A novel phospholipid-binding protein from the yeast Saccharomyces cerevisiae with dual binding specificities for the transport GTPase Ypt7p and the Sec1-related Vps33p

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Introduction

Studies in vitro and in vivo of yeast vacuole membrane fusion have added a great deal to our understanding of Ypt/Rab GTPase-controlled, SNARE-mediated membrane fusion events in eukaryotic cells (reviewed in (Götte and Lazar, 1999; Wickner, 2002)). During the so-called docking step of the fusion event the syntaxin-like vacuolar SNARE Vam3p binds the regulatory Vps33 protein which is part of a large multisubunit protein complex (Sato et al., 2000). This heterogeneous supramolecular aggregate is variously known as HOPS (Seals et al., 2000) or the Vps C complex (Sato et al., 2000). Vps33p is a member of the Sec1 protein family. It shuttles between a membrane-bound and a soluble state (Gerhardt et al., 1998). During the docking step, the HOPS/Vps C protein complex is transferred to the regulatory GTPase Ypt7p (Price et al., 2000). Ypt7p belongs to the highly conserved 11-member family of Ypt/Rab GTPases that are involved in the regulation of membrane recognition/docking processes (Lazar et al., 1997). Each of these GTPases is supposed to perform a similar function at a specific transport step. Ypt7p is required to be present on both membranes determined to fuse with each other. In agreement with its role during vacuolar membrane docking, the membrane-bound fraction of the cellular pool of Ypt7p has been localized to the vacuolar membrane (Haas et al., 1995). A component of the HOPS/Vps C complex, Vps39p (also known as Vam6p), directly interacts with Ypt7p and drives the exchange of GDP for GTP, thereby helping switch the GTPase from the inactive to the active state (Wurmser et al., 2000). Downstream of Ypt7p, two additional monomeric GTPases, Rho1p and Cdc42p (Müller et al., 2001; Eitzen et al., 2001) help regulate vacuoles to dock to each other. The docking step with its multitude of protein factors most likely serves as an identification/proofreading step during the sequence of events from priming to fusion. During the docking process, Ypt7p interacts with Ccz1p, a protein known to be involved in the detoxification of metal ions by the vacuole (Kucharczyk et al., 2001). Details of the molecular events downstream of Ypt7p are summarized in (Mayer, 2001).

By searching for new interaction partners for the GTPase Ypt7p with a two-hybrid system library screen, we have discovered Ydr229w (named Ivy1p), an as yet undescribed protein product of the open reading frame YDR229w (named IVY1 for: Interacting with Vps33p and Ypt7p) was found to interact with both the GTPase Ypt7p and the Sec1-related Vps33p protein. While deletion of IVY1 does not lead to any recognized change in phenotype, overexpression of Ivy1p leads to fragmentation of the vacuole, missorting of the vacuolar enzyme carboxypeptidase Y (CPY) to the exterior of the cell, and an accumulation of multivesicular bodies inside the cell. All effects caused by the overexpression of Ivy1p can be reset by simultaneously raising the amount of Vps33p. This suppression activity of Vps33p suggests that Ivy1p and Vps33p at least partially counteract the action of each other in the cell. The intracellular level of Ivy1p increases in cells approaching stationary growth phase at which part of the protein is located at the rim of the vacuole. In addition to its specific interactions with members of two regulatory protein families, Ivy1p in vitro shows a marked propensity for binding phospholipids with high affinity.
binding partner for Ypt7p. Ivy1p represents a novel class of proteins which were not known to interact with Ras-related transport GTPases. Next to Ypt7p, Ivy1p binds the Sec1-related Vps33p and various phospholipids. To our knowledge, Vac1p is the only yeast protein with a similar set of interactions that has been described to date (Tall et al., 1999). Certain phospholipids have been recognized to fulfill not only structural functions as membrane components but to exert effects as membrane-localized sensor molecules or as signal mediators (Hama et al., 2000; Huijg新娘ts et al., 2000; Odorizzi et al., 2000). In particular, lipids of the phosphatidylinositol phosphate group have been recognized as important regulatory molecules, presumably mediating signals or providing points of aggregation for specific proteins (Wera et al., 2001). For the fusion machinery of the yeast vacuole it could be shown that PIP2 (phosphatidylinositol-4,5-bisphosphate) is necessary at two different stages of the fusion event (Mayer et al., 2000). Phosphatidylinositol-4-phosphate (the precursor of phosphatidylinositol-4,5-bisphosphate) is a regulatory molecule itself the intracellular pool of which affects vacuolar morphology (Foti et al., 2001). Finding Ivy1p in this context suggests the involvement of proteins with novel, non-classical lipid-binding domains. The identification of a member of the fusion machinery as a novel-type lipid-binding protein may lead to a deeper understanding of the mechanisms of fusion, in particular the physical interactions of molecules necessary for removing/attachment of proteins to/from the membrane or for stabilising/destabilising distinct membrane regions.

Materials and methods

**Yeast cell growth and genetic manipulation**

Yeast strains used are described in Table I. Yeast culturing, transformation, DNA and protein extraction was done according to Adams et al. (1997). To overcome growth inhibition in galactose-containing media, 0.02% sucrose (from a sterile 20% stock solution) was added where necessary.

Yeast epitope tagging was carried out as described by De Antoni and Gallwitz (2000). Epitope-tagging modules were obtained with short flanking homolog (SFH) primers working on the template plasmids described by De Antoni and Gallwitz (2000). Primers for IVY1-tagging and YPT7 and YPT7(T22N) and YPT7(Q68L) mutant alleles were cloned by PCR from genomic DNA. An NdeI restriction site including the start codon was inserted into the different plasmids.

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**Plasmid generation and cloning**

**YPT7** and **YPT7(T22N) and YPT7(Q68L)** mutant alleles were cloned by PCR from genomic DNA. An NdeI restriction site including the start codon was inserted into the different plasmids. **Restriction sites in parentheses indicate the fragments harbouring the corresponding genes that were inserted into the different plasmids.**

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codon was introduced by using the oligonucleotide TL 1 (5'-CTAGAGACATATGTTAATTTGATGATAC-3'). PCR fragments were subcloned into pBluescript KS II and transfected after sequencing into two-hybrid or/and expression vectors employing restriction sites common to both plasmids. An Nco I restriction site was introduced at the 5' end of the coding regions of the IVY1 gene with the help of TL 167 (5'-CAGGAAAGAGAAATCGGCATGGGTACAAATACCG G-3') for cloning into the high expression vectors of the pYX series (R&D Systems, Abingdon, UK). VPS33 expression vectors pRS325-VPS33 and pEGKT-VPS33 were kindly provided by Martin Gütte. Plasmids generated for this study are described in Table II.

**PCR**
Polymerase chain reactions were carried out in the following buffer: 10 mM Tris-HCl, pH 8.3; 50 mM KCl; 2.5 mM MgCl₂. The reaction mixture contained variable amounts of template DNA plus 0.2 mM deoxy nucleoside triphosphates (dNTPs), 1 μM of each primer and heat-stable polymerase. A mix of 5 parts Taq polymerase (Perkin-Elmer, Wellesley, MA, USA) and 1 part Deep Vent DNA polymerase (NewEnglandBiob, Beverly, MA, USA) was used. Genomic DNA from the EUROFAN (http://mips.gsf.de/proj/eurofan/) reference strain FY 1679 was used as template in all cases except when non-genomic DNA (e.g. plasmid DNA) was of the essence.

**Fusion protein production and immobilization**
Glutathione-S-transferase (GST) fusion proteins have been produced in *E. coli* DH5α IQ (from GIBCO BRL, Invitrogen, Karlsruhe, Germany) or in the *S. cerevisiae* cl 3-ABYS86 strain. Induction of bacterial cultures was done with 0.1 mM IPTG for 1 to 4 hours, depending on the protein of interest. Fusion protein expression in yeast was done by transfecting yeast cells after washing with sterile water into galactose-containing medium. Induction time of yeast cultures expressing GST-fusion proteins was 12–48 hours, depending on the protein of interest. Harvest and purification of the fusion proteins was identical with bacterial and yeast cells and carried out as described in the manufacturer’s manual (Pharmacia, Sweden). Buffer of choice for the experiments described in this article was PBS (prepared according to Sambrook et al. (1989)), supplemented with protease inhibitors.

**Overexpression of Ivy1p**
To achieve strong overproduction of Ivy1p in the cell, IVY1 was cloned by PCR and inserted into pYX212, pYX213, pYX242, and pYX243 expression vectors (R&D Systems) as an Nco I-Sac I fragment for cloning into the high expression vectors of the pYX series (R&D Systems) as an Nco I-Sac I fragment for cloning into the high expression vectors of the pYX series (R&D Systems) for cloning into the high expression vectors of the pYX series (R&D Systems). FY 1679 was used as template in all cases except when non-genomic DNA (e.g. plasmid DNA) was of the essence.

**Results**
A two-hybrid cDNA library screen was performed to find novel proteins that physically interact with the transport GTPase Ypt7p. To select for proteins with a putative preference for the GDP-bound form of the GTPase, the mutated form Ypt7(T22N)p was used for the bait fusion with the DNA-binding part of the Gal4 protein. This bait fusion protein was tested against a yeast cDNA library (kindly provided by Bruce Horazdovsky, Dallas, USA). After a thorough analysis of the obtained two-hybrid system candidate clones, the gene YDR229w (named IVY1 for: Interacting with Vps33p and Ypt7p) was singled out as the only new protein faithfully interacting with the GTP-binding protein Ypt7p. IVY1 encodes a 435-amino-acid polypeptide chain (Fig. 1).

All isolated clones harbouring this cDNA contained a 5'-truncated form of the gene YDR229w lacking the first 34 codons. The *Saccharomyces cerevisiae* genome does not contain any homologue of IVY1 or any other gene coding for a protein of any significant similarity to Ivy1p. Ivy1p therefore represents a novel protein type, hitherto unknown to interact with GTP-binding proteins of the Ras superfamily.

A computer search for homologues from other organisms has not yielded any protein with a significant degree of similarity in primary structure to Ivy1p in higher eukaryotes. However, a gene coding for a 398-amino-acid protein with sequence homology to IVY1 does exist in the fungus *Neurospora crassa* (accession numbers: CAD11793; emb AL 442164; NCBJ23; N. crassa DNA linkage group II, BAC clone B2J23).
Ivy1p interacts with the transport GTPase Ypt7p and with the Sec1-related Vps33p

After Ivy1p had been identified as a binding partner for Ypt7p (T22N) and Ypt7p (wild-type), the specificity of this interaction was probed by testing the new protein with a number of related GTPases from baker’s yeast. Positive lacZ assay results were obtained with Ypt7p (T22N), Ypt7p (wild-type), and Ypt7p (Q68L) (Fig. 2). A faint reaction was also seen with Ypt10p. Even if it was reproducible, it was not followed up because of the weakness of the signal. Unequivocally negative lacZ reactions were reproducibly achieved with the transport GTPases Ypt51p, Ypt52p, Ypt6p, Ypt6p (Q/L), Ypt31p, and Sec4p. A strong, positive lacZ reaction was obtained with Ivy1p fused to both parts of the Gal4 protein: Ivy1p also interacts with itself (Fig. 2). After Ivy1p was established to interact specifically with the vacuolar transport GTPase Ypt7p, a number of specific proteins, components of the vacuolar membrane fusion machinery and respective counterparts as controls, were tested for their two-hybrid system interactions with Ivy1p. The only protein from this group which tested positive in the lacZ assay was the vacuolar Sec1-related Vps33p (Fig. 2). The endosomal Sec1 homologue Vps45p consistently tested negative. Ivy1p did not show lacZ activation in combination with the vacuolar syntaxin-like protein Vam3 (t-SNARE) in either its full-length version containing the C-terminal transmembrane domain or a truncated version lacking the transmembrane domain (Vam3(AC)). The negative lacZ assay outcome of this interaction test was borne out by testing the endosomal syntaxin-like SNARE Pep12p. Furthermore, Ivy1p did not show any interaction in the two-hybrid system with the GTPase-activating protein (GAP) for Ypt7p, Gyp7p, Gyp6p (Ypt6p-GAP), used as a control for this combination, likewise tested negatively.

Affinity-chromatography experiments confirm the two-hybrid system interactions

Fig. 1. Primary structure of Ivy1p (Ydr229w). The translation product of the YDR229w gene consists of 453 amino acids. The codons for the first 34 amino acids were missing in the retrieved two-hybrid cDNA plasmid clones.

Fig. 2. Two-hybrid system interactions of Ivy1p. Ivy1p yields positive lacZ assay results with the transport GTPase Ypt7p (wild-type), the mutant GTPases Ypt7p (T22N) (predominantly in the GDP-bound form) and Ypt7p (Q68L) (predominantly in the GTP-bound form), with itself, and with Vps33p. No lacZ activation is seen with either the Gal4p activation domain or with the Gal4p DNA-binding domain when Ivy1p is fused to the corresponding second half of the Gal4 protein. Ypt52p is shown as an example of a Ypt7p-related GTPase that tests negative.
GTP-bound GST-Ypt1p (as compared to GST-Ypt7p in both Fig. 3B and Fig. 3C) are most probably not meaningful in vivo. As attempts to produce soluble recombinant Ivy1p fusion proteins in bacteria encountered obstacles, GST-Ivy1-HAp and GST-Ivy1p were instead produced in protease-deficient yeast strains. Using this approach, we could cross-prove the interaction with Ypt7p (detected by anti-Ypt7p antisera) and with Vps33p. Vps33p, and its related endosomal counterpart, Vps45p, were tagged with either the HA or the VSV epitope by modifying the chromosomal VPS33 and VPS45 genes according to De Antoni and Gallwitz (2000). By expressing GST-Ivy1p or/and various other GST-tagged proteins (detected by anti-GST antibodies) in cells containing a chromosomal copy of the IVY1-FA gene, it was possible to confirm the interactions known from the two-hybrid fusion experiments, including Ivy1p-Ivy1p self-interaction (Fig. 3C, D). GST-Vps33p could retain Ivy1-HAp while GST-Vps45p could not. In a reciprocal experiment, GST-Ivy1p was able to retain epitope-tagged Vps33-VSVp (Fig. 3D) but was unable to hold back epitope-tagged Vps45-HAp (Fig. 3E). Vps45p did bind to GST-tagged Vac1p, though. Vac1p exhibits a similar set of interactions (a transport GTPase, Ypt51p, the Sec1p/Vps33p-related Vps45p, and a phosphatidylinositol-phosphate, PI-3-P) as Ivy1p (Tall et al., 1999). The specificity of the interaction for the Sec1-family member Vps33p was in this way validated, too (Fig. 3C, E).

**Deletion of IVY1 is phenotypically neutral**

Removal of the chromosomal copy of the entire coding region of the gene IVY1 by the short flanking homology (SFH) PCR replacement method described by Guildner et al. (1996) led to an ivy1 deletion strain (ivy1Δ TAL39/TAL42). Correct and specific removal of the IVY1 coding region was confirmed by analytical PCR and by Southern blotting (not shown). The ivy1 deletion strains in any assayed way behaved exactly like wild-type yeast strains. Growth rates at temperatures from 14 °C to 40 °C were indistinguishable from the parent strain the deletion strains were based on. Growth on various media containing a range of additives did not differ from the parent strains. Particular attention was paid to the vacuole of ivy1Δ cells. Protein transport to the vacuoles (monitored by the trafficking of carboxypeptidase Y, see below) and vacuolar morphology were undisturbed and indistinguishable from wild-type yeast cells. The cells of a ypt7Δ ivy1Δ double knock-out strain...
Immobilized GST-Ivy1p was incubated with liposomes containing phosphatidylcholine alone, phosphatidylcholine/phosphatidylinositol (1:1 ratio) or a mixture of phosphatidylserine and several phosphatidylinositol phosphates. Compared to immobilized GST, GST-Ivy1p showed strong binding to the phosphatidylinositol vesicles, with preference for higher phosphorylated derivatives. To further elucidate the binding specificity of Ivy1p, vesicles were constructed with a 10% (w/w) content of distinct phosphatidylinositides in a background of phosphatidylcholine. Ivy1p bound all phosphatidylinositides tested with a preference for 4-phosphorylated phosphoinositides over 3-phosphorylated phosphoinositides (Fig. 4).

Ivy1p and Vps33p show similar distributions
To get access to the in vivo behaviour of Ivy1p, yeast cells expressing Ivy1-HA and Vps33-VSVp were fractionated as described by Wichmann et al. (1992). A crude extract, cleared of cell wall fragments, served as standard, representing 100% of the cellular protein. Pellets were redissolved after 10000g (P10) and 100000g (P100) centrifugations. The supernatant after the 10000g (S 100) run is cytosol with soluble proteins. Ivy1-HAP is predominantly found in the P10 fraction. A minor portion of Ivy1-HAP is found in soluble form (S100 fraction), and only a minute quantity of the protein is detectable in the P100 fraction. This distribution is similar to the one for Vps33-VSVp. Ivy1p and its binding partner Vps33p behave in similar fashions in this crude fractionation experiment (Fig. 5).

Localization of Ivy1p
The intracellular localization of Ivy1p was studied by indirect immunofluorescence of the epitope-tagged protein. As a reference, the localization of its interaction partner Ypt7p was used since the membrane-bound pool of this GTPase is known to localize to vacuolar membranes (Haas et al., 1995). In yeast cells of exponentially growing cultures (O.D.₆₀₀ approximately 1), no distinct subcellular distribution of the Ivy1p protein could be detected with anti-HA antisera. In cells approaching stationary phase (O.D. 600 approximately 8), a distinct fluorescence distribution at the vacuolar perimeter was found, similar to that of Ypt7p (Fig. 6A). When the expression of the HA-tagged Ivy1p protein was followed by immunoblotting, a steady increase in the amount of the protein up to a point of saturation was found in high-cell-density cultures (Fig. 6B). In exponentially growing yeast cultures, the amount per cell of Ivy1-HAP is either not sufficient for detection by indirect immunofluorescence or the protein changes its distribution when the cellular protein content rises, or newly synthesized Ivy1-HAP in stationary cells localizes to the perimeter of the vacuole.

Ivy1p senses the absence of its binding partners
The striking difference in electrophoretic behaviour of Ivy1p derived from wild-type versus ypt7 mutant cells (seen in Figure 6B) led to the investigation of Ivy1p in various mutant genetic backgrounds (Fig. 7). Wild-type cells show HA-tagged Ivy1p as a pair of distinct bands migrating very close to each other (Fig. 7, best seen in vam3 mutant cells in which Ivy1p behaves as in a wild-type genetic background). A band shift towards faster migration in a denaturing SDS-polyacrylamide gel was seen in cells having no Ypt7p at all (ypt7A) or synthesising the mutated form Ypt7(T22N)p. A slight difference in mobility can be distinguished between these two mutant

**Ivy1p binds phospholipids**

From the analogy with the protein Vac1p in terms of being able to bind a specific transport GTPase and its cognate Sec1 homologue, the conjecture was made that Ivy1p might exhibit further similarities with Vac1p. Vac1p is known to bind the signal lipid phosphatidylinositol-3-phosphate (Tall et al., 1999). We therefore tested whether Ivy1p can bind lipids, too. By a liposome-based capture assay (Mayer et al., 2000), we found that Ivy1p binds certain phospholipids with high affinity.
Samples were retrieved at an O.D. 600 of about 0.15 (saturation point is reached in cells of an optical density at 600nm of electrophoresis. The amount of Ivy1-HAp increases steadily until a number of cells in each sample, was separated by denaturing gel culture at regular intervals, and the total protein normalized for the aliquots of cells were drawn from a wild-type (wt) and a panel the vacuole (Fig. 6). In aged cells of stationary or near stationary cultures, Ivy1-HAp is not readily localizable. Only a diffuse signal all over the cell is seen (centre panel). In aged cells of stationary or near stationary cultures, Ivy1-HAp shows in indirect immunofluorescence a punctate staining at the rim of the vacuole (right panel), reminiscent of the localization of Ypt7p. Aliquots of cells were drawn from a wild-type (wt) and a ypt7Δ yeast culture at regular intervals, and the total protein normalized for the number of cells in each sample, was separated by denaturing gel electrophoresis. The amount of Ivy1-HAp increases steadily until a saturation point is reached in cells of an optical density at 600 nm of about 6. Samples were retrieved at an O.D.600 of about 0.15 (a), 1 (b), 2.5 (c), 5 (d), and 8 (e).

Fig. 6. Localization of Ivy1p and growth phase-dependent expression. (A) By indirect immunofluorescence, the GTPase Ypt7p was shown to localize predominantly to the vacuole (left panel). In yeast cells of exponentially growing cultures, Ivy1-HAp is not readily localizable. Only a diffuse signal all over the cell is seen (centre panel). In aged cells of stationary or near stationary cultures, Ivy1-HAp shows in indirect immunofluorescence a punctate staining at the rim of the vacuole (right panel), reminiscent of the localization of Ypt7p. (B) Aliquots of cells were drawn from a wild-type (wt) and a ypt7Δ yeast culture at regular intervals, and the total protein normalized for the number of cells in each sample, was separated by denaturing gel electrophoresis. The amount of Ivy1-HAp increases steadily until a saturation point is reached in cells of an optical density at 600 nm of about 6. Samples were retrieved at an O.D.600 of about 0.15 (a), 1 (b), 2.5 (c), 5 (d), and 8 (e).

Fig. 7. Ivy1p senses the absence of its proteinaceous binding partners. Total protein extracts from various strains containing chromosomally HA-epitope-tagged IVY1-HA genes were separated by SDS-PAGE. Wild-type cells (and vam3Δ, ypt51Δ and ypt10Δ deletion/disruption mutant strains) show an anti-HA antibody-reactive double band (best seen in the vam3Δ cells). In strains which harbour a YPT7 mutant allele or which are deleted for genes encoding translation products that interact with Ivy1p, only a single, faster moving anti-HA-reactive band is seen.

cell lines, suggesting that the state of Ivy1p differs between the ypt7 null mutant and the one harbouring a functionally impaired Ypt7 protein. A band shift similar to the one in ypt7Δ mutants is found in cells without Vps33p (vps33Δ). Deletion of other genes coding for Ypt GT-Pases has no effect on the mobility of denatured Ivy1p (ypt10Δ and ypt51Δ). Deletion of the VAMS3 gene, coding for a vacuolar SNARE protein, does not affect Ivy1p. Ivy1p specifically reacts to the presence/absence of its binding partners Vps33p and Ypt7p. In the case of Ypt7p, Ivy1p is sensitive not only to the presence or absence of the GTPase but, in addition, to either the functional status of Ypt7p (Ypt7(T22N)p is bound to GDP and inhibited in nucleotide exchange) or its overall functionality (Fig. 7). Cells expressing the preferentially GTP-bound mutant protein Ypt7(G68L)p (which is inhibited in its GTP-hydrolytic capacity) invariably gave blurred Ivy1p bands that did not allow a statement with regard to reactivity of Ivy1p towards this mutant form. The band shift in the ypt7Δ mutant could be remedied by adding back and expressing Ypt7p from a plasmid (not shown). The molecular nature of the modification underlying the band shift(s) is not known. Phosphatase treatment of wild-type protein extracts did not lead to a ypt7- or vps33-like change in Ivy1-HAp. Phosphorylation/dephosphorylation is therefore unlikely to be the molecular cause of this effect. Conformational change can be ruled out, too, as the alterations in electrophoretic mobility were detected on SDS-containing denaturing gels. Degradation has been ruled out as only one distinct band and no further degradation products have ever been observed in the mutant strains.

Overexpression of Ivy1p leads to a disruption of vacuolar morphology which can be suppressed by raising the amount of Vps33p

When we expressed GST-Ivy1p in yeast cells, a conspicuous morphological change was seen after long induction times to maximize fusion protein yield. A majority of cells in stationary expression cultures showed fragmented vacuoles, a phenotypic trait frequently seen in yeast cells deleted for genes encoding products which play a role in vacuolar transport pathways or vacuolar inheritance (see (Götte and Lazar, 1999) for review). Therefore, yeast cells were transformed with a 2-micron multi-copy plasmid containing the IVY1 coding region plus 400 base pairs 5′ of the start codon, most likely representing the complete native promoter. Transformed cells did not show any morphological alterations visible with a light microscope, irrespective of the growth phase of the culture. The IVY1 coding region (with and without a 3′-extension for HA epitope tags) was then cloned into 2-micron high-expression plasmids with either the constitutive triosephosphate isomerase (TPI) promoter or the inducible GAL1 promoter. Cells containing plasmids with the IVY1 gene constitutively expressing large amounts of Ivy1p (with or without HA epitopes) driven by the TPI promoter showed fragmented vacuoles (Fig. 8). The same effect could be achieved with GAL1 promoter-driven Ivy1p expression but vacuolar fragmentation was only seen after considerable times of induction (more than 24 hours). A sufficiently high amount of the protein had to be synthesized before the morphological alteration could be observed. As cells constitutively expressing large quantities of Ivy1p do not show any change in growth rates as compared to wild-type cells and cells expressing the epitope-tagged protein behaved no different from ones making the unmodified Ivy1 protein, most experiments were carried out with cells expressing Ivy1-HAp protein constitutively (pTL89-transformed cells). All effects described for cells overexpressing Ivy1p were observed in the same way (indistinguishable in both appearance and extent) in cells synthesizing Ivy1-HAp. Ivy1-HAp was employed to confirm the production and measure the extent of the plasmid-driven (over)expression in addition to the normal (background) amount of Ivy1p (or Ivy1-HAp) produced by the
The fragmented vacuole phenotype seen by light microscopic inspection of cells overproducing Ivy1p validated ultrastructural analysis to search for additional structural alterations below the resolution threshold of the light microscope. The striking finding in cells overproducing Ivy1p was a massive accumulation of multivesicular bodies (MVBs). MVBs are believed to represent the late endosomes of eukaryotic cells (Piper and Luzio, 2001; Sachse et al., 2002), the compartment which is functionally the closest neighbour of the vacuole (lysosome). Excess amounts of Ivy1p greatly enhance the accumulation or formation of MVBs/late endosomes. The observed MVBs are of different sizes with typical diameters resembling the size of the vacuolar fragments. The MVBs do contain vesicles of various sizes but a given MVB preferentially harbours vesicles of a distinct size. A mix of vesicles differing in size was occasionally found but was not the norm (Fig. 9).

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**Overexpression of Ivy1p leads to the missorting of vacuolar proteins which can be suppressed by raising the amount of Vps33p**

As morphological alterations at the vacuolar level were induced by overexpression of Ivy1p rather than by deletion of the *IVY1* gene, we tested whether delivery of proteins to the vacuole is also affected in Ivy1p-overexpressing cells. A missorting of vacuolar proteins is usually found to go hand in hand with changes in vacuolar morphology. Two lines of experimental evidence proved that cells strongly overexpressing Ivy1p indeed missort soluble luminal vacuolar enzymes such as CPY.

In an enzymatic in vivo overlay assay developed by Horazdovsky et al. (1994), cells with a defect in vesicular transport to the vacuole shed into their periplasms a fusion protein consisting of the vacuolar sorting signal of CPY and the sucrose-cleaving enzyme invertase (pro-CPY-invertase). Enzyme-driven oxidation of glucose released by the secreted invertase fusion protein leads to a colour reaction if pro-CPY-invertase gets beyond the plasma membrane. A *ypt7* deletion strain (TAL2; a class B vps mutant) served as standard in this assay. “Wild-type” cells of the reporter strain BHY11 were transformed with plasmids containing the *IVY1* coding region under the control of either the TPI promoter (pTL89) or the inducible GAL1 promoter (pTL90). Wild-type cells with the unmodified plasmid pYX212 showed no colour development, while TAL3 *ypt7Δ* cells show the expected colour reaction of a *vps* mutant. BHY11 cells with pTL89 show a colour reaction that is similar to that of *ypt7Δ* cells in strength and speed of the colour appearance. Cells overexpressing Ivy1p missort proteins with a vacuolar targeting signal. Reporter cells (BHY11) with pTL90 do not show any colour development when expression of the plasmid-borne *IVY1* copies is not induced by galactose (Fig. 10A).
The results obtained in the overlay colour assay were confirmed by in vivo labelling of cells with radioactive amino acids and subsequent immunoprecipitation with anti-CPY antisera (Fig. 10B). Cells strongly overexpressing Ivylp (+ pTL89, high expression Ivylp) show a retardation of the proteolytic processing of the portion of the reporter protein that stays inside the cell, indicating that the Golgi-derived p2-pro-CPY did not reach the vacuolar compartment where it would have been activated by partial proteolysis by proteinase A (Pep4p). Some of the unprocessed pro-enzyme is missorted to the outside of the cell where it could be recovered by immunoprecipitation at time 0 from the medium surrounding the protoplasts used in this experiment. The (frequently observed) absence of detectable extracellular p2-pro-CPY at later times (45 minutes in this case) is most probably due to degradation of the extracellular pro-enzyme (irrespective of added protease inhibitors) rather than an inability of the antibodies to precipitate it (Fig. 10B). Part of the protein retained inside the cell is aberrantly processed (*, ‘X’-CPY; Fig. 10B; cf. (Wichmann et al., 1992)). After adding a 2-micron multi-copy plasmid containing VPS33, cells constitutively expressing Ivylp in high amounts behave like untransformed wild-type cells (strain TAL 45). The additional Vps33 protein in these cells completely suppresses the missorting of vacuole-destined CPY in TAL 45 + pTL89 cells (Fig. 10B). This result reflects the suppression of the vacuole fragmentation by additional VPS33 as described above (see Fig. 8).

**Discussion**

The gene YDR229w, named IVY1, of *Saccharomyces cerevisiae* codes for a protein that specifically interacts in vitro with the GTPase Ypt7p, the Vps33 protein, and, in a more promiscuous way, various phospholipids with different degrees of affinity. Both proteinaceous binding partners of Ivylp serve important roles during the docking step of vacuolar fusion. In addition, Ivylp has the ability to self-aggregate into either homooligomers or supramolecular structures containing multiple copies of it.

The propensity of Ivylp to bind Ypt7p over other Ypt proteins, is strong but not perfect (cf. Fig. 3). In the living cell, the functional context in which Ivylp exerts its effect is specified not only by its binding to Ypt7p, though, but also by other interactions. The results obtained in vitro show that binding of Ivylp to Ypt7p may encompass structural elements common to all members of the highly related Ypt GTPase family. Physiologically meaningless interaction with Ras-like GTPases other than Ypt7p may be avoided by subcellular compartmentalization of protein distributions or by blocking of binding sites through competitive binding of other, compartment-specific factors.

Deletion of the genes for Ivylp – interacting proteins (YPT7, VPS33) leads to strong defects in protein trafficking to the vacuole and in vacuolar morphology (cf. (Wichmann et al., 1992; Banta et al., 1990; Wada et al., 1990)). Deletion of *IVY1* has proved phenotypically neutral under all conditions tested. *IVY1* is dispensable for normal growth under non-competitive laboratory conditions. No protein sorting defect and no alterations in vacuolar morphology could be observed in haploid ivyl null mutants. The absence in *S. cerevisiae* of any gene that shows even a weak homology to *IVY1* suggests that the inconspicuousness of the ivyl deletion does not result from functional redundancy. Although its precise role in normal cellular physiology is still obscure, it can be surmised that the presence of a functional copy of the non-redundant *IVY1* gene increases the fitness of the organism, although presumably in a subtle way (Thatcher et al., 1998).

In cells of exponentially growing cultures, the level of Ivylp HAp detectable with anti-HA antisera is apparently low, suggesting that in unmodified cells the Ivylp content is equally low under these conditions. The low amount of Ivylp in vigorously growing cells impeded detection by indirect immunofluorescence. It can therefore not be said if the low amount of protein at this stage localizes in the same way as the increased amount in stationary cells and is just below the threshold of detection, or whether Ivylp is distributed differently in exponentially growing cells. In cultures approaching the stationary phase, the Ivylp content of the cells rises. In the latter case it is possible to localize the protein to the vacuolar membrane by indirect immunofluorescence. This
Overexpression of Ivy1p leads to p2-CPY misorting. (A) In vivo overlay assay of pro-CPY-invertase secretion. yp7 deletion cells and wild-type BHY 11 cells are used as controls. Wild-type cells (transformed with unmodified plasmids) show no colour development while yp7A cells (containing insert-free plasmids to allow growth on selective media) show a colour reaction due to misrouted pro-CPY-invertase into their periplasm. Colour develops after approximately 15 minutes due to an enzyme-catalyzed oxidation of the leuko-dye O-dianisidine. BHY11 cells with a multi-copy plasmid constitutively expressing large amounts of Ivy1p (+ pTL89, bottom; IVY1 expression driven by the TPI promoter) show a colour reaction. Strong overexpression of Ivy1p leads to the misorting of soluble proteins containing a vacuolar sorting signal in their polypeptide chain. BHY 11 reporter cells containing IVY1 on a multi-copy plasmid under the control of the GAL1 promoter (+ pTL90, right side) do not show any colour development when IVY1 expression is not induced. An ivy1Δ strain (TAL43) tested by this overlay secretion assay behaved like the isogenic wild-type strain (not shown). (B) Radioactive labelling and immunoprecipitation. Cells were subjected to immunoprecipitation with anti-CPY antibodies after in vivo labeling with radioactive amino acids. After a 15-min pulse, time 0 samples were removed and the remaining protoplasts incubated for another 45 min. Wild-type cells show a normal processing of pro-CPY. Cells overexpressing Ivy1p (+ pTL89) show a marked retention inside the cells of the p2-proform of CPY after 45 min. Part of the non-matured p2-proenzyme is exported from the cell by Golgi-derived vesicles (e = extracellular). This portion is seen at time 0; the secreted CPY at 45 min chase time was probably degraded. Additional, plasmid-borne VPS33 copies inside cells overexpressing Ivy1p suppress the retarding effect on CPY maturation and restore the wild-type condition. i = intracellular, e = extracellular. 0 = time zero, end of pulse, start of chase period. 45 = situation after 45 minutes of chase after addition of an excess of non-radioactive cysteine/methionine. p1 = core-glycosylated, endoplasmic reticulum pro-CPY; p2 = fully glycosylated, Golgi apparatus pro-CPY; m = matured, Pep5p—processed vacuolar CPY; * = “X”-CPY, an aberrant, presumably partially degraded, form of carboxypeptidase Y (cf. Wichmann et al., 1992)).

All morphological and biochemical Ivy1p overexpression-mediated effects can be completely reversed by raising the amount of Vps33p. To suppress the overexpression effects of Ivy1p, it is sufficient to provide the cells with a 2-micron multi-copy plasmid harbouring the VPS33 gene in its native form. No rescue could be achieved by raising the amount of Ypt7p in the same way. From this remediation effect of Vps33p it follows that the overexpression effects seen by greatly increasing the amount of Ivy1p are threshold effects.

The idea that a certain “threshold” amount of Ivy1p needs to be present in a cell to confer the observed mutant phenotype is further supported by the fact that many but not all cells of a multi-copy transformed yeast cell culture exhibit the overexpression effects. As the cellular copy number is not precisely set in the case of 2-micron multi-copy plasmids, so that the individual plasmid copy number can vary by quite large a margin, the absence of the mutant features in some cells is explicable and, indeed, expected. The threshold theory of the Ivy1p action is supported by the observation that the overexpression effects are not seen until the liquid culture reaches a late, stationary growth phase after a shift to inducing galactose medium when IVY1 expression is driven by an inducible GAL1 promoter.

The fact that Ivy1p binds to Vps33p suggests an interpretation of the effects seen when Ivy1p is overexpressed and their perfect suppression by increased amounts of Vps33p. Vacular fragmentation, missorting of vacuolar proteins and the accumulation of multivesicular bodies could be due to the sequestration of the cellular Vps33p pool by the excess Ivy1p. By binding a critical portion of the cellular Vps33p, Ivy1p in this model would remove an important component of the vacuolar docking/fusion HOPS protein complex. Deletion of VPS33 leads to a vps class C mutant phenotype characterized by a severe fragmentation of the vacuole, missorting of pro-CPY to the periplasm and a severe growth retardation (Wada et al., 1990; Banta et al., 1990). The severity of the Ivy1p overexpression effects is more akin to vps class B mutants, though, and fragmentation of vacuoles is less pronounced than in vps32A cells. Either Ivy1p does not scavenge the entire Vps33p pool, so that a residual fusion activity remains, or its action on the vacuolar fusion machinery does not affect all pathways leading to this organelle in the same way as the deletion of VPS33 does, or, most probably, the mode of action of Ivy1p contains additional levels.

This last possibility is lent support by the striking appearance of a vast number of multivesicular bodies. The generation or accumulation of multivesicular bodies inside cells overexpressing Ivy1p is a novel feature, and a massive accumulation of membrane-enclosed entities of this or a similar sort has, to our
knowledge, never before been described in the published scientific literature on yeast cells. Multivesicular bodies (MVBs) are believed to represent a particular type of endosome in which ubiquitinated membrane proteins are delivered to the vacuole for degradation. MVBs are thought to form by invagination of endosomal membranes which pinch off to yield the vesicles found inside multivesicular bodies (Reggiori and Pelham, 2001). The massive accumulation of these organelles in yeast cells overproducing Ivy1p suggests that Ivy1p either interferes with endosome-vacuole fusion in a particular way (as MVB accumulation is not seen in vps/vam mutants in which vacuolar fusion is disabled or severely inhibited – or has escaped attention), or Ivy1p serves a role in an as yet undefined, novel pathway involving MVB-vacuole fusion. Both the vps and vam mutant phenotypes in cells with enlarged amounts of Ivy1p can successfully be suppressed when the cellular content of Vps33p is enhanced at the same time. Ivy1p and Vps33p are, at least to some degree, functionally reciprocal.

Phospholipids of the phosphoinositide group have only recently been found to be important regulatory substances of membrane fusions. As Ivy1p strongly binds to members of this group of lipids, it could be responsive to phosphoinositol-4,5-bisphosphate, which Ivy1p is likely to meet at the vacuolar membrane, and which Mayer et al. (2000) have found to be important in two chronologically distinct steps of vacuolar membrane fusions. It is this lipid that in vitro shows the highest affinity for Ivy1p (Fig. 4). It is possible that the Ivy1p-lipid interaction facilitates membrane interaction in the first place which is then followed by interaction with Ypt7p and Vps33p. We cannot exclude that Ivy1p in addition reacts to another, not yet defined, lipid signal. It is intriguing, though, that the GTPases Cdc42p and Rho1p are required for vacuolar fusion downstream of Ypt7p (Müller et al., 2001; Eitzen et al., 2001). Cdc42p had previously been shown to bind directly to PI-4,5-bisphosphate (Zheng et al. 1996). In addition, phosphoinositide lipids have been found to bind to guanine nucleotide exchange factors (GEFs) and GTPase-activating proteins (GAPs) via interaction with the PH (pleckstrin homology) domain found in these proteins (Welch et al., 2002; Krugmann et al., 2002; Han et al., 1998). Rho1p is activated by the PH domain-containing GEF Rom2p in a Ypt7p-dependent way (Wickner, 2002). The vacuolar fragmentation seen upon overexpression of Ivy1p could therefore be the result of its lipid-binding capacity. Excess amounts of Ivy1p could be envisaged to sequester the vacuolar PI-4,5-bisphosphate, inhibiting the action of Cdc42p and/or Rom2p by stripping them of their regulatory lipid. PI-4,5-bisphosphate was shown by Zheng and colleagues (1996) to activate Cdc42p directly through the stimulation of GDP-release. It is not known if other Ras-related GTPases such as Ypt7p can be activated in the same way. In addition, a detailed picture of the interactions at the cellular membrane of mammalian Cdc42 bound to its guanine nucleotide exchange factor Dbs, including PI-4,5-bisphosphate, has recently been described (Rossman et al., 2002), suggesting an allosteric function for the phosphoinositide in the GTPase-GEF interaction. From this working hypothesis a model for the physiological role of Ivy1p can be derived: if PI-4,5-bisphosphate were rate-limiting for the progression of vacuolar membrane fusion, the increasing amounts of Ivy1p in aging cells could serve a regulatory role in fine-tuning or shutting down delivery of vesicular carriers/late endosomes to the vacuole in non-dividing, stationary yeast cells. Many of the Ypt GTPases are still devoid of described proteinaceous activating factors, i.e. guanine-nucleotide exchange factors. Maybe lipids – alone or as co-factors of proteins – fulfill this function for some of the Ypt proteins. The presumptive GEF for Ypt7p, Vps39p (a member of HOPS), in vitro displays a rather low activity towards its cognate GTPase (Wurmser et al., 2000). Maybe its activity in vivo is enhanced by additional interactions with other proteins or/and lipids.

This model is able to explain the effects seen when Ivy1p is overexpressed. How increasing the amount of Vps33p (part of the Ypt7p-binding HOPS complex) facilitates the relieve of the detrimental effects of excess Ivy1p is mechanistically not fully understood at the moment. Maybe Vps33p contributes to the action of Rom2p, thereby short-circuiting the need for phospholipids. Despite the severity of the respective deletion-strain phenotype (vps, class C), the role of Vps33p in the HOPS complex is still poorly understood.

In addition to the interactions described in this publication, Ivy1p has been found in two-hybrid system studies to interact physically with a number of yeast proteins other than the ones described here. The interactions are listed at the internet site: http://www.proteome.com/databases/YPD/reports/YDR229 W.html. To our knowledge the physiological relevance of these interactions has not been confirmed. The possibility cannot be ruled out that some of these interactions may be artifactual. Ivy1p is said to interact with: Rvs167p (Ydr388w, a protein which seems to be involved in endocytic vesicle trafficking (Bon et al., 2000); Crm1p (Ygr218w; Exportin); Yjl066p (function unknown); Yj017p (function unknown); Cod2p (Yln041c; a component of the Sec34p-Sec35p complex involved in vesicular transport to the Golgi apparatus). Except for Rvs167p, all of these interactions were found in high-throughput two-hybrid screens, an experimental approach the “in-vivo validity” of which has to be borne out through confirmatory studies. The interaction of Ivy1p with the protein Rvs167p in the two-hybrid system as described by Bon et al. (2000), could be confirmed by us. In an affinity chromatographic experiment HA-tagged Rvs167p was retained by GST-Ivy1p (not shown). If this interaction is of relevance in the living cell has to be determined.

In summary, our results suggest a role for Ivy1p in the vacuolar fusion machinery of the yeast cell as a antagonist for Vps33p. In this way it might add an additional level of regulation to the HOPS-controlled docking step of vacuolar membrane fusion.

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