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3	Freeze tolerance of the yeast Torulaspora delbrueckii: cellular and biochemical
4	basis
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1 Abstract

The freeze-stress responses to prolonged storage at - 20 °C in *Torulaspora delbrueckii* PYCC5323 were investigated. In this yeast no loss of cell viability was observed for at least 120 days during freezing at - 20°C, whereas a loss of 80% was observed in a commercial baker's yeast after 15 days. In the former strain, freeze resistance was dependent on an adaptation process. The primary cell target of freeze stress was the plasma membrane, preservation of it's integrity being related with a lower increase of lipid peroxidation and with higher resistance to H₂O₂, but not with intracellular trehalose concentration.

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1 1. Introduction

2 Frozen-dough technology is well established in the baking industry, making it easier for bakers to 3 supply oven-fresh bakery products to consumers and improving labour conditions. However, storage 4 of frozen bread-dough may lead to the loss of baker's yeast cell viability as well as of its baking 5 capacity, and consequently to economic losses. Thus, bread-making industry keeps a high demand 6 for yeast strains with improved freeze resistance [1,2]. Most research on this field has focused on 7 strains of Saccharomyces cerevisiae, which is the species currently used as baker's yeast. In this 8 species, tolerance to freezing has generally been correlated with the intracellular trehalose 9 concentration, but no direct correlation has been found above a threshold value [3-6]. Prior to the 10 frozen storage, once the yeast cells are mixed with flour the fermentation takes place and a rapid loss 11 of stress resistance occurs [7]. This has also been associated with the degradation of intracellular 12 trehalose [8]. However, it has been shown that retention of high trehalose levels in fermenting cells 13 does not prevent the loss of fermentation capacity during freezing, and that other factors - not yet 14 identified - are required for the maintenance of freeze stress resistance [9, 10]. Accumulation of other 15 solutes such as amino acids and glycerol, and expression of aquaporins were also reported to 16 increase freezing resistance [11-15]. In addition, oxidative damage has been considered to be a factor 17 underlying freeze-thaw damage, since an oxidative burst has been predicted to occur during thawing 18 [16]. In agreement with this, Park et al. found that respiratory ability and functional mitochondria are 19 necessary to confer full resistance to freeze-thaw stress [17]. It has also been shown that freeze 20 tolerance is correlated with tolerance to H₂O₂, and free radicals were detected in S. cerevisiae after 21 the freeze-thaw process [5, 18].

The strain PYCC5323 of *Torulaspora delbrueckii*, isolated from traditional corn and rye bread dough from the North of Portugal, besides presenting dough-raising capacity, growth rates and biomass yields similar to commercial baker's yeast, displays high freeze and osmotic tolerance [19-21].. Therefore this yeast emerges as a powerful candidate for the bread making industry, and the elucidation of such a peculiar behaviour reveals to be of great interest.

In this work the freeze-stress responses to prolonged storage at - 20 °C in *Torulaspora delbrueckii* PYCC5323 were investigated. The results obtained were compared to the ones of a commercial
 baker's strain of *S. cerevisiae*. The following cellular and biochemical parameters were analysed: cell

1 viability, plasma membrane integrity, oxidative damages, intracellular trehalose content and

2 trehalase(s) activity.

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4 **2. Materials and methods**

5 2.1. Microorganisms, growth and freezing conditions

6 The strains used were Torulaspora delbrueckii PYCC5323, isolated from homemade corn and rye 7 bread dough, and Saccharomyces cerevisiae PYCC5325, isolated from commercial compressed 8 baker's yeast - both supplied by the Portuguese Yeast Culture Collection, Faculdade de Ciências e 9 Tecnologia, Universidade Nova de Lisboa, Caparica, Portugal. Stock cultures were maintained on 10 glucose-yeast extract-peptone-agar (2% (wt/vol) glucose, 1% (wt/vol) peptone, 0.5% (wt/vol) yeast 11 extract, 2% (wt/vol) agar), at 4 °C. Yeast strains were grown on yeast extract-peptone-sucrose (YPS) 12 medium containing 2% (wt/vol) sucrose, 4% (wt/vol) peptone, 2% (wt/vol) yeast extract, 0.2% (wt/vol) 13 KH₂PO₄, and 0.1% (wt/vol) MgSO₄.7H₂O, at 30 °C. Peptone (ref. 0118-17) and yeast extract (ref. 14 0127-17) were from Difco (Becton Dickinson, Sparks, MD, USA) and sucrose from Merck (E. Merck, 15 Darmstadt, Germany).

16 For freezing assays, cells were harvested at initial stationary phase (24 hours of culture, 2.4-17 2.7x10⁸ cells ml⁻¹ for *T. delbrueckii* and 1.1-1.5x10⁸ cells ml⁻¹ for *S. cerevisiae*), washed twice with deionised water, and suspended in a quarter of the initial volume in sterile water to an A_{640} of 12-20. 18 19 Aliguots (5 ml) of cells were transferred into 15-ml polycarbonate tubes, centrifuged and the pellet 20 (400-500 mg of wet weight) suspended in 500 µl of the storing medium: LF medium or as described in 21 results. LF medium is a liquid medium formulated to simulate the fermenting ability of yeast in bread 22 dough, with the following composition: 1% (wt/vol) glucose, 1% (wt/vol) sucrose, 3% (wt/vol) maltose, 23 0.25% (wt/vol) (NH₄)₂SO₄, 0.5% (wt/vol) urea, 1.6% (wt/vol) KH₂PO₄, 0.5% (wt/vol) Na₂HPO₄.12 H₂O, 24 0.06% (wt/vol) MgSO₄, 22.5 ppm nicotinic acid, 5.0 ppm pantothenic acid, 2.5 ppm thiamine, 1.25 ppm 25 pyridoxine, 1.0 ppm riboflavin and 0.5 ppm folic acid [22]. The samples were then frozen at -20 °C for 26 different time periods (cooling rate approximately 3 °C min⁻¹) [5], and then thawed at 30 °C for 2 min. 27 For fast freezing, 15-ml polycarbonate tubes containing the cell suspensions were directly immersed in 28 liquid nitrogen (cooling rate approximately 200 °C min⁻¹) [5]. For pre-fermentation treatments, cells 29 were subjected to a fermentation period before freezing as follows: the pellets in the 15-ml polycarbonate tubes were suspended in LF medium to a final OD_{640 nm} of 0.3-0.5, and incubated for 30

120 min, at 30 °C. After this, the suspension was centrifuged and the pellet was suspended in LF
 medium and frozen as described above.

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4 2.2. Extraction and assay of trehalose

5 Cells were harvested by centrifugation, washed twice with cold deionised water and sampled for 6 dry weight contents. Trehalose was extracted from cold cell pellets with 5% (wt/vol) trichloroacetic acid 7 (Merck, Darmstadt, Germany) for 45 min with occasional shaking. Cells were then centrifuged at 735 8 g, for 10 min. Extraction was repeated once more, and supernatants from the two extractions were 9 combined and used for the determination of trehalose by high-performance liquid chromatography. 10 The apparatus used was a Gilson chromatograph (132-RI Detector) equipped with a carbohydrate H^{+} column (SS-100, H⁺, Hypersil) which was maintained at 30 °C. A solution of H₂SO₄ (0.0025 M) was 11 12 used as the mobile phase at a flow rate of 0.45 ml min⁻¹. The relative values (%) of intracellular 13 trehalose concentration after different periods of freezing were calculated by dividing the values of 14 intracellular trehalose concentration obtained for the frozen samples, by those obtained for the 15 unfrozen samples.

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17 2.3. Trehalase activity and protein assay

18 Pellets containing 75-100 mg (wet wt) of washed cells were suspended in 1 ml ice-cold 50 µM 19 MES (4-morpholineethanesulfonic acid) (Boehringer Mannheim, GmbH-Germany)/KOH buffer, pH 7, 20 containing 50 µM CaCl₂. Cells were broken by vortexing with 500 µl of glass beads (0.5 mm diameter), 21 for four periods of 1 min, with 1 min intervals on ice between them. The crude enzyme extract was 22 centrifuged for 3 min at 13 200 g, at 4 °C. The supernatant was dialysed overnight at 4 °C against 10 23 mM Mes/KOH buffer, pH 7, containing 50 µM CaCl₂. Trehalase was assayed as described previously 24 [23]. The glucose liberated was determined by glucose oxidase/peroxidase method (Glucose GOD -25 Perid, Boehringer Mannheim, GmbH-Germany). Protein determination was carried out according to 26 Lowry et al. [24]. Specific activity of trehalase was expressed as units (U, nmol glucose released per 27 min) per mg protein.

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29 2.4. Measurement of cell viability and membrane integrity

30 The viability of yeast cells was determined by counting CFU. For this assay, yeast cell

31 suspensions were washed twice with deionised water and, after convenient dilution, spread on YPDA

1 medium plates. The plates were incubated for 48 hours at 30 °C before counting. The relative values 2 (%) of viable cells after different periods of freezing were calculated by dividing the values of CFU 3 counts obtained for the frozen samples, by those obtained for the unfrozen samples. 4 Membrane integrity was analysed by flow cytometry using the membrane exclusion dye, 5 propidium iodide (PI). In these assays, cells with preserved membrane integrity are not permeated by 6 propidium iodide (PI⁻ cells) while those that lost their membrane integrity do incorporate the 7 fluorochrome (PI⁺ cells) [25]. Cell suspensions (about 10⁷ cells ml⁻¹) were incubated for 10 min in the 8 dark with a 20 µg/ml PI solution (ratio of 2:1, respectively) and injected on a Partec PAS III flow 9 cytometer equipped with an argon-ion laser emitting a 15 mW beam at 488 nm . From each sample 10 2x10⁴ cells were analysed. Control suspensions of membrane-disrupted cells were prepared by boiling 11 cell suspensions. The relative values (%) of PI⁻ cells after different periods of freezing were calculated 12 by dividing the values of PI cells obtained for the frozen samples, by those obtained for the unfrozen 13 samples.

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16 2.5. Oxidative stress evaluation

For evaluating the effect of pre-treatment with the radical scavenger N-tert-butyl-α-phenylnitrone (PBN) (Aldrich Chem. Co. Milwaukee, WI 53201), cells were grown for 24 hours to initial stationary phase in YPS medium at 30 °C. PBN was added to the culture at the final concentrations of 0.5, 5.0 and 15 mM, and cells were cultured for 30 min before being harvested.. PBN was removed by washing cells twice with deionised water and cells were frozen in LF medium as described above. In assays where a pre-fermentation was performed, PBN was added to the fermenting cell suspensions 30 min before the end of the fermentation period.

For treatment with oxidising agents, cells were grown to mid-exponential (OD_{640 nm} 0.5-1.0) and
initial stationary phase (24 hours of culture), harvested and washed twice with deionised water.
Subsequently, these cells were suspended in water in order to achieve a concentration of OD_{640nm} 0.60.7, and 0.1 ml of this suspension was spread on solid YPDA plates (50 mm diameter). A paper disc
(6 mm diameter - BBL, 231039, Becton Dickinson, Sparks, MD, USA) containing 10 μl of one oxidising
agent, was laid on each inoculated plate. The oxidants used were: menadione at concentrations of
0.05, 0.10, 1.0 mM; diamide at concentrations of 0.3, 3.0 M; hydrogen peroxide at 1.1 and 11 M, all

purchased from Sigma Chemical Co (St. Louis, MO). The plates were incubated at 26 °C during two
 days, after which the diameters of the inhibition halos around the paper disks were measured. The
 value obtained was the average of two perpendicular diameters, excluding the disc diameter [26].

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5 2.6. Thiobarbituric acid (TBA) reaction for lipid peroxide analysis

For TBA-reactive substances (TBARS) quantification, pellets containing 350-400 mg (wet weight)
of cells were washed with ice-cold deionised water and suspended in 0.75 ml ice-cold sodium
phosphate (E. Merck, Darmstadt, Germany) buffer, pH 7.2. Cells were broken by vortexing with 500 μl
of glass beads (0.5 mm diameter) for six periods of 1 min with 1 min intervals on ice between them.
TBARS were then determined according to Buege and Aust [27]. The TBARS concentration was
expressed as μmol malondialdehyde per mg protein.
2.7. Reproducibility of the results

All the experiments were repeated at least three times, and the data reported are mean values and SD. When statistical analyses were performed, the significance was tested by analysis of variance (Anova, Microsoft Excel 2000).

17

18 **3. Results**

19 3.1. Cell viability and membrane integrity along freeze storage

20 To characterise the freeze resistance of T. delbrueckii PYCC5323, we first studied cell viability, 21 assessed as colony-forming units (CFU), during freezing at -20 °C for up to 120 days as described in 22 materials and methods (Fig. 1A). For a comparative analysis, a commercial baker's yeast, S. 23 cerevisiae PYCC5325, was used as a reference strain (Fig. 1B). In T. delbrueckii no loss of cell 24 viability was observed for the entire storage period, whereas a loss of 80% was obtained in S. 25 cerevisiae after 15 days. In both species, viability loss was faster when cells were stored in freezing 26 medium without sugars, even when glycerol was added to keep the osmotic pressure. These results 27 are consistent with previous reports showing that the presence of the disaccharide trehalose in the 28 extracellular medium has a protective effect in cell viability during freezing [28]. 29 To elucidate whether the loss of membrane integrity along freezing was directly conditioning cell

30 viability, membrane damage was monitored by flow cytometry using propidium iodide (PI) cell staining.

The decrease of cells with preserved plasma membrane (PI⁻ cells) was much less pronounced in *T*. *delbrueckii* (Fig. 2 A) than in *S. cerevisiae*. As shown in Fig. 2B, there was a direct correlation between the percentage of PI⁻ cells and the percentage of CFU counts for both *T. delbruekii* and *S. cerevisiae* yeast strains. This was evident either in cells subjected to a short fermentation period (120 min) or not subjected to fermentation before freezing. The results show that, independently of the physiological state of the cells, membrane integrity is directly conditioning cell viability, expressed as CFU counts, and therefore the plasma membrane seems to be one of the first freezing targets.

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9 3.2. Cell oxidative stress responses during the freeze-thaw process

10 The results described above indicate that T. delbrueckii displays a higher freeze resistance when 11 compared with S. cerevisiae, which is mainly due to plasma membrane integrity. To examine whether 12 this capacity to preserve plasma membrane integrity was correlated with oxidative stress resistance, 13 we directly assessed oxidative damage in the membranes of both T. delbrueckii and S. cerevisiae 14 during frozen storage by measuring TBA-reactive substances (TBARS). The TBARS test quantifies 15 lipid peroxides in the thiobarbituric acid derivatized form. As shown in Table 1, for both yeast strains, 16 frozen cells presented higher amounts of products of lipid peroxidation than control cells (cells before 17 freezing). However, the percentage increase in TBARS levels was significantly enhanced (P < 0.05) 18 much earlier in S. cerevisiae (approximately 61% for the fifth day), which was associated to its rapid 19 decrease in cell membrane integrity. As shown in the previous section, the presence of glycerol did 20 not protect cells of T. delbrueckii during freezing. Therefore we tested T. delbrueckii under these 21 conditions to avaliate the increase in the percentage of TBARS. The results showed that the values of 22 percentage increase of TBARS in T. delbrueckii frozen in a medium with glycerol, for 60 and 84 days, 23 were similar to those observed for S. cerevisiae (respectively 64% and 83%). Moreover, samples of T. 24 delbrueckii frozen under conditions where no loss of cell viability was observed, presented much lower 25 percentage increase in TBARS production (Fig. 1, Table 1).

The role of oxidative stress during freezing, was examined by using the oxygen radical scavenger PBN. In *S. cerevisiae* PYCC5325, pre-incubation of cells with PBN (1 or 5 mM) resulted in higher levels of membrane integrity preservation for the first days of frozen storage, but no protective effect by PBN was observed in cells frozen after a period of pre-fermentation (results not shown). This result might reflect the different physiological status of the cells and it is conceivable that oxidative stress is

1being carried over by other factors [17]. Next, the oxidative stress responses of both strains to2hydrogen peroxide (H₂O₂, toxicity mainly due to hydroxyl radicals), menadione (a superoxide-3generating agent) and diamide (thiol-oxidizing drug) were evaluated. The results obtained in a disk4diffusion assay are shown in Fig. 3. *T. delbrueckii* was more sensitive to menadione (P < 0.001) and5diamide (P < 0.05) while more tolerant to H₂O₂ (P < 0.01) when compared with *S. cerevisiae*.6Therefore, like it was previously reported for *S. cerevisiae*, in *T. delbrueckii* freeze resistance is also7correlated with H₂O₂ resistance [5].

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9 3.3. Intracellular trehalose content during the storage period

10 The results described above pointed to the possibility that oxidative stress was not the only 11 condition influencing the cell freeze resistance, mainly in cells frozen after a short period of pre-12 fermentation. Intracellular trehalose accumulation has been described to protect cells from oxygen 13 radicals and also to be involved in the stabilisation of the plasma membrane structure during freezing 14 [29-31]. Hence, it was also investigated whether trehalose was involved on the higher capacity of T. 15 delbrueckii to maintain plasma membrane integrity compared to S. cerevisiae. Immediately before 16 freezing, the values of the intracellular trehalose content were high and similar in both species: 109.34 17 \pm 8.72 and 112.79 \pm 20.97 mg g⁻¹ dry wt for S. cerevisiae and T. delbrueckii, respectively. During the 18 entire freezing storage period, these values were kept high and constant in T. delbrueckii but 19 decreased quickly in S. cerevisiae. For both species the relative values of the intracellular trehalose 20 content during freezing followed closely the percentage of PI cells (Fig. 4). In S. cerevisiae, the 21 decrease in the intracellular trehalose content was accompanied by an increase of the extracellular 22 trehalose content and of the medium OD_{260nm} (used as a measure of leakage of cell contents) 23 (Fig.4B). At the end of the assay, the amount of trehalose found in the media was about the same as 24 the one lost from the cells, indicating that total trehalose amount (intracellular + extracellular) remained 25 unchanged. Thus, cell leakage appears to be responsible for the decrease in trehalose content of the 26 cells, which is consistent with the observed loss of plasma membrane integrity. In addition, a decrease 27 in the intracellular trehalose content was observed in cells that were subjected to a short fermentation 28 period before freezing (cells more sensitive to freeze stress), which was similar for both strains (from 29 109.34 ± 8.72 to 9.79 ± 0.93 mg g⁻¹ dry wt for *S. cerevisiae* and from 112.79 ± 20.97 to 17.16 ± 6.24 mg g^{-1} dry wt for *T. delbrueckii*, during a 120 min fermentation period). 30

1 These results seemed to indicate that a similar activation pattern of trehalase(s) by glucose was 2 present in both yeast strains. To address this point, trehalase(s) activities were monitored in cell free 3 extracts prepared from cells in the absence of glucose (stationary phase cells) as well as after a 4 glucose pulse. A similar behaviour was presented by both strains with a two to three-fold increase in 5 activity almost immediately after glucose addition (from 15.0 ± 0.4 to 32.1 ± 1.3 mU/mg protein for S. 6 cerevisiae and from 11.1 ± 0.2 to 31.0 ± 0.7 mU/mg protein for T. delbrueckii, respectively before and 7 after glucose addition). The results are in agreement with the mobilisation of trehalose observed 8 during the pre-fermentation period.

- 9 The results described above are consistent with the conclusion that the higher freeze resistance 10 displayed by *T. delbrueckii* can not be attributed to higher intracellular trehalose contents.
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12 3.4. Cell adaptation during the freeze period

13 We evaluated the ability of T. delbrueckii and S. cerevisiae to adapt to freezing by inhibiting 14 protein synthesis with cycloheximide. The results obtained when cells were frozen at - 20 °C, showed 15 that for T. delbrueckii, contrasting with S. cerevisiae, the presence of cycloheximide in the freezing 16 medium increased the loss of cell viability throughout frozen storage (Fig. 1). To assess if 17 cycloheximide could have a toxic effect in T. delbrueckii cells, being responsible for the observed 18 decrease in cell viability, a cell suspension of this yeast was incubated in LF medium with 19 cycloheximide, and the number of CFU counts was determined without previous freezing the 20 suspension. No differences were found between the number of CFU counts estimated for this cell 21 suspension and for the control without cycloheximide. These results indicated that at the concentration 22 tested, cycloheximide was not having a toxic effect. In addition, when cells of T. delbrueckii were 23 frozen at a much faster rate, in liquid nitrogen (-196 °C), and subsequent storage at -80°C, loss of cell 24 viability rapidly increased, CFU counts dropping aproximatly 50% after one day of frozen storage. 25 Together these results suggest that cell viability of T. delbrueckii is dependent on de novo protein 26 synthesis, and therefore that yeast cells can adapt in a slow freezing process, most probably during 27 the initial cooling period. In accordance with these findings, the capacity of T. delbrueckii cells to adapt 28 might be essential for the yeast high freeze tolerance phenotype.

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1 4. Discussion

2 Our findings clearly indicate that the strain T. delbrueckii PYCC5323 exhibits a high freeze 3 tolerance, when compared with the baker's yeast S. cerevisiae, which reinforces previous reports 4 claiming for its useful exploitation in baking industry. Evidence was presented that the primary cell 5 target of freezing stress is the plasma membrane and that the capacity to preserve membrane integrity 6 displayed by T. delbrueckii PYCC5323 is correlated with higher resistance to lipid oxidative damage. 7 Hydroxyl radicals appear to be the agents responsible for cell membrane damage in freeze stress in 8 this yeast. This is the opposite of a previous report on S. cerevisiae where superoxide radicals were 9 considered as the agents responsible for cell damage in these stress conditions [18]. To ascertain the 10 key role of these radicals during freezing we are developing a catalase null mutant in T. delbrueckii 11 PYCC5323. Loss of cell viability seems to be correlated with the percentage increase in TBARS levels 12 and not with their absolute values, suggesting that a higher content in unsaturated fatty acids could 13 allow the cell to cope with higher absolute levels of lipid oxidation without compromising membrane 14 integrity. This hypothesis would also agree with previous work, showing that freeze-tolerant yeast 15 strains have larger amounts of unsaturated fatty acids when compared with sensitive strains [32]. 16 Previous studies have pointed to a lack of capacity in S. cerevisiae to adapt to cold stress [17], 17 although evidence for cold-induced expression changes associated with improved cryoresistance has 18 also been provided more recently [33, 34]. According to our results, in T. delbrueckii, contrasting with 19 S. cerevisiae, the surviving capacity (evaluated by cell viability) is dependent on de novo protein 20 synthesis. An adaptation process during slow freezing appears to be determinant for the yeast high 21 freeze tolerance phenotype. In the light of these observations, the yeast response to freeze stress 22 seems to be strongly dependent on the yeast strain, culture and freeze conditions. However, further 23 studies will be necessary to clarify the molecular basis of cell adaptation to cold/freeze. In the case of 24 the strain T. delbrueckii PYCC5323, the observed behaviour is particularly relevant in view to its 25 utilisation for frozen dough production, and implies that the dough should be frozen at a slow, rather 26 than at a fast rate.

In addition and noteworthy from a methodological point of view, the results regarding the
correlation observed along freezing between the loss of membrane integrity and cell proliferative
capacity, validate the application of flow cytometry and the use of the fluorochrome PI as a measure of
viability of cells subjected to freeze stress. Therefore, and contrary to other stress conditions [25], the

1 assessment of PI⁻ cells by flow cytometry, as a method to determine cell viability either in T. 2 delbrueckii or S. cerevisiae, can replace the more laborious and time consuming determination of CFU 3 counts. 4 5 Acknowledgements 6 We are grateful to Professor Victor Costa for laboratory support and for critical reading of the 7 manuscript. Cecília Alves-Araújo was supported by a PhD grant (PRAXIS XXI/BD/21543/99) from 8 Fundação para a Ciência e a Tecnologia, Portugal. 9 10 References 11 [1] Randez-Gil, F., Sanz, P. and Prieto, J.A. (1999) Engineering baker's yeast: room for improvement. 12 TIBTECH 17, 237-244. 13 [2] Randez-Gil, F., Aguilera, J., Codón, A., Rincón, A. M., Estruch, F. and Prieto, J. A. (2003) 14 Baker's yeast: challenges and future prospects. In: Functional Genetics of Industrial Yeasts, De 15 Winde JH (ed). Springer-Verlag, Heidelberg; 57-97. 16 [3] Attfield, P.V., Raman, A. and Northcott, C. (1992) Construction of Saccharomyces cerevisiae 17 strains that accumulate relatively low concentrations of trehalose, and their application in testing 18 the contribution of the disaccharide to stress tolerance. FEMS Microbiol. Lett. 94, 271-276. 19 [4] Yokoigawa, K., Murakami, Y. and Kawai, H. (1995) Trehalase activity and trehalose content in a 20 freeze-tolerant yeast, Torulaspora delbrueckii, and its freeze-sensitive mutant. Biosci. Biotechnol. 21 Biochem. 59(11), 2143-2145. 22 [5] Lewis, J.G., Learmonth, R.P., Attfield, P.V. and Watson, K. (1997) Stress co-tolerance and 23 trehalose content in baking strains of Saccharomyces cerevisiae. J. Ind. Microbiol. Biotechnol. 18, 24 30-36. 25 [6] Sano, F., Asakawa, N., Inoue, Y. and Sakurai, M. (1999) A dual role for intracellular trehalose in 26 the resistance of yeast cells to water stress. Cryobiology 39, 80-87. 27 [7] Hino, A., Tacano, H. and Tanaka, Y. (1987) New freeze-tolerant yeast for frozen dough 28 preparations. Cereal Chem. 64(4), 269-275.

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Fig. 1. Relative values (%) of colony forming units (CFU) in stationary growth phase cells of *T*. *delbrueckii* (A), and *S. cerevisiae* (B) during frozen storage at –20 °C, for 120 and 30 days, respectively. Cells were grown in YPS medium during 24 hours at 30 °C, harvested, washed with water and suspended in different storage media: (□) LF medium; (●) LF medium + 0.01% cycloheximide; (○) LF medium without sugars + 0.16 M glycerol; (△) Sugars solution (1% glucose + 1% sucrose + 3% maltose).

Fig. 2. Relative values (%) of negative propidium iodide cells (PI⁻ cells) of *T. delbrueckii* (close symbols) and *S. cerevisiae* (open symbols) during storage at -20 °C (A) and correlation between relative values (%) of PI⁻ cells and colony forming units (CFU) (B). Cells grown in YPS medium during 24 hours at 30 °C (initial stationary growth phase), harvested, washed with water and suspended in LF medium were frozen directly (\blacktriangle , Δ) or after a pre-fermentation period of 120 min (\bullet , \odot), as described in material methods. In Fig. 2A, for some results error bars are within the data point labels.

Fig. 3. Effect of oxidising agents on the radial growth of *T. delbrueckii* (black bars) and *S. cerevisiae* (open bars). Cells were grown in YPS medium at 30 °C, to mid-exponential growth phase (B) or to initial stationary growth phase (A), and 0.1 ml of a suspension (OD_{640} nm 0.6-0.7) of these cells was spread on YPDA plates. A paper disc, 6 mm diameter, containing 10 µl of one oxidising agent was laid on each inoculated plate. The plates were incubated during two days, after which the diameters of the inhibition halos were measured. The values shown are those obtained for the highest concentration of oxidant tested.

Fig. 4. Cells of *T. delbrueckii* (A) and *S. cerevisiae* (B) were grown in YPS medium during 24 hours at 30 °C, harvested, washed with water, suspended LF media and frozen at -20 °C. During freezing samples were taken and assessed for: (\bigcirc) relative values (%) of negative propidium iodide cells (PI⁻ cells); (\blacktriangle) relative values (%) of intracellular trehalose concentration (T_{in}); (\square) extracellular trehalose concentration (T_{out} , g/l); (\blacklozenge) medium optical density at 260 nm, used as a measure of leakage of cell contents (OD_{260 nm}) The relative values (%) were estimated as described in material and methods.