A plant proton-pumping inorganic pyrophosphatase functionally complements the vacuolar ATPase transport activity and confers bafilomycin resistance in yeast

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INTRODUCTION

The occurrence of an electrochemical proton gradient across the internal membranes of eukaryotic cells is one of the known requirements for the correct function of the endo- and exo-cytic pathways [1]. Moreover, pH gradients are necessary not only for the correct functioning of the vesicle transport machinery, but also for many other organelar functions, such as, among others, dissociation of ligand–receptor complexes in receptor-mediated endocytosis or glycosylation of proteins in the Golgi apparatus [2]. A specific class of proton-pumping ATPases, V-ATPases (vacuolar ATPases or V-type ATPases), play an essential role in the generation of these positive-intraluminal proton gradients [1,3]. V-ATPases were originally described in the vacuolar membrane of plants and fungi (hence their name), thereafter they were found in many single-membrane organelles of eukaryotic cells. In addition, they have also been reported to occur at the plasma membrane of certain mammalian cell types [2]. All in all, these multi-subunit proteins fulfill important roles in diverse scenarios, both physiological and in disease, including cancer [3,4]. Not surprisingly then, there is an active search for specific inhibitors of these pumps that could be used both in basic research and in therapeutic applications. Bafilomycin A1 was the first reported specific inhibitor of V-ATPases [5], it consists of a large macrocyclic lactone ring which places it in the macrolide antibiotic group. Unlike other eukaryotes, Saccharomyces cerevisiae cells can grow either in the presence of V-ATPase inhibitors [6] or when a null mutation occurs in one of the genes encoding certain subunits of the V-ATPase complex [1]. However, they show a range of conditional-lethal phenotypes and thus become very sensitive to neutral/alkaline pH values [6,7], high extracellular calcium concentrations [8] and certain metal cations, such as zinc [9]; actually, the combination of sensitivity to high pH and extracellular calcium is widely accepted as the most characteristic Vma-deficient phenotype [1].

V-ATPases are very complex multi-subunit pumps that contain both membrane-bound and peripheral components. This paradoxical complexity, in comparison with other transporters, has been justified by proposing moonlighting functions of some subunits or even whole domains of the V-ATPases [10,11]. However, to study non-transport functions of V-ATPases is a complicated task if organelle acidification is not taken over by some other protein(s). Strikingly, this situation may be the natural scenario in plant cells and many protists where V-ATPases co-exist with alternative proton pumps. However, to date, no work has proven that V-ATPases can be unambiguously substituted by any other proton pump in vivo.

H⁺-PPases [proton-translocating PPases (inorganic pyrophosphatases)] are a type of homo-oligomeric membrane-embedded proteins containing a single highly hydrophobic subunit type that couples the energy obtained in the hydrolysis of PPI (inorganic pyrophosphate) to the transport of protons.

Key words: chimaeric membrane protein, green fluorescent protein, heterologous expression, proton-translocating inorganic pyrophosphatase, Saccharomyces cerevisiae, vacuolar-type ATPase.

Abbreviations used: ACMA, 9-amino-6-chloro-2-methoxy acridine; DIC, differential interference contrast; DTT, dithiothreitol; GFP, green fluorescent protein; H⁺-PPase, H⁺-translocating inorganic pyrophosphatase; PPase, inorganic pyrophosphatase; PPI, inorganic pyrophosphate; V-ATPase, vacuolar H⁺-ATPase; yEGFP, yeast enhanced GFP.

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across biological membranes [12–14]. They are the structurally simplest class of primary proton pumps known to date, and was the last one established, in the middle of the 1980s [12,13]. H+-PPases are reportedly located in cell membrane vesicles and acidicalcosomes of prokaryotes and diverse acidic single-membrane organelles (acidocalcosomes, lysosomes, vacuoles) of certain eukaryotes: protists and plants [13,15–17]. H+-PPases seem to co-localize with V-ATPases in those eukaryotes where they occur [18], thus, one point of discussion is the physiological significance of this apparent functional redundancy and whether H+-PPases contribute in varying degrees to the generation of proton gradients under diverse physiological scenarios.

We recently reported that translational fusions with certain N-terminal signal peptides enhance the expression levels and/or alter the subcellular distribution of diverse H+-PPases heterologously expressed in S. cerevisiae. When this approach was applied to the vacuolar H+-PPase AVP1 from Arabidopsis thaliana, a protein expressed at moderate levels in yeast in its natural form [19], its subcellular distribution was altered; thus, chimaeric proteins constructed by fusing the N-terminal signal sequence of the H+-PPase from the protist Trypanosoma cruzi [20] to AVP1 were shown to preferentially accumulate in intracellular acidic vesicles [21]. In the present study, we show that the expression of such an internal membrane-targeted chimaeric AVP1 derivative confers yeast resistance to the antibiotic macrolide bafilomycin A1, a specific V-ATPase inhibitor. Further experiments performed with a yeast vma mutant defective in its V-ATPase activity demonstrated that the H+-PPase acts by functionally complementing the transport function of the former proton pump. These results raise again the question of the presence of both proton pumps in certain organelles, such as the plant vacuole. Potential applications of this system in order to elucidate the roles of V-ATPases and/or H+-PPases are also discussed.

EXPERIMENTAL

Yeast strains

The YPC3 mutant strain was generated from the S. cerevisiae haploid strain W303-1A (MATa, ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1, ural3-1) by the single-step transplacement procedure as described previously [21]. This mutant has the IPP1 gene coding for the cytosolic soluble PPase under the control of the yeast galactokinase (GAL1) promoter. YPC4 was generated from S. cerevisiae mutant RS-1144 (W303-1A vma1::LEU2) following the same procedure. Mutant RS-1144, provided by Professor Ramón Serrano (Universidad Politécnica de Valencia, Spain), was derived from strain W303-1A by disrupting its VMA1 (TFP1) gene with the yeast LEU2 cassette [22]. The VMA1 gene of S. cerevisiae is translated by protein splicing into two different proteins: the 69 kDa catalytic subunit of the vacuolar H+-ATPase and a 50 kDa DNA endonuclease [23]. Vacuolar protease-deficient mutant BJ5457 was provided by Professor Andrés Aguilara (Universidad de Sevilla, Sevilla, Spain). This mutant carries mutations both in the PEPl gene and the PRBl gene, coding for the yeast vacuolar proteases PrA (Proteasein A) and PrB (Proteasein B) respectively [24]. Yeast strains used in the present study are described in Supplementary Table S1 at http://www.BiochemJ.org/bj/437/bj4370269add.htm.

Transformation of YPC3 and YPC4

S. cerevisiae mutant strains YPC3 and YPC4 were transformed with plasmids pRS699b (negative control), pAVP1, pTcAVP1, and pTcGFPAVP1 by using the method described by Schiestl and Gietz [25]. All plasmids are 2 μm-based derivatives of the URA3-containing Escherichia coli/S. cerevisiae shuttle plasmid pRS699

[26], as described previously [21] (Supplementary Table S2 in [21]). These plasmids bear the yeast PMA1 promoter for constitutive expression of inserts [26]. Three different constructs were made using the sequence coding for the K+-dependent H+-PPase from the higher plant A. thaliana (AVP1): pAVP1, carrying an intron-less open reading frame construct of AVP1; pTcAVP1, a 5′-in-frame fusion of the sequence coding for the N-terminal signal peptide of the T. cruzi H+-PPase (TcVP) with that of AVP1; and pTcGFPAVP1, an in-frame fusion where the sequence coding for yEGFP [yeast enhanced GFP (green fluorescent protein)] is placed between the coding sequence of the TcVP N-terminal signal peptide and that of AVP1. In addition to these, plasmids pIPP1, bearing the whole open reading frame encoding the cytosolic soluble PPase from yeast (IPP1), and pVMA1, bearing the genomic sequence of VMA1 coding for the A catalytic subunit of the yeast V-ATPase, were constructed using the above-mentioned expression plasmid backbone (Supplementary Figure S1 at http://www.BiochemJ.org/bj/437/bj4370269add.htm). YPC3 and YPC4 cells were initially grown at 30°C in galactose-containing synthetic medium [27] devoid of histidine (YPC3) or histidine and leucine (YPC4); transformants were selected by growing cells on 2% agar plates in culture medium without histidine and uracil (YPC3) or without histidine, leucine and uracil (YPC4). Sorbitol (0.5 M) was added to the plates where transformants of YPC4 were selected as an osmotic protectant.

Phenotype complementation tests

Complementation studies were performed by pre-inoculating 2 ml of galactose-containing selective medium with transformed cells from the plates and growing overnight at 30°C with agitation (200 rev/min). The following day, 2 ml of glucose-containing selective medium or YPD medium [1% (w/v) yeast extract/2% (w/v) peptone/2% (w/v) glucose] were inoculated with 20 μl (1:100 dilution) of cells previously grown on galactose and these were allowed to grow as described above. This treatment is necessary to bring down the PPase activity associated with Ipp1p [28]. After overnight growth on glucose, 10-fold serial dilutions of the cultures were made in sterile water and 5 μl drops of each dilution were spotted on to YPD and YPGal [1% (w/v) yeast extract/2% (w/v) peptone/2% (w/v) galactose] agar plates containing 50 mM Tris/Mes adjusted to pH 5.5, 7.5 or 8. CaCl2 (100 mM) or ZnCl2 (4 mM) was added to the plates at pH 5.5 where indicated. Plates were typically grown at 30°C for 2–5 days.

Fluorescence microscopy

Vacuolar lumen acidification was assessed by fluorescence staining with quinacrine as described previously [29]. Cells were visualized after 10 min of fluorephore accumulation with a fully automated Leica DM6000B microscope (Leica Microsystems) with FITC green fluorescence filters (excitation filter 480/40 nm, dichromatic mirror 505 nm, suppression filter 527/30 nm), a 100 x objective and equipped with a cooled CCD (charge-coupled device) camera (ORCA-AG, Hamamatsu Photonics). Vacuolar membrane labelling was carried out with the vital lipophilic dye FM4-64 as described previously [29]. Cells were visualized 30 min after incubation with the dye as described above, except that Texas Red filters (excitation filter 560/40 nm, dichromatic mirror 595 nm, suppression filter 645/75 nm) were used. In all cases, cells were also viewed using DIC (differential interference contrast) microscopy to observe cell morphology.

Isolation of yeast vacuolar membranes

Colonies of transformed YPC4 cells were collected from a plate and liquid-grown up to stationary phase in galactose-containing
selective medium; then, 4 ml of stationary cultures were used to inoculate 400 ml of YPD adjusted to pH 5.5 with 50 mM Tris/Mes and grown overnight to early exponential phase (A660 was approximately 1). Yeast vacuoles were obtained from these cultures using the method described by Roberts et al. [30]

Isolation of total membrane fractions from yeast
Preparation of membrane fractions from YPC4 cells transformed with the different plasmids were obtained by a modification of a method described previously [21]. Yeast colonies were collected from a plate and liquid-grown up to stationary phase in galactose-containing selective medium; then, 4 ml of stationary cultures were used to inoculate 400 ml of YPD. After overnight growth, cells were sedimented by centrifugation at 700 g for 10 min, washed thoroughly with water, resuspended in 5 ml of ice-cold buffer A [25 mM Tris/HCl (pH 8), 10% glycerol, 4 mM 2-mercaptoethanol, 2 mM DTT (dithiothreitol), 2 mM EDTA, 10 mM MgCl2, 1 mM benzamidine, 2 mM ε-aminocaproic acid and 1 mM PMSF], and homogenized by rigorous shaking with glass beads. The homogenate was diluted up to 25 ml with buffer B [10 mM Tris/HCl (pH 7.6), 10% glycerol, 2 mM DTT and 1 mM EDTA] and centrifuged for 10 min at 700 g to remove beads and debris. The resulting supernatant was centrifuged for 30 min at 40,000 rev./min (Beckman 60Ti rotor) thus yielding a total membrane pellet. This pellet was homogenized in stripping buffer [60 mM Tris/HCl (pH 8), 12% glycerol, 0.72 M KCl and 1.2 mM CaCl2] [31] and centrifuged for 20 min at 40,000 rev./min (Beckman 60Ti rotor). This pellet was subsequently washed with buffer B and centrifuged [40,000 rev./min (Beckman 60Ti rotor) for 20 min]. The final pellets were resuspended and homogenized in 1 ml of buffer B.

PPase activity and H+-translocation assays
Membrane-associated PPase activity was measured by spectrophotometric detection of released phosphate [32]. Proton-translocation activities were assayed by monitoring the fluorescence quenching of ACMA (9-amino-6-chloro-2-methoxy acridine) [33]. The statistical significance of the differences observed between sets of activity data was determined by Student’s unpaired t tests using the application available at http://graphpad.com/quickcalcs/ttest1.cfm.

Western blotting and total protein estimation
Western blot analysis was carried out after SDS/PAGE as described previously [34]. For immunodetection of H+-PPases, an affinity-purified polyclonal antibody raised in rabbit against the membrane-bound PPase (TVP) from the bacterium Thermotoga maritima [35] was used at a 1:200 dilution. For Vma1p, GFP and Pma1p immunodetection, commercial monoclonal antibody 8B1 (Molecular Probes), a polyclonal ‘Living Colors’ antibody from BD Biosciences and a polyclonal antibody provided by Professor Ramón Serrano (Universidad Politécnica de Valencia, Valencia, Spain) were used respectively, all of them at a 1:2000 dilution. For Vma1p immunodetection, commercial monoclonal antibody 8B1 (Molecular Probes), a polyclonal ‘Living Colors’ antibody from BD Biosciences and a polyclonal antibody provided by Professor Ramón Serrano (Universidad Politécnica de Valencia, Valencia, Spain) were used respectively, all of them at a 1:2000 dilution. The total protein concentration was estimated using the Bradford method [36] with ovalbumin as a standard.

RESULTS
Chimaeras of AVP1 bearing the N-terminal domain of TcVP induce bafilomycin resistance in YPC3 cells
YPC3 is a yeast mutant whose IPP1 gene coding for the essential cytosolic PPase is under the control of the galactokinase (GAL1) promoter, therefore these cells are non-viable when glucose is present as a carbon source. It has been shown previously that plant and bacterial membrane-bound H+-PPases can functionally complement the yeast cytosolic sPPase (Ipp1p), especially when fused to certain N-terminal signal peptides [21,28]. Addition of macroside bafilomycin A1 to YPC3 cells transformed with different plasmids bearing constitutively expressed genes coding for soluble or membrane-bound PPases resulted in sensitivity to neutral/alkaline pH and to the divalent cations Ca2+ and Zn2+, when grown on galactose (Figure 1), that is, when Ipp1p is expressed. In contrast, when cells were grown on glucose in the presence of bafilomycin A1, those cells transformed with plasmids pAVP1 and pTcAVP1 expressing internal membrane-targeted chimaeras could grow after 4 days, particularly the latter (Figure 1B).

Chimaeras of AVP1 bearing the N-terminal domain of TcVP reverse the sensitivity of YPC4 Vma1p- cells to neutral/alkaline pH, Ca2+ and Zn2+
In order to fully ascertain whether the observed bafilomycin A1 tolerance was actually brought about by complementation of V-ATPase proton-transport activity, similar experiments to those described above were performed with YPC4 cells. This mutant also has IPP1 under the control of the GAL1 promoter

Figure 1 Drop tests of yeast mutant YPC3 cells grown in the presence of the macroside inhibitor bafilomycin A1 at pH 5.5 and 8.0 (A) and in the presence of 100 mM CaCl2 or 4 mM ZnCl2 at pH 5.5 (B)
but, additionally, it has the gene coding for the catalytic subunit VMA1 of the vacuolar H^+-ATPase disrupted. YPC4, like YPC3, is unable to grow on glucose, but, as expected, when grown on galactose it exhibits the typical range of Vma^− phenotypes: very high sensitivity to neutral/alkaline pH values and to Ca^{2+} and Zn^{2+} cations (Figures 2A and 2B). However, when YPC4 cells were transformed with AVP1 and its chimaeric derivatives, they recovered the capacity to grow on glucose at pH 5.5 and 7.5, especially in the case of the TcGFPAVP1 chimaera. In the presence of Ca^{2+}, only cells transformed with plasmids pTcAVP1 and pTcGFPAVP1 were able to grow, whereas only the latter showed growth in the presence of Zn^{2+}. Again, as in the case of YPC3 with bafilomycin A1, the construct bearing the gene coding for Ipp1p could only support growth on glucose at pH 5.6 and in the absence of divalent cations. On the other hand, transformation of YPC4 with VMA1 resulted in the reversal of the phenotypes associated with the disruption of this gene only in the presence of galactose (Figures 2A and 2B).

Expression of TcGFPAVP1 allows acidification of vacuoles in YPC4 cells

To verify that AVP1 and its derivatives were able to promote acidification of internal compartments, we analysed the accumulation of quinacrine, a vital dye that concentrates in acidic lumina. Figure 3 shows typical patterns of quinacrine accumulation in vacuoles of YPC4 cells transformed with the different constructs tested. No internal acidification could be observed in cells expressing the soluble cytosolic PPase Ipp1p, whereas cells expressing AVP1 and TcAVP1 showed a punctate pattern of green fluorescence suggesting acidification of prevacuolar/endosomal compartments. In contrast with this, cells transformed with the plasmid pTcGFPAVP1 showed a different fluorescence pattern, with a clear vacuolar accumulation of quinacrine just like YPC4 cells transformed with plasmid pVMA1 grown in galactose. The former cells showed a punctate green fluorescence pattern, corresponding to the TcGFPAVP1 polypeptide, when visualized in the absence of quinacrine (see below). It is important to point out that some of the cells transformed with pTcGFPAVP1 exhibited clusters of smaller vacuoles rather than the single large vacuole usually observed in cells expressing Vma1p (see Figure 3). These results are in agreement with the previously presented drop tests (compare with Figures 1 and 2), which show comparatively higher growth of TcGFPAVP1-expressing clones at neutral/alkaline pH values and in the presence of Ca^{2+}/Zn^{2+}, these tolerance phenotypes being strictly dependent on physiologically competent acidic vacuoles.

Expression of all AVP1 derivatives allows normal internalization of the vital dye FM4-64

The vital fluorescent dye FM4-64 binds to the plasma membrane and follows the endocytic pathway to reach the vacuole [29]. FM4-64 internalization has been shown to be severely impaired in vma mutants [37]. Figure 4 shows labelling of vacuole membranes in YPC4 cells after incubation with FM4-64 for 30 min. It can be observed that cells expressing the AVP1
Functional complementation of yeast vacuolar \( H^+ \)-ATPase by a \( H^+ \)-PPase

Figure 4  Endocytic internalization of the fluorescent marker FM4-64 in transformed YPC4 cells

The rate of endocytosis was studied by incubating transformed YPC4 cells grown to early exponential phase with FM4-64 for 15 min, as described previously [28]. Cells were then transferred to YPD and visualized as described in Figure 3 after 30 min of incubation with the fluorescent dye. In (A), the left-hand column shows DIC microscopy of a typical field of transformed YPC4 cells, the middle column shows the pattern of FM4-64 labelling in the same cells, and the right-hand column shows the overlap of both. (B) YPC4 cells transformed with plasmid pTcGFPAVP1. From left-to-right: visualization of the cells by DIC microscopy, with red (Texas Red) and green (FITC) filters, overlap of red and green filter microphotographs, and overall overlap of the two former microphotographs.

Membrane-bound \( H^+ \)-PPases are present in YPC4 vacuoles

In order to check whether the different natural and chimaeric plant \( H^+ \)-PPases expressed in YPC4 cells were localized in the vacuoles, activity assays and immunodetection experiments were carried out with purified vacuole preparations obtained from YPC4 cells transformed with the different plasmids described above. Cells transformed with pTcGFPAVP1 showed a characteristic trace of ACMA fluorescence quenching that was dependent on the addition of PPi, this quenching was reversed by the addition of the \( H^+ /K^+ \) ionophore nigericin. Smaller quenching rates were obtained with cells transformed with pTcAVP1 and pAVP1 (Figure 5A). Vacuoles from cells transformed with pVMA1 did not show PPi-dependent ACMA quenching, just like cells transformed with plasmid pRS699b and pIPP1 (results not shown). When ATP-dependent proton transport was recorded, only vacuoles isolated from cells transformed with plasmid pVMA1 exhibited ACMA quenching (Figure 5A). This quenching was sensitive to bafilomycin A1 (results not shown). Membrane-associated PPase activity levels determined for vacuolar membranes obtained from the different YPC4 transformants were consistent with the \( H^+ \)-translocation results (Figure 5B), thus no PPase activity was detected in samples obtained from cells transformed with
Figure 5  PPi- and ATP-dependent proton-pumping activity measured as quenching of ACMA fluorescence (A), membrane-associated PPi, hydrolysis activity (B) and Western blot analyses (C) of vacuolar membrane preparations obtained from transformed YPC4 cells

(A) Proton-translocation activity was measured by recording the PPi-dependent (left-hand panels) and ATP-dependent (right-hand panels) ACMA fluorescence quenching. The upper panels show typical traces obtained in these experiments. The lower panels show initial rates of fluorescence quenching for the different preparations. Values are means ± S.E.M. for three independent experiments (n = 3) and asterisks indicate differences that are statistically significant (P < 0.005). n.d., not detected. Approximately 30 μg of protein was used for these assays. (B) Specific PPi-hydrolysis activity associated with vacuolar membranes obtained from YPC4 cells transformed with the different plasmids tested. Activity values shown are means ± S.E.M. (n = 3) using preparations obtained from YPC4 cells transformed with pRS699b as a reference. Asterisks indicate differences that are statistically significant (P < 0.005). Open and closed bars represent activities in the absence and presence of potassium chloride respectively. n.d., no detected activity. (C) Immunodetection of AVP1 (upper panel) and yeast Vma1p (middle panel) in vacuolar membranes of transformed YPC4 cells. Approximately 30 μg of protein was loaded per lane. Asterisks indicate bands associated with chimaeric protein TcGFPAVP1 (approximately 120 kDa) and AVP1 (72 kDa).

Immunodetection analyses carried out with total membrane preparations obtained with mutants YPC4 and BJ5457 using an antibody against a bacterial membrane-bound PPase (anti-TVP) showed remarkable results (Figure 6A). Thus, in the case of YPC4, a polypeptide of approximately 72 kDa was detected in all cases, although in cells transformed with pTcGFPAVP1, two more bands of approximately 120 kDa and 64 kDa were clearly recognized by the anti-TVP antibody. Peptide mass fingerprint analysis of the 120 kDa polypeptide by MALDI–TOF (matrix-assisted laser-desorption ionization–time-of-flight) MS showed that it actually corresponded to a fusion between GFP and AVP1 (results not shown). Consistently, an antibody

Vacuolar proteases are involved in the processing of native AVP1 and its chimaeric derivatives targeted to the vacuolar membrane

The possibility that AVP1 and its chimaeras might be subjected to proteolytic processing by vacuolar proteases in vivo was investigated by using the vacuolar-protease-deficient yeast mutant BJ5457 [24]. Immunodetection analyses carried out with total membrane preparations obtained with mutants YPC4 and BJ5457 transformed with plasmids pAVP1, pTcAVP1 and pTcGFPAVP1 using an antibody against a bacterial membrane-bound PPase (anti-TVP) showed remarkable results (Figure 6A). Thus, in the case of YPC4, a polypeptide of approximately 72 kDa was detected in all cases, although in cells transformed with pTcGFPAVP1, two more bands of approximately 120 kDa and 64 kDa were clearly recognized by the anti-TVP antibody. Peptide mass fingerprint analysis of the 120 kDa polypeptide by MALDI–TOF (matrix-assisted laser-desorption ionization–time-of-flight) MS showed that it actually corresponded to a fusion between GFP and AVP1 (results not shown). Consistently, an antibody

plasmids pRS699b, pIPP1 and pVMA1. Samples from cells expressing the different AVP1 chimaeras did show PPase activity that was increased by the addition of K+, a feature of AVP1 [19]. Immunodetection in Western blots of vacuolar membranes of the diverse native and chimaeric plant H+-PPases used in the present study (Figure 5C) gave somewhat unpredicted results despite the use of protease inhibitor cocktails during the isolation procedure, thus a polypeptide of the expected size for the native AVP1 protein (approximately 72 kDa) was obtained in the case of cells transformed with pAVP1, pTcAVP1 and pTcGFPAVP1; additionally, another band of approximately 64 kDa was present in these preparations. The relative amounts of H+-PPases immunodetected in these preparations were, however, in accordance with the estimated activity values. A minor band of approximately 120 kDa, the size expected for TcGFPAVP1, could also be detected only in samples obtained from cells transformed with plasmid pTcGFPAVP1 (compare with Figure 5C).
Functional complementation of yeast vacuolar \( H^+\)-ATPase by a \( H^+\)-PPase

Figure 6 Western blot analysis (A) and membrane-associated PPi hydrolysis activity (B) of total membrane preparations obtained from transformed YPC4 and BJ5457 cells, and fluorescence microscopy visualization of YPC4 and BJ5457 cells transformed with pTcGFPAVP1 (C)

(A) Immunodetection of AVP1 and derivatives with antibodies against a membrane-bound PPase (anti-TVP, upper panel) and GFP (anti-GFP) in total membranes of YPC4 (left-hand panels) and BJ5457 (right-hand panels) cells transformed with plasmids pRS699b, pIPP1, pAVP1, pTcAVP1 and pTcGFPAVP1. Approximately 70 \( \mu\)g of protein was loaded per lane. Asterisks indicate bands associated with chimaeric protein TcGFPAVP1 (approximately 120 kDa) and AVP1 (72 kDa). (B) Specific PPi hydrolysis activity associated with total membranes obtained from YPC4 and BJ5457 cells transformed with the different plasmids tested. Activity values shown are means \( \pm \) S.E.M. (\( n = 3 \)) using preparations obtained from cells transformed with pRS699b as a reference. Asterisks indicate differences that are statistically significant (\( P < 0.005 \)). Open and closed bars represent activities in the absence and presence of potassium chloride respectively. n.d., not detected. (C) YPC4 and BJ5457 cells were transformed with plasmid pTcGFPAVP1 and visualized in a Leica DM6000B microscope. Left-hand column shows DIC microscopy of a typical field of cells, the middle column shows the pattern of green fluorescence distribution (FITC green filter) in the same cells, and the right-hand column shows the overlap of both. White arrowheads indicate vacuolar membranes.

against GFP (anti-GFP) only recognized the 120 kDa band. In the case of mutant BJ5457, the anti-TVP antibody recognized polypeptides of approximately 72 kDa only in preparations obtained with cells transformed with plasmids pAVP1 and pTcAVP1, whereas a single polypeptide of approximately 120 kDa was detected in membranes of cells transformed with plasmid pTcGFPAVP1. The anti-GFP antibody could only detect the latter (Figure 6A).

Analysis of membrane-associated PPi hydrolytic activity showed that, in YPC4, the presence of GFP at the N-terminus of AVP1 increased the specific activity of the resulting protein 3–4-fold with respect to AVP1 and TcAVP1. In mutant BJ5457, the specific PPase activities associated with AVP1, TcAVP1 and TcGFPAVP1 were significantly increased with respect to the same proteins expressed in YPC4. The increases were much higher in the cases of AVP1 (approx. 6-fold) and TcAVP1 (approx. 4-fold) than in the case of TcGFPAVP1 (approx. 1.7-fold) (Figure 6B).

Fluorescent microscopy visualization of YPC4 and BJ5457 cells transformed with plasmid pTcGFPAVP1 remarkably showed different patterns of green fluorescence distribution, thus, in YPC4 cells, a punctate pattern was observed, suggesting a preferential accumulation of green fluorescence in internal membrane systems other than the vacuolar membrane. In contrast, green fluorescence mostly associated with the latter was clearly observed in BJ5457 cells (Figure 6C).
DISCUSSION

In previous studies, we emulated in yeast the cytosolic PP, metabolism of the plant cell, i.e. absence of a soluble cytosolic PPase, so that the PP, generated by the anabolism was efficiently hydrolysed by a membrane-bound PP-dependent proton pump [21,28]. In the present study, we have taken one step forward using a metabolic background in which the ATP-dependent proton pump of yeast internal membranes, the V-ATPase, has been either chemically inhibited or functionally impaired by mutation of a gene encoding an essential catalytic subunit. The results obtained strongly suggest that a distinct type of PP-dependent proton pump can efficiently substitute for the V-ATPase of S. cerevisiae in diverse physiological scenarios. Thus overexpression of the K+-dependent H+-PPase from the higher plant A. thaliana, a protein that occurs along with a V-ATPase in plant tonoplast [18], alleviates the range of phenotypes associated with the lack of the latter. The presence of an appropriate N-terminal domain proved to be important in order to observe the optimal phenotypic complementation in our system. The N-terminal domain with signal sequence features of a protistan H+-PPase, when fused to AVP1, alters the subcellular distribution of the latter so that it becomes preferentially located in internal membranes of yeast [21]. Not surprisingly, as shown in the present study, chimaeric proteins of AVP1, constructed with that signal peptide (TcAVP1 and TcGFPAVP1), functionally substituted for the V-ATPase more efficiently than the native AVP1 protein with no extra N-terminal domain. The difference in favour of TcGFPAVP1 compared with TcAVP1 is probably due to the higher overall expression levels attained by the former (see below). Actually, under non-restrictive conditions towards Vma- defects, TcGFPAVP1 supports growth of YPC4 more efficiently than TcAVP1 (Figure 2A) which means that the former is more efficient at complementing the phenotype associated with the absence of Ipp1p. Addition of GFP at the N-terminus of some membrane proteins has been previously reported either because it stabilizes proteins expressed in heterologous systems or because the presence of GFP at the C-terminus yields polypeptides with no activity [21,38,39]. In fact, both assumptions are true for the plant H+-PPase AVP1 expressed in yeast, as the present study and our previous study [21] demonstrates.

The results with both bafilomycin-treated YPC3 and YPC4 cells also showed that no version of heterologously expressed AVP1 was able to complement Vma- phenotypes in galactose, even though the corresponding genes are under the control of the constitutive PMA1 promoter. Similarly, when the parental strains (bafilomycin-treated W303-1A and RS-1114) were transformed with the same plasmids no complementation of the V-ATPase by AVP1 was observed either on galactose or on glucose (results not shown). These results suggested that AVP1 cannot pump protons when the soluble PPase is present in the cytosol, probably due to the competition for the substrate, therefore it was necessary to bring down the activity of the latter to negligible levels in order to have the former efficiently working in vivo. YPC3 and YPC4 were designed and generated so that this could be accomplished by transferring the cells from galactose to glucose and allowing them to grow for several hours. Actually, this reflects the metabolic scenario in cells of plant green tissues, where sPPases are restricted to energy-linked organelles (mitochondria and chloroplasts) and, probably, nuclei [17], although it must be mentioned that other PP,-utilizing enzymes involved in central carbon metabolism, such as a PP-dependent phosphofructokinase, are also present in the plant cell cytosol [40]. All in all, these results suggest that the presence of active soluble PPases is incompatible with H+-PPase functions, and helps to explain why the former are not found in the cytosol of eukaryotic cells if H+-PPases are present in endomembranes.

Acidification of internal compartments of YPC4 cells was studied with the fluorescent probe quinacrine, which accumulates in acidic vacuoles. TcGFPAVP1 was able to acidify the vacuole in YPC4 cells, although it also seemed to induce some fragmentation of this compartment. This has already been reported for YPC3 cells and may be related to the accumulation of this heterologous chimera protein in the internal compartments of yeast cells [21]. It should be noted in this respect that a more or less severe vacuolar fragmentation takes place in yeast under several, and not necessarily pathological, scenarios. In contrast, cells transformed with AVP1 and TcAVP1 showed a punctate pattern of quinacrine accumulation which probably means acidification of prevacuolar/endosomal vesicles. This pattern was more significant in the case of TcAVP1, probably due to the effect of the addition of the N-terminal signal domain added to the latter that favours accumulation of this chimera protein in internal membranes of yeast [21]. In the presence of quinacrine, cells expressing TcGFPAVP1 only exhibited the green fluorescence associated with the dye; however, these cells were also expected to show green fluorescence due to the GFP. Consequently, they were also visualized in the absence of quinacrine showing, in this case, a punctate pattern of fluorescence (see below). This suggests that quinacrine fluorescence masks that of GFP in our system when both are monitored simultaneously.

Results with FM4-64 are consistent with those obtained with quinacrine in the case of TcGFPAVP1; however, they further suggest that full acidification of the vacuole may not be absolutely essential to have the endocytic system operational at the right pace, acidification of the prevacuolar/endosomal compartments being sufficient. The fact that the green fluorescence observed in YPC4 cells transformed with plasmid pTcGFPAVP1 does not necessarily co-localize with the red fluorescence associated with acidic vesicles further suggests that either TcGFPAVP1 polypeptide is not significantly translocated to the vacuolar membrane or the GFP moiety of this chimera has been lost, presumably by proteolysis.

Biochemical analysis of vacuolar membrane preparations from YPC4 transformed with the different plasmids tested showed that AVP1 and all of its chimaeric derivatives were heterologously expressed and located, at least partially, in the vacuoles, as demonstrated by the data of membrane-associated H+-translocating and PPase activities, as well as by immunodetection analysis of vacuolar membrane preparations. Activity data were consistent with the phenotypes observed, thus TcGFPAVP1 showed higher PPase and H+-translocating activities than TcAVP1 which, in turn, produced higher activities than the native AVP1. Immunodetection with the anti-TVP antibody revealed that TcAVP1 and TcGFPAVP1 gave higher expression levels in this compartment than AVP1. This is consistent with previous reports regarding the role of the N-terminal signal domain of T. cruzi H+-PPase on the subcellular distribution of AVP1 [21]. On the other hand, a significant processing of all of the proteins examined was also found; actually, only a tiny amount of a polypeptide of approximately 120 kDa, corresponding to a fusion of GFP and AVP1, was detected, and another band of approximately 64 kDa was consistently observed in all cases. This evidence suggested that vacuolar proteases may be involved in the cleavage of AVP1 and its derivatives, with a concomitant degradation of the GFP in the case of TcGFPAVP1. In order to study this situation, the vacuolar-deficient yeast mutant BJ5457 [24] was transformed with plasmids pAVP1, pTcAVP1 and pTcGFPAVP1 and total membrane preparations obtained with the resulting strains were analysed and compared with preparations obtained with YPC4.
cells transformed with the same plasmids. Immunochemical analysis strongly suggested that vacuolar proteases are involved in the processing of AVP1 and chimaeric derivatives observed in mutant YPC4. These results, along with the membrane-associated PPase activity data, further support the idea that, although AVP1 and TcAVP1 can be found in YPC4 vacuoles, a high proportion of these polypeptides is subjected to proteolysis. This effect is much smaller in the case of TcGFPAVP1, thus suggesting that the combination of the T. cruzi H-PPase N-terminal signal peptide and yEGFP is more efficient than the signal peptide alone at altering the subcellular distribution of the resulting chimera. Fluorescence microscopy supported the data obtained by immunodetection and activity assays, thus, unlike in mutant BJ5457, the green fluorescence corresponding to TcGFPAVP1 in mutant YPC4 showed a pattern not clearly associated with the vacuolar membrane. There are two contributions to this effect according to our results: on the one hand, a significant fraction of TcGFPAVP1 is targeted to other internal membranes and, on the other hand, the polypeptide located in the vacuolar membrane is subjected to proteolysis so that most of the GFP moiety is degraded. Therefore, the role of vacuolar proteases provides an explanation of why the fluorescence associated with GFP is not located in the vacuolar membrane and further suggests that the proton pump responsible for the acidification of the vacuole in YPC4 cells transformed with pTcGFPAVP1 is actually a native-like AVP1 protein and not the chimaeric derivative.

Taken together our results show that the PP-dependent proton pump AVP1 can alleviate the most characteristic phenotypes associated to the lack of V-ATPase in yeast, especially when the former is preferentially targeted to internal membranes by attaching an appropriate signal peptide to its N-terminus [21]. In the opinion of the authors, this is the first direct proof that a H-translocating PPase can sustain growth and significant acidification of intracellular lumina in the absence of any V-ATPase activity.

These results raise another question related to the occurrence of two proton pumps in the same subcellular location in certain organisms, one of them being an extremely complex molecular machine (the V-ATPase), and the other a much simpler membrane pump. There are numerous recent data supporting that V-ATPases may play more roles than the acidification of internal compartments, such as the participation of the V$_s$ domain in membrane fusion [41] and others [42]. On the other hand, the V-ATPase complex may offer many more possibilities of membrane fusion [41] and others [42]. On the other hand, the V-ATPase complex may offer many more possibilities of membrane fusion [41] and others [42].

In any case, the H-PPases probably imply an adaptive advantage to the organisms where they occur because they offer an alternative pump to generate proton gradients in internal organelles by using an abundant by-product of anabolism, PP, [43]. This is likely to be a critical issue under conditions that require the optimization of cellular energy usage. In this respect, it is worthwhile pointing out that the H-PPase seems to be the main tonoplast pump in other plant systems, such as grape berry cells [44].

The yeast system described in the present work may have biotechnological applications, thus it is a good experimental system to accomplish studies of random mutagenesis aimed at obtaining enhanced versions of AVP1 and other PP-dependent ion pumps. On the other hand, YPC4 transformed with diverse H-PPases might be a useful system to check for possible roles other than the acidification of internal organelles proposed for fungal and animal V-ATPases, and it may be a suitable system to study the effect of H-PPases inhibitors in vivo, as these proton pumps occur in micro-organisms responsible for diseases such as malaria, leishmaniasis, Chagas disease and sleeping sickness [15,45]. Finally, this experimental system envisages novel ways of engineering cell tolerance to macrolides, a class of drugs of increasing relevance in therapeutics of a range of diseases in which cellular proton homeostasis may play crucial roles, such as cancer.

**AUTHOR CONTRIBUTION**

José Pérez-Castielva carried out most of the experimental work and wrote the paper; Agustín Hernández generated the YPC4 mutant and corrected the paper; Rocío Drake performed preliminary studies; and Aurelio Serrano supervised the work and wrote the final version of the paper.

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SUPPLEMENTARY ONLINE DATA

A plant proton-pumping inorganic pyrophosphatase functionally complements the vacuolar ATPase transport activity and confers bafilomycin resistance in yeast

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Figure S1 Schematic diagram showing the constructions used for plasmidic expression in yeast of AVP1, AVP1-derived chimaeric H⁺-PPases, IPP1 and the vacuolar ATPase subunit VMA1

Table S1 Yeast strains utilized in this article and their parental strains

<table>
<thead>
<tr>
<th>Strains</th>
<th>Genotype</th>
<th>References</th>
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<td>Parental strains</td>
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<td>W303-1A, RS-1144</td>
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<td>Strains</td>
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<tr>
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<td>W303-1A (pIPP1:PGAL1-IPP1 HIS3)</td>
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<td>Other strains</td>
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<tr>
<td>BJ5457</td>
<td>MATa pop4::HIS3 prb1-Δ tpi1 ura3-52 leu2-Δ his3-Δ lys2-801 can1</td>
<td>[4]</td>
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