Evaluation of stress tolerance and fermentative behavior of indigenous \textit{Saccharomyces cerevisiae} \\

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Abstract

Sixty six indigenous \textit{Saccharomyces cerevisiae} strains were evaluated in stressful conditions (temperature, osmolarity, sulphite and ethanol tolerance) and also ability to flocculate. Eighteen strains showed tolerant characteristics to these stressful conditions, growing at 42 °C, in 0.04% sulphite, 1 mol L\textsuperscript{-1} NaCl and 12% ethanol. No flocculent characteristics were observed. These strains were evaluated according to their fermentative performance in sugar cane juice. The conversion factors of substrates into ethanol ($Y_{p/s}$), glycerol ($Y_{g/s}$) and acetic acid ($Y_{ac/s}$) were calculated. The highest values of $Y_{p/s}$ in sugar cane juice fermentation were obtained by four strains, one isolated from fruit (0.46) and the others from sugar cane (0.45, 0.44 and 0.43). These values were higher than the value obtained using traditional yeast (0.38) currently employed in the Brazilian bioethanol industry. The parameters $Y_{g/s}$ and $Y_{ac/s}$ were low for all strains. The UFLA FW221 presented the higher values for parameter related to bioethanol production. Thus, it was tested in co-culture with \textit{Lactobacillus fermentum}. Besides this, a 20-L vessel for five consecutive batches of fermentation was performed. This strain was genetically stable and remained viable during all batches, producing high amounts of ethanol. The UFLA FW221 isolated from fruit was suitable to produce bioethanol in sugar cane juice. Therefore, the study of the biodiversity of yeasts from different environmental can reveal strains with desired characteristics to industrial applications.

Key words: alcoholic fermentation, biofuel, fermentation kinetics, UFLA FW221, \textit{Saccharomyces cerevisiae}.

Introduction

Carbon dioxide (CO\textsubscript{2}) released from the burning of fossil fuels, such as petroleum and coal, is considered a major contributor to global warming. There is consequently a need for alternative, carbon-neutral energy sources. Bioethanol, a renewable fuel, can be considered an alternative to fossil fuel use (Cardona and Sánchez, 2007).

Ethanol production by microorganisms has received global attention because it can be obtained from renewable sources. Many new ethanol plants are being built to increase supply, and researchers are investigating ways to improve the yield of ethanol production (Antoni \textit{et al.}, 2007).

In the United States, the most ethanol is produced from corn, while Brazil mainly produces ethanol from sugar cane (Basso \textit{et al.}, 2008). Sugar cane is used as a substrate, and ethanol concentrations of 8-11% (v/v) are achieved within a period of 6-11 h at 30-35 °C. After fermentation, yeast cells are recycled during a production season of 200-250 days (Wheals \textit{et al.}, 1999).

The Brazilian bioethanol industry has grown in the last few decades as a result of the international oil crisis. Sugar cane crop productivity has increased due to genetic improvements in cultivars and ongoing research to increase the efficiency of fermentative processes (Cardona and Sánchez, 2007). Scientific and technological advances, regard-
ing sugar cane varieties, agricultural and fermentation pro-
cess management and engineering, for example, have led to
an increase in the efficiency of Brazilian bioethanol distill-
eries. The selection of new *Saccharomyces cerevisiae*
strains could be a suitable way to amplify fuel production
on an industrial scale (Basso et al., 2008). Industrial fer-
mentation processes impose multiple stressful conditions
(e.g., temperature, ethanol concentration, osmotic pressure
and ionic stress) on yeast that affect its performance and ki-
etics during alcoholic fermentation (Fleet, 2008). Changes
in temperature are by far the most studied stress inducers in
living cells (Aguilera et al., 2007; Babiker et al., 2010).
*Saccharomyces cerevisiae* has been chosen over the centu-
 ries because it is physiologically adapted to these unfavor-
able conditions (Attfield, 1997). The present work pro-
poses to study the behaviour of indigenous *S. cerevisiae* in
stressful conditions and selected strains were subjected to
sugar cane fermentation in order to study the associated ki-
etics parameters. Finally, the selected strain was tested in
col-culture with *Lactobacillus fermentum* to simulate a con-
taminated fermentation and also batch fermentations in
sugar cane juice on a semi-industrial scale were performed.

**Material and Methods**

**Yeast strains**

The indigenous yeast strains used in this study were
isolated from bioethanol distilleries (sugar cane fermenta-
tions) and fruit wine fermentations and belong to the micro-
bial collection of the Microbial Physiology Laboratory/De-
partment of Biology /Federal University of Lavras (UFLA), Brazil. The strain PE-2 (commercialized in Brazil
for bioethanol production) was used as a reference strain. A
total of 66 yeast strains were studied (Table 1).

### Table 1 - *Saccharomyces cerevisiae* strains studied and their respective sources.

| *S. cerevisiae* strain | Source                  | *S. cerevisiae* strain | Source                  |
|------------------------|-------------------------|------------------------|-------------------------|
| UFLA CA751             | Distillery (UFLA/Brazil)| UFLA CA776             | Distillery (UFLA/Brazil)|
| UFLA CA752             | Distillery (UFLA/Brazil)| UFLA CA777             | Distillery (UFLA/Brazil)|
| UFLA CA753             | Distillery (UFLA/Brazil)| UFLA CA778             | Distillery (UFLA/Brazil)|
| UFLA CA754             | Distillery (UFLA/Brazil)| UFLA CA779             | Distillery (UFLA/Brazil)|
| UFLA CA755             | Distillery (UFLA/Brazil)| UFLA CA780             | Distillery (UFLA/Brazil)|
| UFLA CA756             | Distillery (UFLA/Brazil)| UFLA CA781             | Distillery (UFLA/Brazil)|
| UFLA CA757             | Distillery (UFLA/Brazil)| UFLA CA782             | Distillery (UFLA/Brazil)|
| UFLA CA155             | Distillery (UFLA/Brazil)| UFLA CA783             | Distillery (UFLA/Brazil)|
| UFLA CA758             | Distillery (UFLA/Brazil)| UFLA CA784             | Distillery (UFLA/Brazil)|
| UFLA CA759             | Distillery (UFLA/Brazil)| UFLA CA785             | Distillery (UFLA/Brazil)|
| UFLA CA760             | Distillery (UFLA/Brazil)| UFLA CA786             | Distillery (UFLA/Brazil)|
| UFLA CA493             | Distillery (UFLA/Brazil)| UFLA CA787             | Distillery (UFLA/Brazil)|
| UFLA CA761             | Distillery (UFLA/Brazil)| UFLA CA788             | Distillery (UFLA/Brazil)|
| UFLA CA762             | Distillery (UFLA/Brazil)| UFLA CA789             | Distillery (UFLA/Brazil)|
| UFLA CA15              | Distillery (UFLA/Brazil)| UFLA CA790             | Distillery (UFLA/Brazil)|
| UFLA CA76              | Distillery (UFLA/Brazil)| UFLA CA791             | Distillery (UFLA/Brazil)|
| UFLA CA116             | Distillery (UFLA/Brazil)| UFLA CA792             | Distillery (UFLA/Brazil)|
| UFLA FW45              | Fruit wine (UFLA/Brazil)| UFLA CA793             | Distillery (UFLA/Brazil)|
| UFLA FW221             | Fruit wine (UFLA/Brazil)| UFLA CA794             | Distillery (UFLA/Brazil)|
| UFLA FW510             | Fruit wine (UFLA/Brazil)| UFLA CA795             | Distillery (UFLA/Brazil)|
| UFLA CA763             | Distillery (UFLA/Brazil)| UFLA CA796             | Distillery (UFLA/Brazil)|
| UFLA CA764             | Distillery (UFLA/Brazil)| UFLA CA797             | Distillery (UFLA/Brazil)|
| UFLA CA765             | Distillery (UFLA/Brazil)| UFLA CA798             | Distillery (UFLA/Brazil)|
| UFLA CA766             | Distillery (UFLA/Brazil)| UFLA CA799             | Distillery (UFLA/Brazil)|
| UFLA CA767             | Distillery (UFLA/Brazil)| UFLA CA800             | Distillery (UFLA/Brazil)|
| UFLA CA768             | Distillery (UFLA/Brazil)| UFLA CA801             | Distillery (UFLA/Brazil)|
| UFLA CA769             | Distillery (UFLA/Brazil)| UFLA CA802             | Distillery (UFLA/Brazil)|
| UFLA CA770             | Distillery (UFLA/Brazil)| UFLA CA803             | Distillery (UFLA/Brazil)|
| UFLA CA771             | Distillery (UFLA/Brazil)| UFLA CA804             | Distillery (UFLA/Brazil)|
| UFLA CA772             | Distillery (UFLA/Brazil)| UFLA CCA 022           | Distillery (UFLA/Brazil)|
| UFLA CA773             | Distillery (UFLA/Brazil)| UFLA CCA 035           | Distillery (UFLA/Brazil)|
| UFLA CA774             | Distillery (UFLA/Brazil)| UFLA CCA 083           | Distillery (UFLA/Brazil)|
| UFLA CA775             | Distillery (UFLA/Brazil)| UFLA CCA 385           | Distillery (UFLA/Brazil)|
Screening for stressful conditions

Temperature, ethanol, sulfite and osmolarity tolerance

The temperature tolerance test was performed according to Breisha (2010) at 30, 37, and 42 °C. Tolerance to ethanol concentrations was evaluated according to Breisha (2010) at 0, 6, 12, and 18% ethanol. The sulfite tolerance test was carried out according to Valles et al. (2008) in 0.02% and 0.04% (w/v) sodium metabisulfite. Osmolarity tolerance was observed after incubation of the strains for 72 h at 30 °C in YPD medium (10 g L⁻¹ Yeast extract; 20 g L⁻¹ Peptone; 20 g L⁻¹ dextrose; 14 g L⁻¹ agar) supplemented with 0, 0.5, 0.7 and 1 mol L⁻¹ NaCl according to Rep et al. (2000) with modifications. Tolerance to NaCl concentrations was observed by measuring colony size.

Flocculation evaluation by spectrophotometry

Flocculation was determined according to Valles et al. (2008) with some modifications. The yeast strains were inoculated in 5 mL of YPD broth and incubated at 30 °C for 72 h. After incubation, the cell cultures were centrifuged and the cells were resuspended in 5 mL of Helm’s buffer (3 m mol L⁻¹ calcium chloride, 50 m mol L⁻¹ acetate-acetic buffer, pH 4.5). The degree of flocculation of the different strains was determined in terms of the ratio between the optical density at 620 nm of the culture suspension and that obtained 10 min after the Helm’s buffer was added (ODₓ/OD₀ x 100). The following flocculation scale was established: ratio > 90% (no flocculence); ratio between 70% and 90% (low flocculence); ratio between 30% and 70% (medium flocculence); and ratio < 30% (high flocculence).

Screening of S. cerevisiae strains for ethanol and glycerol production

Cells were grown in YPS (10 g L⁻¹ Yeast extract; 20 g L⁻¹ Peptone; 20 g L⁻¹ Sucrose) at 30 °C. After 36 h, cells corresponding to an optical density of 2.0 at 600 nm were used to inoculate (5% v/v) the ethanol production medium (40 g L⁻¹ sucrose; 5 g L⁻¹ yeast extract; 20 g L⁻¹ peptone; 0.5 g L⁻¹ KH₂PO₄; 0.5 g L⁻¹ (NH₄)₂SO₄; 0.5 g L⁻¹ MgSO₄7H₂O). The flasks were incubated at 30 °C for 5 days. The ethanol and glycerol concentrations were evaluated by high-performance liquid chromatography (HPLC), according to Duarte et al. (2009).

Performance of S. cerevisiae strains during alcoholic fermentation

Fermentation assays

The yeast isolates were grown in 250-mL flasks containing 100 mL of YPD medium for 24 h at 30 °C and 200 rpm. After measuring the absorbance at 600 nm, the cell suspension volume was adjusted to obtain an inoculum corresponding to an optical density of 2.0 (approximately 1.0 x 10⁸ cells mL⁻¹) and were inoculated in 100 mL of sugar cane juice at 18 ºBrix and incubated at 30 °C without agitation. The experiment was conducted in duplicate. Samples were collected at intervals of 4 h to determine the concentrations of residual sugar, acetic acid, glycerol and ethanol produced.

Chemical analysis

Sugar contents (sucrose, glucose and fructose), organic acids (acetic acid and succinic acid), glycerol, methanol and ethanol were quantified by HPLC, according to Duarte et al. (2009).

Calibration curves

Yeast strains were grown in 250-mL flasks containing 100 mL of YPD broth for 24 h at 30 °C and 200 rpm. After the 24-h incubation, the yeast cells were centrifuged (6000 g) for 5 min at 20 °C and washed twice with sterile peptone water. The biomass obtained was inoculated into 100 mL of sugar cane juice at 18 °Brix and incubated at 30 °C for 36 h without agitation. After growth, the cells were recovered by centrifugation (6000 g) for 5 min at 20 °C and washed twice with sterile peptone water. The biomass was then resuspended in 30 mL of sterile peptone water (0.1%), of which 15 mL was used to determine the dry weight at 105 °C after 24 h. The remaining 15 mL was used for serial dilutions to determine the absorbance at 600 nm. Calibration curves were obtained by plotting the absorbance values against the dry weight values. The calibration curves were used to determine the initial inoculum concentration and to monitor yeast growth during the fermentation process.

Evaluation of the co-incubation of S. cerevisiae UFLA FW221 and L. fermentum during alcoholic fermentation

The fermentation was performed as previously described. The L. fermentum isolated from distillery belonging to the culture collection of the Microbial Physiology Laboratory/Department of Biology/Federal University of Lavras (UFLA), Brazil was employed in this study. Flasks containing 100 mL of sugarcane juice at 18 °Brix were inoculated separately with (1) L. fermentum (10⁶ cfu mL⁻¹), (2) S. cerevisiae UFLA FW221 (10⁶ cfu mL⁻¹) and (3) with a mixed culture of L. fermentum and S. cerevisiae UFLA FW221 (10⁶ cfu mL⁻¹ bacteria; 10⁸ cfu mL⁻¹ yeast). Samples were microbiological analyzed by plating in YPD medium for yeast and Man Rogosa Sharpe (MRS, Merck, Darmstadt, Germany) medium for bacteria, and by HPLC (Duarte et al., 2009) for chemical contents.

Batch scale fermentation

Fermentation using the selected strain S. cerevisiae UFLA FW221 was carried out in 20-L stainless steel vats during five consecutive batches. The selected S. cerevisiae strain was grown in 1 mL YPD broth at 30 °C on an orbital
Evaluation of the fermentation performance

To determine the fermentation performance, conversion factors were used for calculating the conversion of substrates (g g⁻¹) into ethanol (Yp/s), biomass (Yx/s), methanol (Ym/s), glycerol (Yg/s), succinic acid (Yac/s) and acetic acid (Yac/s), and also volumetric productivity (g L⁻¹ h⁻¹) of ethanol (Qp), biomass productivity (g g⁻¹ h⁻¹) (Pp), conversion efficiency (%) (Ep) and percentage of conversion relative to the theoretical value (conversion (%)) (Duarte et al., 2010).

The equations used in this work are presented below:

\[ Y_{p/s} = \left(\frac{P_f - P_i}{S_i - S_f}\right) / (S_i - S_f); \]
\[ Y_{x/s} = \left(\frac{X_f - X_i}{S_i - S_f}\right); \]
\[ Y_{m/s} = (M_f - M_i) / (S_i - S_f); \]
\[ Y_{ac/s} = (Ac_f - Ac_i) / (S_i - S_f); \]
\[ Y_{g/s} = (g_f - g_i) / (S_i - S_f); \]
\[ P_p = (P_f - P_i) / t_f; \]
\[ P_x = (X_f - X_i) / t_f; \]
\[ E_p = \left(\frac{Y_{p/s}}{0.51}\right) \times 100; \]
\[ \text{conversion} = \left(\frac{Y_{p/s}}{0.51}\right) / 100; \]

where \( P_i \) is the initial concentration of ethanol, \( P_f \) is the ethanol concentration at the end of fermentation, \( S_i \) is the initial substrate concentration, \( S_f \) is the substrate concentration at the end of fermentation, \( X_i \) is the initial biomass concentration, \( X_f \) is the biomass concentration at the end of fermentation, \( g_i \) is the initial glycerol concentration, \( g_f \) is the glycerol concentration at the end of fermentation, \( Ac_i \) is the initial acetic acid concentration, \( Ac_f \) is the concentration of acetic acid at the end of fermentation, \( M_i \) is the initial methanol concentration, \( M_f \) is the concentration of methanol at the end of fermentation, \( P_p \) is the productivity of ethanol, \( P_x \) is the productivity of biomass, \( E_p \) is the conversion efficiency, and \( t_f \) is the total time of fermentation.

Statistical analysis

Cluster analysis was performed by using the software Statistica for Windows, version 6.0 (Statsoft Inc., Tulsa, OK, USA) The binary matrix was constructed with the results of growth (+) or no growth (-) for each level of treatment according to methods described above, regarding ethanol and glycerol screening the production of ethanol higher than 10 g L⁻¹ and glycerol less than 3 g L⁻¹ were considered positive (+) in the binary matrix. Principal component analyses (PCA) were performed using the XLSTAT 7.5.2 software (Addinsoft’s, New York, NY, USA). CO2 production (dCO₂/dt) was calculated using the Origin Pro 8.0 software (OriginLab, Northampton, MA, USA). Analyses of the variance by Scott-Knott test were performed with SISVAR 5.1 software (Ferreira, 2008).

Results

Screening of S. cerevisiae for stress tolerance and ethanol production

Figure 1 shows the grouping of yeasts according to their behavior under different growth conditions, such as different temperatures (30, 37 and 42 °C), NaCl concentrations (0.5, 0.7 and 1 mol L⁻¹), sulfite concentrations (0.02

![Dendrogram obtained by Ward’s hierarchical clustering method performed for different growth conditions (temperature, osmolarity, sulfite and ethanol tolerance and flocculation capacity). G1 = most resistant strains (Growth at 42 °C, in 1 M NaCl, 0.04% (w/v) sulfite and tolerated 12% ethanol. They were not flocculent) and G2 = most sensitive strains.](image)
and 0.04%, w/v), ethanol concentrations (0, 6, 12 and 18%) and flocculation capacity. The 66 yeast strains were classified into two large groups (G1 and G2). The G1 group contains the yeast strains that were able to support the most stressful conditions. The S. cerevisiae strains included in this group were able to grow at 42 °C, in 1 M NaCl, 0.04% (w/v) sulfite and tolerated 12% ethanol. There were no flocculant strains among those in G1. These results demonstrate that these strains were well adapted to adverse conditions.

Ethanol production was also evaluated during the yeast selection process. Strains of Saccharomyces that were resistant for the most stressful conditions and produced more than 10 g L⁻¹ of ethanol were selected to study the fermentation behavior in sugar cane juice.

Fermentative behavior of S. cerevisiae strains in sugar cane juice

A total of 18 non-flocculent strains were selected to produce ethanol from sugar cane juice fermentation. These strains included fruit wine and distilleries isolates. S. cerevisiae PE-2, commercially used in Brazil for ethanol production, was also used as a reference to compare with the indigenous strains selected.

The results obtained from the kinetic parameters were subjected to PCA. Three initial principal components (PC) accounted for 73.68% of the total initial variance. Figure 2 shows the plot of the PCA for the first (PC1) and the second (PC2) principal components, which explains 56.03 and 17.65% of the total variance, respectively. The values for \( Y_{p/s} \) ranged from 0.21 g g⁻¹ to 0.46 g g⁻¹. The strains UFLA FW221, CCA083, UFLA CA798, UFLA CA759, UFLA CA759, UFLA CA93, UFLA FW510 and UFLA CA762 showed higher \( Y_{p/s} \), 0.46, 0.44, 0.43, 0.43, 0.40, 0.39 and 0.39 g g⁻¹, respectively. The industrial strain PE-2 isolated from a distillery and studied by Basso et al. (2008) for ethanol fuel production showed a value of \( Y_{p/s} \) of 0.38 g g⁻¹. This value was lower than the values obtained by indigenous strains evaluated in this work.

The conversion factor of \( Y_{g/s} \) ranged from 0.04 g g⁻¹ to 0.09 g g⁻¹. Strains UFLA CA759, UFLA FW211, UFLA CA798 and CCA083 showed the lowest values of \( Y_{g/s} \) corresponding to 0.05 g g⁻¹ for strains UFLA CA759 and UFLA CCA083 and 0.06 g g⁻¹ for the other two. Acetic acid production was not detected for any of the 18 selected yeasts. The methanol conversion (\( Y_{m/s} \)) was 0.01 g g⁻¹ for all strains except for UFLA CA753, whose value was 0.02 g g⁻¹.

The strains UFLA CA759, UFLA CA798, UFLA FW211 and CCA083 were then tested in fermentation with sugar cane juice, and calibration curves were performed to determine the initial inoculum concentration and to monitor yeast growth during the fermentation process. Figure 3 shows the PCA analysis based on the results obtained during fermentation for the four selected yeast strains. Three initial principal components (PC) accounted for 75.62% of the total initial variance; the first (PC1) and the second (PC2) principal components explained 46.97 and 28.65% of the total variance, respectively.

Table 2 shows the values of the fermentative parameters, the data did not differ significantly (p < 0.05) by Scott-Knott test, however the UFLA FW211 strain showed the highest value of \( Y_{p/s} \) (0.50 g g⁻¹) among all evaluated strains. The parameters \( Y_{ac/s} \), \( Y_{g/s} \) and \( Y_{suc/s} \) were similar among the UFLA CA759, UFLA CA798, UFLA FW211 and CCA083 strains. There was no production of acetic acid; the \( Y_{ac/s} \) value was 0.05 g g⁻¹ for the UFLA CA759 strain and 0.06 g g⁻¹ for the other three strains, and \( Y_{suc/s} \) was 0.01 g g⁻¹ for all four strains. The maximum value of \( Y_{x/s} \) was 0.06 g g⁻¹ for UFLA CA759, and the minimum was...
0.01 g g\(^{-1}\) for the UFLA CA798 strain. The \(Y_{x/s}\) value obtained from UFLA FW211 was 0.04 g g\(^{-1}\).

These results showed that the four \textit{S. cerevisiae} strains UFLA CA759, UFLA CA798, UFLA FW211 and CCA083 were able to adapt and to ferment the sugar cane juice quickly. However, since the strain UFLA FW221 presented the higher value of \(Y_{p/s}\) this strain was selected for further studies.

**Behavior of \textit{S. cerevisiae} UFLA FW221 in co-culture with \textit{L. fermentum}**

Fermentation monitoring was based on the production rate of CO\(_2\) (\(\frac{d\text{CO}_2}{dt}\)), carbohydrate consumption and ethanol production (Figure 4). The rates of CO\(_2\) production ranged from approximately 0.3 to 0.35 g L\(^{-1}\) h\(^{-1}\), where the maximum CO\(_2\) production rate was obtained approximately 12 h after the beginning of the fermentative process. The strain PE-2 was employed as a reference, because it is successfully employed in Brazilian industrial fermentations. According to CO\(_2\) rates, both strains showed similar performance when it was inoculated single or in co-culture with \textit{L. fermentum}. All yeast strains showed a similar behavior for carbohydrate consumption, and ethanol production (Figure 4).

**Batch scale fermentation**

Fermentation was carried out in five consecutive batches using the selected strain UFLA FW221. Pulsed Field Gel Electrophoresis analysis were performed to observe if the yeast is genetically stable during the batches (Figure 5), fluorescence microscopy was performed to assess cell viability (Figure 6), and analyses of ethanol, glycerol, organic acids, trehalose (Table 3) and biomass (Figure 7) were also carried out to observe the fermentative characteristics of the yeasts during the batches. All colonies isolated during the batches had the same chromosomal profile, shown in Figure 5 and were identical to the corresponding inoculated yeast UFLA FW221. Figure 6 shows the fluorescent image of the yeast at the end of the fifth batch of fermentation. It was observed that the inoculated yeast remained viable until the last of the five batches even at high alcohol concentrations (approximately 77.4 g L\(^{-1}\)).

Table 2 shows that high volumes of ethanol were produced during the batches, reaching levels up to 77 g L\(^{-1}\) (approximately 10% v/v) of ethanol. The values of glycerol were approximately 5 g L\(^{-1}\) and succinic acid 1 g L\(^{-1}\). Acetic acid production was not observed.

**Table 2 - Fermentative parameters calculated from four selected \textit{Saccharomyces cerevisiae} strains.**

| Yeast strain | Fermentative parameters |
|--------------|-------------------------|
|              | Yp/s       | Yg/s       | Qp | Yx/s       | Ysuc/s      | % Px     |
| UFLA CA759   | 0.47 ± 0.01\(^a\) | 0.05 ± 0.0001\(^a\) | 2.81 ± 0.03\(^a\) | 0.06 ± 0.004\(^a\) | 0.01 ± 0.0001\(^a\) | 91 ± 2.0\(^a\) | 0.39 ± 0.03\(^a\) |
| UFLA CA798   | 0.46 ± 0.02\(^a\) | 0.06 ± 0.0003\(^a\) | 2.85 ± 0.07\(^a\) | 0.01 ± 0.00001\(^a\) | 0.01 ± 0.0001\(^a\) | 90 ± 5.7\(^a\) | 0.04 ± 0.0001\(^a\) |
| UFLA FW221   | 0.50 ± 0.01\(^a\) | 0.06 ± 0.0002\(^a\) | 2.80 ± 0.04\(^a\) | 0.04 ± 0.002\(^a\) | 0.01 ± 0.0001\(^a\) | 99 ± 2.7\(^a\) | 0.19 ± 0.0001\(^a\) |
| CCA 083      | 0.47 ± 0.01\(^a\) | 0.06 ± 0.0001\(^a\) | 2.85 ± 0.01\(^a\) | 0.04 ± 0.002\(^a\) | 0.01 ± 0.0001\(^a\) | 92 ± 1.3\(^a\) | 0.23 ± 0.01\(^a\) |

Presented values are means of duplicate determinations; \(\pm\) indicates standard deviations from the mean. Mean values (± standard deviation) within the same column followed by different superscript letters differ significantly (\(p < 0.05\)) by Scott-Knott test.
The \textit{S. cerevisiae} strain UFLA FW221 was able to accumulate intracellular trehalose (about 32 μmol trehalose per g of wet weight) at the end of five batches (Table 2).

**Discussion**

In general, it is known that industrial process to produce ethanol is harsh for yeast. Therefore, suitable yeast is an important factor to obtain success in productivity. Stressful conditions affect yeast cell metabolism, leading to loss of cell viability and fermentation ability. Resistance to stress conditions is strain-dependent. The yeasts into G1 group were able to grow in higher temperature (42 °C) and ethanol concentration (12%) and also they were osmo- and sulphite- (1 M NaCl and 0.04%, respectively) tolerant. Thus, they displayed characteristics to tolerate these unfavorable conditions imposed by industrial process. The temperature during the fermentation process in Brazil can reach 40 °C and the maximum ethanol concentration, which increases gradually, reaches approximately 8 to 11% (v/v) at the end of the process (Basso et al., 2008). Hence, the G1 strains are probably able to survive until the end of the high ethanol processes. The ability to adapt to changes in the osmolarity of the surrounding medium is fundamental to life, and the accumulation of compatible solutes to decrease intracellular water potential is an adaptation strategy employed by all cell types (Rep et al., 2000). Thus, yeast osmotolerance may be an important factor in ethanol production. Since sulfite and sulfite-generating compounds have long been used as antimicrobial agents in alcoholic fermentation (Walker, 1998), sulfite tolerance in yeast is another desired trait for bioethanol production from sugar cane juice.

Yeast flocculation capacity is considered an important characteristic for some fermentative industrial processes (Campos et al., 2010; Schwan et al., 2001; Stewart and Russell, 1981). However, in the bioethanol industry, flocculent yeast cells can obstruct the pipes and reduce cell-substrate contact, which increases the fermentation

**Table 3** - Concentrations (g L⁻¹) of total sugar, ethanol, glycerol, organic acids and intracellular trehalose contents detected in sugar cane juice fermentation after five batches with the UFLA FW221 strain.

| Sugar cane juice | 1          | 2          | 3          | 4          | 5          |
|------------------|------------|------------|------------|------------|------------|
| Residual Total sugar (g L⁻¹) | 169.27 | 3.51 ± 0.62 | 2.83 ± 0.28 | 8.69 ± 1.56 | 13.66 ± 3.20 | 1.91 ± 1.03 |
| Ethanol (g L⁻¹)    | ND         | 67.00 ± 8.68 | 76.50 ± 1.81 | 77.38 ± 5.43 | 71.36 ± 6.36 | 73.41 ± 3.21 |
| Glycerol (g L⁻¹)   | ND         | 5.57 ± 0.40 | 5.84 ± 0.15 | 5.67 ± 0.57 | 5.16 ± 0.44 | 5.16 ± 0.20 |
| Succinic acid (g L⁻¹) | 0.40 | 1.29 ± 0.01 | 1.33 ± 0.02 | 1.20 ± 0.14 | 1.11 ± 0.14 | 0.97 ± 0.07 |
| Citric acid (g L⁻¹) | 0.15 | 0.79 ± 0.02 | 0.81 ± 0.10 | 0.70 ± 0.11 | 0.68 ± 0.07 | 0.58 ± 0.01 |
| Acetic acid (g L⁻¹) | ND         | ND         | ND         | ND         | ND         | ND         |
| Trehalose (μmol glucose g wet weight⁻¹) | - | 25.39 ± 3.01 | 20.34 ± 4.90 | 20.42 ± 2.28 | 25.55 ± 4.45 | 31.99 ± 0.87 |

ND = not detected.
time and the amount of residual sugar at the end of the fermentative process (Basso et al., 2008).

The strains into G1 group which produced more than 10 g L$^{-1}$ of ethanol were tested in sugar cane fermentations. The Figure 2 present that all yeast strains grouped on the positive axis of PC1 displayed greater correlations among the $Y_{p/s}$ conversion (%), $E_t$ and $Q_p$ parameters (Figure 2), which indicate that these yeasts are able to convert sugar cane juice into ethanol. The strains UFLA CA798, UFLA FW221, CCA083, UFLA FW510, UFLA CA93, UFLA CA762 and UFLA CA759 were closely related to the $Y_{p/s}$, $Q_p$ and $E_t$ parameters (Figure 2). The increase of these parametric values is related to the high yield of ethanol production (Andrietta et al., 2008).

The parameters related to the conversion of a substrate into secondary products, such as $Y_{g/s}$, $Y_{ac/s}$ and $Y_{m/s}$, are in the negative axis of PC1 and were low for all 18 selected strains.

It is known that glycerol formation is coupled to yeast growth and is also formed in response to stressful conditions (Walker, 1998). Glycerol is formed by yeasts at the beginning of fermentation. This period corresponds to the start of glycerolpyruvic fermentation, which is the only way that yeasts can ensure the reoxidation of the NADH$^{+}$/H$^+$ coenzyme by reducing dihydroxyacetone to glycerol (Ribereau-Gayon et al., 2006). Low levels of glycerol synthesis can be associated with an active metabolism for ethanol formation and implies better yeast fermentation performance.

Because the strains UFLA CA798, UFLA FW221, CCA083, UFLA FW510, UFLA CA93, UFLA CA762 and UFLA CA759 showed the best results in the last experimental step (higher $Y_{p/s}$ and lower $Y_{g/s}$ values), they were cultured in starvation medium to observe asc or pseudohyphae formation. The strains UFLA FW510, UFLA CA93 and UFLA CA762 showed pseudohyphae formation (data not shown), which is an undesirable characteristic in the ethanol industry. Cells exhibiting hyphae or filamentous morphologies also differ in their expression and accumulation of undesirable cell constituents and in their ability to be disrupted or fractionated (Gibbs et al., 2000).

The strains UFLA CA759, UFLA CA798, UFLA FW211 and CCA083 were selected for a second fermentation. Even though all four strains did not differ significantly ($p < 0.05$) by Scott-Knott test and all of them are able to convert sugar in ethanol in a great ratio, the strain FW221 exhibited the higher rate of conversion of substrate into product (99% of the theoretical value) and a lower conversion of substrate into secondary products. This strain showed a value of 0.50 g g$^{-1}$ to $Y_{p/s}$. Ortiz-Muñiz et al. (2010) investigated the kinetics of ethanol fermentation using S. cerevisiae ITV-01 strain isolate from molasses in different conditions to determine the optimum fermentation conditions. They found in the optimal conditions that this strain obtained ethanol yield of 0.41 g g$^{-1}$.

The parameters which represent conversion of carbohydrates into secondary products were low for all strains studied. The conversion factor for converting substrate into biomass ($Y_{c/s}$) represents the growth capacity of a strain in the fermentation medium. The strain UFLA FW221 presented a value of 0.04. It means that this strain was capable to use carbohydrates to convert part in ethanol and part in biomass and there were a good balance between these parameters. Marini et al. (2009) studied indigenous S. cerevisiae to produce cachaça from sugar cane and they found values of $Y_{c/s}$ greater than 0.09 g g$^{-1}$ and $Y_{p/s}$ ranging from 0.25 to 0.40 g g$^{-1}$. These values indicate that the strains to produce ethanol convert less carbohydrates in biomass and more in ethanol than cachaça strains. These results showed that the fruit fermentation isolate UFLA FW211 presented a high capacity for growth in sugar cane juice and produced a high volume of ethanol. Nowadays, suitable yeasts have been selected from distilleries process because they are resistant to stress conditions impose by industrial fermentation process (Basso et al., 2008). However, this work has showed the importance of studying microbial diversity from different habitats. This study presented an isolate from fruit fermentation with excellent fermentative characteristics in sugar cane juice.

In fermentation assay with mixed culture of L. fermentum and S. cerevisiae strains UFLA FW221 and PE-2, L. fermentum did not affect the fermentation performance of the yeasts (Figure 4). It was observed that in pure and co-cultures the carbohydrate consumption and ethanol production were similar (Figure 4b). According to CO$_2$ measurements (Figure 4a), the sugar cane fermentation was not affected negatively by L. fermentum presence for both yeasts PE-2 and UFLA FW221. It is known that Lactobacillus species can contaminate ethanol plants and cause losses in productivity, since they may compete for nutrients (Wheals et al., 1999). Thus, the studied yeast strains which presented the same fermentative behavior in pure or co-culture with Lactobacillus is desirable, considering the initial population of $10^6$ CFU mL$^{-1}$ and $10^6$ cfu mL$^{-1}$.
for yeast and bacteria, respectively. Besides this, it could avoid the over-use of antibiotics products to control the bacteria growth. Since, Yokoya and Oliva-Neto, (2001) showed that some biocides used in industrial fuel alcoholic fermentation in Brazil could affect yeast activity.

The growth of \textit{S. cerevisiae} strains were not affected by the \textit{L. fermentum} presence. However, the bacterial growth decline in the yeast presence (Figure 4c). In the literature, we can find some reports of yeast and bacterial interaction (Meignen \textit{et al.}, 2001). Nobre \textit{et al.} (2007) reported that the lactic acid bacteria (LAB) \textit{L. fermentum} reduced the viability of \textit{S. cerevisiae}. In the alcoholic fermentation of corn mash, when the mash was inoculated with the \textit{S. cerevisiae} and \textit{L. fermentum} at the same time, bacterial growth still occurred but was reduced by 94%. The reduction in bacterial growth may be due to alcohol produced by the yeast which can exert inhibitory effects on the multiplication of lactobacilli (Thomas \textit{et al.}, 2001). The results observed in batch fermentations showed that the UFLA FW221 strain remained genetically stable in the five batches and presented high viability. These data are important for the selection of industrial strains because yeasts are recycled during long fermentation periods.

The strain UFLA FW221 was able to accumulate a great amount of trehalose approximately 32 \(\mu\)mol trehalose per g of wet weight. The mechanism by which the yeast protects against the stress imposed by the industrial process includes trehalose accumulation, the synthesis of molecular chaperones and the synthesis of antioxidative enzymes. Trehalose has been used as a good indicator of the ability of yeasts to withstand stressful conditions (Tanghe \textit{et al.}, 2006). Furthermore, trehalose metabolism has been suggested to be an important physiological criterion for yeast selection in biotechnological processes.

This study screened 66 indigenous \textit{S. cerevisiae} strains and selected strains as potential candidates for use in industrial bioethanol processes. The behaviours of the four selected strains UFLA CA759, UFLA CA798, UFLA FW211 and CCA083 were studied in sugar cane juice. The UFLA FW221 strain isolated from fruit fermentation showed a higher \(Y_{ps}\) value (0.50 g g\(^{-1}\)), corresponding to 99% of the theoretical value. This strain was subject to batch scale fermentation. It was observed that UFLA FW221 was suitable for ethanol production, producing approximately 10% (w/v) ethanol from sugar cane juice in 24 h. These results showed that despite being isolated from fruit fermentation, the UFLA FW221 strain demonstrated an excellent fermentative performance in sugar cane juice, showing the importance of studying microbial biodiversity for applications in biotechnology.

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