Metabolic engineering of *Saccharomyces cerevisiae* for improvement in stresses tolerance

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**ABSTRACT**

Lignocellulosic biomass is an attractive low-cost feedstock for bioethanol production. During bioethanol production, *Saccharomyces cerevisiae*, the common used starter, faces several environmental stresses such as aldehydes, glucose, ethanol, high temperature, acid, alkaline and osmotic pressure. The aim of this study was to construct a genetic recombinant *S. cerevisiae* starter with high tolerance against various environmental stresses. Trehalose-6-phosphate synthase gene (*tps1*) and aldehyde reductase gene (*ari1*) were co-overexpressed in *nth1* (coded for neutral trehalase gene, trehalose degrading enzyme) deleted *S. cerevisiae*. The engineered strain exhibited ethanol tolerance up to 14% of ethanol, while the growth of wild strain was inhibited by 6% of ethanol. Compared with the wild strain, the engineered strain showed greater ethanol yield under high stress condition induced by combining 30% glucose, 30 mM furfural and 30 mM 5-hydroxymethylfurfural (HMF).

**KEYWORDS**

Aldehyde reductase; ethanol production; neutral trehalase; *Saccharomyces cerevisiae*; Trehalose; Trehalose-6-phosphate synthase

**INTRODUCTION**

Lignocellulosic biomass and agricultural residues are most widely used for bioethanol production. Acid hydrolysis is commonly used as a pretreatment to convert lignocellulosic materials into sugars. During acid hydrolysis of lignocellulosic materials, furfural and HMF are generated by dehydration of pentoses and hexoses, respectively, and both furfural and HMF are representative aldehyde inhibitors. Aldehydes are toxic to yeasts in inhibition of cell growth and production of ethanol. Beside aldehydes inhibitors, yeasts need to overcome various stresses during converting sugars to ethanol, such as acid left from pretreatment, high osmotic pressure of hydrolysates, high ethanol concentrations at the last stage of fermentation, high temperature for accelerating fermentation, and process interruptions. Survival under various stress conditions is an important feature required for yeast to produce ethanol efficiently.

*Saccharomyces cerevisiae* has been wildly used as a starter in bioconversion of ethanol. It has natural ability to convert furfural and HMF to furfuryl alcohol and 2,5-bis-hydroxymethylfuran, respectively. Reduction of aldehydes, mainly furfural and HMF in the lignocellulosic substrate, helps to improve the cell growth and subsequent ethanol production. Many researches demonstrated that over-expression of the aldehyde dehydrogenase/reductase genes (*ari1*, *ald6* and *adh6*) in *S. cerevisiae* not only could eliminate the toxicity of aldehydes but also improve tolerability.

Trehalose, a non-reducing disaccharide, works as an energy source in yeasts, bacteria, fungi, invertebrates, insects and plants. Many research demonstrated that the higher trehalose accumulated in the yeast cell the higher tolerance against various environmental stresses, for instance ethanol stress, heat stress, saline stress, and various other environmental stresses. In *S. cerevisiae*, concentration of trehalose is regulated by synthesis enzymes and hydrolysis enzymes. Enzymes involved in trehalose synthesis including trehalose-6-phosphate synthase (TPS, encoded by *tps1*), trehalose-6-phosphate phosphatase (TPP, encoded by *tps2*) and 2 other proteins, TSL1...
On the other hand, trehalose is hydrolyzed into glucose by neutral trehalases (encoded by nth1 and nth2) and acid trehalase (encoded by ath1).

In our previous study, an engineering S. cerevisiae (named as SCTΔN) was constructed by overexpression of tps1 and disruption of nth1. The engineered strain showed greater ethanol yield than the wild type strain when the medium contained more than 15% glucose (under glucose stress). In the present study, aldehyde reductase gene (ari1) was further overexpressed in SCTΔN and the tolerance of present strain was great improved. This study is the first to report the co-expression of tps1 gene and ari1 gene in S. cerevisiae to protect cells against not only environmental stresses but aldehyde inhibitors as well.

Results

Expression of trehalose-6-phosphate synthase and aldehyde reductase

In our previous study, SCT (S. cerevisiae with tps1 gene overexpression) and SCTΔN (nth1 gene of SCT was disrupted) were constructed from S. cerevisiae BCRC 21685 (SC). Concentration of intracellular trehalose was in the order of SCTΔN>SCT>SC. Compared to SC, SCTΔN showed greater viability and ethanol productivity under 15% of ethanol and 10% of glucose, respectively.

To create a starter for production of bioethanol, ari1 DNA (encode for aldehyde reductase gene) of SC was inserted in the sense orientation, downstream of the GAP promoter of pGAPZaC to create pGAPZC-ari1. Both SCT and SCTΔN were served as the host for homologous recombination of pGAPZC-ari1 in their gnomic DNA, and PCR and western blot were applied to confirm the transformates, named as SCTA and SCTAΔN, respectively.

Fig. 2 showed the DNA electrophoresis of PCR products for verification of recombinant ari1 DNA. As shown in the Fig. 2A, SCT contained a DNA fragment about 1.5 kb (tps1 gene); while two DNA fragments, about 1.5 kb (tps1 gene) and 1.0 kb (ari1 gene), presented in SCTA and SCTAΔN. Wild strain (SC) did not generate any PCR product. Results of western blot (Fig. 2B) showed one protein band about 56 kDa (trehalose-6-phosphate synthase) in SCT, and 2 protein bands about 56 kDa (trehalose-6-phosphate synthase) and 38 kDa (aldehyde reductase) in SCTA and SCTAΔN. Again, no protein band appeared in SC. Results of PCR and western blot approved successful recombination and expression of ari1 in SCTA and SCTAΔN.

Figure 1. Multiple insertion events in yeast genome by using expression vectors pGAPZC-tps1 and pGAPZC-ari1.
Trehalose concentration under ethanol stress conditions

In our previous study, we found that overexpression of *tps1* gene could increase the intracellular trehalose content in yeast cells.\(^25\) In the present study, the effect of *ari1* gene overexpression on the intracellular trehalose content in yeast cells were evaluated (Table 3). When yeasts were suffered from environmental stress induced by ethanol (10 and 15%) for 1 h, all yeasts increased their intracellular trehalose contents with the increase of ethanol concentration. The intracellular trehalose contents of SCT (*S. cerevisiae* with overexpression of *tps1* gene) and SCTA (*S. cerevisiae* with co-overexpression of *tps1* and *ari1* genes) did not show significant difference under conditions of with/without ethanol stress (*p* > 0.05). SCTA\(\Delta N\) obtained significant higher intracellular trehalose content than SCT (*p* < 0.05), as well as SCTA\(\Delta N\) was compared with SCTA. High intracellular trehalose content in SCTA\(\Delta N\) and SCTA\(\Delta N\) was due to the effect of *nth1* deletion. SCTA\(\Delta N\) showed significantly higher intracellular trehalose content than SCT\(\Delta N\) (*p* < 0.05). Overexpression of *ari1* did not increase the trehalose content in SCT, but it did increase the trehalose content in SCTA\(\Delta N\).

Gene expression of *tps1* and *ari1* in SCTA\(\Delta N\) under stress conditions

SCTA\(\Delta N\) had the highest intracellular trehalose content, therefore, its gene expression level of *tps1* and *ari1* was analyzed and compared with wild strain, SC. Compared with SC, SCTA\(\Delta N\) expressed significant higher level of *tps1* (*p* < 0.05), no matter of ethanol concentration in growth medium (Fig. 3A). Ethanol stress induced expression of *tps1* in a range of 1.2 to 1.25 fold, however, SCTA\(\Delta N\) did not expressed *tps1* differently under 10% and 15% ethanol (*p* > 0.05). On the other hand, *tps1* expression of SC did not be affected by ethanol under tested conditions, and all in a very low level compare with that of SCTA\(\Delta N\). Similar phenomenon was observed when yeasts were incubated with various concentration of HMF (0, 10 and 30 mM). The gene expression of *ari1* was induced to the same level with the present of HMF at 10 and 30 mM in SCTA\(\Delta N\) (Fig. 3B). Expression of *ari1* was also induced by 10 mM of furfural but not 30 mM of furfural (Fig. 3C).

When incubated in YPD broth for 1 h, SC and SCTA\(\Delta N\) did not show significant difference in their viability (Fig. 3D, 3E and 3F), however, the difference emerged under stress conditions induced by ethanol (Fig. 3D), HMF (Fig. 3E) and furfural (Fig. 3F). SCTA\(\Delta N\) showed significantly higher survival than SC when it was confronted with environmental stresses (*p* < 0.05).

Overexpression of *tps1* resulted in increasing synthesis of trehalose in yeast cells, which functioned as a very efficient defense mechanism against ethanol stress. Overexpression of *ari1* resulted in increasing activity of aldehyde reductases in yeast cells, which reduced the toxicity of aldehydes toward yeast cells.

Effects of ethanol concentrations on cell growth

To further understand the effect of ethanol on the growth of SCTA\(\Delta N\), cells were cultured in YPD medium containing 0 to 18% (v/v) ethanol and incubated at 30°C for 24 h (Fig. 4). Without ethanol stress, SCTA\(\Delta N\) did not grow better than wild strain, SC (*p* > 0.05). However, when the medium contained 6% or more ethanol, SCTA\(\Delta N\) showed superior growth over SC (*p* < 0.05). The growth of SCTA\(\Delta N\) was inhibited by ethanol concentrations greater or equal to 14%, while 6% ethanol could slow the growth of SC. When the media contained 14 to 18% ethanol, SC showed no growth in 24 h. When SCTA\(\Delta N\) was incubated under 10 and 15% ethanol for 1 h, its viability decreased but its intracellular trehalose content increased (Fig. 3D and Table 3). SCTA\(\Delta N\) showed higher value of OD\(_{600}\) at YPD containing 6–10% ethanol than that of at YPD without adding ethanol (Fig. 4) indicating that under ethanol stress conditions higher trehalose was
accumulated and played a dual role as stress protectant and carbohydrate energy storage.

**Capacities of furfural and HMF reduction**

To compare the capacity to reduce aldehydes, SC and SCTAΔN were incubated independently under the challenge of furfural (10 mM, 30 mM) or HMF (10 mM, 30 mM) in YPD broth. Both SC and SCTAΔN completely degraded 10 mM of furfural in 12 h. When the medium contained 30 mM of furfural, furfural was totally reduced by SCTAΔN in 24 h; however, 8.64 ± 0.35 mM of furfural was remained after incubation with SC for 96 h (Fig. 5A). Both SCTAΔN and SC reduced 10 mM of HMF efficiently (Fig. 5B). When HMF was increased to 30 mM, SC needed more than 96 h to accomplish reduction and SCTAΔN almost finished reduction in 24 h. Compared with SC, SCTAΔN showed greater capacity on reduction of furfural or HMF ($p < 0.05$).

SC and SCTAΔN showed similar growth rate when medium contained 10 mM of furfural or HMF.

![Figure 3](image-url)
Growth of SC was decreased by 30 mM of furfural or HMF, while SCTA \( \Delta N \) was not affected under the same condition (Fig. 5A, B).

The above results indicated that overexpression of \( \text{ari1} \) gene in engineered strain (SCTA \( \Delta N \)) enhanced reduction of aldehydes and increased cell viability as well.

**Ethanol production under high stress conditions**

The ethanol productivity of SCTA\( \Delta N \) was compared with that of SC under high stress conditions induced by combined effects of 30% glucose, 30 mM furfural and 30 mM HMF (Fig. 6A). After incubation for 96 h, both SCTA\( \Delta N \) and SC produced maximum amount of ethanol, and the ethanol concentration were 110.59 \( \pm \) 0.28 g/L and 44.03 \( \pm \) 1.43 g/L, respectively. At the end of processes, 12.70 \( \pm \) 0.10% glucose was remained in fermentation broth using SC as starter, whereas the glucose in medium was almost used up by SCTA\( \Delta N \) (1.67 \( \pm \) 0.03% residual glucose). Biomasses of SC and SCTA\( \Delta N \) reached their peaks at 24 h incubation, and SCTA\( \Delta N \) showed better growth in the high stress conditions than SC.

After incubation for 96 h, HMF in fermentation broth was reduced to 8.03 \( \pm \) 0.12 mM and 1.05 \( \pm \) 0.03 mM by SC and SCTA\( \Delta N \), respectively (Fig. 6B). On the other hand, furfural was almost completely reduced by SCTA\( \Delta N \) in which broth contained 0.69 \( \pm \) 0.03 mM of furfural. Significant amount of furfural (14.21 \( \pm \) 0.83 mM) was not reduced by the SC. Intracellular trehalose of SC was gradually increased during growth and reached to 58.54 \( \pm \) 0.74 mg/g. SCTA\( \Delta N \) had the highest trehalose accumulation (134.05 \( \pm \) 1.18 mg/g) after incubation for 48h.

The time course of theoretical ethanol yields for SC and SCTA\( \Delta N \) was shown on Fig. 6C. These two strains converted glucose into ethanol continuously during 96 h. Ethanol yields of SCTA\( \Delta N \) were significantly higher than that of SC ( \( p < 0.05 \)). SCTA\( \Delta N \) achieved a great efficiency in ethanol production (72.14 \( \pm \) 0.18%), while, SC got 28.72 \( \pm \) 0.93% of ethanol yield.

**Discussion**

Single insertion event (SCT with insertion of \( tps1 \) gene) is much more likely to happen but the occurrence of multiple insertion may be 1–10% that of single insertion events. To locate these “jack-pot” clones of transformants with multiple insertions of \( tps1 \) gene and \( \text{ari1} \) gene (SCTA and SCTA\( \Delta N \)), we screened more than hundreds of Zeocin-resistant transformants.

Trehalose is a nonreducing disaccharide, in which 2 glucose are linked with an \( \alpha,\alpha-1,1 \) glycosidic linkage.
Trehalose functions as a protector for cell membranes and cellular proteins under environmental stresses such as heat, cold, desiccation, dehydration, ethanol, oxidation and anoxia. In yeast, trehalose level is low under favorable growth condition but is induced by environmental stresses. However, Ratnakumar and Tunnacliffe suggested that there was no consistent relationship between intracellular trehalose concentration and desiccation tolerance in *S. cerevisiae*. In our previous study, SCTΔN (*S. cerevisiae* with overexpression of *tps1* gene and deletion of *nth1* gene) showed not only the highest accumulated intracellular trehalose but the greatest tolerance under stress conditions compare with SCT (*S. cerevisiae* with overexpression of *tps1* gene) and wild strain. Consistent with those results, engineered strain SCTΔN (co-overexpression of *tps1* gene and *ari1* gene, and deletion of *nth1* gene) constructed in this study showed higher trehalose content than SCTΔN under favorable/stress condition. Compared with SCTΔN, SCTΔAN exhibited greater viability under stresses induced by ethanol or high glucose osmosis confirming the defense mechanism of intracellular trehalose.

Furfural and HMF, the most important aldehyde inhibitors in lignocellulosic hydrolysates, are known to affect the growth and fermentation performance of *S. cerevisiae*. However, *S. cerevisiae* has the natural ability to reduce furfural and HMF to their corresponding less toxic alcohols by multiple dehydrogenases/reductases. Researchers stated that, overexpression of the dehydrogenase/reductase genes (*ari1*, *adh7*, *ald6* and *adh6*) increased enzyme activities for furfural and/or HMF reduction and simultaneously improved the tolerability of yeast against inhibitors.

Quantitative real-time PCR data confirmed the overexpression of *tps1* and *ari1* in engineered strain SCTΔN. As compare with wild strain (SC), SCTΔN exhibited significantly higher cell survival under stresses of ethanol, furfural and HMF. The engineered strain (SCTΔN) showed significantly higher ethanol tolerance which might be due to overexpression of *tps1* that results in more trehalose accumulation. Similarly, overexpression of *ari1* in SCTΔN increased the enzyme activities for furfural and/or HMF reduction and improves the tolerability toward them.

When the medium contained 10 mM furfural and/or HMF; cell growth and reduction capacities of SC and SCTΔN were almost similar (Fig. 5A, B). As the concentrations were increased to 30 mM; SCTΔN showed significantly greater growth and furfural and/or HMF reduction capacity than the wild strain SC (*p* < 0.05). Overexpression of *ari1* (aldehyde reductase) improved the inhibitor tolerance and cell viability of engineered strain. Liu and Moon stated that aldehyde reductase gene is involved in the detoxification of aldehyde inhibitors.

Under the stresses of 30% glucose, 30 mM furfural and 30 mM HMF engineered strain SCTΔN showed greater performance. Compared with SC, SCTΔN showed better capacity on converting glucose to ethanol, reducing HMF and furfural to less toxic compounds, and generating intracellular trehalose. The high accumulation of intracellular trehalose and

![Figure 5. Furfural (A) or HMF (B) degradation capacities and biomass of wild strain (SC) and recombinant strain (SCTΔN). Data are presented as the means ± SD (n = 3). *Significantly higher than the other group. One-way ANOVA, Student’s t test, p < 0.05. SC at 10 mM furfural/HMF; SC at 30 mM furfural/HMF; SCTΔN at 10 mM furfural/HMF; SCTΔN at 30 mM furfural/HMF; viability of SC at 10 mM furfural/HMF; viability of SC at 30 mM furfural/HMF; viability of SCTΔN at 10 mM furfural/HMF; viability of SCTΔN at 30 mM furfural/HMF.](image_url)
overexpression of \textit{ari1} in SCTA\textDelta N provided the cells high tolerance against the high environmental stresses induced by combining stresses of glucose, HMF and furfural.

Overexpression of \textit{tps1} (trehalose-6-phosphate synthase) and removal of \textit{nth1} (neutral trehalase) improved the intracellular trehalose level and ethanol tolerance. A positive correlation was observed between cell viability and trehalose concentration because intracellular trehalose prevents the ethanol-induced electrolyte leakage in yeast cells. \textsuperscript{31} Moreover, overexpression of \textit{ari1} (aldehyde reductase) improved the enzyme activities for furfural and/or HMF reduction and inhibitor tolerance. Yeast strain with overexpression of aldehyde reductase gene showed not only more tolerant response to furfural and HMF, but earlier recoveries and a better growth as compare with wild type yeast strain.\textsuperscript{12} All these aspects of the strain improvement played an important role in cell viability as well as ethanol production under several different stress conditions.

The engineered strain (\textit{tps1} gene overexpression, \textit{ari1} gene overexpression and \textit{nth1} gene deletion) presented in this study exhibited increased viability and intracellular trehalose content with high ethanol yield under glucose, furfural, HMF and ethanol stress conditions. This strategy of strain improvement might provide a preferable reuse of starters as immobilized cells and even provide applications in continuous processes.

\textbf{Conclusion}

This study describes the construction of stress tolerant yeast strain to increase intracellular trehalose concentration and to improve its tolerability toward the aldehyde inhibitors. The strain was improved by overexpression of \textit{tps1} gene, \textit{ari1} gene and the entire deletion of \textit{nth1} gene. During this study we found that engineered strain was more tolerant to environmental stresses and aldehyde inhibitors. Compared to the wild strain, the engineered strain exhibited not only higher trehalose accumulation, but cell viability as well as increased furfural and/or HMF reduction capacity. The novel strain constructed in this study will be promising for bioethanol production from lignocellulosic materials and agricultural residues.

\textbf{Materials and methods}

\textit{Strains, vectors, and media}

All the strains and vectors used in this study are listed in \textit{Table 1}. \textit{Escherichia coli} TOP10F' was purchased.
from Novagen Inc. (Wisconsin, USA) and *S. cerevisiae* (BCRC 21685) was purchased from Bioresources Collection and Research Center, Food Industry Research and Development Institution, Shinchu, Taiwan. The expression vector pGAPZαC, purchased from Invitrogen (Carlsbad, CA, USA), was used for *ari1* gene overexpression study. Recombinant vectors were multiplied in *E. coli* TOP10F'.

*E. coli* and yeast were maintained and cultivated in Luria-Bertani (LB) (10 g/L peptone, 10 g/L NaCl, and 5 g/L yeast extract) medium at 37°C and YPD (10 g/L yeast extract, 20 g/L peptone, 20 g/L dextrose) medium at 28°C, respectively. To select the Zeocin resistant transformants, low salt LB (10 g/L peptone, 5 g/L NaCl, and 5 g/L yeast extract) plates were supplemented with 25 mg/L Zeocin (Invitrogen Corp., Carlsbad, California, USA) and YPD plates were supplemented with 100 mg/L Zeocin. All parental strains and engineered strains were maintained in 30% glycerol at −80°C. Vectors within the host cells were extracted using Gene-spin miniprep plasmid purification kit (Protech Technology, Taipei, Taiwan). Yeast genomic DNA (gDNA) was prepared according to the manufacturer’s protocol (Genomic DNA purification kit, BioKit, Miaoli, Taiwan).

**Primers**

Table 2 lists the primers used in this study. These primers were designed according to the website of NCBI (National Center for Biotechnology Information) for nucleotide sequences of *tps1* gene (GenBank ID: NM_001178474.1), *ari1* gene (GenBank ID: NM_001181022.3), *taf10* gene (GenBank ID: NM_001180474.3) and genomic DNA of *S. cerevisiae*.

**Genetic manipulation**

Genomic DNA of *S. cerevisiae* wild strain was used as the template to amplify *ari1* gene by PCR with primers of Ari1-F and Ari1-R. DNA thermal cycler (Labcycler Gradient, SENSQUEST, Gottingen, Germany) was used to obtain the PCR products. All PCR products were electrophoresed, observed by ethidium bromide (EtBr) staining and purified with a Clean/Gel Extraction Kit (BioKit, Miaoli, Taiwan). The BstBI-XhoI fragment carrying the *ari1* gene was digested and

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**Table 1. Strains and vectors used in this study.**

| Strains or vectors           | Properties or product                                                                 | Reference or source            |
|-----------------------------|---------------------------------------------------------------------------------------|--------------------------------|
| *E. coli* TOP10F'            | F' [lac*Δ*10(Tet*)] mcrA (mrr-hsdRMS-mcrBC) B0lacZM15 lacX74 recA1araD139 (ara-leu)/7697 galU galK rpsL (Str') endA1 nupG | Invitrogen, Carlsbad, CA       |
| *S. cerevisiae* (BCRC 21685) | Wild type                                                                             | Bioresources Collection and Research Center, Taiwan |
| *Saccharomyces cerevisiae* with overexpression of *tps1* gene (SCT) | Carrying pGAPZC-tps1 vector                                                        | Divate et al., 2016            |
| *Saccharomyces cerevisiae* with overexpression of *tps1* and *ari1* gene (SCTA) | Carrying pGAPZC-tps1 vector and pGAPZC-ari1 vector                                    | This study                     |
| *Saccharomyces cerevisiae* with overexpression of *tps1* gene and deletion of *nth1* gene (SCT ΔN) | Carrying pGAPZC-tps1 vector and gene disruption cassette                             | Divate et al., 2016            |
| *Saccharomyces cerevisiae* with overexpression of *tps1*, *ari1* genes and deletion of *nth1* gene (SCT ΔN) | Carrying pGAPZC-tps1 vector, pGAPZC-ari1 vector and gene disruption cassette         | This study                     |
| pGAPZC-ari1                  | Expression Vector, GAP promoter, *Sh ble* gene (Zeocin™ resistance gene)              | Invitrogen, Carlsbad, CA       |
| pGAPZC-ari1                  | pGAPZCαC vector carrying a *ari1* gene                                                | This study                     |

**Table 2. Oligonucleotides used in this study.**

| Primer   | Sequence 5’ − 3’ | Restriction site | Purpose                          |
|----------|------------------|------------------|----------------------------------|
| Ari1-F   | TCGTTCGAAAAAATGGCGACTACTGACACACTGTTTTTCGTTTCTGTCTG | BstBI            | *ari1* gene amplification        |
| Ari1-R   | TCGTCGTGATTGATGTTGATGTTGATGGCGCTCTATTCTAACTCTCC | XhoI             |                                   |
| VTF      | TCGAAGAAGAGATTGTGAATACACGTATCTA               | —                | Verification of *tps1* or *ari1* gene insertion |
| VAF      | TCGAAAAAATGGCGACTACTA | —                |                                   |
| VR       | AGTGATGATGATGATGATGAGG   | —                |                                   |
| qTps1-F  | TTTGGTGGTCCCAAGACGGCTT | —                | quantitative real-time PCR       |
| qTps1-R  | GGAGGAGGGAGGTCTCTCGT  | —                |                                   |
| qAri1-F  | TTTGGTGGTCCCAAGACGGCTT | —                |                                   |
| qAri1-R  | CGTCTCCTCCTCTGTTTAGT  | —                |                                   |
| qTaf10-F | TCCAGAAGCAGGCTCTCTCCG | —                |                                   |
| qTaf10-R | TGCCCTGAAATAGCTGCTCTCG | —                |                                   |

The underlined bases encode restriction site and italicized bases encode 6xHis-tag.
ligated into the expression vector, pGAPZaC which was pre-digested with same restriction enzymes. The resulting vector was designated as pGAPZC-ari1. To amplify pGAPZC-ari1, recombinant vector was transformed into CaCl2-treated E. coli TOP10F’ according to Hanahan and Meselson.32

SCT (S. cerevisiae with tps1 gene overexpression) and SCTA ΔN were constructed in our previous study25 and further used in this study for ari1 gene overexpression. Prior to transformation, pGAPZC-ari1 was linearized by AvrII enzyme. Yeast transformation was performed by electroporation method according to the manufacture’s protocol (MicroPulser electroporation apparatus, Bio-Rad Laboratories, Hercules, CA, USA). To generate SCTA and SCTA ΔN, pGAPZC-ari1 was transformed into SCT and SCTA ΔN, respectively (Fig. 1). The colonies were selected by plating on YPDS plates (10 g/L yeast extract, 20 g/L peptone, 20 g/L dextrose, 1 M sorbitol) containing 100 mg/L Zeocin. Transformants were confirmed by PCR using VTF (for tps1 gene) or VAF (for ari1 gene) as forward primers and VR as reverse primer in same PCR reaction tube.

Table 3. Intracellular trehalose content of yeast cells under non stress and stress conditions.

| Ethanol concentration | Intracellular trehalose content of cell (mg/g dry weight) under stress | Reference or source |
|-----------------------|-----------------------------------------------------------------------|---------------------|
| SC                    | 9.2 ± 0.2±Δξ                                                         | Divate et al., 2016 |
| SCT                   | 12.2 ± 0.6±Δξ                                                         | Divate et al., 2016 |
| SCTA                  | 14.3 ± 2.1±Δξ                                                         | This study          |
| SCTA ΔN               | 24.9 ± 1.4±Δξ                                                         | Divate et al., 2016 |
| SCTA ΔN               | 33.1 ± 1.1±Δξ                                                         | This study          |

SC: Wild type Saccharomyces cerevisiae
SCT: S. cerevisiae with overexpression of tps1 gene
SCTA: S. cerevisiae with co-overexpression of tps1 and ari1 genes
SCTA ΔN: S. cerevisiae with overexpression of tps1 gene and nth1 gene deletion
SCTA ΔN: S. cerevisiae with co-overexpression of tps1 and ari1 genes and nth1 gene deletion

Measurement of trehalose concentration

Frozen culture (1 mL) was activated and grown in YPD broth at 30°C with shaking (150 rpm). To measure the intracellular trehalose concentration under ethanol stress, the culture was transferred into 100 mL of YPD broth to achieve an OD620 value of 0.3 and incubated in a 500 mL baffled Erlenmeyer Flask at 30°C with shaking (150 rpm) for 24 h. After incubation, cells were collected by centrifugation and transferred into fresh YPD broth (100 mL) containing ethanol (0%, 10%, and 15%) and incubated at same conditions for another 1 h.

After stress treatment, cells were collected by centrifugation at 7000 × g for 5 min and dried at 100°C for 12 h. To extract trehalose, a pellet of 40 mg was mixed with 2 mL ethanol (99.5%) and incubated in a boiling water bath for 1 h. For HPLC analysis of trehalose, extracts were suspended in 0.5 mL of acetonitrile: water (1:1). Acetonitrile: water (7:3) was used as the mobile phase. The HPLC system was equipped with a refractive index detector and a Lichrocart® 250–4 Purospher® Star NH2 column (5 µm) (Merck, Darmstadt, Germany).35

Blots were probed with Anti-His tag, Clone His.H8 (Millipore, Schwalbach, Germany; 1:5000 dilution), and visualized with a 1:5000 dilution of peroxidase-conjugated AffiniPure Goat Anti-Mouse IgG (Jackson immunoresearch, West Grove, PA, USA). Chemiluminescence detection was performed with Immobilon Western Chemiluminescent HRP Substrate (Millipore, Billerica, MA) and detected with a CCD-camera (Fusion-SL 3500.WL; Peqlab Biotechnologie, Erlangen, Germany).

Protein extraction, SDS-polyacrylamide gel electrophoresis (PAGE), and Western blot analysis

Frozen culture (1 mL) was activated and grown in YPD broth at 30°C with shaking (150 rpm). Yeast pellet (100 mg) was used for proteins extraction.33 SDS-PAGE was performed according to Laemmli.34 For western blotting, protein bands on the gel were transferred to polyvinylidene fluoride (PVDF) membrane (Perkin Elmer, Foster City, CA, USA) using Mini-Trans-Blot system (BioRad Laboratory, Inc., USA).
**Quantification of gene expression by real-time reverse transcription PCR (RT-qPCR)**

Frozen culture (1 mL) was activated and grown in YPD broth at 30°C with shaking (150 rpm). For stress treatment study, the culture was transferred into fresh YPD medium and incubated under the same conditions. When the OD<sub>620</sub> reached approximately 1.0, cells were collected by centrifugation and transferred into fresh YPD broth (100 mL) containing ethanol (0%, 10%, 15%), furfural (0 mM, 10, mM, 30 mM) or HMF (0 mM, 10 mM, 30 mM) in a 500 mL baffled Erlenmeyer Flasks. Cells were incubated at 30°C with shaking (150 rpm) for another 1 h. Stress tolerance was expressed as percentage of survivors.

For gene expression study, Cells were collected by centrifugation at 7000 × g for 5 min and used for RNA isolation. Total RNA was extracted with TRIzol reagent (Life Technologies, Inc., Grand Island, NY) according to the manufacturer’s instructions. RNA concentration and quality were assessed spectrophotometrically. Total RNA (1 μg) was subjected to reverse transcription using the iScript<sup>™</sup> cDNA synthesis kit (BioRad, Hercules, CA, USA). Real-time PCR was performed with iQ SYBR Green Supermix according to the manufacturer’s instructions (BioRad, Hercules, CA, USA). The Mini-Opticon<sup>™</sup> system (Bio-Rad, Hercules, CA, USA) was used to quantify the expression levels of tpsl and ari1. Amplifications were performed under the following conditions: 95°C for 3 min, 40 cycles at 95°C for 10 s and at 57.8°C for 30 s, and final extension at 95°C for 10 s. Gene expression levels of tpsl and ari1 were normalized to an internal control taf10 gene. Analyses were performed with the Bio-Rad CFX manager 2.1 software.

**Ethanol tolerance**

Ethanol tolerance was determined as mentioned in our previous study. Cells were adjusted to an OD<sub>620</sub> value of 0.05 with YPD containing 0–18% ethanol and incubated in the wells of a 96-well flat bottom polystyrene microtiter plate (Costar, Corning Inc., Corning, NY, USA) sealed with a gas-permeable sealing membrane (Breathe Easy membrane, Sigma Chemical Co., St Louis, MO, USA) at 30°C. The growth of yeasts was monitored by measurement of OD<sub>620</sub> with a microplate reader (Fluostar optima, BMG Labtech, Germany).

**Furfural and HMF reduction capacities**

Frozen culture (1 mL) was activated and grown in YPD broth at 30°C with shaking (150 rpm). To measure furfural and HMF reduction capacities, the culture was transferred into 100 mL of YPD broth containing furfural or HMF (10 mM and 30 mM) to achieve an OD<sub>620</sub> value of 0.3 and incubated in a 500 mL baffled Erlenmeyer Flask at 30°C with shaking (150 rpm). Samples were withdrawn at the indicated time intervals for determination of furfural and HMF concentrations and viable cell count.

Supernatant was collected by centrifuging at 10,000 × g for 10 min and filtering through a 0.45 μm membrane. The HPLC system was equipped with a refractive index detector and ICSep ICE-COREGEL 87H3 column (Transgenomic, Omaha, USA). The mobile phase was 5 mM H<sub>2</sub>SO<sub>4</sub> with a flow rate of 0.8 mL/min.

**Ethanol production capacity under stress conditions**

Frozen culture (1 mL) was activated and grown in YPD broth at 30°C with shaking (150 rpm). Cells were collected and centrifuged at 4000 × g for 5 min. The collected cells (3 × 10<sup>7</sup> cells/mL) were transferred into 250-mL Erlenmeyer flasks containing 100 mL of fermentation broth (1%, yeast extract, 2% peptone, 30% glucose, 30 mM furfural, and 30 mM HMF). The flasks were incubated statically at 30°C and samples were withdrawn at the indicated time intervals and centrifuged at 10,000 × g for 10 min, filtered through 0.45 μm membranes and used for analysis of glucose, ethanol, furfural and HMF concentration by HPLC. Pellet was used for trehalose extraction as mentioned above. The HPLC system was equipped with a refractive index detector and ICsep ICE-COREGEL 87H3 column (Transgenomic, Omaha, USA). The mobile phase was 5 mM H<sub>2</sub>SO<sub>4</sub> with a flow rate of 0.8 mL/min. The ethanol yield was calculated according to the following equation:

\[
\text{Ethanol yield (\%) = \left[ \frac{\text{gm ethanol produced}}{\text{gm glucose in medium} \times 0.511} \right] \times 100}
\]

**Viable cell count**

Sample was serially diluted as required and spread on YPD agar plates and plates were incubated at 30°C for 72 h. Viable cells were counted as colony forming units (CFU)/mL sample.
**Statistical analysis**

Trehalose concentration, ethanol concentration, glucose concentration, furfural concentration, HMF concentration, gene expression and biomass values were evaluated by one-way ANOVA, followed by Duncan’s new multiple-range test to determine the differences among means or Student’s t-test when only 2 groups were compared using the Statistical Analysis System (SAS institute, Cary, NC, USA). A significance level of 5% was adopted for all comparisons.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

**Funding**

This research was supported by the Ministry of Science and Technology, R.O.C. Taiwan (NSC 100–2313-B-126–001-MY3). Its financial support is greatly appreciated.

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