Expression of GUP1 and GUP2, Saccharomyces cerevisiae glycerol active transport genes

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Introduction

Two highly homologous genes related to a phenotype of salt stress tolerance were identified in *Saccharomyces cerevisiae*. Upon salt stress, a strain lacking the capacity to sinthesize glycerol (*gpd1gpd2*) is able to increase intracellular levels of glycerol by taking it up from the medium. Accordingly, the presence of small amounts of glycerol in the medium decreased osmosensitivity of this strain. Based on these findings yeast genomic fragments cut on these findings, yeast genomic fragments cut out from a mTn-*lacZ/LEU2*-mutagenized plasmid library were used to transform a *gpd1gpd2* strain with subsequent secreening for decreased osmotolerance of the transformants. This



Characterization of glycerol transport activity

Previous experiments of glycerol transport on conditions (fig. 2). Only the combination of gup1 wild type strains have shown that the active and gut1 deletions eliminated the saturable transport activity was absent in glucose-grown glycerol uptake and further deletion of the GUP1 cells, being this transport detectable in cells homologue (GUP2, YPL189w) did not change the grown with non-fermentable carbon sources such uptake results. The phenotype in ethanol-grown as ethanol (fig. 2).

attributed to a residual uptake pulled by an active symport described before (Lages and Lucas, 1997). catabolism of glycerol under derepressing growth

cells from radioactive glycerol uptake and Glycerol kinase (fig. 1), encoded by GUT1, extracellular alcalinization of cell suspensions interfered on transport measurements, being the upon addition of glycerol (not shown), suggested saturation kinetics present in *gup1* deleted strains that *GUP1* is the gene responsible for the proton

Regulation of glycerol transport activity

As mentioned above, glycerol active transport protonophore sensivity characteristic of symports was found to be under glucose repression. (not shown). Furthermore, the Km value was very However, the mutant screening that lead to GUP1 close to the one attributed to GUP1 activity identification was performed in gpd1gpd2 genetic (1mM). background growing on media containing glucose One of the possibilities to explain these results grown in this medium (fig. 3).

Surprisingly, *gpd1gpd2* strain displayed a strong suggest that *GUP1*-dependent and *GUP2*-uptake which, once determined as a total kinetic dependent glycerol uptake in glucose-grown cells study revealed a Vmax more than two times are tightly controlled and need the extreme higher than wild type cells grown in ethanol (figs. conditions of strong osmotic stress combined 2 and 3). This transport activity displayed proton with glycerol synthesis impairment to be uptake, and accumulation capacity, and detectable.

as carbon and energy source supplemented with was to attribute to GUP1 homologue the NaCl as osmotic stress agent and glycerol for responsability for this active uptake. As shown compatible solute purposes. Therefore, the in fig. 3, $gup\ddot{Z}$ deletion affects glycerol transport glycerol transport study was extended to cells by abolishing any measurable transport in a gpd1gup1 genetic background. These results

Optimization of RT-PCR

The expression of GUP1 and GUP2 was determined exponential phase was determined for each by relative quantitative RT-PCR. Previously, PCR analysis. By stoping PCR at this point, the conditions were optimized in order to fluorescence intensity of the bands is proportional quantitatively detect mRNA differences among to the amount of target molecules at the begining strains and growth conditions tested. Exponential of PCR. Once optimization has been established, phase of amplification was determined for *GUP1* the linearity of the method was tested using (fig. 4 and 5), GUP2 (fig. 6 and 7) and the internal different amounts of DNA to perform PCR (fig. standard 18S rRNA (figs. 8 and 9). A number of 10 and 11). amplification cycles corresponding to mid-

Relative quantification of mRNA's

for both genes (figs. 12 and 13). In wild type cells, highest levels of GUP1 mRNA mutant (fig. 12). were observed by growth on glucose as sole Contrarily, for GUP2, transcription is the lowest carbon and energy source. In cells grown on in glucose-grown cells, pointing to a probable glycerol, the transcription was the lowest of all, involvement in salt stress response and ehanol indicating a possible effect of extracellular utilization. In gpd1gpd2 mutant, mRNA levels are glycerol as inhibitor of transcription. higher for glucose-grown cells in the presence Unexpectedly, the double mutant gpd1gpd2 of NaCl and glycerol, which correlates with the unable to synthesize glycerol, exhibited lower high transport activity measured in this condition.

According to microarrays data available at the levels of mRNA in glucose-grown cells than the Saccharomyces Genome Database, we have wild type. On the other hand, glycerol apparently observed a decrease in mRNA levels during the increases *GUP1* transcription, as can be seen by shift from fermentative to respiratory metabolism the difference in mRNA levels between salt stress with and without added glycerol in the double

Conclusions

Taking together the results of the transport and provided its presence in the medium. assays and RT-PCR, we had detected GUP1 and GUP2 mRNA's for every growth between mRNA levels and transport conditions, suggesting post-transcriptional regulation of glycerol transport activity. Nevertheless, a match between mRNA levels and active transport activity was observed for gpd1gpd2 double mutant cells grown on glucose in the presence of NaCl and glycerol, for which the highest values were determined (figs. 3 and 13). Hence, *GUP2* is apparently involved in response to salt stress when simultaneous expression and the involvement of GUP2 impairment of glycerol synthesis occurs in salt stress response in yeast.

As for *GUP1*, no correlation was detected activity, suggesting that the major steps of regulation of *GUP1* expression lie downstream transcription.

In silico analysis of the promoter region of both genes had shown the presence of a stress response element (STRE - AG₄) at position -292 only on *GUP2*. So, the results presented here provide good evidence for different mechanisms for regulation of



Legend

YPD ferm - fermentative metabolism in YPD (OD600<1) YPD shift - shift to respiratory metabolism in YPD (OD600=2) YPD resp - respiratory metabolism in YPD (OD600>3) YPD S - YPD supplemented with NaCl 1M YPD SG - YPD supplemented with NaCl 1M and glycerol 15mM YPEt - YP with ethanol as carbon and energy source YPGly - YP with glycerol as carbon and energy source

References lolst et al. 2000. Molecular Microbiology. 37: 108-124 Nelissen et al. 1997. FEMS Microbiol Rev. 21: 113-134 Lages and Lucas. 1997. Biochim Biophys Acta. 1322: 8-18