The Cytoplasmic Tail of Invariant Chain Regulates Endosome Fusion and Morphology

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The major histocompatibility complex class II associated invariant chain (Ii) has been shown to inhibit endocytic transport and to increase the size of endosomes. We have recently found that this property has a significant impact on antigen processing and presentation. Here, we show in a cell-free endosome fusion assay that expression of Ii can increase fusion after phosphatidylinositol 3-kinase activity is blocked by wortmannin. In live cells wortmannin was also not able to block formation of the Ii-induced enlarged endosomes. The effects of Ii on endosomal transport and morphology depend on elements within the cytoplasmic tail. Data from mutagenesis analysis and nuclear magnetic resonance-based structure calculations of the Ii cytoplasmic tail demonstrate that free negative charges that are not involved in internal salt bridges are essential for both interactions between the tails and for the formation of enlarged endosomes. This correlation indicates that it is interactions between the Ii cytoplasmic tails that are involved in endosome fusion. The combined data from live cells, cell-free assays, and molecular dynamic simulations suggest that Ii molecules on different vesicles can promote endosome docking and fusion and thereby control endosomal traffic of membrane proteins and endosomal content.

INTRODUCTION

Invariant chain (Ii) is a type II transmembrane glycoprotein known to have several important functions in antigen presentation (for a review see Germain, 1994; Nordeng et al., 1998). In the endoplasmic reticulum (ER) Ii trimers transiently associate with major histocompatibility complex (MHC) class II αβ 10 dimers. MHC class II molecules are transported from the ER to the endocytic pathway where they encounter endocytosed antigens. Two Leu-based-sorting signals in the cytoplasmic tail are independently sufficient for efficient endosomal targeting of Ii (Pieters et al., 1993; Bretnes et al., 1994; Odorizzi et al., 1994). During intracellular transport of αβ-Ii complexes, Ii is sequentially degraded from the luminal side (Blum and Cresswell, 1988). Partially degraded Ii remains trimeric and still associated with MHC class II, suggesting that the luminal region is not required for maintaining the nonameric αβ-Ii complex (Amigorena et al., 1995; Newcomb et al., 1996).

At high levels of expression, Ii has been reported to accumulate in enlarged endosomal compartments in transfected cells (Pieters et al., 1993; Romagnoli et al., 1993). In addition, Ii expression has been shown to cause a delayed endocytic transport of both antigens and membrane proteins in transfected cells (Romagnoli et al., 1993; Gorvel et al., 1995; Gregers, Nordeng, Gilje, Sandlie, and Bakke, unpublished data). These results demonstrate that Ii can modify the structure and function of the endosomal compartment, and this could influence antigen processing and presentation. A morphological study of Ii-induced large endosomal vesicle(s) (ILEV) in a human fibroblast cell line demonstrated that enlarged compartments containing early endosomal markers appeared as single membrane vesicles, whereas those containing late markers were generally filled with internal membranes (Stang and Bakke, 1997). Chimeric proteins consisting of the Ii tail fused to reporter molecules that formed tetramers did not induce vacuolation (Pieters et al., 1993; Bretnes et al., 1994). In addition, Ii constructs unable to trimerize were localized in normal-sized endosomes, showing that the luminal region of Ii was involved in Ii-
induced vacuolation (Gedde-Dahl et al., 1997). Nuclear magnetic resonance (NMR) studies of a synthetic Ii tail peptide have shown that it takes up an α-helix (Motta et al., 1995) and that this peptide at high concentration forms a trimeric aggregate (Motta et al., 1997). Together, these data indicate that Ii trimers are formed by interactions both in its cytoplasmic tail and in the luminal domain.

It has previously been reported that ILEV were not sufficient to avoid vacuolation. ILEV: large endosomal structures.

Table 1. Ii induced vacuolation

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Summary of Ii constructs used in this study. Whereas alanine substitutions had no effect on ILEV formation, mutations that changed the overall charge of the N-terminus (D(6)→R or D(3)→R) was sufficient to avoid vacuolation. ILEV: large endosomal structures.

* Motta et al., 1995.

MATERIALS AND METHODS

Recombinant cDNA Constructs

The cDNA fragment encoding the p33 form of human li (Ii,33) has been described previously (Bakke and Dobberstein, 1990). An overview of mutants used in this study is given in Table 1. The alanine (Aα)-scanned mutants (nos. 4–7 and 11) have been described previously (Motta et al., 1995). Point mutations or deletions in the cytoplasmic tail of li were introduced by polymerase chain reaction (PCR) mutagenesis using li,33 in the expression vector pSV51L (Bakke and Dobberstein, 1990) as template. The forward primers contained the point mutations or deletion and a Kozak translation initiation sequence for proper initiation of translation (Kozak, 1987). The PCR reactions were performed using Vent DNA polymerase (New England Biolabs, Beverly, MA) for primer extension. The final PCR products were cloned into pcDNA3 (Invitrogen, Carlsbad, CA). All mutants were verified by sequencing.

Iiwt and IiD6R in the heavy metal-inducible expression vector pMEP4 (Invitrogen) has been described (Nordeng and Bakke, 1999). The C-terminal endosome-binding domain of early endosomal antigen 1 fused to green fluorescent protein (EEA1-GFP, McBride et al., 1999) was a gift from Dr. H. Stenmark (Oslo, Norway).

Cell Culture

Madin-Darby canine kidney strain II (MDCK) and COS-1 cells were grown in complete medium: DMEM (Bio Whittaker, Walkersville, MD) supplemented with 9% fetal calf serum (Integro, Zaandam, Holland), 2 mM glutamine 25 U/ml penicillin, and 25 μg/ml streptomycin (all from Bio Whittaker) in 6% CO2 in a 37°C incubator.

Antibodies and Ligands

The high production in serum-free medium hybridoma BU45-HPSFI was kindly provided by Dr. S. Buus (Copenhagen, Denmark) and produces the mouse mAb BU45, which recognizes the luminal C-terminal part of human li (Wright et al., 1990). The hybridoma was grown in protein-free hybridoma medium II supplemented with 100 U/ml penicillin and 100 μg/ml streptomycin (all purchased from Life Technologies, Paisly, Scotland). BU45 is a mouse immunoglobulin (Ig) M mAb specific for the luminal C-terminal part of human li (Binding Site, Birmingham, UK), and the mouse mAb VicY1 recognizes the N terminus of the li tail (Quaranta et al., 1984) and was a gift from Dr. W. Kapp (Vienna, Austria). Goat anti-mouse antibody coupled to Alexa 488 was obtained from Mo-
Immunofluorescence Confocal and Video Microscopy

Transfected cells were transiently transfected by the DEAE-dextran method essentially as described by Huylebroeck et al. (1988). MDCK cells stably transfected with IiWT and IiWT-2A have been described (Nordeng and Bakke, 1999; Gregers, Nordeng, Gjilje, Saudlie, and Bakke, unpublished data). MDCK cells were transfected with EEA1-GFP by the DNA-calcium phosphate procedure, yielding stable clones as described elsewhere (Wigler et al., 1979). Clones expressing moderate levels of EEA1-GFP were chosen to avoid interference with the endosomal fusion machinery (McBride et al., 1999). Resistant clones were selected in the presence of 500 μg/ml G418 (Genetics, Nippon Becton Dickinson, San Jose, CA). Approximately 100% positive clones expressing EEA1-GFP were then double-transfected with Ii and selected with 300 μg/ml hygromycin B. Resistant clones were induced with 25 μM CdCl2 for 16 h to induce Ii expression. Double-positive clones were identified by immunofluorescence microscopy.

NMR Spectroscopy

NMR spectra were recorded on a DRX-500 spectrometer (Bruker, New York, DE) equipped with a triple resonance probe head and pulsed field gradients. Proton chemical shift assignments were performed using conventional two-dimensional experiments as described in Motta et al., 1997. Data processing was performed using Aurelia 2.0 software (Bruker Analytic GMBH, Reinstetten, Germany) running on Silicon Graphics computers (Mountain View, CA).

Structure Calculations

Model building and preliminary calculations were performed on a United Atom (UA) model (Weiner et al., 1984) with the Sybyl 6.5 package (Tripos, St. Louis, MO). The solvation effects were roughly included using a distance-dependent dielectric constant ε = r. In the UA model, distance restraints, as previously derived from NMR spectra of the wild-type 27-amino acid cytosolic Ii tail, were included as the C-C or C-N distance increasing the upper limits calculated for interproton distances of 1 Å. Semiparabolic penalty functions were used with force constants in energy minimization (EM) and room temperature molecular dynamics (MD) simulations. The Cα atoms were restrained at 1 kcal mol⁻¹ Å⁻² for distance restraints. In simulated annealing (SA) and preliminary MD runs, a time step of 1 fs, with no restraint on bond length applied, was used. SA calculations were performed with different lengths (from 10–250 ps), temperature (maximum values from 500–800 K, applied for 25–75% of the total time, cooling rates from 1–5 K ps⁻¹), and restraint force constant time profiles. After analysis of energy and violations, the best 40 structures were energy minimized, solvated in a periodic box of 5120 methanol molecules, and subjected to a 500-ps run of restrained MD simulations, followed by 1 ns of unrestrained MD run.

EM and MD simulations with solvent were performed with the Sander module of the Amber 4.1 package using the Amber/OPLS force field (Jorgensen, 1986; Jorgensen and Tirado-Rives, 1988). A time step of 2 fs, with rigid restraining of all bond lengths (SHAKE algorithm, Ryckaert et al., 1977) and periodic boundary conditions were applied. All MD simulations were performed in the isothermal-isobaric ensemble (300 K, 1 atm), with a solvent box that initially extended 8 Å from the most external solute atom on each side of the box. The final analyses were performed with specific modules of the Amber 4.1, Sybyl 6.5, and Molmol (Koradi et al., 1996) packages. Molecular structures have been drawn with Molmol and rendered with the POV-ray software.
Quantification of Cell-free Fusion of Endocytic Vesicles

Cell-free endosome fusion was quantified as previously described (Gruenberg et al., 1989; Bomsel et al., 1990; Gorvel et al., 1991). Briefly, 80% confluent cells were incubated for 30 min with 1.8 mg/ml biotinylated horseradish peroxidase (BHRP) or 3.6 mg/ml avidin. After washing with PBS/bovine serum albumin, cells were homogenized in homogenization buffer (250 mM sucrose, 3 mM imidazole, pH 7.4) by six passes through a 1.25-mm, 22-G needle, and postnuclear supernatants were prepared. Aliquots (50 μl) of the two postnuclear supernatants were combined in the presence of 50 μl of rat liver cytosol (10 mg/ml protein), adjusted to 12.5 mM HEPES (pH 7.4), 1.5 mM MgOAc, 3 mM imidazole (pH 7.4), 1 mM dithiothreitol, 50 mM KOAc, and complemented with 8 μl of an ATP-generating system (1:1:1 mixture of 100 mM ATP brought to pH 7.0 with KOH, 800 mM creatine phosphate, and 4 mg/ml creatine kinase) and 8 μl of a 1-mg/ml biotin-insulin stock solution (Bremnes and Bakke, unpublished data).

RESULTS

ILEV Formation Depends on the Ii Cytoplasmic Tail N Terminus but Not on Specific Amino Acid Residues

Transient expression of Ii in transfected COS cells causes the formation of enlarged endosomes (Figure 1). We have used this phenotype, defined as spheres with a diameter >1 μm, as an assay to study the structural requirements for vacuolation. The N-terminal 11 first amino acids of the Ii cytoplasmic tail has previously been shown to be required for ILEV formation (Pieters et al., 1993). A panel of Ii constructs with mutated cytoplasmic tails were therefore transiently expressed in transfected COS cells and subjected to immuno-fluorescence microscopy (Table 1). To identify the minimal region of the Ii cytoplasmic tail required for vacuolation, we first expressed two additional deletion mutants. IiΔ2–8 (no. 2) did not induce vacuolation, whereas expression of IiΔ2–9 (no. 3) did, but less extensively than Iiwt. This indicated that information for vacuolation was contained within the six first residues in the cytoplasmic tail of Ii, denoted as the Ii tailpiece. To investigate whether single amino acids within this region were required for vacuolation, we substituted all amino acids in positions 2–8 with Ala residues. Immunofluorescence and phase contrast microscopy analysis showed that all mutants induced vacuolation to similar extents as Iiwt (Table 1, nos. 4–7 and 11).

From a previous study it was reported that the sorting signal in the N-terminal region of the cytoplasmic tail, Leu7→Ile8, was not required for ILEV formation as long as the second signal was kept intact (Bremnes et al., 1994). The second Leu signal may also be replaced with tyrosine-based endosomal sorting signals (from the transferrin receptor or TGN 38) without affecting vacuolation, showing that vacuolation is not dependent on a specific endosomal sorting signal (Bremnes and Bakke, unpublished data).

Ii harbors four charged amino acids within the N-terminal tailpiece: three aspartic (Asp) residues and one arginine (Arg) giving a net negative charge in this region of –2. In addition, previous studies of Ii trafficking (Pond et al., 1995) have demonstrated a role for charged amino acids within the tailpiece of Ii. As noted in Table 1, Ala substitutions, except for construct 11, did not change the overall charge of the tailpiece and all still induced ILEV formation. The Δ2–5 truncation, although presenting a single negative charge in the tail, also formed ILEV. These data prompted us to ask whether the charge distribution rather than the specific amino acid sequence would determine ILEV formation. We studied the role of the charged residues by introducing amino acids that altered the overall charge distribution in the tailpiece or by inverting charges at specific positions. We observed that when the total charge of the N-terminal tailpiece was negative (nos. 3–10) we always observed vacuolation. In contrast, a total positive charge (nos. 17–21) always prevented ILEV formation. However, when the tailpiece had...
For construct 5 (IiR5A,D6A), Ala substitution cancels the wild-gously, the structure found for constructs 4 and 7 is helical. Although the tailpiece charge is now 0. Construct 14 (IiD3R) is that spans the region from Asp 3 to Leu 17 with a kink on.

We have reported that an Ii tail peptide takes up an -helix and the substitution induced vacuolation (Table 1) without affecting the structure of hexapeptides corresponding to constructs 14, 15, and Asp6 to Arg (no. 15) substitutions were sufficient to prevent vacuolation (Table 1; Figure 1). Similar results were obtained when Lys was introduced in position 3 or 6, suggesting that positive charges in these positions prevented vacuolation. Likewise, when the Asp in position 3 or 6 were replaced with another acidic residue, Glu, ILEV still formed (data not shown), indicating that the Asp residue per se was not required, but rather the presence of a negative charge appeared to be necessary. When both positions 3 and 6 were exchanged simultaneously for Ala residues (no. 16), vacuolation did not occur, indicating that Asp3 alone was not sufficient for vacuolation.

Although endosomal distribution of the constructs was confirmed by immunofluorescence microscopy (Figure 1), the inability of some mutants to form ILEV could in principle be due to inefficient transport to endosomes. Hence, transfected COS cells were subjected to metabolic labeling, immunoprecipitation, and Endo H treatment. Cells expressing Iiwt, IiD3R, and IiD6R all gained Endo H resistance, indicating that a fraction of the proteins was successfully transported out of the ER to the TGN (Figure 2). However, a control construct, IiD2R/D3R/D6R (no. 21) containing two double-Arg ER retention motifs (Schutze et al., 1994), was sensitive to Endo H, indicating that most of this construct did not reach the TGN. These data demonstrate that, except for IiD2R/D3R/D6R, mutations in the cytoplasmic tailpiece did not significantly affect the intracellular distribution of the resulting Ii constructs.

Predicting Vacuolation: The Importance of Available Negative Charges Not Involved in (i, i ± 3) Salt Bridges within the Ii Tailpiece

MD simulations of some of the above constructs helped to explain the mutation data. For the calculations we relied on the structure found for the wild-type 27-amino acid peptide. We have reported that an Ii tail peptide takes up an α-helix that spans the region from Asp3 to Leu17 with a kink on Pro15 (Figure 3A; Motta et al., 1995) and that it can form a triple-stranded α-helical bundle (Figure 3B) at high concentration (Motta et al., 1995). Constructs with internal Ala substitution induced vacuolation (Table 1) without affecting the helix and the −2 tailpiece charge. Figure 3C shows MD simulations of the 27-amino acid peptide corresponding to constructs 6 (Ii2A2A2A), for which a helix is observed. Analogously, the structure found for constructs 4 and 7 is helical. For construct 5 (Ii5A5A5A5), Ala substitution cancels the wild-type Asp3−Asp6−capping box (Forood et al., 1993), thus shortening the helix (Figure 3D). However, the tailpiece charge and the ILEV formation are not altered (Table 1), and the same is true for construct 11 (Ii11A11A11A11, Figure 3E). Although the tailpiece charge is now 0.

Construct 14 (IiD3R) is very interesting because it bears a neutral tail, for which vacuolation is not observed (Table 1), but MD simulations indicate the presence of a helix (Figure 3F). Finally, although vacuolation is not observed, a short helix, similar to that observed for construct 5 (Figure 3D), was predicted for constructs 17–21, all showing a positive tailpiece.

The above data rule out that vacuolation is linked to the helix, because its different length does not affect ILEV formation. What appears to be relevant is the presence of an available negative charge at either position 2, 3, or 6 not engaged in an i ± 3 electrostatic interaction. The electrostatic interactions observed in the Iiwt and constructs with a neutral tailpiece are reported in Table 2. It is interesting to observe that for constructs 14–16 all the negative charges within the tail are engaged in (i, i ± 3) salt bridges with Arg, whereas in constructs 11–13 at least one negative charge is available for further interactions (Table 2). Accordingly, whereas the former constructs do not vacuolate, the latter do form large vesicles.

It could be argued that the above electrostatic interactions are essentially due to restraints imposed on the N terminus by the presence of a helix. We therefore studied the solution structure of hexapeptides corresponding to constructs 14, 15, and 16 (MDQQRD-NH2, MDDQRR-NH2, and MDAQRA-NH2, respectively) in their amidated form to avoid the electrostatic contribution of the C terminus. NMR spectra indicated the presence of strong nuclear Overhauser effects between backbone amide protons of residues 3–4 and residues 4–5, as well as an effect between the α proton of Asp5 and the amide proton of Arg5 in all hexapeptides. They strongly suggest the presence of a type I β-turn comprising Asp5−Arg5 for all of them. Refinement of the structure by MD simulations highlighted the presence of a hydrogen bond between the Asp2 side chain and the amide proton of Gln4 backbone NH, as well as with the side chain of Arg5. Therefore, the isolated neutral tailpiece assumes a confor-
information that favors the network of electrostatic interactions present in wild-type Ii, independently of the secondary structure that follows. Altogether, the mutagenesis analysis and MD data show that ILEV formation does not depend on specific amino acids but on free negative charges within the Ii cytoplasmic tail, which are available for interactions required for vacuolation.

Vacuolation Depends on the Ii Expression Level

Ii-induced endosomal vacuolation would require that a substantial amount of Ii is transported to endosomes. We have shown above that the intracellular routing of Ii and the Ii constructs could not account for the differences in the ability to induce ILEV. However, it was also necessary to study in more detail ILEV formation in response to the expression level. To this end, MDCK cells were stably transfected with Iiwt or Ii D6R under the control of an inducible metallothionein promoter. Using this system, we previously demonstrated that protein expression was proportional to the concentration of CdCl2 in the 0–40 µM range (Nordeng and Bakke, 1999). The MDCK cells were incubated with various concentrations of CdCl2 overnight, metabolically labeled, and lysed, and Ii proteins were immunoprecipitated and subjected to SDS-PAGE. The amount of protein immunoprecipitated from the lysate was quantified as described in MATERIALS AND METHODS, and the protein expression level was related to the CdCl2 concentration. Cells were also grown on coverslips and incubated with the same concentration of CdCl2 as above and labeled for Ii. Figure 4 shows that the fraction of cells containing ILEV, the average number of ILEV per cell, as well as the average size of ILEV increased with augmented Iiwt expression. In contrast, endosomes were normal-sized when IiD6R was expressed at high levels similar to Iiwt. This shows that one specific positively charged amino acid in the Ii cytoplasmic tail is sufficient to impair the ability of Ii to induce ILEV.

Ii-induced Endosome Vacuolation and Fusion IsInsensitive to Wortmannin

The giant size of the ILEV depends on membrane supply, most likely obtained by endosome fusion, and we have found that ILEV are generated from a population of early endosomes (Bakke and Nordeng, Nordeng, Gregers, Kongsvik, Méresse, Gorvel, Jourdan, Motta, and Bakke, unpublished data). To study the role of early endosome fusion for ILEV formation, CV1 cells were double-transfected with Iiwt and rab5lle133, a dominant negative mutant that inhibits early endosome fusion (Gorvel et al., 1991). Whereas the normal phenotype of rab5lle133 is early endosome fragmentation, we found by confocal microscopy that ILEV were still formed in double-transfected cells, and conversely, IiD6R did not inhibit vacuolation induced by the constitutively active rab5lle133 mutant. This suggested that the Ii-induced vacuolation was uncoupled from the rab5-regulated endosome fusion. To study this further, we used wortmannin, a specific inhibitor of phosphatidylinositol 3 (PI3)-kinase that has been reported to block fusion of early endosomes in cell-free assays (Jones and Clague, 1995; Jones et al., 1998). We first studied Ii-induced vacuolation in the presence of wortmannin in live MDCK cells expressing inducible Iiwt or IiD6R. We chose to use IiD6R as a model for cells with normal-sized vesicles, because this mutant has been shown not to interfere with the transport capacity of Ii itself and MHC class II
Fusion efficiency was reduced to 50% in both control cells and cells transfected with IiD6R, whereas, the fusion efficiency was reduced by only 10% using endosomes from Iiwt-transfected cells. When Iiwt endosomes were fused with Hage endosomes in the presence of wortmannin, there was a 35% reduction in efficiency. The data show that Ii can complement for wortmannin in endosome fusion. The above data thus indicate that Ii is able to increase endosome fusion via a mechanism, which is able to complement for fusion processes inhibited by wortmannin.

DISCUSSION
The Ii-induced endosomal vacuolation and retention observed in transfected cells hints for a previously undescribed cell biological function of Ii. In this study we have attempted to elucidate the structural requirements for and the mechanism behind this effect of Ii expression. We have...
shown that the negative charges provided by acidic residues in the cytoplasmic tail of Ii are a prerequisite for ILEV formation. Acidic residues upstream of the membrane-distal Leu-based sorting signal of Ii have previously been shown to be necessary for this motif to work efficiently in endosomal transport (Motta et al., 1995; Pond et al., 1995; Simonsen et al., 1998a). However, the Leu-based motif per se is not required for vacuolation, because Ii constructs with the Leu7Ile8-based motif inactivated and the Met 16Leu17-based signal replaced by Tyr-based motifs were also vacuologenic (Nordeng, Gregers, Kongsvik, Mørresse, Gorvel, Jourdan, Motta, and Bakke, unpublished data).

Pond et al. (1995) found that when the eight N-terminal residues of the Ii tail, comprising the Leu\textsuperscript{7}Ile\textsuperscript{8} signal and the upstream acidic residues, were replaced by the corresponding region from the lysosomal integral membrane protein II, M6PR, or CD3\textgamma, vacuolation still occurred. However, the Leu-based motif per se is not required for vacuolation, because Ii constructs with the Leu\textsuperscript{7}Ile\textsuperscript{8}-based motif inactivated and the Met\textsuperscript{16}Leu\textsuperscript{17}-based signal replaced by Tyr-based motifs were also vacuologenic (Nordeng, Gregers, Kongsvik, Mørresse, Gorvel, Jourdan, Motta, and Bakke, unpublished data).

We substituted the Asp residues in position 3 or 6 with Arg in full-length Ii, changing the net charge in the tailpiece from \( /H_{11002}^2 \) to zero, and we found that these single mutations were sufficient to avoid vacuolation. Intracellular trafficking of the mutants and wild-type Ii was compared by immunofluorescence microscopy and by quantification of the fractions of newly synthesized proteins that gained Endo H resistance, and there is no indication from the results that the inability of the mutants to form ILEV was due to impaired transport to the endocytic pathway. Moreover, structural data suggest that vacuolation can be interpreted in terms of \((i, i \pm 3)\) electrostatic interactions present in the Ii tailpiece. This was...
applied as a control to the above chimeric Ii constructs reported by Pond et al. (1995), and we found that for neutral tailpieces one may explain ILEV formation using the above hypothesis on electrostatic interactions.

Pond et al. (1995) sought to reconstitute large vesicle formation by replacing one or two residues with Asp and obtained two positive (MDQIKRL and MSDKKRL) and one neutral (MDIKKWL) tail. In agreement with our observations, no large vesicles were observed for positively charged tails. More interesting, for the neutral one no large vesicles were formed (Pond et al., 1995), as it can be predicted by our electrostatic hypothesis. This is due to the presence of double (i, i ± 3) electrostatic interactions (between residues 2-5 and 3-6). The results show that Ii-induced vacuolation depends on available negative charges in the Ii N terminus rather than specific amino acids residues.

It was previously reported that Ii constructs with deletions in the luminal domain, which abolished trimerization, did not cause vacuolation (Gedde-Dahl et al., 1997). We did not observe ILEV in cells transfected with chimeric constructs in which the Ii tail was fused to the transmembrane domains of either the transferrin receptor, which is dimeric (Jing and Trowbridge, 1987), or the neuraminidase, which is a tetramer (Brennes et al., 1994). This suggests that the trimeric conformation of Ii is required for vacuolation. The luminal domain of Ii is sequentially degraded in endosomes, but the tails remain trimeric (Amigorena et al., 1995; Newcomb et al., 1996). Double-staining of ILEV with antibodies against the luminal and the cytoplasmic domains of Ii has demonstrated that the cytoplasmic tails are recognized by antibodies in “late” ILEV after the luminal domain has been degraded (Stang and Bakke, 1997).

It has previously been shown that Ii is retained within the endocytic pathway (Loss and Sant, 1993) where it induces a delayed endocytic transport of both antigens and membrane proteins (Romagnoli et al., 1993; Amigorena et al., 1995; Gorvel et al., 1995). Use of the protease inhibitor leupeptin showed that MHC class II accumulated in late endosomes in association with an Ii-derived fragment, Lip10 (Maric et al., 1994; Amigorena et al., 1995; Brachet et al., 1997). Lip10 harbors an intact cytoplasmic tail and this led to the suggestion that a signal for endosomal retention was present in the cytoplasmic tail of these fragments, which retained MHC class II until Ii processing was complete. It was further suggested that the retention signal likely was constituted of the two Leu-based signals (Maric et al., 1994). However, in a recent study we have shown that expression of Ii5(1-25) but not Ii5(26-100) caused endocytosed material destined for the TGN or lysosomes to be retained in endosomes (Gregers et al., unpublished data), suggesting that endosomal retention is not dependent on the two Leu-based signals. Instead, available negative charges in the N terminus of Ii seem to be decisive.

In a previous study, we have reported the structure of a synthetic peptide corresponding to the cytoplasmic tails of Ii(4-10) determined in solution (Motta et al., 1997). At high concentrations, this peptide forms as a trimeric α-helical bundle. Interestingly, these trimers were almost coplanar and antiparallel, arranged in an up-down-up orientation, suggesting that different trimers might interact with each other. This trimeric association, which preserves the secondary structure of the monomer, was shown to be mostly determined by the pattern of polar and hydrophobic residues. NMR spectroscopy was also applied to compare the solution structure of the synthetic peptides corresponding to the cytoplasmic tails of Ii(4-10) and Ii(26-35), and they were found to have almost the same overall shape (Motta et al., 1997). When the concentration of the Ii(26-35) tail peptide was increased in methanolic solution, the peptide started to form an aggregate. Interestingly, in similar conditions, the Ii(26-35) tail peptide did not aggregate, even after being refrigerated for several months. The absence of aggregation correlates with the lack of noticeable differences in NMR spectra at various concentrations (Nordeng, Gregers, Kongsvik, Meresse, Gorvel, Jourdan, Motta, and Bakke, unpublished data), suggesting that internal double-salt bridges (Asp2-Arg5 and Asp3-Arg6) alter the charge distribution within the tailpiece, thus disfavoring the trimeric association described for the native peptide.

The morphological and structural evidence prompted us to ask for a possible connection between available negative charges in the cytoplasmic tail of Ii, the intrinsic aggregating ability of tails pointing in opposite directions, and the endosomal vacuolation and delayed endocytic trafficking. We have previously suggested that interactions between Ii molecules on separate membranes could promote fusion between endosomes (Motta et al., 1997). In this scenario, membranes of different endosomes are brought together by trans interactions between protruding tails. Parth et al. (1992) have reported that chemical cross-linking of endosomes in live cells causes vacuolation and partially impaired sorting. Moreover, there are several examples in the literature describing vacuolation in response to increased endosomal fusion (Barbieri et al., 1994; Stenmark et al., 1994). We have searched for a connection between the structural and the morphological data by applying a cell-free fusion assay for endosome membranes containing Ii(4-10) or Ii(26-35). Data from these experiments show that in the cell-free system Ii increased endosome fusion by 20%, and it is possible that this could account for the observed effects on the endosomal pathway.

In vitro homotypic fusion between early endosomes has been reported to be almost completely blocked by the PI3-kinase inhibitor wortmannin (Jones and Clague, 1995). In our cell-free assay, wortmannin inhibited fusion between endosomes by ~45%. The reason for the lower efficiency of wortmannin in our assay could be that we used a mixture of early and late endosomes. However, fusion between endosomes from cells expressing Ii(4-10) was inhibited by only ~10% when wortmannin was included in the fusion mixture. Ii(26-35), on the other hand, was not able to restore fusion under such conditions. This shows that in the presence of Ii(4-10) endosome fusion is not dependent on the PI3-kinase and suggests that this effect depends on free negative charges within the cytoplasmic tail N terminus.

There are a number of recently discovered proteins that serve to link vesicles with their targets (for reviews, see Pfeffer, 1999; Waters and Pfeffer, 1999). One of these, EEA1, has been shown to bind PI3-phosphate on early endosomes (Mills et al., 1998; Simonsen et al., 1998b; Christoforidis et al., 1999). We observed by live cell microscopy that, although EEA1-GFP rapidly dissociated from endosome membranes upon addition of wortmannin, Ii-induced vacuolation was not sensitive to this treatment. In the presence of Ii(4-10), endosome fusion was insensitive to wortmannin, whereas NEM
always blocked fusion. This suggests that Ii can complement for PI3-kinase activity in a step before the NEM-sensitive docking of endosomal membranes. When Ii<sub>wt</sub>-positive endosomes were mixed with endosomes containing Ii<sub>D6R</sub>, the wortmannin-induced block of endosome fusion was not restored. One possible explanation for this observation is that Ii<sub>wt</sub> tails could recruit a cytoplasmic factor involved in homotypic endosome fusion and that Ii<sub>D6R</sub> fails to recruit such a factor. Another explanation could be that the cytoplasmic tail of Ii<sub>wt</sub> is not able to form stable antiparallel interactions with tails of Ii<sub>D6R</sub>. The latter model is supported by NMR calculations showing that electrostatic interactions between the II tails depend on available negative charges, which the N-terminal tail of Ii<sub>D6R</sub> does not provide. Thus, although it is possible that relevant cytoplasmic factors could be recruited specifically by the cytoplasmic tail of Ii, we provide evidence that the aggregating property of the II cytoplasmic tail is involved in the regulation of endosome fusion.

The PI3-kinase together with rab5 recruit EEA1 to early endosomal membranes, and it has been suggested that the EEA1 molecules on separate membranes can tether endosomes together by coiled-coil interactions. This is believed to be an early step in the linkage process ultimately causing early endosomes to fuse (for a review, see Pfeffer, 1999; Waters and Pfeffer, 1999). Our data show that in vitro the cytoplasmic tail of Ii can partially complement for endosome tethering by EEA1. Pfeffer (1999) has defined vesicle tethering as involving loose links that extend over distances &gt;25 nm, whereas docking molecules promote rather stable interactions that hold two membranes within a 5- to 10-nm distance. Interactions between antiparallel II tails are expected to be quite loose but would not hold membranes with a distance of much more than 5 nm because the tails are quite short (30 amino acids in a helical conformation, including a kink). Hence, II does not fall into one or the other category, but we suggest that II may serve to stabilize membrane topology in a conformation that favors SNARE interactions and subsequent vesicle fusion.

II tails have been found on both early and late endosomes (Stang and Bakke, 1997), and interactions between the tails could therefore cause endosome populations to mix. Consequently, vectorial processes such as transport and proteolysis would be disordered. This theory is supported by recent experimental data showing that high expression of II<sub>wt</sub> but not II<sub>D6R</sub> caused endocytosed material destined for the TGN or lysosomes to be retained in endosomes (Gregers et al., unpublished data). Moreover, antigen presentation of an II-dependent epitope was shown to be enhanced in cells expressing II<sub>wt</sub> compared with II<sub>D6R</sub>. Even though it remains to be elucidated whether, and under which circumstances, II is able to regulate its own intracellular environment in vivo, our data support new aspects of regulation of endocytic transport and endosome size and morphology.

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