous basolateral targeting signal (Casanova et al., 1991) makes it more plausible that the structural characteristics of the peptide determined here are biologically relevant. Third, this signal is presumably recognized by another protein(s) that forms part of the sorting apparatus. Binding of this protein may alter the conformation of basolateral signal. In that case it will be extremely interesting to determine the bound conformation.

We also examined the effect of two Ala substitution mutations on the three-dimensional structure of the sorting signal peptide. For one of the mutations, R657-A, there was no detectable change in the β-turn and nascent helix regions of the peptide (Fig. 6 b). For technical reasons, the structural determination of the highly charged region (residues 653-657) of the peptide is less certain. If indeed there are no changes in the backbone structure of the signal, this might suggest that the side chain of the R657 is directly involved in recognition by a sorting molecule. In any case an important conclusion that can be drawn from these data is that the β-turn is not sufficient for basolateral sorting.

In contrast another mutation, V660-A, appears to destabilize the tight-turn but has no effect on the stability of the nascent helix conformation in the COOH-terminal region of the peptide, as shown by no significant alterations in the NH-NH$_{\text{H}^+}$ NOE intensities (calculations not shown). This result is similar to observations by Bansal and Gierasch and others on mutations in endocytosis signals that appear to destabilize the β-turn, but differs in that the nascent helix is not also destabilized (Bansal and Gierasch, 1991; Backer et al., 1992; Lehmann et al., 1992). In these situations, one does not know if the mutation acts by altering the secondary structure and/or by changing a residue that is directly involved in recognition of the signal. Although more work may be necessary to determine whether a β-turn is required for basolateral sorting, our data show that V660 is essential for both basolateral sorting and for the stability of the turn.

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