1	Microbial diversity of type I sourdoughs prepared and back-sloped with wholemeal and refined soft (Triticum
2	aestivum) wheat flours
3	
4	Short version of the title: biodiversity of type I sourdoughs
5	
6	Manuela Taccari, Lucia Aquilanti*, Serena Polverigiani, Andrea Osimani, Cristiana Garofalo, Vesna Milanović, and
7	Francesca Clementi
8	
9	Dipartimento di Scienze Agrarie Alimentari ed Ambientali (D3A), Università Politecnica delle Marche, via Brecce
10	Bianche, 60131 Ancona, Italy
11	
12	
13	
14	
15	
16	
17	
18	
19	
20	
21	
22	
23	
24	
25	
26	
27	
28	
29	* Corresponding author at: Dipartimento di Scienze Agrarie, Alimentari ed Ambientali, Università Politecnica delle
30	Marche, via Brecce Bianche, 60131, Ancona, Italy. Tel.: +39 071 2204959; fax.: +39 071 2204988. E-mail address:
31	<u>l.aquilanti@univpm.it (L. Aquilanti)</u>

32 Abstract: Type I sourdough fermentations were carried out for 20 days with daily back-slopping under laboratory and 33 artisan bakery conditions using one wholemeal and two refined soft wheat (Triticum aestivum) flours, preliminarily subjected to chemical and rheological testing. The bacterial and yeast diversity and dynamics were investigated though 34 plate counting and a combined culture-dependent and -independent PCR-DGGE approach; pH, total titrable acidity and 35 36 concentration of key organic acids (phytic, lactic and acetic) were investigated through conventional tests. The three 37 flours differed for both chemical and technological traits. Regarding the establishment and stabilization of microbial 38 consortia during the continuous sourdough propagation, a microbial succession was seen, with atypical sourdough 39 species detected at day 0 (such as f.i. Lactococcus lactis and Leuconostoc holzapfelii/citreum group for bacteria, or 40 Candida silvae and Wickerhamomyces anomalus for yeasts) progressively replaced by taxa more adapted to the 41 sourdough ecosystem (Lactobacillus brevis, Lb. alimentarius/paralimentarius, Saccharomyces cerevisiae). In the 42 mature sourdoughs, a quite different species composition was seen. As sourdoughs propagated with the same flour at 43 laboratory and artisan bakery level were compared, an influence of both the substrate and the propagation environment 44 on the sourdough microbial diversity was assumed.

45

46 Practical Applications: Foods manufactured with wholemeal flours boast numerous health benefits, mainly due to 47 their high content in bran fibre; however, these flours are poorly exploited in bread-making due to their scarce leavening 48 ability and their richness in anti-nutrients (phytates). The use of wholemeal sourdoughs might represent a valid 49 alternative to bakers' yeast for the future development of technologically and nutritionally superior raw materials for all 50 cereal foods, such as bread, breakfast and snack products manufacture.

51

52 Keywords: High rate extraction rate flour, high-fibre sourdough, high-fibre bread-making, sourdough microbiota, PCR53 DGGE.

- 54
- 55
- 56
- 57
- 58
- 59
- 60
- 61
- 62

## 63 Introduction

64 Soft wheat (*Triticum aestivum*) is the most widely used cereal for the production of bakery products and, in particular,
65 bread. The latter can be manufactured with both high and low extraction rate flours; a 100% extraction rate flour
66 corresponds to wholemeal flour containing all the caryopsis components, whereas low extraction rate flours are refined,
67 whiter flours, commonly classified as type 00 or 0, which are progressively deprived of increasing amounts of bran and
68 germ, and thus B vitamins, fat, iron and other minerals.

69 Refined flours have a higher aptitude to leavening and are preferred to wholemeal flours for the manufacture of some 70 baked leavened products, such as special occasion cakes (i.e. Panettone, Pandoro or Colomba) (Garofalo and others 71 2008; Garofalo and others 2011). On the contrary, wholemeal flours have a lower gluten content, which explains the 72 production of doughs with a lower stability, resistance and extensibility (Azizi and others 2006), but are richer in 73 phytates (Buddrick and others 2014), which sequester and, hence, prevent the intestinal absorption of some essential 74 minerals, such as calcium, zinc, iron and magnesium (Kumara and others 2010). However, wholemeal flours are richer 75 in bran fiber, which confers not only improved sensory and nutritional properties to bread but also health benefits, such 76 as the reduction of blood cholesterol and glycemic index, and the promotion of weight loss by increased satiety 77 (Stevenson and others 2012). Previous studies clearly demonstrated that the use of high extraction rate flours also leads 78 to a reduction in the bread staling rate (Venkateswara and others 1985; Azizi and others 2006).

79 In ancient times unrefined flours were typically processed with sourdough (Poutanen and others 2009). Sourdough is a 80 biological ecosystem, composed of lactic acid bacteria (LAB) and yeasts, which may be deliberately added to doughs 81 made of flour and water or spontaneously selected during sourdough continuous propagation (De Vuyst and others 82 2014). The use of sourdough has undoubtedly a great potential for the manufacture of leavened bakery products with 83 increased sensory properties, nutritional value, and prolonged shelf-life (Gobbetti and others 2014). Doughs started with 84 sourdoughs are characterized by more balanced biochemical (f.i. fermentation quotient, profile of volatile compounds, 85 etc.) and rheological (f.i. structural stability) traits than doughs started with the sole baker's yeast, where alcoholic 86 fermentation end-products largely predominate (Cossignani and others 1996). In the case of doughs prepared with 87 wholemeal flours, the use of sourdough appears particularly advantageous for the activation of endogenous phytases 88 during prolonged fermentation (Buddrick and others 2014), which directly implies the improvement of bread mineral 89 bioavailability (Lopez et al., 2003). Leavening and bread-making trials conducted with flours characterized by low 90 technological aptitudes, such as barley (Zannini and others 2009; Mariotti and others 2014) or even gluten-free flours 91 (Arendt and others 2007) demonstrated the usefulness of sourdough technology for the improvement of bread qualities. 92 In the past decades, this technology was successfully explored for the improvement of volume and shelf-life of high-

93 fibre breads (Salmenkallio-Marttila and others 2001; Katina and others 2005) and a new type of sourdough, consisting

94 of wheat bran pre-fermented with selected yeasts and lactic acid bacteria, was supplemented to bread doughs, with95 encouraging results.

96 The composition of the sourdough microbiota plays a main role for taste and quality of baked goods. For this reason, 97 scientific research focuses largely on the characterization of microbiota of sourdoughs fermented under different 98 conditions, on selection of suitable starter cultures for the baking industry, and on metabolic properties of sourdough 99 microorganisms. Despite intense research conducted in the last decades, it is still unclear to which extent the sourdough 90 microbiota is affected and selected by the kind of substrate, the process parameters (f.i. temperature and number of 91 refreshments, dough yield, etc.), and the establishment of diverse microbial interactions.

102 This study deals with the question how the composition of the sourdough microbiota is influenced by the substrate and 103 the sourdough propagation conditions. More specifically, the focus was laid on both the flour extraction rate and the 104 back-sloping environment. Based on the above premises, this study was aimed at investigating the biochemical (pH, 105 total titratable acidity, concentration of phytic acid (phytate), lactic acid and acetic acid, and fermentation quotient) and 106 microbiological traits of spontaneously fermented sourdoughs prepared and propagated daily at artisan and laboratory 107 scale with wholemeal soft wheat (Triticum aestivum) flour, with a view to their further exploitation in high-fibre bread-108 making. Conventional sourdoughs produced with refined flour from soft wheat grains processed by roll milling or stone 109 grinding were also analyzed, in parallel. Preliminary chemical, farinograph, alveograph, and micro-visco-amylograph 110 tests were also conducted on the three flours whereas the evolving microbial diversity of the high-fibre and 111 conventional type I sourdoughs were investigated through a combined PCR-DGGE approach, relying on the analysis of 112 the bacterial and yeast DNA extracted directly from the (sour)dough samples and the bulks of colonies harvested from 113 selected dilution agar plates used for viable counting.

114

## 115 Materials and Methods

116

# 117 Composition and traits of the three flours

118 Three soft wheat (*T. aestivum*) flours designed for making bread were used in this study. Wheat was cultivated during 119 the growing season 2013-2014, at the farmers' cooperative "Il Biroccio" (Filottrano, Ancona, Italy). A wholemeal 120 (hereafter referred to as " $F_{A"}$ ) and two type 0 (hereafter referred to as " $F_B$ " and " $F_C$ ") flours were obtained through steel 121 roller milling ( $F_A$  and  $F_B$ ) or stone grinding ( $F_C$ ). The grains were thoroughly cleaned by winnowing, sieving and 122 sorting. Flours were kept in sealed containers at -20 °C before use.

- 123
- 124 Chemical analyses

125 For each flour, the following chemical analyses were performed according to international standard methods: moisture

126 (UNI EN ISO 712:2010); ash (UNI-ISO 2171:2010); falling number (UNI EN ISO 3093:2007); and protein (UNI EN

127 ISO 20483:2007). Commercial kits were used for the determination of phytic acid (phytate/total phosphorous) (K128 PHYT 12/12, Megazyme, Wicklow, Ireland).

129

130 Rheological analyses

The three flours were subjected to farinograph (UNI 10790:1999), alveograph (UNI EN ISO 29971:2008), and microvisco-amylograph tests (UNI 10872:2000), according to international standard methods, using the following equipment:
Chopin alveograph (alveo-consistograph with alveolink NG, Chopin, Villeneuve-la-Garenne, France), Brabender
farinograph (Farinograph-E, Brabender, Duisburg, Germany), Brabender micro-visco-amylograph (Micro visco-amylograph, Brabender, Duisburg, Germany).

136

137 Propagation of type I sourdoughs

138 Type I sourdough preparation was started as follows: 100 mL of tap water and 150 g of the flours F<sub>A</sub>, F<sub>B</sub> or F<sub>C</sub> were 139 mixed to produce 250 g of dough [dough yield (dough weight  $\times$  100/flour weight), 166], according to a traditional 140 protocol (Zannini and others 2009) without the use of starter cultures or baker's yeast. The doughs were kneaded 141 manually for 5-10 min until the correct consistency was obtained and fermented for 24 h in glass containers covered 142 with a lid. The resulting sourdoughs were propagated daily over a period of 20 days according to the back-slopping 143 procedure, where 100 g of the sourdough from the day before was used as the inoculum [40% (w/w)] of a new mixture 144 of flour (100 g) and tap water (50 mL) (dough yield, 156). Sourdough propagation was performed at both artisan bakery 145 and laboratory level, at room temperature; in the first case, it was performed at an artisan bakery in the Marche region 146 (central Italy), where the sourdough technology had never been used before, and the manufacture of bread is based on 147 the use of compressed bakers' yeast. Sourdoughs propagated at laboratory and artisan bakery level with the three 148 flours F<sub>A</sub>, F<sub>B</sub> and F<sub>C</sub> are hereafter referred to as "S<sub>A</sub>-l", "S<sub>B</sub>-l", and "S<sub>C</sub>-l" and "S<sub>A</sub>-b", "S<sub>B</sub>-b", "S<sub>C</sub>-b", respectively.

Sampling was performed on the first unfermented doughs ( $t_0$ ) and the sourdoughs immediately before each daily refreshment step; for each type of sourdough, a total of 21 samples were collected, cooled to 4 °C and analyzed in duplicate within 2 h of collection. At each refreshment step all samples were subjected to the measurement of pH and total titratable acidity (TTA), whereas the sole samples collected after 0 ( $t_0$ ), 5 ( $t_5$ ), 10 ( $t_{10}$ ), and 20 ( $t_{20}$ ) days of daily

back-slopping were subjected to both plate counting of presumptive LAB and yeasts and PCR-DGGE analyses.

154 Organic acids content of laboratory and artisan bakery sourdoughs was determined at the end of the 20-day propagation.

156 Viable counts of presumptive lactic acid bacteria (LAB) and yeasts

157 LAB and yeast cell counts were determined by mixing 10 g of each dough and sourdough sample with 90 mL of a 158 sterile aqueous peptone solution (0.1% peptone, w/v), followed by homogenization with a Stomacher apparatus (400 159 Circulator, International PBI, Milan, Italy) at 260 rpm for 3 min. Presumptive lactobacilli were counted on modified de 160 Man Rogosa Sharpe (mMRS) agar, containing 1% maltose (w/v) plus 5% fresh yeast extract (v/v) and Sourdough 161 Bacteria (SDB) medium modified according to Vogel et al. (1994), (hereafter referred to as mSDB), the latter being 162 elective for Lactobacillus sanfranciscensis (Foschino and others 2004). Both media were supplemented with 163 cycloheximide (0.1 g L<sup>-1</sup>) to inhibit the growth of yeasts. Plates were incubated in sealed jars with the Anaerogen system (Oxoid, Basingstoke, UK) at 30 °C for 48 h. Yeasts were counted on Wallestein Laboratory Nutrient (WLN) 164 165 medium added with chloramphenicol (0.1 g L<sup>-1</sup>) to inhibit the growth of bacteria and incubated at 25 °C for 72 h. The 166 results of the viable counts were expressed as means of the Log of colony forming units (cfu) per gram of sample  $\pm$ 167 standard deviation. After counting, plates spiked with the highest (-6, -7 or -8) and lowest dilution (-2) were used for 168 bulk formation, following the procedure previously described by Garofalo and others (2015).

169

Determination of pH, total titratable acidity (TTA), organic acids (phytic, lactic and acetic acid), and fermentationquotient (FQ)

172 The pH measurements on doughs and sourdoughs were carried out with a pH meter equipped with a solid electrode 173 (HI2031, Hanna Instruments, Padua, Italy) which was inserted directly into the sample. Total titratable acidity (TTA) 174 was measured on 10 g of dough or sourdough homogenized with 90 mL of distilled water using a Stomacher apparatus 175 (International PBI). TTA was expressed as the amount (mL) of 0.1 N NaOH needed to achieve a final pH of 8.5. Lactic 176 and acetic acids were determined in the water-soluble extracts of the 20-day back-slopped sourdoughs, according to the 177 following procedure. Ten grams of each sourdough were homogenized with 90 mL of Tris-HCl 50 mM buffer (pH 8.8). 178 After incubation at 25 °C for 30 min under stirring the suspension was centrifuged at 4.000 rpm for 10 min at 4 °C, and 179 the supernatant was analyzed using the commercial kits K-ACETIC 02/11 and K-DLATE 12/12 (Megazyme). The 180 fermentation quotient (FQ) was determined as the molar ratio between D,L-lactic and acetic acids. Phytic acid 181 (phytate/total phosphorous) was determined using the commercial kit K-PHYT 12/12 (Megazyme).

182

183 Bacterial and fungal DNA extraction from (sour)doughs and cell bulks

184 Microbial DNA extraction from the dough and sourdough samples was performed using the PowerFood<sup>™</sup> Microbial 185 DNA Isolation Kit (Mo Bio Laboratories, Carlsbad, USA) following the kit manufacturer's instructions. DNA 186 extraction from the LAB and yeast bulk cell suspensions was performed according to the procedures described by Hynes and others (1992) and Makimura and others (1999), respectively. The procedure indicated by Hynes and others
(1992) was slightly modified as described by Osimani and others (2015). All the DNA suspensions were subjected to
optical readings at 260, 280 and 234 nm with a UV-Vis Shimadzu UV-1800 Spectrophotometer (Shimadzu
Corporation, Kyoto, Japan) for the assessment of the DNA quantity and purity.

191

192 Amplification reactions

Two microliters of each microbial DNA suspension (adjusted to 25 ng/ $\mu$ L) were amplified in 25  $\mu$ L reaction volume using bacterial universal primer sets (338f<sub>GC</sub>- 518r), targeting the variable regions V3 of the 16S rRNA gene, as previously described (Osimani and others 2015). Two microliters of each microbial DNA suspension (adjusted to 25 ng/ $\mu$ L) were amplified in 25  $\mu$ L reaction volume using universal primers for the eumycetes (NL1<sub>GC</sub>-LS2), as previously described (Osimani and others 2015). All the PCR reactions were performed in the thermal cycler My Cycler (Bio-Rad Laboratories, Segrate, Italy); aliquots (5  $\mu$ L) of the PCR products were routinely checked on 2% agarose (w/v) gels, prior to PCR-DGGE analysis.

200

201 Polyacrylamide gel preparation and DGGE runs

PCR products were analyzed using the DGGE Bio-Rad D-code<sup>™</sup> apparatus (Bio-Rad Laboratories). DGGE runs were performed in 8% polyacrylamide (acrylamide/bis-acrylamide mix 37.5:1, w/v) gels with a 30-60% urea-formamide (w/v) gradient. Gels were subjected to a constant voltage of 130 V for 4 h at 60 °C. After electrophoresis, gels were stained in 1X TAE buffer containing SYBR Green I (1× final concentration; Molecular Probes, Eugene, OR, USA) and photographed under UV transillumination, using the Complete Photo XT101 system (Explera, Jesi, Italy).

207

208 Band sequencing and identification

209 For DGGE band sequencing, bands of interest were excised by gel cutting tips, resuspended in 50 µL of sterile water, 210 and incubated overnight at 4 °C to allow diffusion of DNA from the gel cuts. Five microliters of the eluted DNA were 211 re-amplified using the conditions described above, with the forward primer without the GC clamp and sent to Beckman 212 Coulter Genomics (Hope End, Takeley, United Kingdom) for purification and sequencing. The sequences obtained in 213 FASTA format were compared with those deposited in the GenBank DNA database (http://www.ncbi.nlm.nih.gov/) 214 using the basic blast search tools. An identity of the partial 16S and 26S rRNA gene sequences to type strain sequences 215  $\geq$  97% was used as the criterion for species identification. Due to the slowly diverging nature of the 16 rRNA gene 216 (Felis and Dellaglio 2007), a few sequences could not be univocally assigned to a single species, but to a microbial 217 group, including strictly correlated species, namely the Lactobacillus graminis/curvatus group, the Lactobacillus

alimentarius/paralimentarius group, the Lactobacillus pentosus/plantarum/paraplantarum group, the Weissella confusa/cibaria group, and the Leuconostoc holzapfelii/citreum group. Similarly, a discrimination between phylogenetically closely related species, namely Kazachstania unispora, Kazachstania servazzii, Kazachstania aerobia was not achieved through the analysis of the D1 region of yeast DNA, as previously reported (Garofalo and others 2015).

223

#### 224 Statistical analysis

225 Chemical and rheological properties of three flours were analyzed by one-way ANOVA. All the data referring to dough 226 and sourdough microbial communities (viable counts of LAB and yeasts) and biochemical traits (pH, TTA, 227 concentration of phytic acid, lactic acid and acetic acid, FQ) were subjected to two-way ANOVA focusing on both the 228 influence of the flour and the sourdough propagation environment, and on their cross interaction. Due to the presence of 229 a cross interaction for several parameters a Tukey's HSD test at P < 0.05 was performed for mean separation of all 230 possible combinations of the two factors. Principal component analysis (PCA) using a correlation matrix was carried 231 out to visualize the effects of the flour and the sourdough propagation environment on the above parameters. Finally, a 232 two way cluster permutation test was performed using all the data referring to microbial communities and biochemical 233 traits, and used to overlay confidence ellipses on the PCA score plots. For cluster analysis the Ward's method was 234 applied. All the statistical analyses were performed using JMP 9 (SAS Institute Inc., Cary, NC).

235

# 236 Results and Discussion

237

# 238 Chemical and rheological traits of the three soft wheat flours

The results of chemical and rheological analyses carried out on the three flours are shown in Table S1 in the supplementary material. With the exception of moisture, for all the chemical traits investigated,  $F_A$  differed from  $F_B$  and  $F_C$ . Analogously, differences were seen in some rheological properties of wholemeal ( $F_A$ ) and type 0 flours ( $F_B$  and  $F_C$ ), namely consistency, water absorption, dough development time, mixing tolerance index after 10 min, maximum viscosity, falling number, phytic acid.

It has previously been reported that the reduced moisture of high extraction rate flours is mainly dependent on their content in wheat bran, the latter being characterized by a lower moisture with respect to aleuronic particles (Azizi and others 2006). However, in this study, no significant reduction in such a quality parameter was seen in  $F_A$  compared with  $F_B$  and  $F_C$ . According to what was reported by Azizi and others (2006), even the significantly lower falling number of  $F_A$ , which indicates a higher alfa-amylase activity, and the higher ash and protein content of this flour might be directly imputable to its higher bran content. It has previously been reported that the alfa-amylase activity of wheat flour depends on its extraction rate and quality, with wholemeal flour, and especially the bran fraction, having the highest enzyme activity (Martinez-Anaya 2003). As expected,  $F_A$  was also characterized by a higher phytic acid content than the refined flours  $F_B$  and  $F_C$ . In fact, it is known that the flour refining process reduces the levels of phytate (Buddrick and others 2014), which are concentrated in the aleuronic layers of grains (Poutanen and others 2009).

254 As far as the rheological traits of the three flours are concerned, a high water absorption, which significantly differed 255 from that of  $F_B$  and  $F_C$ , was seen in  $F_A$ . The occurrence of a linear correlation of this trait with the flour extraction rate 256 has previously been elucidated by Azizi and others (2006), who reported an increase in this parameter in high extraction 257 rate flours, which are generally characterized by coarser particle sizes. Even the weaker resistance of  $F_A$  to mechanical 258 shocks (revealed by a significantly lower mixing tolerance index after 10 min) and the markedly higher dough 259 development time might be attributed to the higher content in wheat bran, which translates into a reduction in the gluten 260 network and quality. In terms of acidification, the results collected demonstrated that just two refreshment steps were 261 needed to significantly reduce the pH of all the sourdoughs to mean values < 4.5, whereas a higher variability was seen 262 in both the maximum increase in TTA during the 20-day propagation and the number of refreshments needed to reach 263 TTA values > 8.0. It has previously been proposed that sourdoughs achieve their maturity, in terms of constant 264 acidification and leavening capacity, within 5 to 7 days of continuous propagation (Ercolini and others 2013).

265

# 266 Microbial viable counts and acidification kinetics during sourdough propagation

267 After 5 days of propagation at laboratory level, the mean viable counts of presumptive LAB of S<sub>A</sub>-l, S<sub>B</sub>-l, and S<sub>C</sub>-l 268 reached values  $\geq$  9.2 Log cfu/g, irrespective of the growth medium used (mSDB or mMRS), whereas at the same 269 sampling time, the artisan bakery sourdoughs were characterized by mean cell numbers of presumptive LAB > 9.0 Log270 cfu/g ( $S_A$ -b on mMRS;  $S_A$ -b,  $S_B$ -b, and  $S_C$ -b, on mSDB) and > 8.0 Log cfu/g ( $S_B$ -b and  $S_C$ -b on mMRS), respectively 271 (Figure 1). The viable cell counts estimated by plating on mSDB stayed almost constant during subsequent propagation 272 in all the sourdoughs analyzed, whereas those estimated by plating on mMRS showed variable fluctuations, consisting 273 in a slight decrease in laboratory sourdoughs (S<sub>A</sub>-l, S<sub>B</sub>-l, and S<sub>C</sub>-l) and, conversely, a slight increase in artisan bakery 274 sourdoughs (S<sub>A</sub>-b, S<sub>B</sub>-b, and S<sub>C</sub>-b).

No yeasts could be enumerated in all the initial doughs ( $t_0$ ) prepared at laboratory level, considering a detection limit of 2 Log cfu/g, whereas doughs prepared at artisan bakery level contained a significantly (P > 0.001) higher (at least 1.0 log unit) yeast cell density with respect to presumptive LAB (Figure 1). Nevertheless, after 20 days of daily propagation, all the sourdoughs were characterized by a ratio between LAB and yeasts stabilized at ca. 100:1 (Figure 1). In all the sourdoughs analyzed, a pH drop below 4.5 was seen between the 1<sup>st</sup> and the 2<sup>nd</sup> day of propagation, whereas differences were seen in the refreshment step, where pH values  $\leq$  4.0 were reached (remaining almost constant until the end of the 20-day propagation period), namely: days 10, 3 and 7 in S<sub>A</sub>-l, S<sub>B</sub>-l and S<sub>C</sub>-l, respectively, and days 5, 4 and 4 in S<sub>A</sub>-b, S<sub>B</sub>-b and S<sub>C</sub>-b, respectively (Figure 1).

Differences were also seen between the sourdoughs propagated at laboratory and artisan bakery level in the significant maximum increase in TTA reached during the 20-day propagation, namely:  $\geq 20$  mL in S<sub>A</sub>-l and S<sub>A</sub>-b, and  $\geq 11$  mL in S<sub>B</sub>-l, S<sub>B</sub>-b, S<sub>C</sub>-l and S<sub>C</sub>-b, respectively. Analogously, there were also variations in the propagation day when the first significant increase in sourdough TTA was recorded, namely: days 2, 1 and 2 in S<sub>A</sub>-l, S<sub>B</sub>-l and S<sub>C</sub>-l, respectively, and day 2 in S<sub>A</sub>-b, S<sub>B</sub>-b and S<sub>C</sub>-b (Figure 1).

A mean fermentation quotient (FQ) > 5 was seen in all the 20-day back-slopped sourdoughs analyzed, with the exception of  $S_B$ -l, which was characterized by a mean molar ratio of lactic and acetic acid of 1.92. In the two laboratory sourdoughs  $S_B$ -l and  $S_C$ -l, this molar ratio reached levels as high as 13.74 and 12.55, respectively.

291

#### **292** Biochemical characteristics of the mature 20-day back-slopped sourdoughs

293 The values of pH, TTA, concentration of phytic acid, lactic acid and acetic acid, and FQ of the mature high-fibre and 294 conventional type I sourdoughs are shown in Table 1. The mean pH and TTA values recorded at the end of the 20-day 295 propagation were, with a few exceptions, in the typical ranges of 3.6-3.8 and 8-13, respectively (Brummer and Lorenz 296 1991). The significantly higher TTA observed in the sourdoughs propagated with wholemeal flour ( $S_A$ -l and  $S_A$ -b) is 297 feasibly correlated to the higher ash content of  $F_A$  compared with  $F_B$  and  $F_{C_A}$  which has been reported as having an 298 influence on sourdough buffering ability (Mariotti and others 2014). As regards the content in lactic acid and acetic 299 acid, mean values notably higher than those reported for wheat sourdoughs, ranging from 600 to 800 mg/100 g 300 sourdough and from 80 to 160 mg/100 g sourdough, respectively (Barber and others 1992; Hansen and Hansen 1994), 301 were seen. Fermentation quotients exceeding 1 (and in most cases higher than 5) were also seen in all the 20-day back-302 slopped sourdoughs, thus suggesting a predominance of the species with an obligately homofermentative or 303 facultatively heterofermentative metabolism.

Although the phytic acid content of flour and mature sourdough is not directly comparable, at the end of the 20-day continuous propagation, a trend toward the reduction of this parameter was seen. More in detail, a complete hydrolytation of this compound was observed in  $S_B$  and  $S_C$ , whereas a noticeable decrease was seen in  $S_A$  sourdoughs propagated with wholemeal flour ( $F_A$ ), containing the highest amount of this anti-nutrient. Sourdough fermentation has been found to decrease the amount of phytates, either through a mere pH drop to values < 5.5, sufficient to activate the endogenous phytates, or through the phytase activity of LAB and yeasts (Poutanen and others 2009; Mariotti and others 2014). In particular, in whole-wheat flours, the solubilization of minerals due to fermentation has been shown to be
strictly dependent on the bran particle size, being effective in finely milled bran, but not in coarse bran (Lioger and
others 2007).

313

314 Statistical correlations between propagation environment, flour, microbial communities and biochemical traits

The two way ANOVA highlighted a cross interaction of the sourdough propagation environment and the type of flour on TTA (P = 0.0008), acetic acid content (P < .0001), FQ (P = 0.0014), yeast (P = 0.0075) and LAB viable counts (the latter assessed on mMRS) (P < .0001). Both the variability factors considered had a significant effect on all the chemical/microbiological traits analyzed; the only exceptions were represented by phytic acid content, which was not influenced by the sourdough propagation environment, and viable counts of LAB (assessed on mSBD), not influenced by the type of flour (Table 1).

321 PCA indicated eigenvalues of 4.24, 3.44 and 0.79 for the first 3 components, respectively, thus suggesting the adoption 322 of two principle components. These two components accounted for 47.1% and 38.3% of total variance of the data, 323 respectively. The first principle component was dominated by pH, TTA, and content of phytic acid and lactic acid. The 324 second component had the highest scores for acetic acid, FQ, LAB and yeast viable counts. The PCA analysis indicated 325 that the type of flour strongly influenced PC1 whereas the sourdough propagation environment had less influence. 326 Although the correlation with PC2 of the type of flour in the sourdoughs produced at laboratory level was minimal, it 327 was magnified in the sourdoughs propagated at the artisan bakery (Figure 2). According to the permutation test, 328 sourdoughs were distributed into two major clusters (A and B). Irrespective to the propagation environment, cluster B 329 included the sourdoughs refreshed with the flour F<sub>A</sub>, characterized by the highest values of pH, lactic acid content, 330 TTA, and phytic acid content. Cluster A included sourdoughs S<sub>B</sub> and S<sub>C</sub>, whose further clustering (A<sub>I</sub> and A<sub>II</sub>) was 331 influenced by the propagation environment. Acetic acid content and LAB viable counts assessed on mMRS were higher 332 in the sourdoughs propagated at laboratory level whereas FQ and LAB viable counts assessed on mSBD were lower 333 (Figures 2 and 4).

334

**335** PCR-DGGE analyses

336 The results of PCR-DGGE analyses are shown in Figure 3 and in Tables S2 to S5, in the supplementary material.

337 Overall, multiple genera (*Lactobacillus, Leuconostoc, Weissella, Lactococcus,* and *Staphylococcus*) were identified in 338 both the high-fibre and refined-flour sourdoughs with the double PCR-DGGE approach used; these genera correspond 339 to those commonly retrieved in *T. aestivum* flour (Ercolini and others 2013). In almost all the mature sourdoughs 340 analyzed (with the exception of  $S_{c}$ -b) the closest relatives to *Lb. brevis* and the *Lb. alimentarius/paralimentarius* group were stably detected in both the mMRS and mSDB bulks, irrespective of the flour and, hence, the flour extraction rate or the propagation environment. *Lb. brevis* is an obligate heterofermentative species, which is frequently isolated from sourdoughs (De Vuyst and others 2009). This species has been used with some success for the pre-fermentation of wheat bran destined for the manufacture of wheat bread supplemented with bran (Salmenkallio-Marttila and others 2001), thus clearly demonstrating the adaptability and even competitiveness of such a microorganism in high-fibre cereal environments. *Lb. paralimentarius* is another common sourdough inhabitant which has first been isolated from this peculiar ecosystem (Cai and others 1999).

The detection of both these species at the highest dilutions suggests their occurrence at cell densities  $\geq 6$  Log cfu/g. Although newly-developed sequencing techniques, also referred to as next generation sequencing (NGS) methods, have recently been applied to the study of complex food ecosystems, a recent investigation exploiting and comparing NGS and PCR-DGGE clearly demonstrated how this latter technique still represents a valid tool for the profiling of the sourdough core microbiota (Ercolini and others 2013).

In the present study, an apparently flour-dependent occurrence of *Weissella* spp. was seen in  $S_A$ -l and  $S_A$ -b, and in  $S_B$ -l and  $S_B$ -b, whereas for other LAB, a greater influence of the propagation environment might be assumed, as was the case for the sourdoughs refreshed in the laboratory, which were characterized by the establishment of the closest relatives to the *Lb. graminis/curvatus* group (detected in  $S_A$ -l,  $S_B$ -l and  $S_C$ -l) and *Lb. plantarum/pentosus/paraplantarum* group (detected in  $S_B$ -l and  $S_C$ -l).

358 Together with *Lb. fermentum*, which occurred in the sole laboratory sourdough  $S_{c}$ -l, the *Lb.* 359 *plantarum/pentosus/paraplantarum* group is recognized as a highly adapted sourdough-typical LAB (Van der Meulen 360 and others 2007; Weckx and others 2010), stably co-dominating with *Lb. fermentum* in laboratory wheat sourdough 361 fermentations carried out under semi-aseptic conditions (Vogelmann and others 2009).

362 Finally, for the occurrence of other bacteria, a contribution of both parameters might be hypothesized, as in the case of 363 the Leuc. holzapfelii/citreum group and Leuc. mesenteroides/pesudomesenteroides, which were exclusively detected in 364 S<sub>A</sub>-l, and of *Pediococcus pentosaceus* and the two minority taxa *Leuc. holzapfelii/citreum* group and *Staphylococcus* 365 pasteuri detected in the sole S<sub>C</sub>-b. Insight into the impact of both these parameters on the sourdough microbiota is 366 readily available in the literature; several investigations have proved the influence of the substrate (Vogelmann and 367 others 2009; Zannini and others 2009; Ercolini and others 2013), whereas a number of other studies comparing the LAB 368 and yeast diversity of sourdoughs back-slopped at laboratory and artisan bakery level pointed out the effect of the 369 bakery environment (equipment, air, staff) on the establishment of an in-house dominant microbiota (Scheirlinck and 370 others 2007; Van der Meulen and others 2007; Vrancken and others 2010; Weckx and others 2010; Minervini and

371

others 2012).

Regarding the yeast dynamics, to our knowledge, this is the first study investigating the spontaneously developing yeast
microbiota in Italian sourdoughs in controlled laboratory fermentations without the addition of any starter culture.

374 A simpler evolution with respect to LAB was observed, and an overall low number of yeast species was identified, in 375 agreement with what was previously reported in an investigation aimed at assessing the yeast species composition of 376 artisan bakery and spontaneous laboratory sourdoughs (Vrancken and others 2010). In terms of yeast cell densities, in 377 both the laboratory and artisan bakery sourdoughs analyzed in this study, a progressive increase was seen in viable 378 counts to Log cfu/g ranging from mean values of 7.47 to 7.97; however, at day 0, an initial yeast population markedly 379 lower (up to 4 orders of magnitude) with respect to artisan bakery sourdoughs was found in the sourdoughs prepared in 380 the laboratory, feasibly due to a very poor yeast contamination of this latter propagation environment. In a further 381 research study, carried out on seven sourdoughs propagated at artisan bakery and laboratory level, after 20 and 80 days 382 of continuous back-slopping, most of the laboratory sourdoughs showed a yeast cell density markedly lower than that of 383 artisan bakery sourdoughs, not exceeding 3 Log cfu/g (Minervini and others 2012).

In the present study, a clear dominance of *S. cerevisiae* was seen along the whole 20-day propagation, irrespective of the back-slopping environment or the type of flour. This finding is in agreement with what was very recently found by Ercolini and others (2013) and other authors investigating the yeast ecology of this peculiar ecosystem (Stolz 2003; Garofalo and others 2011), thus confirming the high adaptation of this yeast to wheat sourdoughs.

388 However, for the other yeasts identified, interesting remarks can be made by comparing the sourdoughs prepared with 389 the same flour and propagated at laboratory or artisan bakery level; in more detail, except for S<sub>B</sub>-b, the co-dominance of 390 members of the K. unispora/servazzii/aerobia group with S. cerevisiae was established in the sole artisan bakery 391 sourdoughs  $S_A$ -b and  $S_C$ -b and the laboratory sourdoughs  $S_A$ -l. To date, K. unispora has rarely been documented as 392 existing in a sourdough ecosystem, namely once in Belgian artisan wheat sourdough (Huys and others 2013) and twice 393 in rye sourdoughs (Bessmeltseva and others 2014). In SA-b P. kudriavzevii, formerly known as Issatchenkia orientalis, 394 the anamorph Candida krusei, was also found to stably occur with the previous two species from days 5 to 20 of daily 395 propagation. Interestingly, highly phytase-active stains ascribed to P. kudriavzevii have previously been isolated from 396 two Lithuanian rye sourdoughs and a Danish rye sourdough made with wholemeal flour (Nuobariene and others 2012), 397 thus corroborating the positive association of this yeast with whole grain foods containing considerable amounts of 398 phytic acid.

399

## 400 Conclusions

401 To the authors knowledge this is the first report on the exploitation of the back-slopping technique for the production of402 type I sourdoughs with wholemeal flour. In general, the results collected demonstrated the suitability of this flour for

403 this specific purpose, encouraging further research for its future exploitation in high-fibre bread-making. In particular, 404 wholemeal sourdoughs offer interesting challenges for the quality improvement of high-fibre baked leavened goods, as 405 the positive modification of texture, flavour, nutritional value and shelf-life and the overcome of the deleterious effect 406 of wheat bran on bread volume.

When the sourdoughs propagated at artisan bakery and laboratory level are comparatively evaluated irrespective of the type of flour, the overall findings support, on the one hand, what was previously reported about the role of the bakery environment (equipment, air, or staff) on the contamination of sourdough batches, while on the other hand they underline the equally important influence of the substrate on the adaptation capacity of microorganisms (in terms of growth rate and competitiveness).

412

# 413 Author Contributions

Conceived and designed the experiments: L. Aquilanti, M. Taccari. performed the experiments: M. Taccari, V.
Milanović. Analyzed the data: L. Aquilanti, F. Clementi, S. Polverigiani, A. Osimani. Contributed
reagents/materials/analysis tools: S. Polverigiani, C. Garofalo, F. Clementi. Wrote the paper: M. Taccari, L. Aquilanti.

417

# 418 Acknowledgements

This study was financially supported by the Marche Region - *Servizio Agricoltura, Forestazione e Pesca* - in the context of the project "*P.S.R. Marche 2007/2013 - Progetti di Macrofiliera, Filiera Locale e Microfiliera-Misura 1.2.4-Cooperazione per lo sviluppo di nuovi prodotti, processi e tecnologie*". The authors wish to thank the farmers' cooperative "*Il Biroccio*" (Filottrano, Ancona, Italy) for supplying of the soft wheat flours and the propagation of the artisan bakery sourdoughs. They are also grateful to Dr. Catia Governatori of the "Agenzia Servizi Settore *Agroalimentare delle Marche*" (ASSAM) for the chemical, farinograph, and alveograph analyses.

425

### 426 References

- 427 Arendt EK, Ryan LAM, Dal Bello F. 2007. Impact of sourdough on the texture of bread. Food Microbiol 24:165-174.
- Azizi MH, Sayeddain SM, Payghambardoost SH. 2006. Effect of flour extraction rate on flour composition, dough
   rheological characteristics and quality of flat bread. J Agric Sci Tech 8:323-330.
- 430 Barber B, Ortola C, Barber S, Fernandez F. 1992. Storage of packaged white bread. III. Effects of sourdough and
- addition of acids on bread characteristics. Z Lebensm Unters Forsch 194:442-449.
- 432 Bessmeltseva M, Viiard E, Simm J, Paalme T, Sarand I. 2014. Evolution of bacterial consortia in spontaneously started
- 433 rye sourdough during two months of daily propagation. PLoS One 9:1-12.

- 434 Brummer JM, Lorenz K. 1991. European developments in wheat sourdoughs. Cereal Foods World 36:310-314.
- 435 Buddrick O, Jones OAH, Cornell HJ, Small DM. 2014. The influence of fermentation processes and cereal grains in
- 436 wholegrain bread on reducing phytate content. J Cereal Sci 59:3-8.
- 437 Cai Y, Okada H, Mori H, Benno Y, Nakase T. 1999. *Lactobacillus paralimentarius* sp. nov., isolated from sourdough.
- 438 Int J Syst Bacteriol 49:1451-1455.
- 439 Cossignani L, Gobbetti M, Damiani P, Corsetti A, Simonetti MS, Manfredi G. 1996. The sourdough microflora.
- 440 Microbiological, biochemical and bread making characteristics of doughs fermented with freeze-dried mixed starters,
- freeze-dried wheat sourdough and mixed fresh-cell starters. Z Lebensm Unters Forsch 203:88-94..
- 442 De Vuyst L, Vrancken G, Ravyts F, Rimaux T, Weckx S. 2009. Biodiversity, ecological determinants, and metabolic
  443 exploitation of sourdough microbiota. Food Microbiol 26:666-675.
- 444 De Vuyst L, Van Kerrebroeck S, Harth H, Huys G, Daniel HM, Weckx S. 2014. Microbial ecology of sourdough
  445 fermentations: diverse or uniform? Food Microbiol 37:11-29.
- Ercolini D, Pontonio E, De Filippis F, Minervini F, La Storia A, Gobbetti M., Di Cagno R. 2013. Microbial ecology
  dynamics during rye and wheat sourdough preparation. Appl Environ Microbiol 79:7827-7836.
- 448 Felis G, Dellaglio F. 2007. Taxonomy of lactobacilli and bifidobacteria. Curr Issues Intestinal Microbiol 8:44-61.
- Foschino R, Gallina S, Andrighetto C, Rossetti L, Galli A. 2004. Comparison of cultural methods for the identification
  and molecular investigation of yeasts from sourdoughs for Italian sweet baked products. FEMS Yeast Res 4:609-618.
- 451 Garofalo C, Aquilanti L, Clementi F. 2011. The biodiversity of the microbiota of traditional Italian sourdoughs. In:
- Almeida M.T. (Eds.), *Wheat: Genetics, Crops and Food Production* (pp. 366-392). Hauppauge, NY: Nova Science
  Publishers, Inc.
- 454 Garofalo C, Osimani A, Milanović V, Aquilanti L, De Filippis F, Stellato, Di Mauro S, Turchetti B, Buzzini P, Ercolini
- 455 D, Clementi F. 2015. Bacteria and yeast microbiota in milk kefir grains from different Italian regions. Food Microbiol
  456 49:123-133.
- 457 Garofalo C, Silvestri G, Aquilanti L, Clementi F. 2008. PCR-DGGE analysis of lactic acid bacteria and yeast dynamics
  458 during the production processes of three varieties of Panettone. J Appl Microbiol 105:243-254.
- Gobbetti M, Rizzello CG, Di Cagno R, De Angelis M. 2014. How the sourdough may affect the functional features of
  leavened baked goods. Food Microbiol 37:30-40.
- 461 Hansen A, Hansen B. 1994. The influence of wheat flour type on the production of flavour compounds in wheat
- 462 sourdoughs. J Cereal Sci 19:185-190.
- 463 Hynes WL, Ferretti JJ, Gilmore MS, Segarra RA. 1992. PCR amplification of streptococcal DNA using crude cell
- 464 lysates. FEMS Microbiol Lett 94:139-142.

- 465 Katina K. 2005. Sourdough: a tool for the improved flavour, texture and shelf-life of wheat bread. Academic
- 466 Dissertation. VTT publications 569, VTT (Technical Research Centre of Finland), Espoo, Finland (2005), pp. 13-75.
- Kumara V, Sinha AK, Makkara HPS, Becker K. 2010. Dietary roles of phytate and phytase in human nutrition: a
  review. Food Chem 120:945-959.
- Lioger D, Leenhardt F, Demigne C, Remesy C. 2007. Sourdough fermentation of wheat fractions rich in fibres before
  their use in processed food. J Sci Food Agr 87:1368-1373.
- 471 Lopez HW, Duclos V, Coudray C, Krespine V, Feillet-Coudray C, Messager A, Demigné C, Rémésy C. 2003. Making
- bread with sourdough improves mineral bioavailability from reconstituted whole wheat flour in rats. Nutrition19:524-530.
- 474 Makimura K, Tamura Y, Mochizuki T, Hasegawa A, Tajiri Y, Hanazawa R, Uchida K, Saito H, Yamaguchi H. 1999.
- Phylogenetic classification and species identification of dermatophyte strains based on DNA sequences of nuclear
  ribosomal internal transcribed spacer 1 regions. J Clin Microbiol 37:920-924.
- 477 Mariotti M, Garofalo C, Aquilanti L, Osimani A, Fongaro L, Tavoletti S, Hager A-S, Clementi F. 2014. Barley flour
- 478 exploitation in sourdough bread-making: a technological, nutritional and sensory evaluation. LWT-Food Sci Technol479 59:973-980.
- 480 Martinez-Anaya MA. 2003. Associations and interactions of micro-organisms in dough fermentations: effects on dough
  481 and bread characteristics. In: Kulp, K., and Lorenz, K., (Eds.), *Handbook of dough fermentations* (pp. 63-195).
- 482 Marcel Dekker Inc, New York,
- 483 Minervini F, Lattanzi A, De Angelis M, Di Cagno R, Gobbetti M. 2012. Influence of artisan bakery-or laboratory-
- 484 propagated sourdoughs on the diversity of lactic acid bacterium and yeast microbiotas. Appl Environl Microbiol 78:
  485 5328-5340.
- 486 Nuobariene L, Hansen AS, Arneborg N. 2012. Isolation and identification of phytase-active yeasts from sourdoughs.
  487 LWT-Food Sci Technol 48:190-196.
- Osimani A, Garofalo C, Aquilanti L, Milanović V, Clementi F. 2015. Unpasteurised commercial boza as a source of
   microbial diversity. Int J Food Microbiol 194:62-70.
- 490 Poutanen K, Flander L, Katina K. 2009. Sourdough and cereal fermentation in a nutritional perspective. Food Microbiol
  491 26:693-699.
- 492 Scheirlinck I, Van der Meulen R, Van Schoor A, Vancanneyt M, De Vuyst L, Vandamme P, Huys G. 2007. Influence
- 493 of geographical origin and flour type on diversity of lactic acid bacteria in traditional Belgian sourdoughs. Appl
- 494 Environ Microbiol 73:6262-6269.

- 495 Salmenkallio-Marttila M, Katina K, Autio K. 2001. Effects of bran fermentation on quality and microstructure of high496 fibre wheat bread. *Cereal Chem* 78:429-435.
- 497 Stevenson L, Phillips F, O'Sullivan K, Walton J. 2012. Wheat bran: its composition and benefits to health, a European
  498 perspective. Int J Food Sci Nutr 63:1001-1013.
- Stolz P. 2003. Biological Fundamentals of Yeast and Lactobacilli fermentation in bread dough. In: Kulp, K., Lorenz,
  K., (Eds.), *Handbook of dough fermentations*, (pp. 23-43). Marcel Dekker, New York.
- 501 Van der Meulen R, Scheirlinck I, Van Schoor A, Huys G, Vancanneyt M, Vandamme P, De Vuyst L. 2007. Population
- dynamics and metabolite target analysis of lactic acid bacteria during laboratory fermentations of wheat and spelt
   sourdoughs. Appl Environ Microbiol 73:4741-4750.
- Venkateswara Rao G, Indrani D, Shurpalekar SR. 1985. Effect of milling methods and extraction rate on the chemical,
   rheological and breadmaking characteristics of wheat flours. J Food Sci Tech 22:38-42.
- 506 Vogel RF, Böcker G, Stolz P, Ehrmann M, Fanta D, Ludwig W., Pot B, Kersters K, Schleifer KH, Hammes WP. 1994.
- 507 Identification of Lactobacilli from sourdough and description of *Lactobacillus pontis* sp. nov. Int J Syst Bacteriol
  508 44:223-229.
- 509 Vogelmann SA, Seitter M, Singer U, Brandt MJ, Hertel C. 2009. Adaptability of lactic acid bacteria and yeasts to
- sourdoughs prepared from cereals, pseudocereals and cassava and use of competitive strains as starters. Int J Food
  Microbiol 130:205-212.
- 512 Vrancken G, De Vuyst L, Van der Meulen R, Huys G, Vandamme P, Daniel HM. 2010. Yeast species composition
- 513 differs between artisan bakery and spontaneous laboratory sourdoughs. FEMS Yeast Res 10:471-481.
- 514 Weckx S, Van der Meulen R, Allemeersch J, Huys G, Vandamme P, Van Hummelen P, De Vuyst L. 2010.Community
- dynamics of bacteria in sourdough fermentations as revealed by their metatranscriptome. Appl Environ Microbiol
  76:5402-5408.
- 517 Zannini E, Garofalo C, Aquilanti L, Santarelli S, Silvestri G, Clementi F. 2009. Microbiological and technological
  518 characterization of sourdoughs destined for bread-making with barley flour. Food Microbiol 26:744-753.
- 519
- 520
- 521
- 522
- ---
- 523
- 524
- 525



Figure 1-Viable counts of presumptive lactic acid bacteria and yeasts, assessed on selective media mMRS ( $\rightarrow$ ), mSDB ( $\neg \rightarrow$ ) and WLN( $\rightarrow$ ) and kinetics of acidification (pH  $\neg \neg \neg$  and TTA  $\neg \diamond \neg$ ) of wholemeal (S<sub>A</sub>) and type 0 (S<sub>B</sub> and S<sub>C</sub>) sourdoughs daily propagated under laboratory (labeled "l") and artisan bakery (labeled "b") conditions for 20 days; numbers on the  $\chi$  axis indicate days of propagation. Day 0 corresponds to the first water and flour dough sampled and analyzed prior to fermentation, whereas days 1 to 20 correspond to sourdough sampled immediately before each daily refreshment. Means of two independent experiments ± standard deviations are shown.



**Figure 2-**Score and loading plots of first and second principal components based on microbial community and biochemical characteristics data means from sourdoughs produced with different flours (A, B and C) and in two different propagation environments (laboratory, "l" and artisan bakery, "b"). Clusters A<sub>I</sub>, A<sub>II</sub> and B are indicated by dotted ellipses and correspond to those derived from the cluster analysis shown in Fig. 4.







Presence or absence of strains LD. Plantarumparapharatumpertosis group 20 10 W. confusacioaria group Sc Padiococcus spp 20 Leve holtapati i otreun aroup S. S. Pasteuri Lactobacitus sup. 20 S. Cerevisiae 10 P. Kudriavcevii K. unisporalservatziliserobia 5 SA 0

585

- 586
- 588

**Figure 3-**Lactic acid bacteria (LAB) and yeast species identified through the combined PCR-DGGE approach in the sourdoughs propagated at artisan ( $S_A$ -b,  $S_B$ -b,  $S_C$ -b) and laboratory ( $S_A$ -l,  $S_B$ -l,  $S_C$ -l) level for 1, 5, 10 and 20 days using a wholemeal ( $F_A$ ) and two type 0 ( $F_B$  and  $F_C$ ) flours. The *z*-axis indicates only the presence or absence of a given

- species, resulting from either the analysis of the DNA extracted directly from the dough and sourdough samples or the
- 593 bulk of colonies harvested from selected dilution plates. (A) Laboratory, (B) Artisan bakery.



Figure 4-Dendrogram obtained from two way cluster analysis based on the microbial community and biochemical
characteristics of sourdoughs propagated with the three flours (F<sub>A</sub>, F<sub>B</sub> or F<sub>C</sub>) at the laboratory ("1") or artisan bakery
("b").

- Table 1-Biochemical properties and microbial viable counts of wholemeal (S<sub>A</sub>) and type 0 (S<sub>B</sub> and S<sub>C</sub>) mature
- 617 sourdoughs after 20 days propagation under laboratory (labeled "l") and artisan bakery (labeled "b")
- 618 conditions.

Mature sourdough	рН	TTA (mL 0.1 N NaOH)	Phytic acid (g/100g)	Lactic <u>acid</u> (g/100 g)	Acetic <u>acid</u> (g/100 g)	FQ	Yeast (Log cfu/g)	LAB (on mMRS) (Log cfu/g)	LAB ( <u>on mSDB</u> ) (Log cfu/g)
ANOVA									
Flour	0.0001*	<.0001*	<.0001*	0.0003*	<.0001*	0.0177*	0.0079*	<.0001*	0.1431
Site	0.0274*	0.0117*	1	0.0041*	<.0001*	<.0001*	0.0193*	<.0001*	<.0001*
Flour x Site	0.1924	0.0008*	1	0.9458	<.0001*	0.0014*	0.0075*	<.0001*	0.0572
Mean separation test									
TREATMENT									
S <sub>A</sub> -b	$4.02 \pm 0.02^{a}$	$16.00\pm0.20^{\rm b}$	$0.12\pm0.00^{\text{a}}$	$17.20\pm0.10^{\mathtt{a}}$	$1.96\pm0.00^{b}$	$5.85 \pm 0.09^{\mathrm{ab}}$	$7.97 \pm 0.02^{a}$	$9.26\pm0.01^{\mathtt{a}}$	$9.19\pm0.03^{\text{b}}$
S <sub>B</sub> -b	$3.79\pm0.01^{\text{cd}}$	$10.15\pm0.15^{\textrm{d}}$	n.d.	$11.60 \pm 0.51^{bc}$	$4.01 \pm 0.08^{a}$	1.92 ± 0.04°	$7.47 \pm 0.11^{b}$	$9.05\pm0.01^{\text{b}}$	$8.93 \pm 0.05^{b}$
Sc-b	$3.87 \pm 0.00^{bc}$	$11.40\pm0.10^{\circ}$	n.d.	11.63 ±0.22 <sup>bc</sup>	$1.18\pm0.10^{\circ}$	$6.61 \pm 0.67^{b}$	$7.81 \pm 0.02^{a}$	$9.00\pm0.00^{\text{b}}$	$9.01 \pm 0.11^{b}$
S <sub>A</sub> -1	$3.95\pm0.05^{\text{ab}}$	$18.10\pm0.10^{\mathtt{a}}$	$0.13\pm0.00^{a}$	14.66 ±0.88 <sup>ab</sup>	$1.43 \pm 0.05^{\circ}$	$6.83 \pm 0.62^{b}$	$7.97 \pm 0.02^{a}$	$8.45\pm0.02^{\text{d}}$	$9.75\pm0.05^{a}$
S <sub>B</sub> -1	$3.72\pm0.00^{d}$	$9.40\pm0.20^{d}$	n.d.	9.35 ± 0.83°	$0.45\pm0.00^{d}$	$13.74 \pm 1.28^{a}$	$7.93 \pm 0.06^{a}$	$8.62 \pm 0.02^{\circ}$	$9.85 \pm 0.00^{\mathrm{a}}$
$S_{C}$ -1	$3.87\pm0.00^{\text{bc}}$	$11.70\pm0.30^{\rm c}$	n.d.	8.94 ± 0.99°	$0.47\pm0.01^{\text{d}}$	$12.55 \pm 1.10^{a}$	$7.78\pm0.00^{ab}$	$7.00\pm0.00^{\rm e}$	$9.65 \pm 0.03^{a}$

620 Values represent means  $\pm$  st. err. (n.= 2). Means followed by a different letter were significantly different according to

621 the Tukey's test (P < 0.05).

**622** FQ: Fermentation Quotient; n.d. not detectable.

623