Identification and assessment of kefir yeast potential for sugar/ethanol-resistance

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Abstract

Biochemical and molecular analysis was used for identification of different kefir yeasts species from Brazil, Canada and the United States of America. The sugar/ethanol-resistant activity of the yeasts was evaluated. Saccharomyces cerevisiae and Kluyveromyces marxianus had the highest growth rates, suggesting biotechnological applications possible for these strains.

Key words: kefir, sugar/ethanol-resistant yeasts, yeasts diversity, rep-PCR, PCR-DGGE.

Kefir is a refreshing, naturally carbonated fermented dairy beverage with a slightly acidic taste, yeasty flavor and creamy texture (Miguel et al., 2010; Magalhães et al., 2011a, 2011b). Previous results have shown that two groups of microorganisms co-exist in milk kefir grains: lactic acid bacteria and yeast (Miguel et al., 2010; Magalhães et al., 2011a, 2011b). The presence of yeast enhances the organoleptic quality of the kefir beverage, promoting a strong and typically yeasty aroma, as well as a refreshing and pungent taste (Magalhães et al., 2011a, 2011b). Yeasts isolated from kefir could be used for research that aims to produce fermented alcoholic beverages and ethanol using other substrates, e.g. whey. Cheese whey is the major by-product of the dairy industry, and its disposal without expensive sewage treatments represents a major source of environmental pollution. Some methods have been proposed for whey valorization by yeast (Dragone et al., 2009; Guimarães et al., 2010) including distilled beverages (Dragone et al., 2009) and kefir-like whey beverages (Paraskevopoulos et al., 2003; Magalhães et al., 2010b, 2011a, 2011b). Seeking this possibility, the aim of this study was to isolate and identify kefir-derived yeast species from different countries (Brazil, Canada, and the United States of America) and to evaluate their potential for sugar/ethanol-resistance. The yeast diversity was also assayed using denaturing gradient gel electrophoresis (DGGE).

The samples of kefir grains were obtained from families (private households) that traditionally consumed the kefir beverage. These families were from locations in three different countries: Lavras (Brazil - BR), Bowmanville (Canada - CA), and Fayette (United States of America - USA). Each kefir grain sample was collected aseptically for phenotypic analysis (50 g of sample) and genotypic analysis (5 g of sample).

Yeasts were enumerated by plating 10 µL of each diluted sample in triplicate using the surface spreading technique described by Magalhães et al. (2010a). Enumeration of yeasts was carried out using the culture media malt yeast glucose peptone extract (MYGP agar, Merck, Whitehouse Station, USA); 0.3% yeast extract, 0.3% malt extract, 0.5% peptone, 1.0% glucose, 2.0% agar) containing 100 mg/L chloramphenicol (Sigma, St. Louis, USA) and 50 mg/L chlortetracycline (Sigma, St. Louis, USA) to inhibit bacterial growth. After spreading, plates were incubated at 28 °C for 120 h and colony forming units (log10 cfu/g) were quantified. Colonies were taken at random for identification from each plate containing isolated colonies (the number of colonies used for identification was equal to the square root of the total number of colonies on that plate). The phenotypic characteristics of all yeast isolates were determined by their morphology and spore formation, as well as their assimilation and fermentation of different carbon sources (Magalhães et al., 2010a). For the selection of yeast isolates, molecular biology-based grouping of the isolates was performed using repetitive extragenic palindromic (rep-PCR) using the primer (GTG)5 (5’-GTG GTG GTG GTG GTG GTG-3’). The screening was performed with all 125 isolates. The cluster was done only by the different profiles obtained from rep-PCR technique. DNA was extracted from the pure cultures as described by Pereira et al. (2010). The
yeast *Saccharomyces cerevisiae* UFLA GF33 was used as the reference strain. The rep-PCR profiles were normalized, and cluster analysis was performed using Bionumerics 2.50 (Applied Maths, Sint-Martens-Latem, Belgium). After the selected isolates by rep-PCR technique were identified by Internal Transcribed Spacer (ITS) region analysis (Magalhães et al., 2011c).

In addition, a culture-independent method was used to analyze each sample. Kefir grains (5 g) from the different countries were transferred into a plastic tube and subjected to DNA extraction using a NucleoSpin Tissue kit (Macherey-Nagel, Düren, Germany) according to the manufacturer’s instructions. Genomic DNA was used as a template for PCR amplification of yeast ribosomal target regions, and the products from the PCR reaction were then analyzed by PCR-DGGE. The primers NS3fGC and YM951r were used for partial ribosomal 18S DNA amplification as described by Magalhães et al. (2010a). Selected bands from the PCR-DGGE gels were excised and reamplified using the same primers without GC clamp. The sequencing of portions of the ITS region from the yeast isolates and 18S rRNA gene amplicons were performed by the Central Laboratory of Molecular Biology - LCBM/UFLA (Lavras, MG, Brazil). Sequence similarity was analyzed using the BLAST database from GenBank (http://www.ncbi.nlm.nih.gov/BLAST/).

One representative microorganism from each identified microbial species and each kefir grain was inoculated in triplicate in 2.0 mL culture medium MYGP medium containing 100 mg/L chloramphenicol and 50 mg/L chlorotetracline to inhibit bacterial growth. The medium was then supplemented with 0, 4, 6, 9 or 12% v/v ethanol (Merck, Whitehouse Station, USA), and the flasks were incubated at 28 °C for 120 h. The ethanol resistance was determined by measuring biomass formation. The yeast *Saccharomyces cerevisiae* UFLA GF33, which is ethanol-resistant and has a high growth rate under these conditions, was used as a control. The ethanol-resistant yeasts were inoculated in triplicate in 2.0 mL of culture medium containing 10 g/L yeast extract (Merck, Whitehouse Station, USA) and 20 g/L peptone (Merck, Whitehouse Station, USA). This medium was supplemented with 100 or 250 g/L (Osho, 2005; Miguel et al., 2010) glucose (Merck, Whitehouse Station, USA) or lactose (Merck, Whitehouse Station, USA). Sugar resistance was determined as described above.

The enumeration values (cfu/g) of the isolated viable yeasts ranged from a minimum value of 7.43 ± 0.01 log_{10} cfu/g in the sample from Canada to a maximum value of 9.57 ± 0.03 log_{10} cfu/g in the sample from Brazil, with the sample from the USA falling in between at 8.38 ± 0.02 log_{10} cfu/g. A total of 125 isolates were obtained from kefir grain samples and grouped for molecular characterization using the (GTG)^5-PCR technique (Figure 1a). The cluster was grouped by Bionumerics 2.50 software (Applied Maths, Sint-Martens-Latem, Belgium). Cluster analysis using (GTG)^5-PCR patterns resulted in the identification of two groups referred to as Y1 and Y2 at the level of 45% similarity. The Y1 group was subdivided into three subgroups (Y1A, Y1A1 and Y1B) with greater than 70% similarity. The Y1A group consists of isolates of the *Saccharomyces* genus. The Y1A1 group is composed of *Saccharomyces cerevisiae* isolated, which are grouped at the level of ~ 96% similarity. This group therefore is grouped with *Saccharomyces bayanus* strain, forming the group Y1A to ~ 79% similarity. These isolates were derived from kefir grains from Brazil, Canada and the USA. Group Y2 contained 14 isolates that demonstrated less similarity with UFLA GF33. These isolates were also present in kefir grains from Brazil, Canada and the USA. The high incidence of band differences between yeast isolates makes the rep-PCR technique efficient for distinguishing between different strains. Isolates representing different branches in the (GTG)^5-PCR fingerprinting analysis were chosen for partial sequencing of the PCR-amplified ITS region.

The 125 yeasts isolated were identified by phenotypic and genotypic techniques as *Saccharomyces cerevisiae* (11 from Brazil, 7 from Canada and 9 from the USA), *Saccharomyces* sp. (3 from Brazil, 3 from Canada and 5 from the USA), *Saccharomyces bayanus* (2 from Brazil), *Zygosaccharomyces mellis* (3 from Brazil), *Dekkera anomala* (3 from Brazil and 1 from Canada), *Kluyveromyces marxianus* (21 from Brazil, 16 from Canada and 19 from the USA), *Pichia membranifaciens* (1 from Brazil), *Pichia guilliermondii* (2 from Brazil and 1 from Canada), *Pichia fermentans* (5 from Brazil and 6 from the USA), *Pichia anomala* (1 from Canada and 1 from the USA), *Candida parapsilosis* (1 from Brazil and 1 from the USA) and *Candida valida* (1 from Canada and 2 from the USA). The species *Saccharomyces cerevisiae*, *Pichia fermentans*, *Saccharomyces bayanus*, *Kluyveromyces marxianus* and *Zygosaccharomyces mellis* have been previously found in kefir grains (Magalhães et al., 2010a, 2011c; Miguel et al., 2011), whereas *Dekkera anomala* has not been reported as a microorganism present in kefir grains. *Dekkera anomala*, may have been introduced to the kefir grains as a contaminant during manipulation or transport. Another factor may be the species *Dekkera anomala* be part of the original microflora of these grains, because different kefir grains may have diverse microbiota (Miguel et al., 2010, 2011; Magalhães et al., 2011c). This species was identified in kefir grain samples from Brazil and Canada. *Saccharomyces* spp. were also identified in this study. Sequencing of the ITS region confirmed that the isolates are closely related to *Saccharomyces cerevisiae* and *Saccharomyces* spp., with 99.2-100% similarity. However, *S. cerevisiae*, *S. bayanus* and *S. pastorianus* are very closely related, so to obtain an unambiguous identification, the isolates were differentiated using species-specific PCR primer pairs homologous to the HO genes of *S. cerevisiae*.
A single amplicon of approximately 40 bp confirmed the identity of the Saccharomyces spp. as S. cerevisiae (data not shown).

In addition to the culture-based methods, the yeast communities from the kefir samples were investigated using DGGE analysis of the PCR-amplified 18S rRNA gene. Kefir grain samples from Brazil had the greatest band diversity according to a statistical comparison using Duncan’s multiple range test (p = 0.05; Figure 1c), suggesting that this sample had the greatest diversity of yeast species. The CA and USA grain samples showed no significant differences in band diversity. To determine the composition of the yeast communities from the kefir samples, the bands were excised from the DGGE gel, and the DNA was

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isolated and sequenced. Figure 1b shows the DGGE profiles of yeast 18S rDNA fragments amplified from kefir grains samples of different countries. The bands were identified based on a search for similar sequences in GenBank as follows: band 1 - *Pichia fermentans*, band 2 — *Pichia guilliermondii*, band 3 - *Candida validaiviana*, bands 4 and 8 - *Zygosaccharomyces* sp., band 5 - *Lachancea meyersii*, band 6 - *Saccharomyces cerevisiae*, band 7 - *Klyveromyces marxianus*, band 9 - *Saccharomyces bayanus* and band 14 - *Kazachatania unispora*. Despite the fact that the *Zygosaccharomyces* sp. and *Candida* sp. could be identified only at the genus level, *Zygosaccharomyces fermentati* and *Candida valdiviana* or *Candida parapsilosis* were the most probable species because they were sporadically isolated from kefir grains in this study. The *Zygosaccharomyces* sp. (bands 4 and 8) appeared in the DGGE gel from the Brazil sample in two different positions. The multiple banding patterns may be due to sequence heterogeneity between multiple copies of the 18S rDNA in this strain (Miguel et al., 2011). *Lachancea meyersii* and *Kazachatania unispora* were not isolated by culture-dependent methods, but had been previously identified in kefir grains by Magalhães et al. (2010a, 2010b). It was not possible to identify band 10, 12 and 13; the bands were excised from the gel, but DNA could not be recovered for sequencing.

Twenty-five kefir yeasts (grouped by rep-PCR) isolated from fermented milk were screened for ethanol and sugar (glucose/lactose) tolerance (Figure 2). Only the isolates of *Saccharomyces cerevisiae*, *Saccharomyces* sp. and *Klyveromyces marxianus* were able to grow at a concentration of 12% (v/v) ethanol. Figure 2 shows the identities of the isolates as well as the percentage of ethanol (v/v) that inhibited the growth of the other isolates. All of the ethanol-tolerant yeasts were tested in different concentrations of both glucose and lactose, and all reached their maximum growth rates at a sugar concentration of 25% (v/v) (Figure 2). *Saccharomyces cerevisiae* and *Klyveromyces*

| Morphotypes yeast MYGP medium | Accession number | Concentrations of Ethanol | Concentrations of Sugar |
|-------------------------------|-----------------|---------------------------|-------------------------|
| **Saccharomyces cerevisiae** UFLA GF 33 reference strain | GI61332350 | + | + | + | + | (GL) | + | (GL) |
| *Saccharomyces cerevisiae* UFLA KFG01, 02, 03 *Saccharomyces* sp UFLA KFG09, 10, 11 Brazil (1), Canada (1), USA (1) | EU649673.1 | + | + | + | + | + | (GL) | + | (GL) |
| *Saccharomyces cerevisiae* UFLA KFG09, 10, 11 Brazil (1), Canada (1), USA (1) | AY796128 | + | + | + | + | + | (GL) | + | (GL) |
| *Saccharomyces bayanus* UFLA KFG29 Brazil (1) | D89889.1 | + | + | + | - | - | n.a. | n.a. |
| *Zygosaccharomyces melliis* UFLA KFG40 Brazil (1) | X90755 | + | + | - | - | - | - | n.a. | n.a. |
| *Dekkera anomala* UFLA KFG32 Brazil (1), Canada (1) | X83820 | + | + | - | - | - | - | n.a. | n.a. |
| *Klyveromyces marxianus* UFLA KFG21, 22, 23 Brazil (1), Canada (1), USA (1) | GM620995 | + | + | + | + | + | (LA) | + | (LA) |
| *Pichia membranifaciens* UFLA KFG121 Brazil (1) | DQ104710 | + | + | - | - | - | - | n.a. | n.a. |
| *Pichia guilliermondii* UFLA KFG91, 92 Brazil (1), Canada (1) | EF197951.1 | + | + | - | - | - | - | n.a. | n.a. |
| *Pichia fermentans* UFLA KFG73 Brazil (1), USA (1) | FJ770542.1 | + | + | - | - | - | - | n.a. | n.a. |
| *Pichia anomala* UFLA KFG56, 57 Canada (1), USA (1) | EU330185 | + | + | - | - | - | - | n.a. | n.a. |
| *Candida parapsilosis* UFLA KFG42 Brazil (1), USA (1) | AY227019 | + | + | - | - | - | - | n.a. | n.a. |
| *Candida validaiviana* UFLA KFG69, 10 Canada (1), USA (1) | FN052300.1 | + | + | - | - | - | - | n.a. | n.a. |

Figure 2 - Screening sugar/ethanol-tolerance kefir yeasts. (+) biomass formation; (-) biomass not formation. Abbreviations: (GL) Glucose; (LA) Lactose; n.a. not analyzed. The values in parentheses refer to the yeast isolates number tested of each country.
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Paraskevopoulou et al., 2003; Pescuma et al., 2008; Dragone et al., 2009; Guimarães et al., 2010; Magalhães et al., 2010b, 2011a, 2011b). Due to the large lactose surplus generated, its conversion to bio-ethanol has long been considered as a possible solution for cheese whey bioremediation. Because of this, emphasis has been given to advances in engineering a consortium of K. marxianus and S. cerevisiae strains for efficient cheese whey-to-ethanol bioprocesses (Dragone et al., 2009; Guimarães et al., 2010). The yeasts isolated in this study (particularly those with high ethanol- and sugar-tolerance, i.e., K. marxianus and S. cerevisiae) suggest biotechnological applications possible for these strains e.g. represent new avenues for the study of the alcoholic fermentation of cheese whey. Another suggestion, the production of fermented alcoholic beverages produced upon whey fermentation by a consortium of K. marxianus and S. cerevisiae strains (identified and analyzed in this study) could be an interesting alternative for cheese whey utilization. Cheese whey fermentation by these yeasts could decrease the high lactose content in cheese whey, producing mainly lactic acid and other metabolites such as aroma compounds contributing to the flavor and texture and increasing carbohydrate solubility and sweetness of the end product (Pescuma et al., 2008). Manufacture of beverages through lactic fermentations can provide desirable sensory profiles and have already been considered an option to add value to cheese whey (Pescuma et al., 2008).

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