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Effects of bioprotective cultures on the microbial community during storage of Italian fresh filled pasta

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36 **ABSTRACT**

37 Filled pasta is a typical Italian product consisting in a thin dough stuffed with a filling containing dairy, meat
38 or vegetable ingredients. When industrially produced, its microbial stability relies on thermal treatment,
39 proper storage temperature and modified atmosphere packaging. Since these processes can strongly affect
40 the traditional features of pasta (mainly flavor and texture), alternative strategies have been investigated.
41 In this research milder heat treatments were applied and, to assure microbial quality and safety, they were
42 combined with the addition of bioprotective cultures (*Lactobacillus rhamnosus* and *Lactobacillus paracasei*)
43 in the filling of Ricotta based *Tortelloni*. Their effects on microbiological patterns during storage at 6°C was
44 studied through culture dependent and independent methods and also the influence on organoleptic
45 profile (through SPME-GC-MS and sensory evaluation) was assessed. The results demonstrated that
46 bioprotective cultures had a relevant quantitative and qualitative effect on the microbiota of *Tortelloni*
47 during storage: indeed, even if they were not dominant, their presence reduced the initial microbiota
48 associated with raw materials and gave a competitive advantage to safer or organoleptically acceptable LAB
49 species, such as leuconostocs. Although these LAB cultures influenced the aroma profile of filled pasta
50 (increase of alcohols, esters and acids), the sensory evaluation confirmed the overall acceptability of the
51 product. The addition of bioprotective cultures can be therefore a helpful strategy to reduce thermal
52 treatments and better maintain the traditional textural and flavor characteristics of this product.

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57 **Key words:** bioprotective cultures; filled pasta; shelf life; metagenomics; lactobacilli

1. INTRODUCTION

Filled pasta is a typical Italian product with different shapes and filling varying with the geographical area of production (Alexander, 2000). The pasta consists in a thin dough made with water, flour and usually eggs that is stuffed with a filling prepared with dairy, meat or vegetable ingredients (Marotta et al., 2018). The artisanal products are usually handmade and locally distributed at refrigerated temperature with a limited shelf life (4-5 days). On the contrary, shelf life of 60-90 days and more can be achieved for industrial production by applying thermal treatments, reducing the water activity (a_w) of the filling, packaging the pasta under modified atmosphere (MAP) and adopting appropriate storage temperature (Marotta et al., 2018; Zardetto & Dalla Rosa, 2015).

According to Italian legislation, “fresh pasta” (filled or not) must have a water content lower than 24% and a_w lower than 0.97 and higher than 0.92 (Decreto Repubblica Italiana, 2001) and it must be stored at temperature not higher than $4\pm 2^\circ\text{C}$.

The microbial stability of this product relies in first instance on thermal treatments, which can be applied following two strategies. In the first case, only a treatment is carried out on the loose product with an injected steam belt pasteurizer. In the second case, this first treatment is followed by a further pasteurization in static chambers after packaging. In addition to the effects on microbial population, both these strategies definitely affect the mechanical and functional properties of the final products (Alampese, Casiraghi, & Rossi, 2008; Zardetto & Dalla Rosa, 2015). After packaging, the shelf life of filled pasta is strictly dependent on the ability of microorganisms survived to the thermal treatments to grow during storage, overcoming the hurdles determined by a_w , MAP composition and storage temperature (Sanguinetti et al., 2011, 2016).

In spite of the great diffusion of this typology of pasta and the increase of its consumption (ISMEA, 2018), there is a relatively scarce literature concerning its microbiological characteristics, mainly focused on the presence of *Enterobacteriaceae*, total mesophilic counts and moulds (Marotta et al., 2018; Ricci, Barone, & Petrella, 2017; Zardetto, 2005).

Tortelloni is a filled pasta produced in Emilia Romagna Region (Italy) since several centuries (Tanara, 1644) with a soft filling obtained mixing Ricotta and Parmesan cheese, eggs, salt, nutmeg and vegetables (beet, spinach or parsley, according to the zone). The productive process, including thermal treatments and a_w lowering, can have a strong impact on the “traditional” features of such type of pasta, inducing to adopt milder conditions to preserve regional traditions and peculiar characteristics of *Tortelloni*. In particular, in order to preserve its softness, often the a_w of the filling cannot be excessively lowered, with the consequent risk of rapid growth of spoiling microflora.

This aspect is in contrast with the shelf life expected for the commercialization of this product (45-60 days). For this reason, other hurdles to the microbial growth must be exploited to satisfy this need while maintaining the traditional organoleptic properties and assuring the hygienic quality of *Tortelloni*.

93 The use of bioprotective cultures is an interesting strategy proposed with the aims of reducing the risks
94 associated with the growth of undesirable and pathogenic microorganisms and prolonging the shelf life of
95 foods (Oliveira, Ferreira, Magalhães, & Teixeira, 2018). Biopreservation consists in the use of natural and
96 selected microflora able to control or inhibit spoiling or pathogenic microorganisms by competition or
97 production of specific antimicrobial molecules such as bacteriocins, organic acids, diacetyl, acetoin, etc.
98 (Ghanbari, Jami, Domig, & Kneifel, 2014). Lactic acid bacteria (LAB) are ideal candidates for biopreservation
99 due to their safe history in foods and their wide range of antimicrobial compound production (Cifuentes
100 Bachmann & Leroy, 2015). Among the potential drawbacks of the use of bioprotective cultures there is the
101 possibility of undesired sensory effects on the food organoleptic profile and the choice of LAB species must
102 take into consideration a low or compatible impact on food flavor.

103 This work was aimed to evaluate the effects of the use of bioprotective cultures on the microbiological
104 patterns of Ricotta based *Tortelloni* during production and storage. Filled pasta was produced in a small
105 factory following the traditional recipe. In addition to traditional microbiological protocols, the microbial
106 community profiling was performed through rDNA-targeting pyrosequencing to test the effects of two
107 bioprotective cultures (*Lactobacillus rhamnosus* or a mixture of *L. rhamnosus* and *Lactobacillus paracasei*)
108 on the spoilage microbiota during filled pasta storage. Further, the influence of the cultures on the
109 organoleptic profile of the product was studied. Finally, a validation trial was carried out with the objective
110 to optimize the use of this strategy for the stabilization of this filled pasta.

112 2. MATERIAL AND METHODS

113 2.1 Filled pasta production

114 Filled pasta samples were produced in a small factory located in Emilia Romagna Region (Italy). The
115 *Tortelloni* (50% filling, thickness of pasta 0.7 mm) were obtained with a traditional recipe following the
116 process reported in Figure 1. In particular, pasta was produced using durum wheat semolina (36% w/w) and
117 soft wheat flour (36% w/w) added with pasteurized egg product (28% w/w). Filling was obtained by mixing
118 Ricotta (70.5% w/w), Parmigiano Reggiano cheese (12% w/w), breadcrumbs (12% w/w), parsley (3.5 %
119 w/w), salt (1.5% w/w) and nutmeg (0.5% w/w). The filling rested 18 hours at refrigerate temperature (4°C)
120 and then underwent to the line production process.

121 The filling pH was about 5.5 ± 0.1 while its a_w was 0.960. After forming, *Tortelloni* were subjected to a heat
122 treatment in one-line steam belt pasteurizer and a subsequent drying and cooling in a tunnel chamber.
123 After cooling, the filled pasta was packed in polyamide/polypropylene (PA/PP) film under modified
124 atmosphere (40% CO₂; 60% N₂). The film presented a water vapor permeability <5 gr/m²/24h (38°C, 90%
125 R.H.) and an oxygen permeability <3 cc/m²/24h (23°C, 0% R.H.). After packaging, the samples were cooled
126 and stored at 6°C for 30 days.

Two different trials were performed. During the first trial, three different samples were obtained: the control, produced without bioprotective cultures, and two samples inoculated with two different commercial bioprotective cultures available on the market, containing *Lactobacillus rhamnosus* (BC1) or a mixture of *L. rhamnosus* and *Lactobacillus paracasei* (BC2). After appropriate hydration according to the manufacturer's suggestions, these cultures were added (cell concentration of about 7 log CFU/g) in the filling during mixing. In the first trial, a mild pasteurization was carried out in a steam belt pasteurizer set for a treatment of 60°C for 3 min in the product inner part. In the second trial (validation), two samples were produced: the control and the sample with the bioprotective culture BC2 (inoculum level 7 log CFU/g), added as reported above. In this case, the thermal treatment was carried out at 70°C for 3 min in the product inner part. In both treatments, the reaching of target temperature was controlled using a data logger (S-Micro, Tecnosoft, Italy) inserted inside the product.

138

139 **2.2 Microbial counts, pH and a_w analysis**

Three replicates of each sample were examined at the different sampling times (before and after pasteurization and during the shelf life) for the enumeration of microbial population. 10 g of samples were 10-fold diluted with 90 ml of 0.9% (w/v) NaCl and homogenized in a Lab Blender Stomacher (Seward Medical, England) for 2 minutes. Decimal dilutions were performed and plated onto selective media: LAB were enumerated on MRS agar (Oxoid, Basingstoke, UK) incubated at 30°C for 48 h in anaerobic conditions, staphylococci were counted by surface-plating on Baird-Parker (added with egg yolk tellurite emulsion) (Oxoid) incubated at 30°C for 48 h, yeast and moulds were grown on Sabouraud Dextrose Agar (Oxoid) plates, added with 200 mg/l of chloramphenicol and incubated at 28°C for 72 h, while *Enterobacteriaceae* were enumerated by pour plating in Violet Red Bile Glucose agar (Oxoid) and incubating plates at 37°C for 24 h. The presence/absence of *Listeria monocytogenes* and *Salmonella* was evaluated according to the methods EN ISO 11290–1 and EN ISO 6579, respectively (2017).

The pH and a_w of Ricotta filled *Tortelloni* were monitored using a pH-meter Basic 20 (Crison Instruments, Barcelona, Spain) and an Aqualab CX3-TE (Labo-Scientifica, Parma, Italy), respectively.

153

154 **2.3 Aroma profile analysis**

For the analysis of volatile compounds, 3 g of samples were placed in 10-ml sterilized vials, added with 33 mg/kg of 4 methyl-2-pentanol (Sigma-Aldrich, Steinheim, Germany) as internal standard and sealed by PTFE/silicon septa. The samples were heated for 10 min at 45°C and the volatile molecules in the headspace were adsorbed with fused silica SPME fiber covered with 85 μ m Carboxen/Polydimethylsiloxane (CAR/PDMS) (Supelco, Steinheim, Germany) for 40 min. Adsorbed molecules were desorbed in the injector for 10 min with the same analytical conditions reported by Montanari et al. (2016), using a an Agilent

161 Hewlett–Packard 6890 GC gas-chromatograph and a 5970 MSD MS detector (Hewlett–Packard, Geneva,
162 Switzerland) equipped with a CP-WAX 52CB (50 m X 0.32 mm X 1.2 µm) fused silica capillary column.
163 Volatile peak identification was carried out by computer matching of mass spectral data with those of
164 compounds included in NIST 2011 mass spectral library (Scientific Instrument Services, Ringoes, NJ, United
165 States).

166

167 **2.4 Sensory evaluation of cooked samples**

168 Sensory evaluation was carried out using an affective test (Sanguinetti et al., 2016) involving 40 untrained
169 consumers (age 22-60), among which 20 females and 20 males. Their acceptance of the sample proposed
170 was based on an intensity scale from 0 to 7 (0, dislike; 7, extremely like) for selected attributes (odour
171 intensity, colour, aroma intensity, salty, sourness, bitterness, cohesiveness, chewiness, overall
172 acceptability). The evaluation was carried out in randomized and balanced order. The samples were served
173 to the panelist after a 5 min immersion in boiling water (100°C) without salt added.

174

175 **2.5 DNA extraction and Sequencing**

176 Total genomic DNA was directly extracted from frozen *Tortelloni* samples by taking approximately 10 g of
177 filled pasta. The samples were dissolved in 90 ml of physiological solution (0.9% NaCl) and were
178 mechanically homogenized in stomacher for 4 minutes at 430 beats per minute. After decanting, 1 ml of
179 the supernatant was collected and subjected to enzymatic treatment towards bacteria (lysozyme) and
180 yeasts (lyticase) at 37°C for 1 h, followed by alkaline lysis with the addition of NaOH and SDS at a final
181 concentration of 0.1 N and 1%, respectively. The extracted DNA was purified by chloroform:isoamyl alcohol
182 24:1 treatment and precipitated in 0.54 volumes of isopropanol. Finally, the purified DNA was resuspended
183 in water and quantified using Qubit 4 Fluorimeter (ThermoFisher Scientific, Waltham, MA USA). The
184 concentration of the DNA samples was normalized and the sequencing was carried out through Illumina
185 MiSeq platform which generated 300 bp pair-end sequencing reads. The library for Illumina sequencing was
186 generated from V3-V4 variable regions of ribosomal 16S rRNA in order to characterize the bacterial
187 population of the samples.

188

189 **2.6 Bioinformatics analysis**

190 FASTQ sequence files from Illumina reads were generated using bcl2fastq2 version 2.18. Initial quality
191 assessment was based on data passing the Illumina Chastity filtering. Subsequently, reads containing PhiX
192 control signal were removed using an in-house filtering protocol. In addition, reads containing (partial)
193 adapters were clipped (up to a minimum read length of 50 bp). A final quality assessment was performed
194 on the remaining reads using the FASTQC quality control tool version 0.11.5.

195 The FASTQ sequences obtained were analyzed using DADA2 version 1.8 (Callahan et al., 2016) by R 3.5.1
196 environment. DADA2 implements a new quality-aware model of Illumina amplicon errors without
197 constructing OTUs (Callahan et al., 2016). DADA2 was run as described in
198 <https://benjjneb.github.io/dada2/tutorial.html> applying the following parameters: trimLeft equal to 30 and
199 truncLen option set to 270 and 200 for the forward and reverse fastq files, respectively.
200 The taxonomic assignment was performed comparing the amplicon sequence variant (ASV) predicted from
201 DADA2 against SILVA database (version 128, <https://www.arb-silva.de/documentation/release-128/>). ASVs
202 belonging to taxa classified as external sample (Davis, Proctor, Holmes, Relman, & Callahan, 2018)
203 contaminations were not included in the composition analysis for microbial population as well as for ASVs
204 with low abundance setting a threshold of relative abundance equal to 0.5%. The assignment at specie level
205 for the remaining ASVs was confirmed comparing the nucleotide sequence obtained from the Illumina
206 sequencing with the 16S rRNA sequence of the available type strains included in the LTP database, which
207 represents all bacterial and archaeal taxa with validly published names (Yilmaz et al., 2014).

208

209 **2.7 Statistical analysis**

210 Three independent samples were analysed for each sampling time, each of which analysed in triplicate. The
211 data were statistically analysed using ANOVA. The Tukey critical difference test was performed to
212 determine differences between samples ($p < 0.05$). The presence of significative differences in the sensory
213 test was tested using a non-parametric Mann-Whitney test ($p < 0.05$).

214

215 **3. RESULTS AND DISCUSSION**

216 **3.1 Microbial counts**

217 In the first experimental trial, a mild pasteurization was carried out with a double aim: *i*) to have a low
218 impact on the characteristic of *Tortelloni* and *ii*) to have a limited effect on the viability of the microbial
219 populations to better evidence the relationships between bioprotective cultures and wild spoilage
220 microorganisms. Table 1 reports the results of the microbial counts before and immediately after the
221 pasteurization and after 7, 15 and 30 days of storage at 6°C. The pH of the product was 5.7 and did not
222 significantly change during storage in the control, while low reductions (5.5-5.6) were observed in the
223 presence of bioprotective cultures (BC1 and BC2) after 30 days. The initial cell concentrations of fresh filled
224 pasta before pasteurization were rather high, represented mainly by *Enterobacteriaceae* (more than 6 log
225 CFU/g). Yeasts concentration was about 4 log CFU/g. Similar cell concentrations were found for
226 staphylococci, but no coagulase positive colony was detected. Wild LAB in the control were 5.4 log CFU/g
227 while in the samples added with bioprotective cultures reached the expected concentration (more than 7
228 log CFU/g).

229 After the mild pasteurization (corresponding to a treatment in the steam belt pasteurizer at 60°C for 3 min)
230 yeasts were subjected to the more severe reduction, since in all the samples they were below the detection
231 limit. *Enterobacteriaceae* and LAB decreased of about 2 log units while staphylococci counts were the less
232 affected by heat (1 log unit reduction or less). These levels are high if compared with other industrial
233 products and are more similar to data reported for artisanal fresh pasta (Ricci et al., 2017), due to the low
234 thermal treatment. Marotta et al. (2018) found that 55% of Italian artisanal fresh filled pasta samples
235 showed high *Enterobacteriaceae* counts (> 4 log CFU/g) while these microorganisms were below the
236 detection limit in industrial samples. Higher thermal treatments (80°C for about 3 min) were able to
237 decrease the concentration of this microbial group of at least 4 log units (Sanguinetti et al., 2011). In
238 addition, Ricci et al. (2017) found a mean value of 4.3 log CFU/g of total mesophilic bacteria in packed
239 industrial products and 5.5 log CFU/g in artisanal ones (unpacked). Chavez-Lopez, Vannini, Lanciotti, &
240 Guerzoni (1998) described a prevalence of *Bacillus* spp. in industrial *Ravioli* with a meat-based filling after
241 the thermal treatment. They suggested that shelf life depended also on the textural and micro-structural
242 changes (particularly protein gelation) induced by the thermal treatment.

243 In the case of *Tortelloni*, the high survival rate determined a rapid multiplication of microorganisms during
244 storage at 6°C. In particular, LAB counts reached 7 log CFU/g in the control after 15 days, while in the
245 presence of bioprotective cultures the counts were about 8 log CFU/g at the same sampling time and
246 further increased after 30 days.

247 In the control *Enterobacteriaceae* grew up to 5.9 log CFU/g, while in the samples added with bioprotective
248 cultures their concentration significantly decreased during storage, particularly in the sample added with
249 BC2. The concentration of coagulase negative staphylococci remained rather constant throughout the
250 storage in the control, while the presence of BC1 and BC2 determined a significant decrease of the
251 concentration of this microbial group. Yeasts were always below the detection limit in the presence of the
252 cultures, while concentrations higher than 2 log CFU/g were observed in the control.

253 Pathogens (*Listeria monocytogenes* and *Salmonella*) were never detected in our samples. Mould growth
254 was never observed in packages. Moulds can be responsible for the spoilage of fresh filled pasta but their
255 growth can be controlled by using MAP packaging. According to Zardetto (2005), concentration of CO₂ up
256 to 15% can stimulate the growth of *Penicillium* species, but higher concentrations rapidly inhibited their
257 multiplication.

258

259 **3.2 Study of microbial community composition through amplicon sequencing and metagenomics**

260 Firmicutes were dominant in all the samples including the filled pasta before treatment (control).
261 Immediately after production, in the samples without the addition of bioprotective cultures, *Streptococcus*
262 and *Lactococcus* were the dominant genera (Figure 2). In particular, among streptococci, the species
263 detected were *Streptococcus parauberis* (27.7% of ASVs); *Streptococcus uberis* group (11.4%), including the

species *S. uberis*, *Streptococcus porcinus* and *Streptococcus pseudoporcinus*; *Streptococcus parasuis* (0.9%) and *Streptococcus salivarius* group (0.8%), including both subspecies of *S. salivarius* and *Streptococcus vestibularis*. Regarding lactococci, *Lactococcus lactis* dominated (18.5% of ASVs) followed by *Lactococcus raffinolactis* group (11.9%), which includes the species *Lc. raffinolactis* and *Lactococcus piscium* and *Lactococcus garviae* group (1.1%), including the species *Lc. garviae* and *Lactococcus formosensis*. Among lactobacilli, the homofermentative and thermophilic species *Lactobacillus delbrueckii* and the *Lactobacillus acidophilus* group, which includes the species *Lt. acidophilus* and *Lactobacillus helveticus*, were the most representative (5.0 and 4.9%, respectively), probably deriving from the whey used for Ricotta production. Among Gram positive, the sample without the addition of bioprotective cultures exhibited the presence of members of the species *Rothia endophytica* (belonging to *Micrococcaceae*) and belonging to the *Bacillus cereus* group (4.9 and 3.0% respectively). Gram negative bacteria were represented by *Hafnia* spp. (1.0%) and member of the genus *Acinetobacter*, which included *Acinetobacter guillouiae* group (0.7%) and *Acinetobacter johnsonii* group (4.1%). As expected, many of these species derive from dairy environment, including lactococci and lactobacilli. In particular, the relevant presence of the species *S. parauberis* and those belonging to the *S. uberis* group (*S. uberis*, *S. porcinus* and *S. pseudoporcinus*) can be related to the use of milk from cows affected by mastitis at some degree. In particular, the species *S. uberis* is implied in recurrent cow clinical mastitis (Jamali et al., 2018). Independently on the initial concentration, these species were able to multiply in the industrial Ricotta used for the filling.

After 15 days of storage in the samples without bioprotective cultures, members of the *S. uberis* group and of the species *S. parauberis* still accounted for a relevant proportion of the total population (3.7 and 20.9% of ASVs), together with *Lc. raffinolactis* group (7.8%). However, the species more represented was those of the *Leuconostoc mesenteroides* group (48.7 % of ASVs), including *Leuc. mesenteroides* and *Leuconostoc pseudomesenteroides*, which were not detected immediately after pasteurization. The addition of bioprotective cultures changed the quantitative composition of the microbiota. *Leuc. mesenteroides* group remained the most relevant species but its relative presence greatly increased and reached 77.8% in the samples BC1 and 83.0% in the samples BC2. Streptococci and lactococci decreased significantly (especially *S. uberis* group and *S. parauberis*) while the presence of *Carnobacterium gallinarum* group ranged from 3.8% to 3.0% of ASVs in both samples. The detection of ASVs associated to the *Lactobacillus casei* group confirmed the presence of the bioprotective cultures, representing the 4.7 and 5.6% of the ASVs in BC1 and BC2, respectively. *L. casei* group is constituted by the phenotypically and genotypically closely related species *L. casei*, *Lactobacillus paracasei*, and *Lactobacillus rhamnosus* (Huang et al., 2018). Even if the bioprotective cultures were not able to dominate the pasta microbiota and probably did not survive the thermal treatment, their addition to filling followed by an 18-hour adaptation before production changed significantly the ratios between microbial species during storage. Bacteriocin production is usually maximum in the mid exponential phase or at the end of growth phase (Beshkova & Frankova, 2012) and it

299 is compatible with an accumulation in the overnight incubation of filling. In other words, heat stable
300 bacteriocins can be present in the filling after production, independently of the viability of the cells
301 responsible for their production, due to the thermal treatment. In addition, the inhibition of mastitis
302 streptococci (*S. agalactiae*) by a bacteriocin producing *L. rhamnosus* strain has already been demonstrated
303 by Ruíz et al. (2012).

304

305 **3.3 Volatilome of filled pasta**

306 The SPME-GC-MS aroma profile of filled pasta immediately after pasteurization and after 30 days of storage
307 is reported in Table 2, where data are express as ratio between peak area of each molecule and peak area
308 of the internal standard (4-methyl-2-pentanol). In general, ketones decreased during the storage in the
309 presence of both bioprotective cultures. 2-pentanone, 2-hexanone and 2-heptanone decreased in the
310 samples added with bioprotective cultures after storage, while they remained rather constant in the
311 control. The presence of bioprotective cultures determined also an increase of 2,3-butanedione (diacetyl)
312 and 3-hydroxy-2-butanone (acetoin) immediately after production, due to the overnight culture adaptation
313 in fillings, but these molecules drastically decreased at the end of storage.

314 Aldehydes generally decreased during storage, especially where the bioprotective cultures were added.

315 This diminution was mainly due to the reduction hexanal, more relevant in the samples BC1 and BC2, and
316 2-butenal.

317 Alcohols markedly increased in all the samples during storage, but their presence was almost doubled when
318 bioprotective cultures were added. The main responsible for this change was ethanol, present in low
319 amounts (about 10) in all the samples after production, and found after 30 days at level of 216 in the
320 control and at values higher than 500 in the samples with the bioprotective cultures. Other alcohols
321 increased their concentration, and namely 1-hexanol, 2-heptanol and 2-nonanol; the increase of these
322 latter molecules can be related to the reduction of the corresponding aldehydes. Phenylethyl alcohol and 1-
323 nonanol were found at the end of storage only in the samples with the bioprotective cultures.

324 Acids, present in similar amounts in all the conditions immediately after pasteurization, were represented
325 by acetic, butanoic, hexanoic and octanoic acids. Acetic acid is produced by heterofermentative LAB as
326 result of heterolactic pathway. However, it is also produced under nutritional stress conditions (scarcity or
327 absence of fermentable sugars) through different pathways starting from pyruvate, which can derive from
328 re-oxidation of lactate or from amino acid metabolism (Montanari et al., 2018; Zotta, Parente, & Ricciardi,
329 2017). During storage acids remained almost constant in the control, whereas they drastically increased in
330 the samples BC1 and BC2.

331 A similar behavior characterized the esters, represented mainly by ethyl hexanoate, ethyl acetate, ethyl
332 octanoate and ethyl butanoate. The esterase activity of LAB has been deeply described in dairy and
333 enological environments (Sumby, Grbin, & Jiranek, 2010). Liu, Holland, & Crow (2003) demonstrated that

LAB esterases were active against short chain fatty acid mono- and diglycerides and not against triglycerides. Esterase activity was demonstrated also in leuconostocs by Pedersen, Ristagno, McSweeney, Vogensen, & Ardö (2013). All the molecules described have already been reported as component of dairy product flavor as result of microbial metabolism (Curioni & Bosset, 2002) and their presence can be, at some extent, compatible with a dairy-based filling. Leuconostocs were the dominant microbial component in *Tortelloni* samples, especially in the filled pasta containing protective cultures, and their role described in the aroma formation of dairy products is compatible with the volatile profile described for *Tortelloni* (Hemme & Foucaud-Scheunemann, 2004).

342

3.4 Validation

The effects of the bioprotective cultures on the spoilage profile and the shelf life of fresh filled pasta were validated in a second trial, applying a higher pasteurization treatment (70°C for 3 min) if compared with the first trial. However, this thermal profile was milder, if compared with those often adopted for this typology of filled pasta (Alampese et al., 2008), with the aim to preserve the characteristic of fresh *Tortelloni*. In this production, only the protective culture BC2 was used, due to its better inhibition effects against enterobacteria and staphylococci observed in the first trial. The results of microbial counts are reported in Table 3. Before pasteurization, LAB showed concentration of 5.7 and 7.0 log CFU/g in the control and in the samples added with the bioprotective culture, respectively. The thermal treatment decreased their presence at about 2 log CFU/g in both samples. However, LAB counts were higher in the samples containing BC2 during storage, reaching counts of about 7 log CFU/g after 30 days of storage at 6°C.

Enterobacteriaceae were below the detection level after the thermal treatment and no growth of this microbial group was observed during storage. By contrast, after 30 days lower counts of coagulase negative staphylococci were observed in the samples containing BC2 (4.2 log CFU/g vs. 5.8 log CFU/g). Also yeast concentration was lower in the presence of the culture BC2 (about 2.0 log CFU/g vs. 4.4 log CFU/g after 30 days).

The sensory analysis carried out on cooked *Tortelloni* immediately after production and after 30 days of storage evidenced the acceptability of both samples of pasta (Figure 3). After production, six of the attributes (colour, salty, sourness, bitterness, cohesiveness and chewiness), as well as the overall acceptability, did not result significantly different, while two (odour intensity and aroma intensity) received significantly higher scores in the filled pasta containing BC2, probably due to the activity of bioprotective culture during the 18 h resting of the filling. At the end of storage all the attributes showed no significant differences between the two samples (Figure 3).

366

4. CONCLUSIONS

368 The addition of bioprotective cultures to the filling of fresh filled pasta had a relevant quantitative and
369 qualitative effect on the bacterial microbiota of the product during storage. Even if the added cultures were
370 never dominant at the end of shelf life, their presence and/or activity during the overnight incubation of
371 filling was sufficient to reduce the initial microbiota associated with raw materials and could drive the
372 evolution of microbial community toward the predominance of safer or more organoleptically acceptable
373 species, such as leuconostocs. Although the presence of these LAB cultures had some important effects on
374 the aroma profile of filled pasta, the overall impact on the sensory attributes after cooking was not
375 perceived by the panel, with the exception of more intense aroma and odour intensity immediately after
376 production. This is attributable to the presence of molecules (such as acetoin and diacetyl) compatible with
377 the Ricotta/cheese based filling of *Tortelloni*. The use of these cultures allowed the application of milder
378 thermal treatments with the aim to better maintain the traditional textural and flavor characteristics of this
379 kind of pasta combined with a higher a_w needed to preserve the softness of the filling.

380

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FIGURE CAPTIONS

Figure 1: Flow sheet of the Ricotta based *Tortelloni* production.

Figure 2: Microbial community composition at species level for *Tortelloni* samples by 16S rRNA gene V3–V4 region sequencing. The taxa were plotted for their relative abundance in the control sample immediately after production, control sample after 15 days, samples inoculated with *Lactobacillus rhamnosus* (BC1) or a mixture of *L. rhamnosus* and *Lactobacillus paracasei* (BC2) after 15 days of storage.

Figure 3: Sensory data of cooked samples produced during the validation test immediately after production and after 30 days of storage at 6°C. Sample C: control; sample BC2: sample added with a mixture of *L. rhamnosus* and *Lactobacillus paracasei*. For each attribute, the presence of an asterisk indicates significant differences between the two samples.

Table 1: Microbial counts (log CFU/g) and pH values during production and storage of *Tortelloni* not inoculated (control) or inoculated with the bioprotective culture BC1 (*Lactobacillus rhamnosus*) or BC2 (*L. rhamnosus* and *Lactobacillus paracasei*). Results are the mean of three independent repetitions. For each microbial group significant differences between products at each sampling point are indicated by different capital letters.

Microbial group	Sample	Before pasteurization	After pasteurization *	T7	T15	T30
LAB	Control	5.43 ^A	3.83 ^A	4.03 ^A	7.08 ^A	7.14 ^A
	BC1	7.39 ^B	5.27 ^B	6.64 ^B	7.90 ^B	8.38 ^B
	BC2	7.28 ^B	5.31 ^B	6.56 ^B	8.03 ^B	8.55 ^B
Staphylococci	Control	3.87 ^{AB}	2.93	2.81 ^A	3.10	3.06 ^A
	BC1	4.12 ^A	3.30	3.77 ^B	3.46	3.10 ^A
	BC2	3.65 ^B	3.17	2.97 ^A	3.34	2.44 ^B
Yeasts	Control	3.96	<1 ^{**}	<1	2.69 ^A	2.48 ^A
	BC1	4.07	<1	<1	<1 ^B	<1 ^B
	BC2	3.98	<1	<1	<1 ^B	<1 ^B
Enterobacteriaceae	Control	6.20	4.07 ^A	4.57	5.91 ^A	4.95 ^A
	BC1	6.39	4.52 ^B	4.21	3.29 ^B	2.57 ^B
	BC2	6.41	4.48 ^B	4.56	2.13 ^C	<1 ^C
pH	Control	5.70	5.88	5.77 ^A	5.85 ^A	5.74 ^A
	BC1	5.74	5.90	5.92 ^B	5.90 ^A	5.45 ^B
	BC2	5.70	5.94	5.98 ^B	5.61 ^B	5.58 ^C

* 60°C for 3 min in the product inner part

** Below detection limit (1 log CFU/g). In order to apply ANOVA these values have been numerically treated as 1.

Table 2: Volatile compounds detected by SPME-GC-MS in the samples immediately after pasteurization and after 30 days of storage at 6°C. Data are expressed as ratio between peak area of each molecule and peak area of the internal standard (4-methyl-2-pentanol). For each sampling time, different capital letters represent statistically significant differences ($p < 0.05$) between samples according to Tukey test of the two-way ANOVA.

Volatile compounds	After pasteurization *			After 30 days of storage		
	Control	BC1	BC2	Control	BC1	BC2
Acetone	6.88 ^A	6.71 ^A	6.04 ^A	6.79 ^A	- ^{**B}	- ^B
2-butanone	4.64 ^A	3.15 ^B	2.22 ^C	2.81 ^B	- ^D	- ^D
2,3-butanedione	1.86 ^A	22.15 ^B	18.30 ^B	1.22 ^A	0.90 ^A	1.04 ^A
2-pentanone	26.83 ^A	27.03 ^A	26.90 ^A	19.07 ^B	2.00 ^C	2.64 ^C
2-hexanone	49.92 ^A	38.16 ^B	33.58 ^B	44.37 ^A	33.75 ^B	27.25 ^B
2-heptanone	89.79 ^A	98.90 ^{AB}	105.19 ^B	99.95 ^{AB}	64.09 ^C	89.66 ^A
2-octanone	1.89 ^A	1.87 ^A	2.24 ^A	1.96 ^A	1.48 ^A	1.81 ^A
3-hydroxy-2-butanone	3.94 ^A	13.12 ^B	11.17 ^B	- ^C	- ^C	- ^C
2-nonanone	34.60 ^A	38.75 ^A	39.16 ^A	40.46 ^A	28.78 ^B	38.17 ^A
2-undecanone	5.57 ^A	5.80 ^A	5.92 ^A	6.81 ^{AB}	6.47 ^A	7.42 ^B
Total ketones	225.92^A	255.64^A	250.72^A	223.44^A	137.47^B	167.98^C
3-methyl - butanal	2.33 ^A	2.15 ^A	2.27 ^A	3.53 ^B	1.65 ^A	1.64 ^A
2-butenal	7.03 ^A	6.33 ^A	5.12 ^A	7.38 ^A	- ^B	- ^B
Hexanal	28.38 ^A	25.17 ^A	23.67 ^A	8.57 ^B	3.92 ^C	4.59 ^C
Nonanal	5.33 ^{AB}	6.99 ^A	6.55 ^A	6.29 ^A	4.53 ^B	6.83 ^A
Decanal	2.96 ^A	3.72 ^A	3.62 ^A	3.91 ^A	2.78 ^A	1.94 ^B
Benzaldehyde	4.94 ^A	6.17 ^{<B}	7.29 ^B	1.98 ^C	1.56 ^C	- ^D
Benzenacetaldehyde	4.19 ^A	4.71 ^A	4.04 ^A	6.21 ^B	5.30 ^{AB}	6.32 ^B
Total aldehydes	55.16^A	55.24^A	52.56^A	37.88^B	19.74^C	21.31^C
Ethyl alcohol	10.17 ^A	8.33 ^{AB}	7.19 ^B	216.58 ^C	571.67 ^D	508.84 ^D
2-pentanol	5.83 ^A	6.54 ^A	6.96 ^A	10.27 ^B	8.86 ^C	9.39 ^B
1-butanol	2.89 ^A	1.94 ^A	1.58 ^A	2.16 ^A	4.35 ^B	3.16 ^{AB}
1,2-cyclopentanediol, 3-methyl	1.71 ^A	1.80 ^A	2.90 ^B	1.33 ^A	2.80 ^B	2.68 ^B
1-pentanol	4.00 ^A	3.52 ^A	3.87 ^A	5.44 ^B	5.20 ^B	5.58 ^B
2-heptanol	7.88 ^A	8.15 ^A	8.03 ^A	8.42 ^A	21.78 ^B	21.63 ^B
3-penten-2-ol	- ^A	2.33 ^B	2.60 ^B	0.93 ^C	0.77 ^A	1.86 ^B
1-hexanol	7.58 ^A	12.33 ^B	11.62 ^B	12.93 ^B	23.92 ^C	23.91 ^C
1-octen-3-ol	1.45 ^A	1.31 ^A	1.26 ^A	2.58 ^B	2.25 ^B	2.58 ^B
1-hexanol-2-ethyl	2.80 ^A	3.33 ^A	3.24 ^A	4.24 ^B	3.16 ^A	4.28 ^B
2-nonanol	2.20 ^A	2.50 ^A	2.47 ^A	2.62 ^A	9.19 ^B	8.99 ^B
1-nonanol	- ^A	- ^A	- ^A	- ^A	2.21 ^B	2.43 ^B
Phenylethyl alcohol	- ^A	- ^A	- ^A	- ^A	3.67 ^B	4.24 ^B
Total alcohols	46.50^A	52.08^A	51.71^A	267.49^B	659.83^C	599.58^C
Acetic acid, ethyl ester	2.81 ^A	1.83 ^{AB}	1.34 ^A	4.54 ^C	46.22 ^D	57.09 ^E
Butanoic acid, ethyl ester	31.69 ^A	24.30 ^B	20.00 ^B	44.22 ^C	53.60 ^D	36.63 ^{AC}
Hexanoic acid, ethyl ester	52.63 ^A	58.12 ^A	66.93 ^B	66.96 ^B	120.58 ^C	97.81 ^D
Heptanoic acid, ethyl ester	1.33 ^A	1.19 ^A	1.30 ^A	1.81 ^A	2.49 ^B	2.58 ^B
Octanoic acid, ethyl ester	7.45 ^A	6.06 ^{AB}	5.43 ^B	11.04 ^C	20.57 ^D	23.68 ^D
Decanoic acid ethyl ester	2.53 ^A	2.30 ^A	2.15 ^A	3.91 ^B	9.34 ^C	7.99 ^C
Pentanoic acid, ethyl ester	2.36 ^A	3.00 ^A	2.98 ^A	- ^B	6.63 ^C	6.05 ^C
Total esters	100.82^A	96.80^A	100.13^A	132.49^B	259.42^C	231.84^C

Acetic acid	16.72 ^A	16.06 ^A	15.68 ^A	14.37 ^A	84.39 ^B	82.63 ^B
Butanoic acid	79.33 ^A	81.13 ^A	80.59 ^A	71.41 ^A	183.63 ^B	165.14 ^B
Hexanoic acid	95.57 ^A	100.27 ^A	104.90 ^A	81.54 ^A	249.19 ^B	229.06 ^B
Octanoic acid	28.00 ^A	38.41 ^{AB}	41.16 ^B	24.57 ^A	71.86 ^C	65.68 ^C
Total acids	219.62^{AB}	235.87^A	242.33^A	191.89^B	589.07^C	542.51^C

* 60°C for 3 min in the product inner part

** Not detected under the adopted conditions. For two-way ANOVA the samples under the detection limit were set as 0.

Table 3: Microbial counts (log CFU/g) and pH during production and storage of *Tortelloni* not inoculated (control) or inoculated with the bioprotective culture BC2 (*Lactobacillus rhamnosus* and *Lactobacillus paracasei*) during the validation test. Results are the mean of three independent repetitions. For each microbial group significant differences ($p < 0.05$) between the two products at each sampling point are indicated by the presence of an asterisk.

Microbial group	Sample	Before pasteurization	After pasteurization ¹	T7	T15	T30
<i>LAB</i>	Control	5.67*	2.44	1.95*	4.47*	6.70
	BC2	7.02	1.95	4.48	5.14	6.94
<i>Staphylococci</i>	Control	3.33	<1 ²	2.45*	3.70*	5.82*
	BC2	3.63	<1	<1	2.15	4.24
<i>Yeasts</i>	Control	2.80	1.32	1.33*	2.66*	4.37*
	BC2	2.41	1.51	<1	1.56	1.97
<i>Enterobacteriaceae</i>	Control	4.68	<1	<1	<1	<1
	BC2	4.51	<1	<1	<1	<1
<i>pH</i>	Control	5.39	5.38	5.73	5.71	5.64
	BC2	5.47	5.43	5.73	5.80	5.66

¹ 60°C for 3 min in the product inner part

² Below detection limit (1 log CFU/g). In order to apply ANOVA these values have been numerically treated as 1.

Figure 1

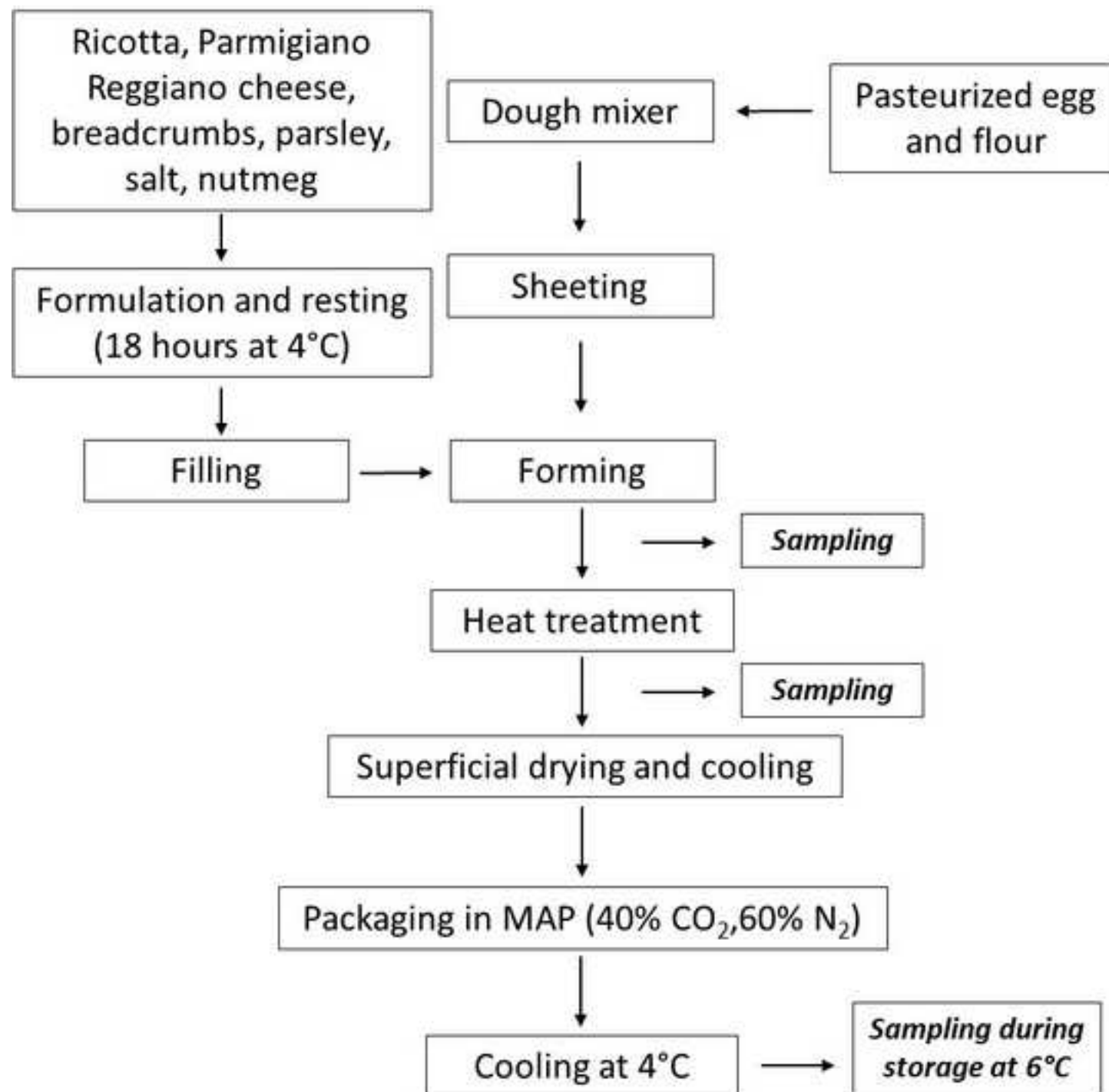


Figure 2

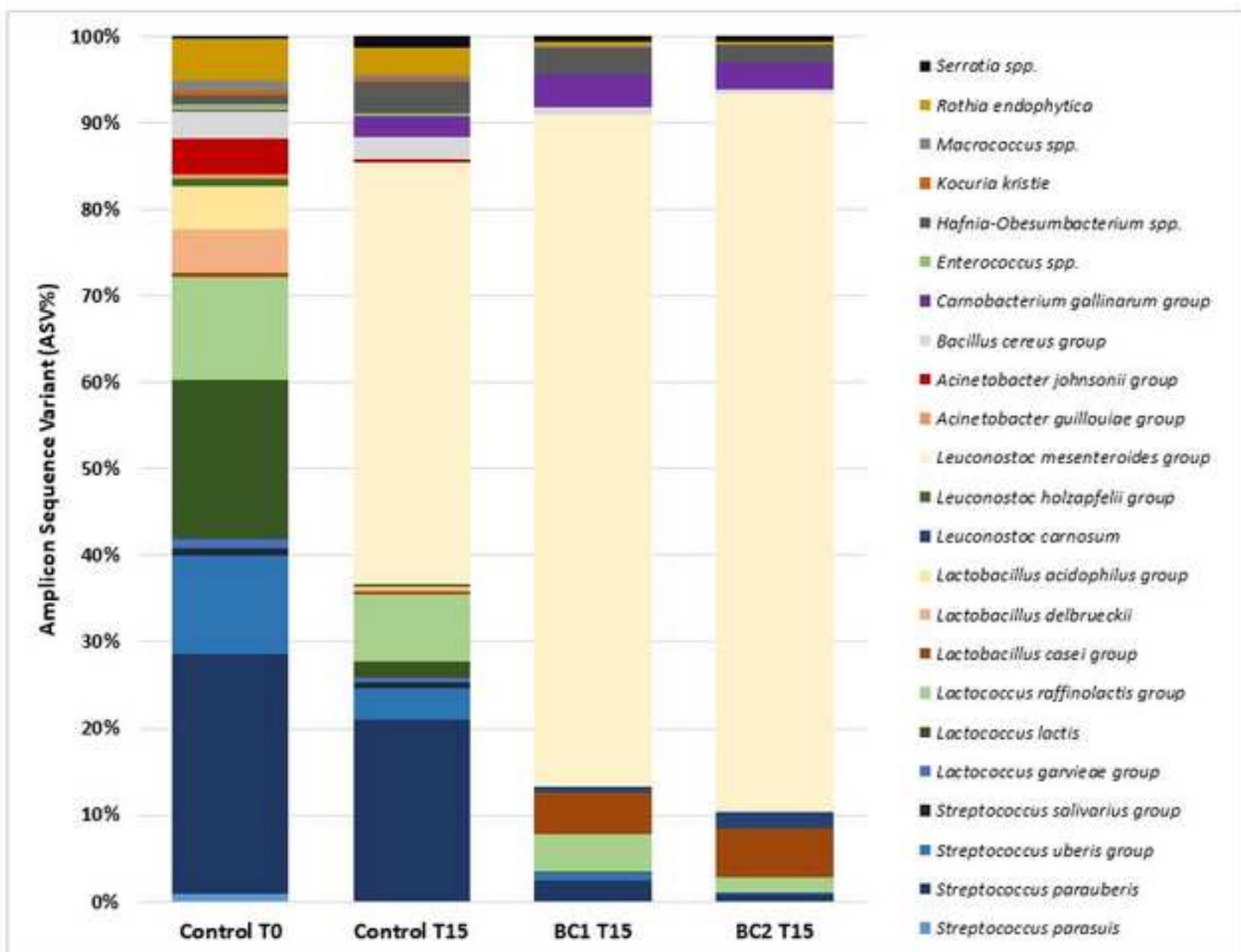


Figure 3

