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Torulaspora delbrueckii in the brewing process: A new approach to enhance bioflavour and to reduce ethanol content

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Abstract: Nowadays, consumers require fermented alcoholic beverages with particular and enhanced flavour profiles while avoiding the health concerns due to high ethanol content. Here, the use of Torulaspora delbrueckii was evaluated for beer production, in both pure and in mixed cultures with a Saccharomyces cerevisiae starter strain (US-05). The yeast interactions were also evaluated. In mixed fermentations with S. cerevisiae, the main analytical characters from T. debrueckii were comparable with those of the S. cerevisiae starter strain, but the beers were characterized by a distinctive overall analytical and aromatic profile. Indeed, there were interactions between S. cerevisiae and T. delbrueckii, with enhanced ethyl hexanoate (0.048 mg 1-1) and ethyl octaonate (0.014 mg 1-1) levels at the 1:20 and 1:10 inoculation ratios, respectively; while phenyl ethyl acetate increased in all mix combinations. The presence of T. delbrueckii resulted in reduced β -phenyl ethanol and isoamyl acetate levels, which are responsible for floral and fruity aromas, respectively. Beer produced with T. delbrueckii pure cultures had a low alcohol content (2.66%; v/v), while also showing a particularly analytical and aromatic profile.

Dear Editor,

We would be glad if you would reconsider the enclosed revised manuscript, entitled, "*Torulaspora delbrueckii* in the brewing process: a new approach to enhance bioflavour and to reduce ethanol content", for inclusion in *Food Microbiology*.

We carefully revised the manuscript following the suggestions of the reviewer 2.

Yours faithfully

Prof. Maurizio Ciani

Response to reviewers

Reviewers' comments:

Reviewer #2: In the highlights the genus names should be written in full: Torulaspora delbrueckii and Saccharomyces cerevisiae

Corrected in the highlights

Line 67: However, the Belgian lambic beer is obtained from the fermentation of Change to: However, the Belgian lambic beers are obtained from the spontaneous fermentation ...

Corrected in the text

Line 136: bottles,adding ... Change to: bottles, adding

we suppose Line 140 the space was already present in previous R2 version

Line 202:showing very closed cell ... Change to:showing very similar cell ...

we suppose line 206 "similar" at place of "closed" already corrected in the previous version R2

Line 211: ...together with maltose residue. Change to: ...associated with higher residual maltose levels.

Changed in the text

Lines 213 and 214: use absolute numbers for attenuation (no decimals)

Corrected in the text

Line 216:an higher consumption ... Change to:a higher consumption ... (as already mentioned before!)

we suppose line 220 already corrected in the R2 previous version

Line 218: a competitive interactions in mixed fermentation even if a large amounts ... Change to: a competitive interaction in mixed fermentation even if large amounts ...

Corrected in the text

Lines 297-298: not clear Do you mean: ...that in mixed fermentations (...) exhibiting a higher consumption of YAN this could be related to the production of aromatic compounds (...)?

Yes, we corrected in the text following your suggestion.

Reviewer #3: This paper presents an interesting study about the application of non-Saccharomyces yeasts in brewing. In particular, the paper studies the influence of Torulaspora delbrueckii on analytical and aromatic profile.

All the comments indicated by previous reviewers have been resolved, and this has significantly improved the paper, specially the introduction.

Highlights

The potential use of a selected *Torulaspora delbrueckii* strain in the brewing process was evaluated.

Mixed fermentation *Torulaspora delbrueckii/Saccharomyces cerevisiae* produced beer with distinctive analytical and aromatic profile.

Torulaspora delbrueckii may be proposed to produce a low alcohol beer.

1	Torulaspora delbrueckii in the brewing process: a new approach to enhance bioflavour
2	and to reduce ethanol content
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19 Abstract

20

Nowadays, consumers require fermented alcoholic beverages with particular and enhanced 21 flavour profiles while avoiding the health concerns due to high ethanol content. Here, the use 22 of Torulaspora delbrueckii was evaluated for beer production, in both pure and in mixed 23 cultures with a Saccharomyces cerevisiae starter strain (US-05). The yeast interactions were 24 25 also evaluated. In mixed fermentations with S. cerevisiae, the main analytical characters from T. debrueckii were comparable with those of the S. cerevisiae starter strain, but the beers were 26 characterized by a distinctive overall analytical and aromatic profile. Indeed, there were 27 interactions between S. cerevisiae and T. delbrueckii, with enhanced ethyl hexanoate (0.048 28 mg l^{-1}) and ethyl octaonate (0.014 mg l^{-1}) levels at the 1:20 and 1:10 inoculation ratios, 29 30 respectively; while phenyl ethyl acetate increased in all mix combinations. The presence of T. delbrueckii resulted in reduced β -phenyl ethanol and isoamyl acetate levels, which are 31 32 responsible for floral and fruity aromas, respectively. Beer produced with T. delbrueckii pure 33 cultures had a low alcohol content (2.66%; v/v), while also showing a particularly analytical and aromatic profile. 34

35

Keywords: beer, *Torulaspora delbrueckii*, mixed fermentation, bioflavour, low-alcohol
content

38

40 **1. Introduction**

41

Over the years, brewers have always tried to find yeast strains that can improve the quality of 42 their beer and provide it with specific sensory notes. The various aroma compounds that 43 characterize different beers styles come from the raw materials of barley, malt and hops. 44 However, yeast has a central role in the brewing process, metabolizing of sugars in the beer 45 46 wort into ethanol, carbon dioxide, and several aroma compounds, including esters, higher alcohols, aldehydes and organic acids (Kyselová and Brányik, 2014; Lodolo et al., 2008; Pires 47 et al., 2014). In particular, in the beer industry, the goal of the use of inoculated yeast is to 48 increase the fermentation efficiency, to develop new beers, and especially to enhance the 49 sensory complexity of the final beer produced (Harrison, 2009). The production of aroma 50 51 compounds through biological methods exploits the metabolic pathways of the yeast, for the promotion of the so-called bioflavour (Cheetham, 1993; Vanderhaegen et al., 2003). This 52 53 approach can include microbial bioconversion of the flavour precursors, use of strains that 54 produce the required compounds, and genetic modification of the yeast (Dequin, 2001; Priefert et al., 2001; Ramachandra and Ravishankar, 2000; Mertens et al., 2015). 55

In winemaking, there has been a re-evaluation of the role of non-Saccharomyces yeast and 56 their use in mixed fermentations, with the aim to enhance the analytical and aromatic profile 57 of the final wine and to reduce the alcohol content (Benito et al., 2011; Ciani et al., 2010; 58 59 Comitini et al., 2011; Contreras et al., 2014; Morata et al., 2012; Quirós et al., 2014; Sadoudi et al., 2012). Within the non-Saccharomyces yeast species, attention has been focused on 60 Torulaspora delbrueckii, as this yeast has shown a positive impact in terms of low production 61 of undesirable compounds, such as acetaldehyde, acetoin and acetic acid, and concomitant 62 enhancement of other desired compounds (Azzolini et al., 2015; Bely et al., 2008; Comitini et 63

al., 2011; Jolly et al., 2014; Loira et al., 2012). The use of non-Saccharomyces yeast has been 64 65 less investigated in the brewing industry, where most beers are brewed with the use of a single yeast strain. However, the Belgian lambic beers are obtained from the spontaneous 66 fermentation of Saccharomyces and Brettanomyces yeasts, with the contribution of lactic acid 67 bacteria and acetic acid bacteria (Bokulich et al., 2012; Spitaels et al., 2014; Vanderhaegen et 68 al., 2003). These mixed fermentations were also used in the production of some weissbier 69 German style beer (Vriesekoop et al., 2012). During the maturation of acidic ale beers, 70 different yeasts belonging to Candida, Torulopsis, Pichia, Hansenula and Criptococcus 71 genera were isolated, but their contribution on the aroma composition was not investigated 72 73 (Vanderhaegen et al., 2003). Other non conventional beers such as Tchapalo, are brewed using Candida tropicalis and S. cerevisiae cultures selected for their ability to ferment 74 sorghum wort (N'Guessan et al., 2010). Regarding to the use of T. delbrueckii strains in 75 76 brewing process, only a few studies have been conducted. King and Dickinson (2000, 2003) reported that T. delbrueckii has the ability to transform hop aroma terpenoids, influencing the 77 78 aroma profile of the final beer. More recently, Tataridis and co-workers (2013) carried out a preliminary study on the use of *T. delbrueckii* strains in the production of "wheat" style beers. 79 These authors found that this species was able to consume maltose more slowly than S. 80 81 *cerevisiae* commercial starter strain, giving more intensity and complexity to the product. In the present study, after a preliminary screening, we evaluated the use of a selected strain of T. 82 delbrueckii in wort fermentation in pure and mixed cultures. The influence on the analytical 83 84 and aromatic profile of beer, as well as the potential of producing a low-alcohol beer with T. delbrueckii was evaluated. 85

86 2. Materials and methods

The 28 yeast strains used in this study belong to the species T. delbrueckii and were obtained 89 from the Yeast Collection of the Department of Life and Environmental Sciences (DiSVA) of 90 91 the Polytechnic University of Marche (Italy). These had been isolated from natural matrices from different environments and in different geographical areas (i.e., Italy, Cameroons) 92 (Table 1). All of the T. delbrueckii strains were identified through 5.8S internal transcribed 93 94 spacer rDNA polymerase chain reaction restriction fragment length polymorphism analysis, and sequencing of the D1/D2 domains of the 26S rDNA gene, as reported by Comitini et al. 95 (2011) and Solieri et al. (2006). The S. cerevisiae commercial strain US-05 (Fermentis, 96 97 Lesaffre, France) was used as the control.

For short-term storage, all of the yeast strains were maintained on YPD medium (1% yeast
extract, 2% peptone, 2% glucose, 1.8% agar; all w/v) (Oxoid, Basingstoke, UK) at 4 °C, and
for long-term storage, in YPD broth supplemented with 80% (w/v) glycerol at -80 °C.

101

102 2.2. Preliminary screening

103 The fermentation of glucose, maltose and sucrose by these 28 *T. delbrueckii* strains was 104 assessed using the Durham test, according to Kurtzman and Fell (1998). The fermentative 105 performance of eight of these *T. delbrueckii* strains that fermented maltose were determined at 106 20 °C in flasks that contained 500 ml malted barley wort under sterile conditions. The main 107 parameters of the fermentation kinetics (fermentation rate, total CO_2 evolved) of these *T. delbrueckii* strains and of the S. *cerevisiae* starter strain on the wort were assayed.

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110 2.3. Fermentation trials

From preliminary screening T. delbrueckii DiSVA 254 was selected and used in the pure and 111 mixed fermentations with the S. cerevisiae US-05 starter strain at different S. cerevisiae to T. 112 delbrueckii ratios (i.e., 1:1, 1:10, 1:20, respectively). A batch of 1,500 l of malted barley wort 113 for the production of American Amber Ale was used in this study. Its main analytical 114 characters were: pH 5.47; specific gravity 12.7 °Plato. The fermentation potential of the 115 selected strain was evaluated in fermentation trials carried out at 20 °C in flasks containing 116 117 500 ml wort under sterile conditions. The flasks were locked with a Müller valve containing sulphuric acid, to allow only CO_2 to escape from the system. 118

Pre-cultures were grown in 10% malt extract at 20 °C for 48 h. The fermentation kinetics 119 120 were monitored by measuring the weight loss of the flasks due to the CO₂ evolution, which was followed to the end of the fermentation (i.e., constant weight for 3 consecutive days). The 121 growth kinetics were monitored using viable cell counts on WL Nutrient Agar (Oxoid, 122 Hampshire, UK) and Lysine Agar (Oxoid, Hampshire, UK) a selective medium unable to 123 support the growth of S. cerevisiae (Lin, 1975), for differentiation of the T. delbrueckii yeast 124 125 from the S. cerevisiae starter strain. The fermentations were carried out in duplicate trials under static conditions. 126

127

128 2.4. Analytical determinations

The specific gravity was measured using a DA-300 specific gravity meter (Kyoto Instruments). The volatile acidity and pH determinations were performed according to the Official European Union Methods (EC, 2000). Ethanol was measured according to the Association of Official Analytical Chemists (1990). Acetaldehyde, ethyl acetate, higher alcohols, and volatile compounds were determined by direct injection into a gas–liquid chromatography system, as reported by Canonico et al. (2014). The free amino nitrogen was determined following a procedure described previously by Dukes and Butzke (1998). Specific
enzymatic kits (Megazyme, Ireland) were used to determine the concentrations of glucose
sucrose, maltose (kit k-masug) and ammonia (kit k-amiar) according to the manufacturer
instructions.

139 2.5. Sensory analysis

At the end of the fermentation process, the beers obtained were transferred into 500-ml 140 bottles, adding 5 g l⁻¹sucrose. The secondary fermentation in the bottle was carried out at 18-141 20 °C for 7-10 days. After this period, the beers underwent sensory analysis (Analytica EBC, 142 1997) on the basis of a list of descriptors related to both the aromatic notes (e.g., floral, fruity, 143 toasty) and the main structural features (e.g., sweet, acidity, flavour, astringency, bitterness, 144 olfactory persistence). A group of six trained tasters were asked to rate each sensory category 145 using a scale from 1 to 10. The results were combined, and the means were subjected to 146 statistical analysis. The data processed in this way were used to construct the Figures to 147 148 provide information on both the contribution of each descriptor to the overall organoleptic 149 quality of the beers, and the significant differences between the beers in relation to each descriptor. 150

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152 2.6. Statistical analysis

153 Analysis of variance (ANOVA) was applied to the experimental data for the main 154 characteristics of the beers. The means were analyzed using the STATISTICA 7 software. 155 The significant differences were determined by the means of Duncan tests, and the results 156 were considered significant if the associated *P* values were <0.05. The results of the sensory 157 analysis were also subjected to Fisher ANOVA, to determine the significant differences with 158 a p-value <0.05.

159 **3. Results**

160

161 *3.1. Sugar fermentation and fermentative performance in the wort*

Initial screening of the 28 *T. delbrueckii* strains was carried out to determine whether they fermented maltose, the most abundant fermentable sugar in the brewing wort (at 50%-60%), to select for their potential use in beer production. The data from this screening with Durham tubes indicated that out of the 28 strains tested, only eight cultures fermented maltose: DiSVA 254, DiSVA 602, DiSVA 603, DiSVA 343, DiSVA 399, DiSVA 413, DiSVA 419 and DiSVA 426. All of the 28 *T. delbrueckii* strains fermented glucose and sucrose.

These eight strains of T. delbrueckii that fermented maltose were evaluated for their 168 fermentative performance on the brewing wort. The parameters of the fermentation kinetics 169 are reported in Table 2. As expected, none of these eight strains showed fermentation 170 parameters comparable to those of the commercial S. cerevisiae starter strain. This behaviour 171 172 is important since the strains with lower fermentation performances in the wort are generally of interest for low-alcohol beer production. However, among the strains tested, T. delbrueckii 173 strain DiSVA 254 showed a good fermentative performance, with the highest fermentation 174 175 rate and final CO₂ production. For this reason, T. debrueckii DiSVA 254 was selected for the subsequent trials in pure and mixed fermentations with the S. cerevisiae starter strain US-05, 176 177 to evaluate the influence of T. debrueckii DiSVA 254 (henceforth: T. debrueckii) on the analytical and aromatic profiles of the beers produced. 178

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182 *3.2.1. Fermentation kinetics*

T. delbrueckii pure cultures and the three *S. cerevisiae* US-05 to *T. delbrueckii* inoculum
ratios of 1:1, 1:10 and 1:20, respectively, were investigated, with *S. cerevisiae* US-05 pure
cultures as the control. These mixed cultures of *S. cerevisiae* US-05 and *T. delbrueckii*showed CO₂ production that was comparable to that of *S. cerevisiae* US-05 alone, irrespective
of the different inoculation ratios (Fig. 1).

The fermentation evolution of the trials inoculated at the 1:1 ratio were consistent with the *S. cerevisiae* US-05 pure cultures. These data revealed that the inoculation of *T. delbrueckii* at the same concentration as *S. cerevisiae* US-05 did not affect the fermentation performance of the starter strain. In contrast, the fermentations carried out with the 1:10 and 1:20 ratios showed slower fermentation kinetics in comparison with *S. cerevisiae* US-05 alone. The *T. delbrueckii* pure culture also showed slow kinetics, which indicated that it did not provide complete wort attenuation.

195 *3.2.2. Biomass evolution*

The growth kinetics of the S. cerevisiae US-05 pure cultures achieved ca. 10^8 CFU ml⁻¹ at 7 196 days of fermentation, and maintained 10^7 CFU ml⁻¹ until the end of fermentation (Fig. 2). In 197 the mixed cultures with the inoculum ratio of 1:1, S. cerevisiae US-05 reached a lower 198 biomass (ca. 10^7 CFU ml⁻¹; Fig 2A), while for the inoculum ratios of 1:10 and 1:20, S. 199 *cerevisiae* US-05 remained at cell concentrations <10⁶ CFU ml⁻¹(Fig. 2B, C). In these trials, 200 the growth kinetics of T. delbrueckii showed biomass evolution compared with the T. 201 delbrueckii pure culture indicating that at these inoculation ratios S. cerevisiae was not 202 competitive with T delbrueckii, while at the 1:1 inoculum ratio, the S. cerevisie competition 203 occurred since the T. delbrueckii growth kinetics were much slower. Therefore, these data 204 indicated that T. delbrueckii at high concentrations (10-fold and 20-fold higher, vs. S. 205

cerevisiae US-05) dominated the process (Fig. 2). With 1:1 inoculation ratio both species
 suffered of the presence of each other showing very similar cell concentrations until the 7th
 day of fermentation. This behavior indicates a high level of competitiveness of *T. delbrueckii* towards *S. cerevisiae* commercial strain in wort.

210

211 *3.2.3. Main analytical profiles*

212 The data for the analytical compositions of the beers produced are reported in Table 3. The mixed cultures with S. cerevisiae US-05 and T. delbrueckii produced beers with ethanol levels 213 that were significantly lower, although essentially comparable, to those of the S. cerevisiae 214 US-05 control (4.51%-4.85%). On the other hand, the inoculations with T. delbrueckii pure 215 cultures showed a large reduction in the ethanol level (2.66%) associated with higher residual 216 217 maltose levels. Consequently, *T. delbrueckii* pure cultures showed a low attenuation (7.51 vs. 2.84-2.96 °P) with the real attenuation of 37%, which was significantly lower than that for the 218 other trials (63%-64%). For the volatile acidity, the mixed cultures produced beers with acetic 219 220 acid levels that were significantly higher than for the *T. delbrueckii* pure cultures (0.15 vs. 0.22-0.29 g L^{-1}). Regarding to assimilable nitrogen (YAN), a higher consumption in mixed 221 fermentations in comparison with pure cultures was showed. This behaviour indicated a 222 competitive interaction in mixed fermentation even if a large amounts of YAN remained. 223

224

225 *3.2.4. By-products and volatile compounds*

The data for the main by-products are reported in Table 4. The *T. delbrueckii* pure culture showed significant reductions for the levels of all of the by-products except for acetaldehyde, in comparison with the *S. cerevisiae* US-05 control. In particular, ethyl acetate is responsible for the fruity and solvent notes in the beer, and this was slightly, but significantly, reduced in the mixed fermentations compared with *S. cerevisiae* US-05 alone (14.65-16.25 vs. 17.57 mg Γ^{-1}), with a significantly greater reduction for the pure *T. delbrueckii* culture (3.46 mg Γ^{-1}). The acetaldehyde level was slightly, but significantly, higher in the beers produced with *T. delbrueckii* alone compared to *S. cerevisiae* US-05 alone (7.50 vs. 5.80 mg Γ^{-1}), and even further enhanced at the 1:1 (10.72 mg Γ^{-1}) and 1:10 (30.48 mg Γ^{-1}) fermentation ratios.

235 For the other analyzed by-products in the mixed fermentations, all of the beers produced by the association of S. cerevisiae US-05 and T. delbrueckii showed different profiles to those 236 produced from the S. cerevisiae US-05 and T. delbrueckii pure fermentations. Indeed, all 237 three mixed fermentations (i.e., 1:1, 1:10, 1:20) showed lower n-propanol levels in 238 comparison with S. cerevisiae US-05 (20.14-26.36 vs. 30.56 mg l^{-1}). For isobutanol, amylic 239 and isomylic alcohols, S. cerevisiae US-05 showed higher levels than the T. delbrueckii pure 240 cultures (24.38 vs. 7.98 mg l⁻¹), while the mixed fermentations showed intermediate levels 241 $(16.04-18.45 \text{ mg l}^{-1}).$ 242

243 The effects of T. delbrueckii on the beers were particularly evident for the main volatile 244 compounds (Table 5). Indeed, the contribution of this non-Saccharomyces yeast was clear for the ethyl butyrate and β -phenyl ethanol levels, where the trend was strictly linked to the 245 inoculum ratios. The mixed fermentation at the inoculum ratio of 1:20 showed ethyl butyrate 246 levels comparable to that for the T. delbrueckii pure cultures (0.185 vs. 0.168 mg l^{-1}), while 247 for the inoculation ratio of 1:1 the ethyl butyrate levels were comparable to S. cerevisiae US-248 05 pure culture (0.339 vs. 0.319 mg l^{-1}). This evolution was also particularly evident for β -249 phenyl ethanol. Indeed, the increasing T. delbrueckii in the mixed fermentations resulted in 250 251 volatile β -phenyl ethanol levels that were lower or higher when compared with the pure cultures of S. cerevisiae US-05 or T. delbrueckii, respectively. As regards the phenyl ethyl 252 acetate, the results showed an increase in mixed fermentations, particularly in 1:1 inoculum 253 ratio. Different behaviors were seen for ethyl hexanoate and ethyl octanoate. Indeed, there 254

were generally higher levels of these compounds in the mixed fermentations, and particularlywith the high inoculation ratio (i.e., 1:20).

257 *3.2.5. Sensory analysis*

The beers obtained by these pure and mixed fermentations underwent sensory analysis, with 258 the data illustrated in Figure 3. All of the beers analyzed showed significant differences for 259 their main aromatic notes. In particular, for the main sensorial descriptors, the data showed 260 261 that the beer obtained with all of the mixed fermentations and for the T. delbreuckii pure cultures were significantly different from those of the S. cerevisiae US-05 starter strain for a 262 variety of the sensorial characteristics. Within the mixed fermentations, with the 1:1 inoculum 263 264 ratio, there was a bouquet with notes that particularly emphasized the cereal, toasted and fullbodied sensorial attributes, while the fruity/ ester notes were poorly pronounced. At the 1:10 265 inoculum ratio, there was a high perception of the alcohol/ solvent, malt, caramel and 266 oxidized/ aged attributes, while those for fruit/ citric were lower and watery. Then for the 1:20 267 268 inoculum ratio, the fruity/ester and hop attributes were enhanced. For the T. delbrueckii pure 269 cultures, these were characterized by fruit/ citric notes and the full-bodied attributes, while the alcohol/ solvent, cereal, caramel, oxidized/ aged and astringency attributes were little 270 expressed. In addition, the beers produced by the T. delbrueckii pure cultures and with the 271 272 inoculum ratio of 1:20 were characterized by a pale yellow colour, clarity, and persistent and compact foam, which are very important features in the assessment of the quality of a beer 273 (data not shown). 274

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²⁷⁶ **4. Discussion**

In the present study, the use of non-conventional yeast to produce beers with distinctive bioflavours and reduced alcohol contents was investigated. Fermented beverages with improved flavours can be profitably obtained through the yeast during the fermentation process (Carlquist et al., 2015). Indeed, among the volatile compounds produced by yeast, the higher alcohols and esters are crucial to the definition of the final quality of a beer (Pires et al., 2014).

284 Of particular interest here was the production of the three esters phenyl ethyl acetate (i.e., floreal, honey, sweet) ethyl hexanoate (i.e., apple, fruit flavours) and ethyl octaonate (i.e., 285 apple, aniseed flavours), which showed increases for the mixed fermentations, possibly 286 287 related to the increase in consumption of YAN. These data were only in part consistent with the results obtained by sensory analysis. Indeed, the panel tests showed that the beers made 288 with T. delbrueckii showed fruity/esters comparable with those obtained with S. cerevisiae 289 US-05 while they were characterized for malt, caramel and hop attributes. This might be 290 explained considering that some esters that are characterized by a powerful odour can act in 291 292 synergy with other compounds, thus influencing the final beer flavor at concentrations below the threshold of perception (Meilgaard, 1975). 293

The formation of β -phenyl ethanol, diethyl succinate, ethyl butyrate and isoamyl acetate, relates directly to the inoculation levels, and thus to the consequent metabolic expression levels of each of the two strains in the mixed fermentations. These results clearly showed that *T. delbrueckii* more significant affects the analytical and aromatic profile of beers when the inoculation ratios for *S. cerevisiae* US-05 to *T. delbrueckii* were at 1:10 or 1:20.

Over the years, several studies have shown that the levels of the aroma compounds can be increased or decreased through modifications to the constituents of the wort, such as the nitrogen content (Hernàndez-Orte et al., 2005, Igyor et al., 2001). In this context, we found

that in mixed fermentations (S. cerevisiae/T.delbrueckii) exhibiting a higher consumption of 302 303 YAN this could be related to the production of aromatic compounds (Carrau et al., 2015). Also the use of commercial enzyme preparations, such as proteases that act on certain 304 305 proteins in the malt can promote increased contents of the higher alcohols (Piddocke et al., 2011; Treimo et al., 2008). However, the results of the present study show that modifications 306 of the volatile profile of beers can be easily obtained without any modifications to the 307 308 brewing process or the use of enzymes, but instead through the use of non-conventional yeast, such as *T. delbrueckii*. 309

310 Another important feature of the use of T. delbrueckii in this brewing process might be the 311 production of beer with a low alcohol content. Indeed, nowadays, as well as considering the aromatic profile of a beer, consumers pay more attention to health-related problems. The 312 biological approaches that have been proposed to reduce the ethanol content in beer mainly 313 rely on the selection of strains with particular properties or on the modification of the brewing 314 yeast through genetic engineering (Remize et al., 1999; Verstrepen et al., 2003). Taking 315 316 advantage of fermentative performance, some practices are based on the use of S. cerevisiae which is also generally used in winemaking. 317

Another yeast species that has been proposed for the production of beer with a low alcohol 318 319 content on an industrial scale is Saccharomycodes ludwigii (Huige et al., 1990). In this regard, however, conflicting data on the volatile profiles of the beers have been reported (De 320 Francesco et al., 2015; Narziss et al., 1992). In the present study, we have shown instead that 321 T. delbrueckii used in mixed fermentations had little or no effect on the ethanol content in 322 comparison with the S. cerevisiae US-05 pure cultures. In contrast, T. delbrueckii in pure 323 cultures showed a substantial, almost 50%, reduction in the ethanol content that was 324 accompanied by a pleasant and valuable sensorial profile. In this regard, other T. delbrueckii 325

strains initially selected in this study as maltose fermenters might be of interest for low-alcohol beer production.

In conclusion, beers produced with T. delbrueckii in mixed fermentations and in pure cultures 328 show important features that are relevant for brewing beer. In particular, the beer obtained 329 using T. delbrueckii pure cultures showed a low alcohol content while maintaining at the 330 331 same time a pleasant and aromatic taste, a lighter colour, and a compact and persistent foam. On the other hand, T. debrueckii in mixed fermentations with S. cerevisiae US-05 can fully 332 convert the fermentable sugars (63- 64% of real attenuation), thus resulting in a final ethanol 333 content that is comparable to that of the control S. cerevisiae US-05 starter strain, while 334 335 showing distinctive analytical and aromatic profiles.

336

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Table 1. *Torulaspora delbrueckii* strains used in this study.

Source of isolation	Geographic origin	Strain code ^a
Winery environment	Sardinia, Italy	315, 130
Soil	Italy	55
Papaya leaves	Cameroon	254, 419, 343, 413, 255
Sugar cane juice	Cameroon	426, 430, 431, 432, 363, 603
Grapes	Italy	258, 259, 260, 261, 313, 606, 607, 608, 609
Fig fruit	Italy	604, 605
Coconut palm	Cameroon	445
Corrosol fruit	Cameroon	602, 399

⁴⁸⁴ ^a Accession number of DiSVA Collection (Department of Life and Environmental Sciences).

DICVA and	Total CO ₂ evolved	Fermentation rate		
DiSVA code	$(g [30 days]^{-1})^{a}$	$(g CO_2 day^{-1})^b$		
254	13.99 ±0.68	0.96 ± 0.05		
602	5.87 ±1.04	0.68 ±0.10		
603	4.92 ±0.37	0.61 ±0.04		
343	5.94 ± 0.80	0.68 ±0.11		
399	4.24 ±0.20	0.51 ±0.03		
413	4.69 ±0.38	0.59 ± 0.01		
419	4.62 ± 0.08	0.55 ± 0.02		
426	4.48 ±0.40	0.61 ±0.06		
S. cerevisiae ^c	20.97 ± 0.58	3.34 ±0.04		

Table 2. Fermentation kinetics parameters of the eight *T. delbrueckii* strains and the *S. cerevisiae* US-05 starter strain for the wort.

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492 Data are means ±standard deviations

493 $^{a}CO_{2}$ g evolved after 30 days of fermentation (in 500 ml wort)

^bFermentation rate: CO₂ g/day (over the first 6 days of fermentation)

495 ^cCommercial strain: *S. cerevisiae* US-05 (Fermentis, Lesaffre, France)

Fermentation	Ratio	Re	esidual sugar ($(\mathbf{g} \mathbf{l}^{-1})^{\mathbf{a}}$	Analytical characteristic						
		Sucrose	Glucose	Maltose	YAN (mg Γ ¹) ^b	рН	Ethanol (% v/v)	Wort gravity attenuation (°P) ^c	Apparent attenuation (%)	Real attenuation (%)	Volatile acidity (g l ⁻¹)
S. cerevisiae US-05	Pure	0.02 ± 0.01	0.01 ± 0.00	0.05 ± 0.01	193.84 ±0.39	4.80 ± 0.03^{a}	4.85 ± 0.00^{a}	2.84 ± 0.00^{b}	78.40 ± 0.00^{a}	63.73 ± 0.00^{a}	$0.22 \pm 0.00^{\circ}$
T. delbrueckii DiSVA254	Pure	0.42 ±0.01	0.00 ±0.00	35.43 ±0.31	174.58 ±3.57	$4.56\pm0.01^{\circ}$	2.66 ± 0.05^{d}	7.51 ±0.00 ^a	45.09 ± 0.00^{b}	36.65 ± 0.00^{b}	0.15 ± 0.02^{d}
S. cerevisiae US-05 +	1:1	0.02 ± 0.01	0.02 ± 0.01	0.33 ± 0.00	141.79±1.96	4.61 ± 0.01^{b}	$4.51 \pm 0.00^{\circ}$	2.96 ± 0.17^{b}	77.44 ± 0.01^{a}	62.76 ± 0.01^{a}	$0.23 \pm 0.01^{\circ}$
T. delbrueckii DiSVA254	1:10	$0.08\pm\!0.00$	0.03 ± 0.00	0.75 ± 0.05	144.32 ±2.41	4.43 ± 0.01^d	4.68 ± 0.03^{b}	2.84 ± 0.00^{b}	$78.40\pm\!\!0.00^a$	63.73 ± 0.00^{a}	0.26 ± 0.00^{b}
	1:20	0.10 ± 0.01	0.04 ± 0.02	1.65 ± 0.10	142.87 ± 2.41	4.38 ± 0.00^{d}	4.59 ± 0.07^{bc}	2.96 ± 0.17^{b}	77.44 ± 0.01^{a}	62.76 ± 0.01^{a}	0.29 ± 0.00^{a}

Table 3. The main analytical characteristics of the beer produced by the pure and mixed fermentations.

^aThe initial composition of the sugars in wort were: Sucrose 6.4 g l⁻¹; glucose 8.5g l⁻¹; Maltose 64.06 g l⁻¹

^bThe initial yeast assimilable nitrogen (YAN) were 251 mg l^{-1}

^cThe wort gravity at the start was 12.7 °P.

Data are means± standard deviation

Data with different superscript letters $(^{a,b,c,d})$ within each column are significantly different (Duncan tests; P <0.05).

Table 4. The main by-products in the beer produced by the pure and mixed fermentations.

Fermentation	Ratio	By-products (mg l ⁻¹)						
		Acetaldehyde	Ethyl acetate	n-propanol	Isobutanol	Amylic alcohol	Isoamylic alcohol	
S. cerevisiae US-05	Pure	5.80 ± 0.45^{d}	17.57 ±0.24 ^a	30.56 ± 0.29^{a}	24.38 ±0.31 ^a	12.69 ± 0.22^{a}	68.53 ± 0.08^{a}	
T. delbrueckii DiSVA254	Pure	$7.50 \pm 0.40^{\circ}$	3.46 ± 0.40^{d}	15.41 ± 0.04^{e}	7.98 ±0.10 ^d	$3.82 \pm 0.04^{\circ}$	32.79 ± 0.42^{e}	
S. cerevisiae US-05 +	1:1	10.72 ± 0.13^{b}	16.75 ±0.06 ^b	26.36 ± 0.29^{b}	18.45±0.32 ^b	10.60 ± 0.13^{b}	$58.69 \pm 0.35^{\circ}$	
T. delbrueckii DiSVA254	1:10	30.48 ± 0.22^a	$14.65 \pm 0.04^{\circ}$	20.14 ± 0.21^d	$16.04 \pm 0.34^{\circ}$	10.61 ± 0.60^{b}	55.65 ± 0.47^d	
	1:20	5.56 ± 0.15^d	16.25 ± 0.16^{b}	$22.59 \pm 0.06^{\circ}$	$16.15 \pm 0.24^{\circ}$	11.29 ± 0.19^{b}	$60.05 \ {\pm} 0.06^{b}$	

Data are means \pm standard deviations

Data with different superscript letters $(^{a,b,c,d,e})$ within each column are significantly different (Duncan tests; P <0.05).

Table 5. The main volatile compounds in the beer produced by the pure and mixed fermentations.

Fermentation	Ratio	o Main volatile compounds (mg l ⁻¹)							
		Ethyl butyrate	Isoamyl acetate	Ethyl Ethyl hexanoate octanoate		Butyric acid	Diethyl succinate	phenyl ethyl acetate	β-phenyl ethanol
S. cerevisiae US-05	Pure	0.319±0.01 ^a	0.346±0.045 ^a	0.037±0.010 ^b	0.007±0.001 ^b	0.158±0.033 ^{ab}	0.016±0.004 ^{ab}	0.001±0.001 ^b	40.77±0.444 ^a
T. delbrueckii DiSVA254	Pure	0.168±0.00 ^c	0.134 ± 0.005^{d}	0.031±0.013 ^b	0.006±0.003 ^b	0.074±0.028 ^{bc}	ND	ND	6.52±0.038 ^e
S. cerevisiae US-05 +	1:1	0.339±0.018 ^a	0.321±0.001 ^{ab}	0.023±0.011 ^c	0.009±0.003 ^{ab}	$0.184{\pm}0.058^{a}$	0.039±0.022 ^a	0.008±0.129 ^a	30.69±0.018 ^{ab}
T. delbrueckii DiSVA254	1:10	0.26±0.03 ^b	0.258±0.017 ^{bc}	0.019±0.004 ^c	0.014±0.001 ^a	0.119±0.021 ^{abc}	0.023±0.010 ^{ab}	0.004±0.001 ^a	27.53±0.035°
	1:20	0.185±0.00 ^c	0.227±0.030 ^c	0.048±0.011 ^a	0.009 ± 0.001^{b}	$0.047 \pm 0.022^{\circ}$	0.010 ± 0.007^{b}	0.003±0.001 ^a	15.48±0.115 ^d

Data are means ±standard deviations

Data with different superscript letters $(^{a,b,c,d})$ within each column are significantly different (Duncan tests; P <0.05).

ND : not detected

Figure captions

Figure 1. Fermentation kinetics of the pure and mixed fermentations. Pure cultures of *S. cerevisiae* (--=-) and *T. delbrueckii* (--=-), and mixed cultures of *S. cerevisiae* + *T. delbrueckii* at 1:1 (-==-), 1:10 (-==-) and 1:20 (-==-).

Figure 2. Growth kinetics of the pure and mixed fermentations. Pure cultures of *S. cerevisiae* (- - -) and *T. delbrueckii* (-- - -) (A-C), and of the mixed fermentation with *S. cerevisiae* (-- -) and *T. delbrueckii* (-- -) individually for the mixed cultures at 1:1 (A), 1:10 (B) and 1:20 (C).

Figure 3. Sensory analysis of the beer produced by the mixed fermentations. From pure cultures of *S. cerevisiae* (- - -) and *T. delbrueckii* (- - -), and mixed cultures of *S. cerevisiae* + *T. delbrueckii* at 1:1 (- - -), 1:10 (- - -) and 1:20 (- - -). *, Significantly different (Fisher ANOVA; p-value 0.05). DMS, Dimethyl sulfide.



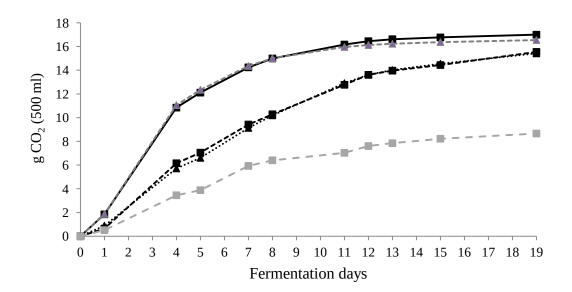
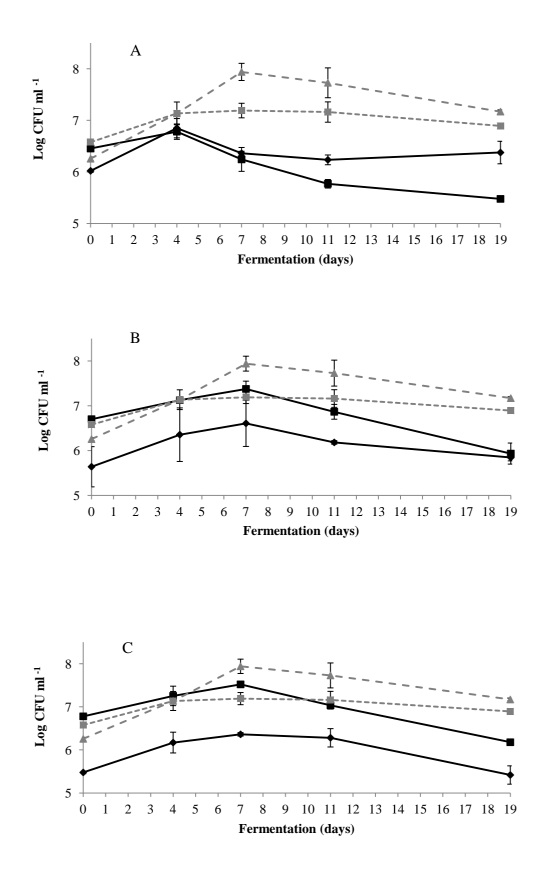




FIG 2



Figure

FIG 3

