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## Genomic Expression Program Involving the Haa1p-Regulon in *Saccharomyces cerevisiae* Response to Acetic Acid

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#### Abstract

The alterations occurring in yeast genomic expression during early response to acetic acid and the involvement of the transcription factor Haa1p in this transcriptional reprogramming are described in this study. Haa1p was found to regulate, directly or indirectly, the transcription of approximately 80% of the acetic acid-activated genes, suggesting that Haa1p is the main player in the control of yeast response to this weak acid. The genes identified in this work as being activated in response to acetic acid in a Haa1p-dependent manner include protein kinases, multidrug resistance transporters, proteins involved in lipid metabolism, in nucleic acid processing, and proteins of unknown function. Among these genes, the expression of *SAP30* and *HRK1* provided the strongest protective effect toward acetic acid. *SAP30* encode a subunit of a histone deacetylase complex and *HRK1* encode a protein kinase belonging to a family of protein kinases dedicated to the regulation of plasma membrane transporters activity. The deletion of the *HRK1* gene was found to lead to the increase of the accumulation of labeled acetic acid into acid-stressed yeast cells, suggesting that the role of both *HAA1* and *HRK1* in providing protection against acetic acid is, at least partially, related with their involvement in the reduction of intracellular acetate concentration.

#### Introduction

THE SACCHAROMYCES CEREVISIAE TRANSCRIPTIONAL AC-TIVATOR Haa1p was first included into a family of copperregulated transcription factors, based on the identification of a putative copper-regulatory domain within its DNA binding domain (Keller et al., 2001). However, unlike its homologous proteins, the function of Haa1p is independent of the copperstatus of the cell (Keller et al., 2001) and a biological role for Haa1p was related with yeast resistance to acetic acid and propionic acid (Fernandes et al., 2005). The expression of the HAA1 gene was shown to lead to the reduction of the duration of the adaptation period of a yeast cell population suddenly exposed to toxic concentrations of these low chain carboxylic acids, by decreasing the loss of cell viability occurring during the phase of latency (Fernandes et al., 2005). More recently, Haa1p was also described as a determinant of yeast resistance to lactic acid (Abbott et al., 2008). Acetic and propionic acids are widely used by food and beverage industries to prevent microbial spoilage often caused by yeasts and other fungi at low pH. However, some strains can adapt and proliferate even in the presence of concentrations of these chemicals, which are close to the legally permitted values, causing serious economic losses (Fleet, 2007). Acetic acid and propionic acid are also bypass products of *Saccharomyces cerevisiae* alcoholic fermentation, and their accumulation in the growth medium to toxic levels contributes to decreased ethanol yields and may lead to stuck and sluggish fermentations (Gibson et al., 2007). Acetic acid is also one of the inhibitory compounds resulting from the pretreatment of lignocellulosic substrates used for bioethanol production (Almeida et al., 2007). It is thus expected that the understanding of the molecular mechanisms underlying yeast resistance to acetic acid and propionic acid may be successfully used to guide the design of more efficient weak acid-based food preservation strategies and of more robust industrial yeast strains.

During the early response of unadapted yeast cell populations to acetic acid, propionic acid, or lactic acid, the rapid transcriptional activation of a small set of genes, previously shown to be transcriptionally regulated in the dependence of *HAA1* expression (Keller et al., 2001), was found to occur (Abbott et al., 2008; Fernandes et al., 2005). These genes include: (1) *TPO2* and *TPO3*, encoding two plasma membrane multidrug resistance (MDR) transporters of the major

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facilitator superfamily involved in the efflux of polyamines (Albertsen et al., 2003); (2) YGP1, encoding a cell wall glycoprotein expressed under nutrient limitation (Destruelle et al., 1994); (3) other genes encoding proteins of unknown or poorly characterized function (Abbott et al., 2008; Fernandes et al., 2005). Among these genes, only the expression of TPO2, TPO3, and YGP1 exerts a moderate protection against acetic acid and propionic acid, but the level of protection is much below the one exerted by Haa1p itself (Fernandes et al., 2005). The protective effect of both TPO3 and HAA1 genes was related with their involvement in the reduction of the intracellular concentration of acetate during acetic acid challenge, being the effect exerted by Haa1p much more evident than the one associated to the expression of its target-gene TPO3 (Fernandes et al., 2005). The much higher susceptibility of  $\Delta haa1$  cells to acetic acid and lactic acid, compared to the susceptibility exhibited by cells devoid of any of its initially described targets (Abbott et al., 2008; Fernandes et al., 2005; Keller et al., 2001), suggests that the Haa1p-regulon active under stress imposed by those weak acids includes other unidentified genes (Keller et al., 2001). Consistent with this idea, the SPI1 gene was recently identified as a novel weak acid-responsive gene dependent on HAA1 expression (Simões et al., 2006). SPI1 encodes a GPI-anchored cell wall protein required for the weak acid-induced remodeling of cell wall structure (Simões et al., 2006) but, contrasting with Haa1p, the protection exerted by Spi1p against acetic acid is negligible, its protective action being more evident toward the more lipophilic weak acids (Simões et al., 2006). In the present study we explored a transcriptomic approach to search for the complete Haa1p-regulon acting in yeast cells following sudden exposure to acetic acid stress. A number of novel genes regulated directly or indirectly by Haa1p, are here described for the first time, some of which play a role in yeast resistance to acetic acid.

#### Materials and Methods

#### Strains and growth media

The parental strain *Saccharomyces cerevisiae* BY4741 (*MATa*,  $his3\Delta 1$ ,  $leu2\Delta 0$ ,  $met15\Delta 0$ ,  $ura3\Delta 0$ ) and the derived mutant strains used in this work were obtained from the Euroscarf collection. Cells were batch-cultured at 30°C, with orbital agitation (250 rpm), in MM4 liquid medium that contains, per liter: 1.7 g yeast nitrogen base without amino acids or NH<sub>4</sub><sup>+</sup> (Difco Laboratories, Detroit, MI, USA), 20 g glucose (Merck, West Point, PA, USA), 2.65 g (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (Merck), 20 mg methionine, 20 mg histidine, 60 mg leucine, 20 mg uracil, all from Sigma (Spain). Yeast Peptone Dextrose (YPD) solid medium, containing, per liter, 20 g glucose, 20 g bactopeptone (Difco), 10 g yeast extract (Difco), and 20 g/L of agar (Iberagar, Barreiro, Portugal), was used to assess viable cell concentration.

#### Acetic acid susceptibility assays

The susceptibility to acetic acid of the parental strain BY4741 and selected deletion mutants was assessed based on the comparison of the growth curves in liquid MM4 medium supplemented with 60 mM acetic acid (acidified to pH 4.0 with HCl). Cells were cultivated in unsupplemented MM4 (pH 4.0) growth medium until mid-exponential phase (standard  $OD_{600 nm}$  0.4) and used to reinoculate (at an initial

 $OD_{600 nm}$  of  $0.05 \pm 0.005$ ) the fresh growth medium supplemented with the acid. Growth was followed by measuring culture  $OD_{600 nm}$ .

# Effect of Haa1p in yeast transcriptional response to acetic acid

The effect of Haa1p in yeast response to acetic acid stress was carried out by comparing the transcriptomes of cells of S. cerevisiae BY4741 and of the mutant BY4741\_ $\Delta$ haa1 in the presence or absence of acetic acid. Cells were cultivated in MM4 growth medium (at pH 4.0) until mid-exponential phase (standard  $OD_{600 nm} 0.8$ ) and then reinoculated into fresh medium either or not supplemented with 50 mM of acetic acid (at pH 4.0). After 30 min of incubation in this medium in the absence (sample A) or presence (sample B) of acetic acid, cells were harvested, immediately frozen in liquid nitrogen, and kept at -80°C until total RNA extraction. Total RNA extraction was performed according to the hot phenol method. Concentration and purity was determined by spectrophotometry and integrity was confirmed using an Agilent 2100 Bioanalyzer with a RNA 6000 Nano Assay (Agilent Technologies, Palo Alto, CA, USA). RNA was processed for use on Affymetrix (Santa Clara, CA, USA) GeneChip Yeast Genome S98 Arrays, according to the manufacturer's One-Cycle Target Labeling Assay. Briefly,  $5 \mu g$  of total RNA containing spiked in Poly-A RNA controls (GeneChip Expression GeneChip Eukaryotic Poly-A RNA Control Kit; Affymetrix) was used in a reverse transcription reaction (One-Cycle DNA synthesis kit; Affymetrix) to generate first-strand cDNA. After secondstrand synthesis, double-stranded cDNA was used in an in vitro transcription (IVT) reaction to generate biotinylated cRNA (GeneChip Expression 3'-Amplification Reagents for IVT-Labeling; Affymetrix). Size distribution of the cRNA and fragmented cRNA, respectively, was assessed using an Agilent 2100 Bioanalyzer with a RNA 6000 Nano Assay. A total of  $15 \,\mu g$  of fragmented cRNA was used in a  $300 - \mu L$  hybridization containing added hybridization controls. Mixture (200  $\mu$ L) was hybridized on arrays for 16h at 45°C. Standard posthybridization wash and double-stain protocols (EukGE-WS2v5) were used on an Affymetrix GeneChip Fluidics Station 400. Arrays were scanned on an Affymetrix GeneChip scanner 3000. To ensure the reliability of the analysis, two biological replicates were used and the replicate data for the same sample was weighted gene-wise using inverse square SE as weights. The scanned arrays were analyzed first with Affymetrix MAS 5.0 software to obtain Absent/Present calls and subsequently with DNA-Chip Analyzer (dChip) 2006 (http://www.dchip.org, Wong Lab, Harvard). The arrays were normalized to a baseline array with median CEL intensity by applying an Invariant Set Normalization Method (Li and Hung Wong, 2001). Normalized CEL intensities of the eight arrays were used to obtain model-based gene expression indices based on a PM (Perfect Match)-only model (Li and Hung Wong, 2001). Genes compared were considered to be differentially expressed if they were called Present in at least one of the arrays and if the 90% lower confidence bound of the fold change between experiment and baseline was above 1.2. For downstream analysis of acetic acid-responsive genes only the genes whose transcriptional change was above or below 1.5-fold the values registered in control cells were considered to be altered. Clustering of the up- or downregulated genes,

based on biological function, was performed using the MIPS Functional Catalogue (http://mips.gsf.de/projects/funct) and the description of gene function is based on the information available in SGD (www.yeastgenome.org).

To confirm some of the results obtained in the microarray analysis carried out, the transcript levels of selected acetic acid-responsive genes were compared by real-time RT-PCR in cells of the parental strain BY4741 and in the *Ahaa1* mutant. For this, cells were cultivated in conditions similar to those used for the microarray analysis and total RNA extraction was performed using the same protocol. Synthesis of cDNA from total RNA samples was performed using the MultiscribeTM reverse transcriptase kit and the subsequent real time RT-PCR step was carried out using SYBR<sup>®</sup> Green reagents 7500 RT-PCR in a Thermal Cycler Block (Applied Biosystems, Bedford, MA, USA). Specific primers for the amplification of the selected genes and of the *ACT1* gene, used as an internal control, were designed using Primer Express Software (Applied Biosystems).

## Extracellular acidification curves promoted by yeast cells in the presence or absence of acetic acid

To compare the *in vivo* active export of protons from cells of the parental strain BY4741 or the  $\Delta hrk1$  mutant in the presence of increasing concentrations of acetic acid, the values of pH of the aqueous cell suspension medium was followed for 10 min. Cells of both strains were grown in MM4 medium (at pH 4.0) and harvested by centrifugation (8,000 rpm, 5 min, 4°C) when the culture reached a standardized  $OD_{600 \text{ nm}}$  of 0.5–0.6, in mid-exponential phase. Cells were washed twice with icecold water and then incubated in sorbitol (20g/liter) for 30 min to deenergize the cells and deactivate plasma membrane H<sup>+</sup>-ATPase (Portillo and Serrano, 1989). After this, cells were washed and resuspended in distilled water to obtain a dense cell suspension ( $OD_{600 nm}$  of  $20.0 \pm 2.0$ ). Acidification experiments were carried out in a water-jacketed cell of 10-mL capacity, at 30°C, containing 3.0 mL of water with acetic acid (0, 0.4, 0.8, and 1.2 mM) and 1 mL of the above-described cellular suspension. The pH of the resulting suspension was adjusted to  $4.0 \pm 0.05$  and 1 mL of 100 g/L glucose (at pH 4.0) was added (to obtain a final concentration of 20 g/L). The variation of extracellular medium pH was followed by potentiometry using a pH microelectrode attached to a pH meter (Metrohm 605).

#### [1-14C]-acetic acid accumulation assays

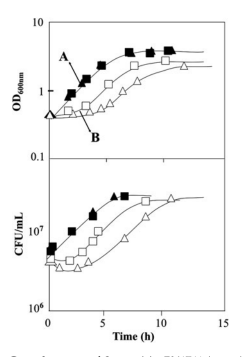
The accumulation ratio of  $[1^{-14}C]$ -acetic acid (defined as the ratio between intracellular and extracellular concentrations of the radiolabeled chemical) in cells of the parental strain BY4741 or the mutant  $\Delta hrk1$  was followed for 30 min during cultivation in MM4 growth medium (at pH 4.0) supplemented with 60 mM of cold acetic acid. Cells used to perform this assay were grown until mid-exponential phase (at the standardized  $OD_{600 \text{ nm}} = 0.5 \pm 0.05$ ) in MM4 growth medium (pH 4.0) and harvested by filtration (Supor®200 membranes,  $0.2 \mu$ m). Following harvesting, cells were washed with fresh MM4 growth medium (pH 4.0) and resuspended in 5 mL of the same growth medium to obtain a cell suspension with  $OD_{600 \text{ nm}} = 0.7 \pm 0.05$ . After 5 min of incubation at 30°C with agitation (150 rev/min) for temperature equilibration, 60 mM of cold acetic acid and  $10 \,\mu\text{M}$ of [1-14C]-acetic acid (sodium salt from GE Healthcare, Piscataway, NJ; 9.25 MBq) were added to the culture medium and incubation proceeded for 30 min. During this period of incubation, the intracellular accumulation of labeled acetic acid was followed by filtering  $200 \,\mu\text{L}$  of cell suspensions, at adequate intervals, through prewetted glass microfiber filters (Whatman GF/C). The filters were washed with ice-cold water and the radioactivity was measured in a Beckman LS 5000TD scintillation counter. Extracellular [1-<sup>14</sup>C]-acetic acid was estimated by measuring the radioactivity of 50  $\mu$ L of the supernatant. The nonspecific [1-<sup>14</sup>C]acetic acid adsorption to the filters and to the cells (<5% of the total bound-radioactivity) was assessed and taken into consideration. To calculate the intracellular concentration of  $[1-^{14}C]$ -acetic acid, the internal cell volume (Vi) of the different strains was considered constant and equal to  $2.5\,\mu\text{L}$  $(mg dry weight)^{-1}$  (Carmelo et al., 1997).

#### Results

# Genome-wide transcriptional response to acetic acid stress controlled by Haa1p

The genome-wide transcription alterations occurring in yeast cells during early response to acetic acid stress and the effect of HAA1 expression in this genomic expression program were examined in this study, based on the DNA microarray profiling of the transcriptomes of unadapted cells of S. cerevisiae BY4741 and of the  $\Delta haa1$  mutant cultivated for 30 min in the absence or presence of acetic acid (50 mM at pH 4.0). The growth curves of the yeast populations examined are shown in Figure 1. During cultivation in acetic acidsupplemented growth medium, a latency period of 2 or 5 h was observed for the parental strain and the  $\Delta haa1$  mutant, respectively, although the reinoculation of the unsupplemented growth medium with the same inoculum caused no detectable growth delay (Fig.1). During the adaptation period, the concentration of the viable cell population was maintained for the parental strain and slightly reduced for the  $\Delta$  haa1 population (Fig. 1). Following adaptation to the acetic acid concentration tested, both populations resumed exponential growth, although with a different delay (Fig.1). These observations suggest that the role of Haa1p in reducing the acetic acid-induced loss of cell viability is essentially exerted during the early adaptation period, as proposed before (Fernandes et al., 2005), therefore justifying the short exposure of 30 min to the acid, during the acute phase of stress, that was selected to perform the microarray experiments. Moreover, an acetic acid concentration that had a slight or moderate effect in the growth of the parental strain and of the  $\Delta haa1$  mutant growth was chosen to avoid a marked deleterious effect of the acid toward the very susceptible mutant population, which could interfere with the identification of the genes, directly or indirectly, regulated by Haa1p.

The concentration of acetic acid tested led to the upregulation (above 1.5-fold) of 112 genes in the parental strain *S. cerevisiae* BY4741 and to the decreased expression (below 1.5-fold) of 83 genes (Supplementaryl Tables S1 and S2 of the supplementary material). The acetic acid-responsive genes were clustered based on the biological process in which they are involved and the most significant functional classes are shown in Supplementary Figure S1 of supplementary

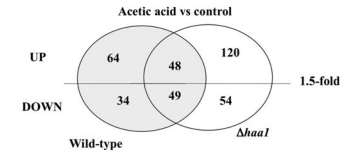


**FIG. 1.** Growth curves of *S. cerevisiae* BY4741 ( $\blacksquare$ ,  $\square$ ) and of the deletion mutant  $\Delta haa1$  ( $\blacktriangle$ ,  $\triangle$ ) in MM4 medium (pH 4.0) ( $\blacksquare$ ,  $\blacktriangle$ ) or in this same basal medium supplemented with 50 mM acetic acid (pH 4.0) ( $\square$ , $\triangle$ ). The arrows indicate the time of cultivation at which cell samples were harvested to compare their transcriptomes. Growth was followed by measuring culture OD<sub>600 nm</sub> and the concentration of viable cells was assessed as the number of colony-forming units/ mL of cell culture (CFU mL<sup>-1</sup>). The growth curves shown are means of, at least, three independent growth curves that gave rise to the same growth pattern.

material. In our dataset of acetic acid transcriptionally activated genes, the functional classes enriched in our dataset are: "Stress response," "Signal transduction," "Regulation of carbohydrate metabolism," and "Drug transport," whereas the "Amino acid catabolism" and "Ion transport" functional classes are the enriched functional classes among the downregulated genes (Fig. S1 of the supplementary material).

In the absence of acetic acid stress, the elimination of *HAA1* gene only had a slight effect on the yeast transcriptome: 11 genes exhibited an increased expression level in  $\Delta haa1$  cells and the level of 15 transcripts was decreased in this mutant strain (using 1.5-fold as the threshold cut level) (Supplementary Table S3 of the supplementary material).

Contrasting with the subtle effect registered in the transcriptome of cells cultivated in the absence of acetic acid, the expression of *HAA1* had a major impact in the alterations occurring in yeast genomic expression during early adaptive response to the acid (Fig. 2) The elimination of *HAA1* gene reduced the transcription of 85 out of the 112 acetic acidactivated genes in the parental strain, corresponding to approximately 80% of the upregulated genes that are dependent of *HAA1* expression. Remarkably, 120 genes were only found to be upregulated (above 1.5-fold) in acetic acid-challenged  $\Delta haa1$  cells but not in the parental strain (Fig. 2), presumably as a result of the higher concentration of acetic acid accumu-



**FIG. 2.** Venn diagram showing the number of genes whose transcription is altered in response to acetic acid in cells of the parental strain *S. cerevisiae* BY4741 and/or in the  $\Delta haa1$  mutant. Genes whose transcript level was increased (or decreased) above the threshold level of 1.5-fold in both strains cultivated in the presence of the acid, compared to the values registered in unstressed cells (control), were selected for this analysis.

lated inside  $\Delta haa1$  cells, compared to the parental strain (Fernandes et al., 2005). The Haa1p-regulated genes could be separated in two classes. The first class includes the 64 genes whose acetic acid-induced transcriptional activation was fully dependent on Haa1p (listed in Table 1 and clustered in Fig. 2 as "wild-type only" activated genes), these genes not being considered upregulated in  $\Delta haa1$  cells, at least above the threshold level of 1.5-fold. The other class includes the 21 genes whose acetic acid-induced transcriptional activation was only partially dependent on Haa1p (listed in Table 2 and clustered in Fig. 2 as genes activated both in "wild-type and  $\Delta haa1''$ ), the transcription of these genes being activated in  $\Delta haa1$  acetic acid-stressed cells (above 1.5-fold) but their transcript levels in the mutant strain is below the one registered in the parental strain. However, only the genes whose transcription was reduced in acetic acid-challenged  $\Delta haa1$ cells by more than 25%, compared to wild-type cells, are shown in Table 2. The full list of genes is available in Supplementary Table S4. The effect of HAA1 expression over the transcription of a few selected genes (TPO2, MSN4, HRK1, HSP30, and PDR12) was confirmed by real-time RT-PCR, thus validating the results of the microarray analysis. The more evident effect of HAA1 elimination in the transcription level of TPO2 (16-fold), MSN4 (7-fold), and HSP30 (6.8-fold) genes (Fig. 3) is also in agreement with the microarray data.

Eleven genes whose transcription was previously reported in the literature to be dependent on the presence of Haa1p-PHM8, TPO2, YRO2, TPO3, YGP1, YER130c, YIR035c, YLR297w, YPR157w, YPR127w, and SPI1 (Abbott et al., 2008; Fernandes et al., 2005; Keller et al., 2001; Simões et al., 2006)were confirmed by our microarray analysis to have a reduced expression in  $\Delta haa1$  acetic acid-challenged cells (Table 1). The remaining 73 genes identified in our work as being activated by Haa1p in response to acetic acid are novel, direct or indirect, targets of this transcription factor. The Haa1p-dependent genes encode proteins belonging to the following functional classes: "Transcription factors," "Multidrug resistance transporters," "Cell wall," "Lipid metabolism," "Regulation of carbohydrate metabolism," "Protein folding," "Carbohydrate metabolism," "Amino acid metabolism," "Nucleic acid processing," and "Uncharacterized biological function" (Fig. 4 and Tables 1 and 2). This last class comprises approximately

	OF 5. CEREVISIAE B	Y4/41 IS ABOLISE	OF 5. CEREVISIAE BY4/41 IS ABOLISHED IN THE ABSENCE OF HAAI EXPRESSION	OF HAA1 EXPRESSIC	N				
		$(mRNA \ wt \ )_{ac}$	(mRNA Δhaa1 ) <sub>ac</sub>	(mRNA wt) <sub>ac</sub>			Induced by	ł by	
Gene or ORF	Biological function	$(mRNA \ wt)_c$	(mRNA Ahaa1) <sub>c</sub>	(mRNA Ahaa1) <sub>ac</sub>	Acetic acia susceptibility	Actld.	Prop.	Sorb.	Lact.
TPO2	Plasma membrane polyamine transporter	14.9	1.3	18.7	+				
70CG IV	of the major facilitator superfamily	c 11	c 7	10 1					
I LN29/W		C.11	7.1	C.UI	-				
SIP3	Zinc-tinger protein of unknown function	7.0	-1.4	8.1 2.1	+				
Y K U Z	Putative protein of unknown function	11.1	1.4	8.7	I				
YAR029w	Unknown function	3.7	-1.1	7.6	Ι				
TOS3	Protein kinase required for phosphorylation and	5.7	1.1	6.2	+				
	activation of Snf1p; glucose metabolism								
$\gamma IR035c$	Unknown function	5.5	1.2	6.2	Ι				
YGP1	Cell wall-related secretory glycoprotein	3.0	1.1	4.8	+				
PCL10	Cyclin involved in regulation	5.0	1.1	4.3	+				
	ot activity of protein kinase Pho85p								
YPR127w	Unknown function	3.3	1.0	4.1	Ι				
DSD1	D-serine dehydratase	3.2	-1.1	4.0	I				
MSN4	Transcription factor activated	6.1	1.4	3.8	Ι				
	in stress conditions								
$\gamma IR096 w$	Putative xvlose and	2.7	-1.1	3.7	Ι				
	arabinose reductase								
CD11	CDI anchored for the sector of	ц -	V L	и r					
JF11	ULI-AUCIOLEU CEIL WAIL PLOTEUL,	1.0	1.1	0.0	I				
			Ţ	Ċ					
HOK2	Giycerol phosphatase involved in alveral hiosvnthesis:	4.8	-1.4	3.4	+				
	induced in response to stress								
$\gamma KR075c$	Unknown function	2.9	,	с. С	I				
CI IR 2	Subinganing (A-bridrowrilsed	ע ייי יי	1.1	0.0	+ +				
7000	optunganue C+11yuoxyase involved in sphingolipid	0.0	7:1	0.0	+ +				
	metabolism								
ICY1	Unknown function	2.7	1.0	3.0	+				
INNI	Inositol monophosphatase	3.4	1.3	2.6	+				
	involved in biosynthesis								
	of inositol								
SAP30	Subunit of a histone	3.2	1.3	2.6	+++++				
	deacetylase complex								
$\gamma NL200c$	Unknown function	2	-1.2	2.4	++				
STF2	Protein involved	2.0	-1.3	2.2	I				
	in the regulation of F <sub>1</sub> F <sub>0</sub> ATPase								
SYC1	Subunit of the APT	2.9	1.1	2.2	+				
	subcomplex of cleavage and								
	рогуацепутации гастог								

(continued)

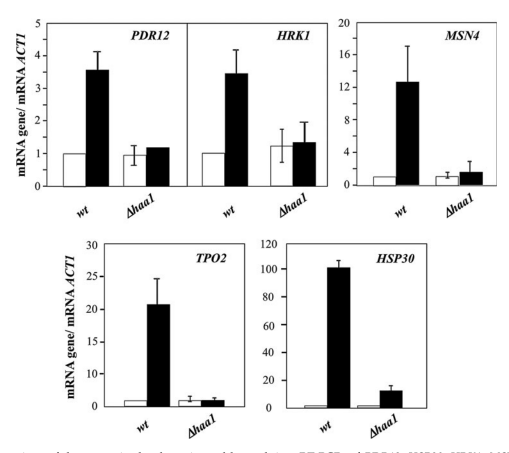
Table 1. List of Genes Whose Acetic Acid-Induced Transcriptional Activation Registered in Cells of *S. cerevisiae* BY4741 Is Abolished in the Absence of *HA41* Expression

RF         Biological function $(mRNA \text{ art })_{ac}$ $(mRNA \text{ art })_{ac}$ $(mRNA \text{ art })_{ac}$ $Aeth_{ac}$			LABLE 1.	IABLE 1. (CUNTINUED)						
ORF         Biological function         (nRVA A what), (nRVA A heat),			(mRNA wt ) <sub>ac</sub>	(mRNA Δhaa1) <sub>ac</sub>	$(mRNA \ wt \ )_{ac}$	E V		Induced by	t by	
600         Unknown function         2.7         1.3         2.2           600         Unknown function         2.0         1.0         2.1           600         Unknown function         2.2         1.3         2.1           610         Unknown function         2.2         1.3         2.1           610         Unknown function         2.2         1.3         2.1           610         Oprotein that interacts with the receptor Grapity, preventably involved         2.2         1.1         2.0           6         Oprotein that interacts with the receptor disorgenase         1.9         1.0         2.0           7.0         Unknown function         2.3         1.4         1.9           7.1         Unknown function         2.3         1.3         1.8           7.1         Unknown function         2.3         1.4         1.7           7.1         Unknown function         2.3         1.4         1.7           7.1	Gene or ORF	Biological function	$(mRNA wt)_c$	(mRNA Ahaa1) <sub>c</sub>	(mRNA Ahaa1) <sub>ac</sub>	Acetic acid susceptibility	Actld.	Prop.	Sorb.	Lact.
%       Unknown function       4.0       1.1       2.1         %       Unknown function       2.2       1.1       2.2         %       Unknown function       2.2       1.1       2.2         %       Unknown function       2.2       1.1       2.2         %       Submut of the heterotimenc       2.2       1.1       2.1         %       Submut of the heterotimenc       2.2       1.1       2.0         %       Submut of the heterotimenc       2.2       1.1       2.0         %       Submut of the heterotimenc       2.2       1.1       2.0         %       Inpected/orbinal represention with the receptor Graph presumably in whether       1.7       1.1       2.0         %       Involved in methonine salvage pathway       2.3       1.4       1.9       2.0         %       Unknown function       2.3       1.4       1.9       2.0 <td>YLR326w</td> <td>Unknown function</td> <td>2.7</td> <td>1.3</td> <td>2.2</td> <td>I</td> <td></td> <td></td> <td></td> <td></td>	YLR326w	Unknown function	2.7	1.3	2.2	I				
4.       Unknown function       2       10       21       23       21         4.0.4.       C protein that interacts with the receptor Grp1 presumably involved       22       11       2       2         6       Divension function       2       11       2	YAR028w	Unknown function	4.0	1.1	2.1	Ι				
44r.a       Unknown function       22       13       2         6       protein that interacts with the receptor Graphy presumably involved       22       11       2         7       G protein that interacts with the receptor Graphy presumably involved       17       11       20         7       Low afinity cycle AMP phosphodiesterase       17       11       20         7       Low afinity cycle AMP phosphodiesterase       17       11       20         7       Low afinity cycle AMP phosphodiesterase       17       11       20         7       Unknown function       23       14       19         7       Unknown function       23       13       18         1       Unknown function       23       13       18         1       Unknown function       23       13       13         1       Unknown function       23       13       13         1       Unknown function       23       13       13       13         1       Unknown function       23       13       13       13         1       Unknown function       23       13       14       15         1       Unknown function       23       13	$\gamma NL024c$	Unknown function	2	1.0	2.1	Ι				
Subunit of the heterotrimetic       22       1.1       2         G protein that interacts with the ceptor Grp1p: presundly involved       1.7       1.1       2.0         G protein that interacts with the ceptor Grp1p: presundly involved       1.7       1.1       2.0         Taw undrived in methionine salvage pathway       2.3       1.4       1.9         Zu       Undrown function       2.3       1.4       1.9         Zu       Undrown function       2.3       1.4       1.9         Zu       Undrown function       2.3       1.4       1.9         Undrown function       2.3       1.3       1.8       1.7         Unknown function       2.0       2.0       1.4       1.7         Unknown function       2.1       2.0       1.7       1.7         Unknown function       2.3       1.4       1.7       1.7         Unknown function       2.0       2.0       1.4       1.7         Unknown function       2.0       2.	YNR034w-a	Unknown function	2.2	1.3	2	Ι				
C protein that interacts with the ceptor Grply: presumably involved in pseudohybal growth       17       11       20         Two stringty cyclic AMP phosphodiesterase       17       11       20         Acireductor diorgenase       19       10       20         Not of mittity cyclic AMP phosphodiesterase       17       11       20         Acireductor diorgenase       19       10       20         Notowed in methionine salvage pathway       23       14       19         Unknown function       23       13       18       13         Le Unknown function       23       13       18       17         Unknown function       23       13       13       18         Distromation function       23       13       13       13         Unknown function       23       13       13       13         Distromation function       23       13       14       17         O collaraced and sorbate anions       23       13       13       13         Distromation function       23       12       17       17         O colutration factor involved       23       13       14       15         O acyltranscrase       22       10       14 <td>GPG1</td> <td>Subunit of the heterotrimeric</td> <td>2.2</td> <td>1.1</td> <td>2</td> <td>I</td> <td></td> <td></td> <td></td> <td></td>	GPG1	Subunit of the heterotrimeric	2.2	1.1	2	I				
70       In preceduably involved         70       In preceduation growth         70       In preceduation growth         70       Involved in methionine salvage pathway         70       Unknown function         70       Unknown function         71       11         72       Unknown function         74       Unknown function         75       Unknown function         76       Unknown function         77       Unknown function         78       Inancriptional repression         78       Unknown function         78       Unknown function         78       Unknown function         79       Unknown function         70       Unknown function         71       12         72       13         73       13         74       13         75       Unknown function         76       Polosphofrance         77       Unknown function         78       Unknown function         79       Unknown function         70       Unknown function         71       13         72       13		G protein that interacts with the								
7u       In periodicyptai gowun         7u       In whithity cyclic AMP phosphodiesterase       17       11       20         7u       Involved in methionine salvage pathway       23       14       19       20         7u       Unknown function       23       14       19       20         1c       Unknown function       23       13       18       19         1c       Unknown function       23       13       18       19         1c       Unknown function       23       13       18       17       13       18         1c       Unknown function       23       13       13       18       17       17         1n       Plasma membrane ABC transporter involved       42       13       13       18       17         1nknown function       21       12       13       13       17       17         1nknown function       Unknown function       23       14       17       17         1nknown function       Unknown function       23       11       17       17       17         1nknown function       Inknown function       23       10       16       11       17         1nknown fu		receptor Grp1p; presumably involved								
7a       Low atmitty cyclic AMP phosphodiesterase       1.7       1.1       2.0         7a       Low atmitty cyclic AMP phosphodiesterase       1.9       1.0       2.0         7a       Unknown function       salvage pathway       2.3       1.4       1.9       2.0         1       Transcriptional repression involved in response       2.2       1.4       1.9       1.9         1       Unknown function       2.3       1.3       1.8       1.9       1.9       1.9         1       Unknown function       2.3       1.3       1.3       1.8       1.9       1.7       1.9       1.7       1.9       1.7       1.1       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7		in pseudohyphal growth	ļ							
7u       Unknown function       1.9       1.0       20         7u       Unknown function       aslvage pathway       2.3       1.4       1.9         1       Transcriptional repressor involved in response       2.3       1.4       1.9       1.9         1       Transcriptional repressor involved in response       2.3       1.4       1.9       1.9         1       Unknown function       2.3       1.3       1.8       1.3       1.8         1       Unknown function       2.1       1.2       1.3       1.8       1.9       1.7         1       Unknown function       2.1       1.2       1.3       1.8       1.7         1       Disknown function       2.1       1.2       1.3       1.8       1.7         1       Disknown function       2.1       1.2       1.1       1.7       1.7         1       Disknown function       2.2       1.1       1.1       1.7       1.7       1.7         1       Disknown function       2.2       1.1       1.1       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7       1.7	PDET	Low atinitity cyclic AMP phosphodiesterase	1.7	1.1	2.0	I				
7umicroved in methonine salvage patiway7uUnknown function1cUnknown function1cUnknown function1cUnknown function1cUnknown function1cUnknown function1cUnknown function1dUnknown function </td <td>ADII</td> <td>Actreductone dioxigenase</td> <td>I.9</td> <td>1.0</td> <td>2.0</td> <td>I</td> <td></td> <td></td> <td></td> <td></td>	ADII	Actreductone dioxigenase	I.9	1.0	2.0	I				
70       Unknown function       2.3       1.4       1.9         17       Transcriptional repressor involved in response       2.2       1.4       1.9         16       Unknown function       2.3       1.3       1.8       1.9         17       Unknown function       2.3       1.3       1.8       1.7       1.8       1.7       1.8       1.7       1.8       1.7       1.9       1.7       1.9       1.7       1.6       1.6       1.6       1.6       1.6 <td< td=""><td></td><td>involved in methionine salvage pathway</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>		involved in methionine salvage pathway								
Transcriptional repressor involved in response       2.2       1.4       1.9         1       Unknown function       2.3       1.3       1.8         1       Unknown function       2.3       1.3       1.8         1       Unknown function       2.1       1.2       1.8         1       Unknown function       2.1       1.2       1.8         1       Unknown function       2.1       1.2       1.8         1       Influx of benzoate and sorbate autions       2.8       1.4       1.7         1       Unknown function       2.8       1.4       1.7         1       Unknown function       2.8       1.4       1.7         1       Unknown function       2.0       1.2       1.7         1       Unknown function       2.0       1.2       1.7         1       Unknown function       2.1       1.0       1.6       1.7         1       O-acyltransferase, presumably       1.9       1.4       1.6       1.7         1       Metersorien of C2/M phase genes       2.2       1.0       1.6       1.6         1       Metersoriense of C2/M phase genes       2.2       1.0       1.6       1.6	YNL217w	Unknown function	2.3	1.4	1.9	I				
1c       Unknown function       2.3       1.3       1.8         1c       Unknown function       2.1       1.2       1.8         1c       Unknown function       2.1       1.2       1.8         1c       Unknown function       2.1       1.2       1.8         1c       Inknown function       2.1       1.2       1.8         1c       Unknown function       2.8       1.4       1.7         1c       Unknown function       2.0       1.2       1.8         1c       Unknown function       2.0       1.2       1.7         1c       Transcription factor involved       2.1       1.0       1.7         1c       Transcription factor involved       2.2       1.0       1.7         1c       Transcription factor involved       2.2       1.0       1.6         1c       Oracyltransferase; presumably       0.       0.44       1.6       1.6         0       Oracyltransferase; presumably	NRG1	Transcriptional repressor involved in response	2.2	1.4	1.9	++				
1       Unknown function       2.3       1.3       1.8         1       Unknown function       2.1       1.2       1.8         1       Instam membrane ABC transporter involved       4.2       1.3       1.8         1       In efflux of beracate and sorbate anions       2.8       1.4       1.7         1       In efflux of beracate and sorbate anions       2.8       1.4       1.7         1       Unknown function       2.0       1.2       1.8         1       Unknown function       2.0       1.2       1.7         1       Unknown functor 2-kinase       2.1       1.0       1.7         1       Transcription factor involved       2.2       1.0       1.7         1       Transcription factor involved       2.2       1.0       1.6         1       O-acyltransferase; presumably       1.0       1.6       1.6         1       Unknown function       1.4       1.6       1.4       1.6         1<		to alkaline pH and glucose catabolic repression								
Unknown function       2.1       1.2       1.8         Plasma membrane ABC transporter involved       4.2       1.3       1.8         In efflux of benzoate and sorbate anions       2.8       1.4       1.7         Unknown function       2.8       1.4       1.7         Unknown function       2.0       1.2       1.8         Unknown function       2.0       1.2       1.7         Unknown function       2.0       1.2       1.7         Glutaredoxin enzyme       2.0       1.2       1.7         Glutaredoxin enzyme       2.1       1.0       1.7         Glutaredoxin enzyme       2.1       1.0       1.7         Glutaredoxin enzyme       2.1       1.0       1.7         Acyl-coenzymeA: ethanol       2.2       1.0       1.6         O-acyltansferase; presumably       1.9       1.4       1.6         O-acyltansferase; presumably       1.9       1.4       1.6         Unknown function       1.9       1.4       1.6         O-acyltansferase; presumably       1.9       1.4       1.6         Unknown function       1.9       1.4       1.6         Unknown function       0       1.9       1.4	$\gamma PL071c$	Unknown function	2.3	1.3	1.8	Ι				
Plasma membrane ABC transporter involved       4.2       1.3       1.8         in efflux of benzoate and sorbate anions       2.8       1.4       1.7         Unknown function       2.0       1.2       1.7         Glutaredoxin enzyme       2.0       1.2       1.7         Glutaredoxin enzyme       2.0       1.2       1.7         Glutaredoxin enzyme       2.0       1.2       1.7         Graphofructo-2-kinase       2.1       -1.4       1.7         Transcription factor involved       2.2       1.0       1.7         Acyl-coenzymeA: ethanol       2.2       1.0       1.6         Acyl-coenzymeA: ethanol       2.2       1.0       1.6         O-acyltransferase; presumably       1.9       1.4       1.6         Novolved in lipid metabolism       1.9       1.4       1.6         Nuknown function       1.9       1.4       1.6         Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         confored uctase complex which       conforeductase complex which       1.6       -1.1       1.5         Component of autophagosomes       1.6       -1.1       1.5       1.6       -1.1       1.5         nin intracellular traffic	MUKI	Unknown function	2.1	1.2	1.8	I				
n filux of benzoate and sorbate anions       2.8       1.4       1.7         Unknown function       2.0       1.2       1.7         Clutaredoxin enzyme       2.0       1.2       1.7         6-phosphofructo-2-kinase       2.1       -1.4       1.7         Transcription factor involved       2.1       -1.4       1.7         in the expression of G2/M phase genes       2.2       1.0       1.6         Acyl-coenzymes/thanol       2.2       1.0       1.6         Acyl-coenzymes/thanol       2.2       1.0       1.6         Acyl-coenzymes/thanol       2.2       1.0       1.6         Acyl-coenzymes/thanol       1.9       1.4       1.6         Acyl-coenzymes/thanol       1.9       1.4       1.6         Acyl-coenzymes/thanol       1.9       1.4       1.6         Acyl-coenzymes/thanol       1.9       1.4       1.6         O-acyltransferase; presumably       1.9       1.4       1.6         involved in lipid metabolism       1.9       1.4       1.6         o-acyltransferase; presumably       1.9       1.4       1.6         o-acyltransferase; producturelocism       1.6       1.4       1.6         o-acyltransferase;	PDR12	Plasma membrane ABC transporter involved	4.2	1.3	1.8	I				
0       Unknown function       2.8       1.4       1.7         6       Phosphofructo-2-kinase       2.0       1.2       1.7         6       Phosphofructo-2-kinase       2.1       -1.4       1.7         6       Fanscription factor involved       2.1       -1.4       1.7         6       Transcription factor involved       2.1       -1.4       1.7         7       Transcription factor involved       2.1       1.0       1.7         7       Transcription factor involved       2.2       1.0       1.7         7       Acyl-coenzymeA:ethanol       2.2       1.0       1.6         7       O-acyltransferase; presumably       1.9       1.4       1.6         6       Putative protein of       1.9       1.4       1.6         9       Subunit of the ubiquol-cytochrome       1.6       -1.1       1.5         6       Untative protein of       1.6       -1.1       1.5         7       0       Subunit of the ubiquol-cytochrome       1.6       -1.1       1.5         8       Component of autophagosomes       1.6       -1.1       1.5       1.6         9       Component of autophagosomes       1.6       -1.1		in efflux of benzoate and sorbate anions								
Glutaredoxin enzyme       2.0       1.2       1.7         6-phosphofructo-2-kinase       2.1       -1.4       1.7         Transcription factor involved       2.1       -1.4       1.7         in the expression of G2/M phase genes       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         O-acyltransferase; presumably       2.2       1.0       1.6         Novolved in lipid metabolism       1.9       1.4       1.6         Mutown function       1.9       1.4       1.6         unknown function       1.6       -1.1       1.5         Subunit of the ubiquol-cytochrome       1.6       -1.1       1.5         c oxidoreductase complex which       c oxidoreductase complex which       1.6       -1.1       1.5         comprises part of the mitochondrial       respiratory chain       1.6       -1.1       1.5         and Cvt vesicles; involved       in intracellular trafficking       1.6       -1.1       1.5	TMA10	Unknown function	2.8	1.4	1.7	I				
6-phosphofructo-2-kinase       2.1       -1.4       1.7         Transcription factor involved       1.0       1.7         in the expression of G2/M phase genes       2.2       1.0       1.7         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       0       2.1       1.4       1.6         Acyl-coenzymeA:ethanol       1.9       1.4       1.6       1.6         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         0       Subunit of the mbiqunol-cytochrome       1.6       -1.1       1.5 <td>GRX8</td> <td>Glutaredoxin enzyme</td> <td>2.0</td> <td>1.2</td> <td>1.7</td> <td>I</td> <td></td> <td></td> <td></td> <td></td>	GRX8	Glutaredoxin enzyme	2.0	1.2	1.7	I				
Transcription factor involved in the expression of G2/M phase genes Acyl-coenzymeA:ethanol       2       1.0       1.7         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       2.2       1.0       1.6         Acyl-coenzymeA:ethanol       1.9       1.4       1.6         O-acyltransferase; presumably involved in lipid metabolism       1.9       1.4       1.6         0       Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       Subunit of the ubidumol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       Component of autophagosomes and Cyt vesicles; involved in intracellular trafficking       1.6       -1.1       1.5	PFK27	6-phosphofructo-Ź-kinase	2.1	-1.4	1.7	Ι				
in the expression of G2/M phase genes Acyl-coenzymeA:ethanol O-acyltransferase; presumably involved in lipid metabolism (6c Putative protein of unknown function 0 Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain (7) Component of autophagosomes and Cyt vesicles; involved in intracellular trafficking	FKH2	Transcription factor involved	7	1.0	1.7	I				
Acyl-coenzymeA:ethanol       2.2       1.0       1.6         O-acyltransferase; presumably       0 acyltransferase; presumably       1.9       1.4       1.6         0 acyltransferase; presumably       1.9       1.4       1.6       1.6         0 butative protein of unknown function       1.9       1.4       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5       1.6       1.1       1.5         0       Component of autophagosomes       1.6       -1.1       1.5       1.6       1.1       1.5         0       Intracel		in the expression of G2/M phase genes								
Ö-acyltransferase; presumably involved in lipid metabolism1.91.41.66cPutative protein of unknown function1.91.41.60Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain1.6-1.11.50Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain1.6-1.11.50Subunit of the ubidunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain1.6-1.11.50Subunit of autophagosomes and Cyt vesicles; involved in intracellular trafficking1.6-1.11.5	EEB1	Acyl-coenzymeA:ethanol	2.2	1.0	1.6	Ι				
<i>l</i> ( <i>c</i> Putative protein of unknown function       1.9       1.4       1.6         0       Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome c oxidoreductase complex which comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       Subunit of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       In intracellular trafficking       1.6       -1.1       1.5		Ó-acyltransferase; presumably								
6c       Putative protein of unknown function       1.9       1.4       1.6         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         0       comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         0       component of autophagosomes       1.6       -1.1       1.5         and Cvt vesicles; involved       in intracellular trafficking       1.6       -1.1       1.5		involved in lipid metabolism								
0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         c oxidoreductase complex which       comprises part of the mitochondrial       1.6       -1.1       1.5         comprises part of the mitochondrial       respiratory chain       1.6       -1.1       1.5         component of autophagosomes       1.6       -1.1       1.5         and Cvt vesicles; involved       in intracellular trafficking       1.6       -1.1       1.5	YLR346c	Putative protein of	1.9	1.4	1.6	Ι				
0       Subunit of the ubiqunol-cytochrome       1.6       -1.1       1.5         c oxidoreductase complex which comprises part of the mitochondrial respiratory chain       1.6       -1.1       1.5         component of autophagosomes       1.6       -1.1       1.5         and Cvt vesicles; involved in intracellular trafficking       1.6       -1.1       1.5		unknown function								
c oxidoreductase complex which comprises part of the mitochondrial respiratory chain Component of autophagosomes 1.6 –1.1 1.5 and Cvt vesicles; involved in intracellular trafficking	QCR10	Subunit of the ubiqunol-cytochrome	1.6	-1.1	1.5	I				
comprises part of the mitochondrial respiratory chain Component of autophagosomes 1.6 –1.1 1.5 and Cvt vesicles; involved in intracellular trafficking		c oxidoreductase complex which								
respiratory chain Component of autophagosomes 1.6 –1.1 1.5 and Cvt vesicles; involved in intracellular trafficking		comprises part of the mitochondrial								
Component of autophagosomes 1.6 –1.1 1.5 and Cvt vesicles; involved in intracellular trafficking		respiratory chain								
and Cvt vesicles; involved in intracellular trafficking	ATG8	Component of autophagosomes	1.6	-1.1	1.5	I				
in intracellular trafficking		and Cvt vesicles; involved								
		in intracellular trafficking								
Unknown function 1.7 1.4	YER188w	Unknown function	1.7	1.4	1.5	I				
	This table wa	This table was prepared based on the comparative analysis of the transcriptome of cells of the parental strain BY4741 (wt) and of the mutant BY4741 Ahaa1 (Ahaa1) cultivated in MM4 medium (pH 4.0)	tome of cells of the p	arental strain BY4741 (w	/t) and of the mutant BV	(4741 Ahaal (Ahaa	1) ailtimetod	/////: E		,

supplemented with 50 mM acetic acid. The magnitude of the transcriptional activation registered in response to acetic acid stress in the parental strain (wt<sub>ac</sub>/wt<sub>c</sub>) is shown as well as the susceptibility toward acetic acid stress in the parental strain,  $(m_{tac}/wt_c)$  is shown as well as the susceptibility toward acetic acid stress in the parental strain, (++,+) acetic acid of Haa1p-target genes (-susceptibility identical to the parental strain, (++,+) is also indicated of mutants devoid of Haa1p-target genes (-susceptibility identical to the parental strain, (++,+) is also indicated (shaded in gray) based on results reported from Aranda and del Olmo (2004). Mira et al. (2009), Schuller et al. (2004), and Abbot et al. (2008), respectively. Genes are sorted based on the ratio of mRNA gene level between the wild-type and the  $\Delta lna1$  mutant.

		$(mRNA wt)_{ac}$	(mRNA Δhaa1 ) <sub>ac</sub>	$(mRNA \ wt \ )_{ac}$	F::: V		Induced by	ed by	
Gene or ORF	Biological function	$(mRNA wt)_c$	$(mRNA \ \Delta haa1)_c$	(mRNA Ahaa1) <sub>ac</sub>	Aceric acta susceptibility	Actld.	Prop.	Sorb.	Lact.
$\gamma PR157w$	Unknown function	32.6	2.2	19.2	Ι				
STP4	Putative transcription factor	12.6	2.2	12.7	Ι				
YDR222w	Unknown function	10.9	1.6	9.5	Ι				
YPL014w	Unknown function	10.3	1.8	7.3	Ι				
PHM8	Unknown function; expression	15.6	2.8	6.3	Ι				
	regulated by phosphate availability								
YER130c	Unknown function	10.9	2.4	4.7	Ι				
TPO3	Plasma membrane polyamine	4.8	1.8	3.7	++				
	transporter of the major								
	facilitator superfamily								
HSP30	Putative negative regulator	86.9	54.1	3.5	Ι				
	of plasma membrane H <sup>+</sup> -ATPase								
	activity; expression								
	induced under stress								
YPC1	Alkaline ceramidase involved in	9.3	2.6	3.4	+				
	sphingolipid and ceramide metabolism								
SSE2	Heat-shock protein involved in protein	6.8	3.1	2.9	+				
	folding; expression induced under stress								
SCM4	Putative regulator of Cdc4p activity required	3.8	2.1	2.7	I				
	for G1/S phase transition								
HSP26	Heat-shock protein involved in	2.7	1.9	2.5	I				
	protein folding; expression								
	induced under stress								
YGR250c	Unknown function	6.5	3.5	2.3	I				
SKS1	Putative serine/threonine protein kinase	4.4	2.3	2.3	I				
	involved in adaptation to glucose starvation								
MTH1	Negative regulator of the glucose-sensing	3.1	1.7	2.1	I				
	signal transduction pathway								
FMP48	Putative mithocondrial protein;	28.2	16.1	2	++				
	unknown function								
RGS2	Negative regulator of glucose-induced	5.2	2.9	2	Ι				
	cAMP signaling pathway								
ECM13	Unknown function	68	3.0	<i>د</i>	Ι				
HRK1	Protein kinase imulicated in activation	5.0	8. <del>-</del>	10	+++				
	of the plasma membrane H <sup>+</sup> . ATPase		2		-				
	in records to alloce metabolism								
10110	Interpoted to glacose includuality	0 C	с С	7					
CUNI		0.4	C.1	1.7	I				
PDR16	Phosphatidylinositol transfer protein	2.3	1.6	1.6	Ι				

TABLE 2. LIST OF GENES WHOSE ACETIC ACID-INDUCED TRANSCRIPTIONAL ACTIVATION REGISTERED



**FIG. 3.** Comparison of the transcript levels, estimated by real-time RT-PCR, of *PDR12*, *HSP30*, *HRK1*, *MSN4*, and *TPO2* genes produced in cells of the parental strain BY4741 (*wt*) or the  $\Delta haa1$  mutant during cultivation for 30 min in MM4 growth medium (at pH 4.0) (white bars) or in this same growth medium supplemented with 50 mM acetic acid (dark bars). *ACT1* mRNA was used as internal control. For each gene the value registered in parental strain control cells (corresponding to cells cultivated in unsupplemented growth medium) were set as 1 and the induction ratios are relative to this control. mRNA values shown are representative of results obtained, at least, in two independent incubation experiments.

40% of the dataset. The functional classes considered enriched in the dataset (*p*-values below 0.01) are "Multidrug resistance transporters," "Carbohydrate metabolism," and "Regulation of carbohydrate metabolism" (Fig. 4).

Using the YEASTRACT database (Teixeira et al., 2006b), the genes transcriptionally activated in response to acetic acid were clustered based on documented regulatory associations with the transcription factors Rim101p, War1p, Msn2p/ Msn4p, and Haa1p that control the so far described regulatory pathways involved in yeast response to weak acid stress (Fernandes et al., 2005; Mira et al., 2009; Schüller et al., 2004). This analysis indicates that Haa1p specifically regulates almost one-half of the acetic acid-induced genes (51 out of 112 genes), whereas the number of genes specifically dependent on Msn2p/Msn4p, Rim101p, or War1p is much lower (Fig. 5). Remarkably, the sole War1p-target gene, PDR12, was also identified among the genes activated by Haa1p in response to acetic acid stress. An overlap between the Haa1p and the Msn2p/Msn4p regulons was also registered 23 acetic acidactivated genes being coregulated by these two regulatory pathways (Fig. 5). Three genes (NRG1, YKR075c, WSC4) are common to both the Haa1p- and Rim101p-dependent regulons (Fig. 5) and six genes (MSN4, YPL014w, JID1, HSP26, MDH2, and HSP30) are shared by the Rim101p, Msn2p/ Msn4p, and Haa1p-regulatory pathways (Fig. 5). The 13 remaining acetic acid-responsive genes have no documented regulatory associations with any of the above referred transcription factors and were clustered as "Others" (Fig. 5). Most of them are documented targets of Yap1p and Hsf1p transcription factors, also involved in yeast response to stress (Hahn et al., 2004; Rodrigues-Pousada et al., 2004).

# Involvement of genes of the Haa1p regulon in yeast tolerance to acetic acid

The susceptibility phenotype of  $\Delta haa1$  cells toward acetic acid is, presumably, due to the lack of activation of Haa1p target genes required for yeast protection against the acid. Among the group of genes whose transcription was initially described by Keller et al. (2001) to be dependent on a functional Haa1p only *TPO2*, *TPO3*, and *YGP1* genes provided protection against acetic acid. However, their protective effect was much less evident than the one exerted by Haa1p (Fernandes et al., 2005), which prompted us to examine the role in yeast tolerance to acetic acid of the novel Haa1p-targets uncovered in the present work. The expression of 20% of these genes, although at different levels, contributed to a reduced duration of latency period observed under acetic acid stress (Fig. 6 and Tables 1 and 2). In the absence of acetic acid, the elimination of these genes had no detectable in the growth

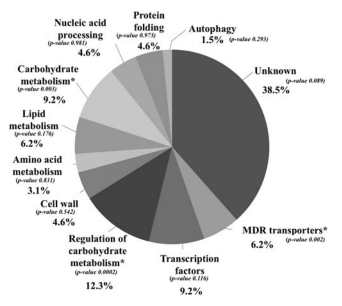


FIG. 4. Clustering, based on biological function, of genes activated in response to acetic acid stress in a Haa1p-dependent way. Genes whose transcript level in yeast cells challenged with acetic acid was reduced by more than 25% in the  $\Delta haa1$  mutant, compared to the levels attained in the parental strain, were grouped according to their biological functions described on MIPS functional catalog. The relative percentage of each functional class was calculated taking into consideration the number of genes in each class compared to the total of Haa1p-regulated genes (85) considered and the *p*-values of each functional class. Enriched classes (*p*-value below 0.01) are marked with\*.

curve (results not shown). The genes of the Haa1p-regulon conferring protection toward acetic acid include: HOR2 and SSE2, two genes of the environmental stress response (Gasch et al., 2000) that encode a glycerol-3-phosphatase involved in glycerol biosynthesis and a heat shock protein, respectively; NRG1, HRK1, TOS3, and PCL10, whose function is related with the regulation of carbohydrate metabolism; and SYC1 and SAP30, which encode two proteins involved in nucleic acid processing. The other genes that are determinants of resistance to acetic acid and that are activated in response to acetic acid stress in the dependence of HAA1 expression encode proteins involved in lipid metabolism. This is the case of YPC1 and SUR2, encoding two enzymes that catabolyze ceramide to the C26-sphingolipid phytosphingosine and INM1 whose product is a monophosphatase involved in inositol biosynthesis.

The elimination of *HRK1* and, to a lower extent, of *SAP30* gene, led to the strongest susceptibility phenotypes to acetic acid (Fig. 6). *HRK1* encodes a protein kinase involved in the activation of plasma membrane H<sup>+</sup>-ATPase activity in response to glucose metabolism (de la Fuente et al., 1997; Goossens et al., 2000) and *SAP30* encodes a component of the Rpd3L histone deacetylase complex (Carrozza et al., 2005). *SAP30* was identified before as a determinant of resistance to propionic acid (Mira et al., 2009) and to a variety of other chemicals (Hillenmeyer et al., 2008). Interestingly, the elimination of *SDS3*, *SIN3*, and *DEP1*, encoding, respectively, two subunits of the Rpd3L complex and a protein associated to this complex (Carrozza et al., 2005), also led to

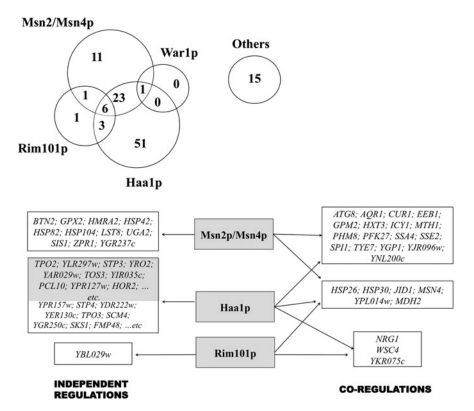
an increase in yeast susceptibility to acetic acid (Supplementary Fig. S2).

# The role of Hrk1p in providing protection against acetic acid involves the reduction of intracellular acetate concentration

Hrk1p was proposed to be a positive regulator of plasma membrane (PM) H<sup>+</sup>-ATPase (Pma1p) activity in response to glucose metabolism, although less important than its functional homologue, the Ptk2p kinase (de la Fuente et al., 1997; Goossens et al., 2000). Taking into account the essential role played by PM-H<sup>+</sup>-ATPase activity in yeast response and resistance to weak acid stress (Holyoak et al., 1997; Viegas et al., 1998), the putative effect of Hrk1p in acetic acid-induced activation of Pma1p was hypothesized. Considering that the result of both the in vivo activity of this H<sup>+</sup>-pump and the passive  $H^+$  influx through the plasma membrane can be roughly assessed based on the acidification rate of the external environment induced by glucose (Portillo and Serrano, 1989), the acidification curves promoted by cells of the parental strain and of the  $\Delta hrk1$  mutant following addition of glucose, either in the presence or absence of acetic acid, were compared (Fig. 7A). In the absence of acetic acid, glucose supplementation triggered the activation of the PM-H<sup>+</sup>-ATPase and the rapid extrusion of H<sup>+</sup> to the external environment was registered (Fig. 7A). Compared with the parental strain, the kinetics of growth medium acidification was inhibited in the  $\Delta hrk1$  mutant, although the effect is mild, probably as the result of Ptk2p activity (Goossens et al., 2000) (Fig. 7A). The presence of increasing concentrations of acetic acid led to the reduction of the rate of extracellular acidification promoted by yeast cells, consistent with the nonspecific permeabilization of plasma membrane induced by the acid and the consequent increase of the H<sup>+</sup>-passive influx (Fig. 7A). Under acetic acid stress, the expression of *HRK1* had little or no detectable effect in the active efflux of protons (Fig. 7A, and other results not shown) indicating that it is unlikely that the role of this kinase in yeast tolerance to acetic acid is exerted through Pma1p activation. Because Hrk1p belongs to a family of protein kinases dedicated to the regulation of plasma membrane transporters (Goossens et al., 2000), it was hypothesized that Hrk1p phosphorylation targets might include putative transporters involved in the reduction of acetate concentration into acetic acid-challenged cells. This mechanism was considered because the concentration of acetate is higher in  $\Delta haa1$  cells (Fernandes et al., 2005), and this effect could not be fully attributed to other candidate targets of Haa1p, specifically to the plasma membrane drug-H<sup>+</sup> antiporters Tpo2p and Tpo3p (Fernandes et al., 2005). Consistent with the hypothesized mechanism, the accumulation of labeled acetic acid in  $\Delta hrk1$  cells attained higher levels than those registered in the parental strain (Fig. 7B).

#### Discussion

The alterations occurring in yeast genomic expression during early response to acetic acid and the involvement of the transcription factor Haa1p in this transcriptional reprogramming is described here for the first time. Results indicate that Haa1p regulates, directly or indirectly, the transcription of approximately 80% of the acetic acid-responsive genes (85



**FIG. 5.** Venn diagram illustrating the crosstalk between the regulatory systems controlled by the weak acid-responsive transcription factors Haa1p, Rim101p, Msn2/Msn4p, and War1p in yeast response to acetic acid stress. The 112 genes found to be upregulated in response to acetic acid (50 mM) in the parental strain were clustered among the referred regulons, based on the existence of documented regulatory associations with the indicated transcription factors, according with the information available in the YEASTRACT database (www.yeastract.com) (Teixeira et al., 2006b). The number of genes in each cluster is indicated and the full or the partial list of genes belonging to each regulon is shown in the schematic representation below the diagram, emphasizing the genes specific of each regulon and the coregulated genes. In the case of the genes specifically dependent on Haa1p, 10 genes were selected from those listed in Tables 1 (indicated in gray background) and 2 (indicated in a white background). These genes exhibit the highest reduction in transcript level in  $\Delta haa1$  cells, compared to the parental strain.

out of 112 genes). This percentage is much higher than the percentage of acetic acid-responsive genes that is regulated by the other weak acid-responsive regulatory systems described so far, controlled by Rim101p (9%), Msn2p/Msn4p (29%), and War1p (the PDR12 gene only), according to the information available in the YEASTRACT database (Teixeira et al., 2006b). These observations sustain the idea that Haa1p is the main player in the direct or indirect control of yeast transcriptional response to acetic acid stress. Approximately 52% of the Haa1p upregulated genes are presumably direct targets of this transcription factor as they exhibit in their promoter region a Haa1p DNA binding motif (our unpublished results). Among them are TPO3, YPR157w, and YGP1 genes, previously proposed as direct Haa1p targets (Keller et al., 2001). The effect of HAA1 deletion over the transcriptome of acetic acid-stressed cells may also result from the indirect effect of Haa1p over the transcription of genes encoding other transcriptional regulators, in particular, MSN4, NRG1, FKH2, STP3, and STP4. The putative Haa1pdependent transcriptional regulatory network presumably involved in the control of the early transcriptional response to acetic acid is proposed in Figure 8, based on the information gathered from the present study and in the literature (Teixeira et al., 2006b). This putative regulatory network reveals that a large number of genes whose transcription is reduced in Δhaa1 cells are documented targets of Nrg1p and Msn4p transcription factors (Fig. 8), suggesting that some crosstalk between these regulons may exist. Msn4p and its close homolog Msn2p are known to mediate yeast transcriptional response to environmental stress and to cooperate with other transcription factors under stress (Bar-Joseph et al., 2003; Gasch et al., 2000; Gorner et al., 1998; Rep et al., 1999; Schüller et al., 2004; Vyas et al., 2001). Nrg1p is a transcriptional repressor involved in glucose-dependent regulation of gene expression, in the negative regulation of filamentous growth, in alkaline pH response, and in the repression of stress-responsive genes (Kuchin et al., 2002; Vyas et al., 2005; Zhou and Winston, 2001). It is likely that under acetic acid stress Haa1p may act over the transcription of Nrg1p-target genes to alleviate the transcriptional repression exerted by this transcription factor, as proposed before to occur with Msn2p/Msn4p (Vyas et al., 2005) and with the Candida albicans transcription factor Mnl1p (Ramsdale et al., 2008). Such mechanism is thought to fine-tune the regulation of Msn2p/ Msn4p- and Mnl1p-dependent genes to prevent unwanted energy expenses (Ramsdale et al., 2008; Vyas et al., 2005).

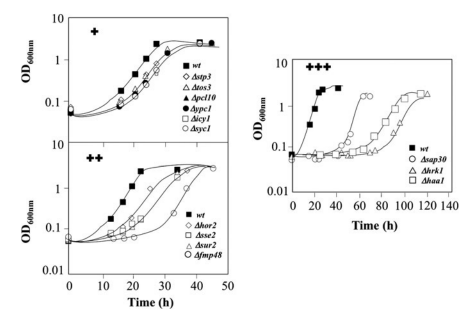


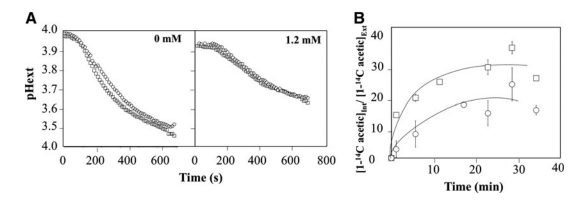
FIG. 6. Comparison of the susceptibility toward acetic acid of mutants devoid of the expression of genes whose transcription is activated in response to the acid in a Haa1p-dependent manner. Cells of the parental strain (wt) and of the indicated deletion mutants were grown until mid-exponential phase in liquid MM4 medium (at pH 4.0) and then used to inoculate the same basal medium either supplemented or not supplemented with 60 mM acetic acid (at pH 4.0; initial  $OD = 0.05 \pm 0.005$ ). Cells were batch cultured at 30°C and growth was monitored based on  $OD_{600 \text{ nm}}$ . Three levels of susceptibility phenotypes were considered based on the level of inhibition of the growth curves of the parental strain and of the different deletion mutants. The growth curves shown are representative of at least four independent experiments that gave essentially the same results.

Remarkably, Nrg1p was identified in this study as a determinant of resistance to acetic acid.

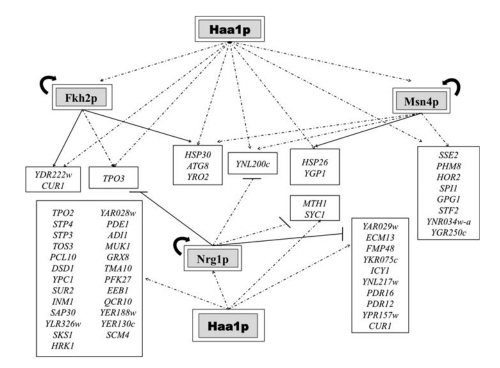
Several genes transcriptionally controlled by *HAA1* expression in response to acetic acid stress were implicated in this study, for the first time, in yeast tolerance to this weak acid but the level of protection exerted could not be directly correlated with the level of transcriptional activation induced by the acid, as reported before in studies dedicated to sorbic

acid, citric acid, and to the lipophilic acid herbicide 2,4dichlorophenoxyacetic acid (Lawrence et al., 2004; Schüller et al., 2004; Teixeira et al., 2006a). For example, the more evident susceptibility phenotypes were observed with mutants deleted for genes whose activation was around twofold only ( $\Delta hrk1$ ,  $\Delta sap30$ , and  $\Delta fmp48$ ).

With the exception of the drug:H<sup>+</sup> antiporters Tpo2p, Tpo3p, and of the cell-wall related proteins Ygp1p and Spi1p,



**FIG. 7.** (**A**) Comparison of the acetic acid-induced inhibition of external medium acidification rate promoted by energized cells of the parental strain BY4741 ( $\Box$ ) and of the deletion mutant  $\Delta hrk1$  ( $\bigcirc$ ). Deenergized cell suspensions of midexponential phase cells of both strains were either exposed or not to acetic acid (0 or 1.2 mM at pH 4.0) and then energized with a pulse of glucose (2%) to assess plasma membrane H<sup>+</sup>-ATPase activity, associated to the external acidification rate of the growth medium. The values shown are means of, at least, three independent experiments that gave rise to similar results. (**B**) Time course of the accumulation ratio of [1-<sup>14</sup>C]-acetic acid (corresponding to the ratio between intracellular (Int) and extracellular (Ext) [1-<sup>14</sup>C]-acetic acid) during cultivation of cells of the parental strain BY4741 ( $\Box$ ) or the  $\Delta hrk1$  mutant ( $\bigcirc$ ) in MM4 growth medium supplemented with 60 mM acetic acid (at pH 4.0). Cells used to perform the assay were grown in unsupplemented MM4 medium (at pH 4.0), harvested by centrifugation and reinoculated in fresh growth medium supplemented with radiolabeled and cold acetic acid as described in Materials and Methods.



**FIG. 8.** Transcriptional regulatory associations that may be governing the Haa1p-dependent response in yeast cells under acetic acid stress. The model is based on the microarray data obtained in this study and on the information available in the YEASTRACT database (December 2009). Genes whose transcription is activated in response to acetic acid in a Haa1p-dependent manner were selected for this analysis. The regulatory network is based on the documented regulatory associations between transcription factors (TF) whose expression is activated by Haa1p (the encoding genes are shown inside double boxes) and the other genes regulated by Haa1p, according with the microarray analysis carried out. Positive regulations between activators and target genes are represented by an arrow ( $\rightarrow$ ) and negative regulations between repressors and target genes are represented by lines (-). Documented associations are based on immunoprecipitation experiments (closed arrows or lines;  $\rightarrow$  or -) or on transcription analysis approaches with no demonstration of the binding of the TF to the promoter region of target genes (dashed arrows or lines,  $- - \rightarrow$  or - -).

previously implicated in yeast tolerance to acetic acid (Fernandes et al., 2005; Simões et al., 2006), all the other genes of the Haa1p regulon that are responsive to acetic acid and determinants of resistance to acetic acid are described here for the first time. Three of these genes encode proteins involved in lipid metabolism: YPC1 and SUR2 involved in sphingolipid metabolism, and INM1 involved in inositol metabolism. Assembly of the vacuolar membrane H<sup>+</sup>-ATPase (V-ATPase) is known to depend on sphingolipid metabolism (Chung et al., 2003), and C26-based sphingolipids also have a role in intracellular trafficking to the plasma membrane of nutrient transporters and of the H<sup>+</sup>-ATPase Pma1p (Chung et al., 2001; Gaigg et al., 2006). Proper vacuolar acidification under acetic acid stress was demonstrated to depend on the signaling molecule phosphatidylinositoide PtdIns(3,5)P2 (Mollapour et al., 2006), produced from inositol. The activities of plasma membrane H<sup>+</sup>-ATPase (PM-ATPase) and vacuolar membrane H<sup>+</sup>-ATPase (V-ATPase) are essential for yeast adaptation and resistance to weak acid-induced stress, being required for maintenance of intracellular pH homeostasis, nutrient compartmentalization, and protein sorting (Carmelo et al., 1997; Fernandes et al., 2003; Makrantoni et al., 2007; Martinez-Munoz and Kane, 2008; Mira et al., 2009; Viegas et al., 1998). Given this, it is likely that the Haa1p-dependent increased transcription of SUR2, YPC1, and INM1 genes in acetic acid-challenged cells may contribute to optimal activity

of these two proton pumps, and thus to increased tolerance towards acetic acid.

Haa1p-mediated resistance to acetic acid was also found to involve the expression of SAP30 gene, which encodes a component of the histone deacetylase Rpd3L complex. Indeed, SAP30 expression provided one of the highest levels of protection against acetic acid identified in our study. The deletion of SIN3 and SDS3 genes, which encode two proteins required for the structural integrity and catalytic activity of the Rpd3L complex and whose expression is not Haa1pdependent, also leads to a higher susceptibility to acetic acid. Surprisingly, the elimination of RPD3 gene, encoding a histone deacetylase of the complex, does not significantly increase yeast susceptibility to acetic acid. It is possible that the reduced susceptibility phenotype of  $\Delta rpd3$  cells to acetic acid results from a functional redundancy between Rpd3p and Hos2p, another histone deacetylase that was recently considered part of the Rpd3L complex (Shevchenko et al., 2008). Acetylation and deacetylation of histones are among the complex mechanisms involved in the regulation of gene expression in eukaryotes by modulating the accessibility of transcription factors to their target promoter regions (Kurdistani and Grunstein, 2003). It is therefore conceivable that, through the Rpd3L complex, Sap30p modulates the yeast transcriptional response to acetic acid by affecting the activity of transcription factors involved in such genomic expression

program. Consistent with this hypothesis, it was recently demonstrated that the Rpd3L complex affects the activity of the general stress responsive transcription factors Msn2p and Msn4p (Alejandro-Osorio et al., 2009), which govern the transcriptional response to acetic acid-induced stress. Interestingly, the activity of Haa1p was also suggested to depend on histone acetylation (Guo et al., 2006) and a protein–protein interaction of Haa1p with the histone deacetylase Hos2p was demonstrated (Beltrao et al., 2009).

The other major determinant of resistance to acetic acid stress that is transcriptionally controlled by Haa1p is the *HRK1* gene. Notably, the elimination of *HRK1* leads to an increased accumulation of acetate into yeast cells challenged with acetic acid, as reported to occur in  $\Delta haa1$  cells (Fernandes et al., 2005). Because Hrk1p belongs to a group of protein kinases thought to be involved in the regulation of the activity of plasma membrane transporters (Goossens et al., 2000), it is possible to speculate that this protein kinase may be involved in the reduction of the intracellular acetate concentration through the activation, by phosphorylation, of putative(s) plasma membrane acetate exporter(s). Due to its negative charge, acetate tends to accumulate in the cytosol following dissociation of the lipophilic acid form, and the active expulsion of the counterion from the cell interior contributes to the cell protection against acetic acid. So far, only the plasma membrane drug:H<sup>+</sup> antiporters Tpo2p, Tpo3p, Aqr1p, and Azr1p were suggested to play a role in acetic acid resistance (Fernandes et al., 2005; Tenreiro et al., 2000, 2002). According to our microarray analysis, the transcription of TPO3, TPO2, and AQR1 genes is activated by acetic acid under the dependence of Haa1p. The AQR1 gene is not listed in Table 2 because the difference in the mRNA levels of this gene produced in the wild-type strain and in  $\Delta haa1$  cells was below 25%, but it is included in Supplementary Table S4. A large scale screening of the yeast kinome, performed using cells cultivated in the absence of stress, in rich growth medium, led to the identification of a few Hrk1p targets although none of them is a plasma membrane transporter (Ptacek et al., 2005). One of the identified Hrk1p targets is a cytosolic enzyme, Gph1p, involved in the catabolism of glycogen, the expression of GPH1 providing protection against acetic acid (our unpublished results). This suggests that other Hrk1p targets relevant for acetic acid tolerance may exist besides plasma membrane transporters. The ongoing analysis of the phosphoproteome of yeast cells under acetic acid stress, either expressing or not HRK1, is expected to bring light into the biological role of Hrk1p as a major determinant of yeast resistance to acetic acid. In previous studies, Hrk1p was proposed to stimulate the activity of the plasma membrane proton pump Pma1p (de la Fuente et al., 1997), but according to our results, there is no positive effect of *HRK1* expression in the *in vivo* activity of PM-H<sup>+</sup>-ATPase under acetic acid stress. However, a possible role for Hrk1p in counteracting the weak acid-induced decrease of plasma membrane electrochemical potential, essential to secondary nutrient uptake and acetate export, is supported by the increased susceptibility of  $\Delta hrk1$ cells to hygromycine B (Goossens et al., 2000), a compound known to disrupt plasma membrane electrochemical potential. Remarkably, Hal4p and Hal5p protein kinases, which modulate the activity of the potassium uptake transporters Trk1p and Trk2p, thereby playing a key role in the maintenance of plasma membrane electrochemical potential, belong to the same family of protein kinases as Hrk1p (Goossens et al., 2000).

Haa1p was also implicated in yeast transcriptional response to lactic acid, propionic acid, and acetaldehyde (Abbott et al., 2008; Fernandes et al., 2005; Mira et al., 2009). The analysis of the published data shows that most of the genes identified in our study as being transcriptionally activated in response to acetic acid, under the direct or indirect dependence of Haa1p, are also upregulated in response to those compounds (Tables 2 and 3), whereas only a small group of these genes is activated in response to the more lipophilic sorbic acid (Tables 2 and 3). These results are consistent with the concept that the Haa1p-dependent regulatory pathway is particularly responsive to less lipophilic chemicals.

In summary, the present study has characterized the key role played by Haa1p and by genes of the full Haa1p-regulon in yeast response to acetic acid. Only 20% of the genes whose transcript level is increased in response to acetic acid under the direct or indirect dependence of Haa1p have a significant role in yeast protection against the acid. These genes are involved in lipid metabolism, in the regulation of gene expression by modulation of chromatin accessibility, and in the reduction of the intracellular acetate concentration. Microbial production of acetic acid or lactic acid using yeasts as cells factories is a very active area of research in the present days (Sauer et al., 2008). Acetic acid is also one of the inhibitory compounds resulting from the pretreatment of lignocellulosics (Almeida et al., 2007) and from yeast alcoholic fermentation, its accumulation contributing to a decreased performance of wine and bioethanol production processes. The present study therefore provides useful insights to guide the engineering of more robust yeast strains suited to overproduce acetic acid or lactic acid and to conduct alcoholic fermentation envisaging bioethanol and wine production. Because the fungicidal action of fluconazole is elicited by short-chain weak acids, in particular by acetic acid (Moosa et al., 2004), and considering that Haa1p and the proteins encoded by its target genes have robust homologs in *Candida glabrata*, the indications emerging from this study might also be useful to develop suitable strategies to control this pathogenic yeast.

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#### **Author Disclosure Statement**

The authors declare that no conflicting financial interests exist.

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