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# Journey to the morpho-textural traits, microbiota, and volatilome of Ciauscolo PGI salami

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1	Journey to the morpho-textural traits, microbiota, and volatilome of Ciauscolo PGI salami
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32 Abstract

33

34 In the present study, microbial dynamics of naturally fermented Ciauscolo PGI salami were studied through culture-35 dependent and metataxonomic analyses. Moreover, physico-chemical and morpho-textural evaluation, together with the 36 analysis of volatile organic compounds were also carried out. Cultures of Latilactobacillus sakei were isolated from the 37 analyzed samples and characterized for: i) acidification activity; ii) enzymatic activity; iii) the production of EPS; iv) 38 production of bacteriocins against Listeria innocua, utilized as surrogate for Listeria monocytogenes. Pseudomonas spp. 39 represented the dominant taxon in the raw materials, whereas L. sakei was the dominant species from  $t_5$ , together with 40 Dellaglioa algida and Leuconostoc carnosum. Among eumycetes, Cladosporium cladosporioides, Debarvomyces 41 hansenii, Kurtzmaniella zeylanoides, and Malassezia restricta/globosa were the most abundant yeasts occurring in all 42 samples. The 44 L. sakei cultures isolated from the analyzed samples showed a suitable acid production capacity, together 43 with the capability to cope with the main environmental stresses occurring in fermented sausages. For most of the isolates, 44 strong aminopeptidase activity (leucine arylamidase and the valine arylamidase) was observed. Moreover, the majority 45 of isolates showed the *in vitro* production of sucrose-dependent exopolysaccharides. Based on these evidence, a few 46 candidate starter or adjuncts cultures, with potential use for product safety and quality improvement, were highlighted, 47 namely L. sakei C5, C7, C11, C31, C45, C48, C53, C55, and C60. In the analyzed samples, 53 volatile substances were 48 fully or tentatively identified; among these, spices-derived components (black pepper and garlic cloves) were constantly 49 detected throughout the whole ripening time.

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51 Keywords: fermented sausages, metataxonomic analysis, *Latilactobacillus sakei*, acidification, enzymatic activity.

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#### 53 1. Introduction

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55 The preservation of meat through drying and fermentation represents the most ancient method to overcome meat decay, 56 as already told in the Homer's Odyssey, ca. 900 B.C., and in some historical texts dating back to the old Roman Empire 57 (Zeuthen, 1995). Nowadays, those that are called charcuterie products represent food delicacies that are highly appreciated 58 by consumers, since they are produced in accordance with ancient traditions. Among charcuterie products, fermented 59 sausages are usually manufactured using minced swine meat (or meat from other animal species as chicken or lamb) and 60 lard, added with salt and spices, and stuffed into animal casings. Sugars, preservatives, antioxidants, and microbial starter 61 cultures can also be added to the meat batter (Cruxen et al., 2019). Then, sausages are left to ferment at specific 62 environmental conditions (temperature and relative humidity) that vary according to the production process.

During fermentation, the sausage is subjected to physico-chemical modifications that are the result of the activity of native
 meat enzymes and microorganisms (Belleggia et al. 2022a). Such modifications contribute to the firmness, cohesiveness,
 and safety of fermented sausages (Cruxen et al., 2019).

In fermented sausages, the microbial succession occurring from the stuffing of the meat batter till the end of ripening produce the transformation of the raw meat into the end product. In more detail, as soon as the meat batter is stuffed into the natural or artificial casing, the naturally occurring adventitious microbiota (e.g., Enterobacteriaceae, *Pseudomonas*, *Campylobacter*, *Psychrobacter*, and *Brochothrix*) is progressively substituted by pro-technological microorganisms as lactic acid bacteria, coagulase-negative cocci, and eumycetes (Belleggia et al., 2020a).

Lactic acid bacteria are among the key pro-technological microorganisms involved in the biochemical, microbiological, and sensory modifications occurring during the production of fermented sausages (Zdolec, Mikuš, & Kiš, 2022). Lactic acid bacteria can naturally be present as minority fraction in the raw materials, or they can deliberately be added to the meat batter as starter or adjunct cultures. In fermented sausages, lactic acid bacteria are primarily responsible for the acidification of the product; moreover, they can exert other biological activities that include the production of bacteriocins and exopolysaccharides (EPS), with positive affect on safety and sensory attributes of the product (Bellegia et al., 2022b). Of note, the metabolic activity of lactic acid bacteria can every so often produce unwanted substances, as biogenic amines

78 (e.g., histamine), with negative impact on the safety of fermented sausages (Fong, El-Nezami, & Po Sze, 2021).

Although the use of lactic acid bacteria as starters or adjuncts for the production of fermented sausages is well established,

80 the selection and characterization of new cultures with biotechnological and safety advantages is still needed (Zdolec et

81 al., 2022). To this end, naturally fermented sausages could represent a source of microbial biodiversity for the selection

82 of lactic acid bacteria with pro-technological features to be used for product improvement (Zdolec et al., 2022).

In Central Italy, a great number of naturally fermented sausages is produced in accordance with ancient traditions, with *Ciauscolo* PGI salami being recognized as one of the most well-known and appreciated products of the Marche Region (Central Italy). *Ciauscolo* PGI salami obtained the Protected Geographical Indication (PGI) status according to Commission Regulation (EC) No 729/2009 of 10 August 2009. The name *Ciauscolo*, also referred to as *Ciavuscolo* or *Ciabuscolo*, originates from the Latin words "*ciabusculum*" or "*cibusculum*" that were used to describe a little portion of

88 food or snack that farmers were used to consume in tiny quantities during breaks and between main meals.

*Ciauscolo* PGI salami consists of a mixture of pork meat resulting from the following cuts used in descending order:
belly, up to a maximum of 70% (w w<sup>-1</sup>); shoulder, up to a maximum of 40% (w w<sup>-1</sup>); trimmings of ham and loin, up to a
maximum of 30% (w w<sup>-1</sup>). The main physical characteristic of *Ciauscolo* PGI salami is softness up to spreadability.

92 To produce *Ciauscolo* PGI salami, the meat cuts are minced using a 2-3 mm plate, together with salt, ground black 93 pepper, wine, and crushed garlic. The addition of sugars (lactose, dextrose, fructose, or sucrose) as well as of L-ascorbic 94 acid, sodium ascorbate, potassium nitrate is allowed. Once prepared, the meat batter is stuffed into pork or bovine intestine 95 casings and ripened for at least 15 days. The final pH of Ciauscolo PGI salami must be higher than or equal to 4.8, with 96 minimum protein content attesting at 15% (w w<sup>-1</sup>) and fat content comprised between 32 and 42% (w w<sup>-1</sup>). According to 97 the production disciplinary, the maximum water/protein ratio allowed is 3.10, whereas the maximum fat/protein ratio 98 allowed is 2.80. The weight of the end product ranges from 400 to 2,500 g, and the diameter ranges between 4.5 and 10 99 cm. Ciauscolo PGI salami has a cylindrical shape with a length comprised between 15 and 45 cm. The slice of Ciauscolo 100 PGI salami is pink, uniform, homogeneous, and free from rancid fractions. The smell of this fermented sausage is delicate, 101 aromatic, and spicy, whereas the taste is savory and delicate, but not acidic. The minimum load of lactic acid bacteria that 102 must be present in the end product is higher than 7 log colony forming units (cfu) g<sup>-1</sup>. Ciauscolo PGI salami can only be 103 manufactured in the geographical area covered by four (Ancona, Ascoli Piceno, Fermo, and Macerata) out of the five 104 provinces of the Marche Region.

105 Although a few studies already dealt with the microbiology of Ciauscolo PGI salami (Aquilanti et al., 2007; Belleggia et 106 al., 2020a; Federici et al., 2014; Silvestri et al., 2007), to the authors' knowledge no research on the characterization of 107 pro-technological traits of autochthonous lactic acid bacteria isolated from this peculiar food matrix has been performed, 108 yet. Similarly, scarce data on its volatilome are available in the scientific literature. Hence, in the present study, different 109 selective growth media and a metataxonomic approach were applied to study the dynamics of the microbiota naturally 110 occurring in Ciauscolo PGI salami manufactured by an artisanal producer located in the Marche Region and to isolate a 111 pool of lactic acid bacteria with potential pro-technological features (starter or adjunct cultures). Moreover, the analysis 112 of volatile organic compounds (VOCs) via Headspace Solid-Phase Microextraction-Gas Chromatography/Mass

113	Spectrometry (HS-SPME-GC/MS) analysis was also carried out. Finally, a pool of isolates ascribed to Latilactobacillus
114	sakei were characterized for: i) acidification activity; ii) key enzymatic activities; and iii) the production of EPS.
115	As reported by EFSA BIOHAZ Panel (EFSA Panel on Biological Hazards, 2018), ready-to-eat foods, including fermented
116	sausages, might represent a risk for the presence of Listeria monocytogenes (Petruzzelli et al., 2010), hence, the L. sakei
117	isolates were also tested for production of bacteriocins against Listeria innocua, utilized as surrogate for L. monocytogenes
118	(ANSES, 2019).
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120	2. Materials and methods
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122	2.1. Ciauscolo PGI salami production
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124	Two independent manufacturing batches (marked as batch 1 and batch 2) of Ciauscolo PGI salami were analyzed. Both
125	batches were manufactured by an artisan producer located in the geographical area of production (Macerata province of
126	the Marche Region), in accordance with the Ciauscolo PGI salami production disciplinary. The first batch was produced
127	in early spring, whereas the second in in late spring of the same year. According to the disciplinary for production of
128	Ciauscolo PGI salami, the meat batter was prepared using the following swine cuts: shoulder 25% (w w <sup>-1</sup> ), belly (including
129	fat) 40% (w w <sup>-1</sup> ), loin 10% (w w <sup>-1</sup> ), and ham 21% (w w <sup>-1</sup> ). Moreover, salt 2.7% (w w <sup>-1</sup> ), ground black pepper ~0.4% (w
130	w <sup>-1</sup> ), ground garlic ~0.1% (w w <sup>-1</sup> ), white wine (20 mL/kg), and 0.05% (w w <sup>-1</sup> ) potassium nitrate (E252) were also added.
131	No starter cultures were added.
132	For each batch a total of 100 kg of meat batter was processed. The meat batter was minced twice with a 3 mm plate and
133	stuffed into swine bowels previously washed in a mixture of water and white wine vinegar 50% (v/v). The drying was
134	performed for 5 days under the following conditions: progressive temperature decrease, from 20 to 14 °C, and concomitant
135	progressive relative humidity (R.H.) decrease from 99 to 60%. Ripening of the sausages was carried out at 15 °C and
136	75% R.H. for 15 days.
137	For each batch, fermented sausages approximately weighing 750 g, with 26 cm length and 6 cm width, were produced
138	(Fig. 1). Collection of samples was performed at 0, 5, 10, 20 days; for each ripening time and batch, three fermented
139	sausages (for a total of 24 sausages) were collected. Samples were aseptically collected using sterile instruments and bags
140	(Nasco Whirl-Pak Easy-To-Close Bags, Fisher Scientific Italia, Rodano, Italy). The samples were rapidly refrigerated
141	and analyzed during the same day of collection.
142	

143 2.2. Physico-chemical analyses

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- A pH meter equipped with an HI2031 solid electrode (Hanna Instruments, Padova, Italy) was used to determine pH values of *Ciauscolo* PGI salami samples through direct insertion of the probe into the food matrix. The data, in duplicate for each sample, were reported as mean ± standard deviation.
- 148 Total titratable acidity (TTA) was determined on 10 g of sample homogenized in 90 mL of distilled water, adjusted to a
- pH value of 8.3 with a 0.1 N solution of NaOH used as a titrant. Percentage (%) TTA of lactic acid equivalents was
- calculated as previously reported by Rampanti et al. (2023). TTA% was reported as mean ± standard deviation of two
  replicates.
- 152 The amounts of acetic acid and lactic acid were assessed using Acetic Acid (Acetate Kinase Manual Format) test kit and
- 153 D-/L-Lactic Acid (D-/L-Lactate) (Rapid) test kit (Megazyme, Bray, Ireland). The data, in duplicate for each sample, were

154 expressed as g  $100g^{-1}$  of the sample and reported as mean  $\pm$  standard deviation.

- 155 The water activity (a<sub>w</sub>) was assessed in accordance with the ISO 21807:2004 standard method using an Aqualab 4TE
- apparatus (Meter Group, Pullman, USA). The data, in duplicate for each sample, were reported as mean ± standard
  deviation.
- Color measurements was performed using a Chroma Meter CR-200 (Minolta, Osaka, Japan) with a D65 illuminant. Color was determined on 2 cm thick slices according to CIE L\*a\*b\* system (L\*, lightness; a\*, redness/greenness; b\*, blueness/yellowness) (Belleggia et al. 2022b). Visual changes during ripening of the sausages were evaluated by longitudinally cutting them (7 mm thickness) and imaging the cross sections with a scanner (ENVY 6200 Series, HP, Palo Alto, CA, United States) (Dreher et al., 2021)
- Cylindrical specimens of meat batter and sausages (height: 15 mm, diameter: 20 mm) were excised and then uniaxially
  compressed twice with a CT3-4500 texture analyzer (Brookfield Engineering Laboratories Inc., Middleboro MA, USA)
  equipped with a 36 mm diameter cylindrical probe (mod. TA-AACC36) at 1.5 mm/s using a non-destructive deformation
- 166 (40%) (Dreher et al., 2021). Specimens were positioned between the load cell and the fixture base table of the instrument.
- 167 A 4500 g load cell was used.
- 168
- 169 2.3. Viable counts

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171 Viable counts were performed by mixing 10 g of each sample with 90 mL of sterile peptone water (1 g L<sup>-1</sup> of 172 bacteriological peptone) homogenized using a Stomacher apparatus (400 Circulator, International PBI, Milan, Italy) for 173 3 min at 260 rpm. After homogenization, ten-fold serial dilutions were prepared, and viable counts of the following 174 microbial groups were evaluated: i) lactic acid bacteria on De Man Rogosa and Sharpe (MRS) agar added with 250 mg

- 175 L<sup>-1</sup> of cycloheximide (250 mg L<sup>-1</sup>) to inhibit the growth of eumycetes, with incubation at 37 °C for 48-72 h; ii) coagulase-
- 176 negative cocci on Mannitol Salt Agar (MSA), with incubation at 37 °C for 48-72 h; iii) Enterobacteriaceae on Violet Red
- 177 Bile Glucose Agar (VRBGA), with incubation at 37 °C for 24 h; iv) eumycetes on Rose Bengal Chloramphenicol Agar,
- 178 with incubation at 25 °C for 72-96 h. The results of two biological and three technical replicates were expressed as the
- 179 log of colony-forming units (cfu) per gram of sample and reported as mean ± standard deviation.
- 180 Finally, a miniVIDAS apparatus (Biomerieux, Marcy l'Etoile, France) was used to assess the presence/absence of *Listeria*
- 181 *monocytogenes* and *Salmonella* spp. through the enzyme-linked fluorescent assay (ELFA) method, in accordance with
- 182 the AFNOR BIO 12/11-03/04 and AFNOR BIO 12/16-09/05 standard methods, respectively (Haouet et al., 2017).
- 183
- 184 2.4. Real-time PCR analysis for the detection of botulinic toxins genes
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- 187 Belleggia et al. (2020a). C. botulinum was analyzed in accordance with the multiplex real-time PCR method of the Italian
- 188 National Reference Centre for botulism (<u>http://old.iss.it/binary/cnrb/cont/CNRB31.010.pdf</u>) for the detection of the target
- 189 genes *bont/A*, *bont/B*, *bont/E*, *bont/F*, *and 4gyrB* (*CP*) that codify for botulinic toxins.
- Briefly, 25 g of *Ciauscolo* PGI salami sample were blended in 225 mL Triptone Peptone Glucose Yeast extract (Microbiol Diagnostici, Italia), incubated in anaerobiosis at 30°C for 96 h and then extracted with 6% Chelex-100 (Biorad, Milan, Italy). The amplification was carried out using the Kit QuantiTect multiplex No Rox (Qiagen) in the Stratagene Mx3005P (Agilent Technologies) thermal cycler and the primers and probes nucleotide sequences are those already reported by Belleggia et al. (2020a). Two different Master mixes were used concurrently, with the following thermal profile: 1 cycle at 95°C for15 min followed by 40 cycles at 94°C for 30 s and 56°C for 90 s.
- 196
- 197 2.5. RNA extraction and cDNA synthesis
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- For each sample homogenate (dilution 10<sup>-1</sup>), prepared as reported above, 1.5 mL aliquot was centrifugated at 16,000 rpm for 10 min. RNA*later* Stabilization Solution (Ambion, Foster City, CA, USA) was used to protect the resulting cell pellets that were subsequently stored at -80 °C. Quick-RNA MiniPrep kit (Zymo Research, Irvine, California, USA) was then used to extract the total microbial RNA from the cell pellets in accordance with manufacturer's instructions. PCR amplification with universal prokaryotic primers (27 f, 1495 r) (Weisburg et al., 1991) was carried out to check for the presence of residual DNA. SensiFAST cDNA Synthesis Kit for RT-qPCR (Bioline, London, UK) was used to synthesize cDNA according to manufacturer's instructions.

<sup>186</sup> The presence/absence of *Clostridium botulinum* was assessed via multiplex real-time PCR as already described by

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## 207 2.6. Amplicon target sequencing and bioinformatic analysis

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The cDNA was used as template to amplify the V3-V4 region of the 16S rRNA gene for bacteria (Klindworth et al.,
2013), whereas the D1 domain of the 26S rRNA gene of Large Ribosomal Subunit (LSU) was targeted for fungi (MotaGutierrez et al., 2019).

A total of 400,802 and 535,306 raw-reads were produced by 16S and 26S amplicon-based sequencing, respectively. To obtain Amplicon Sequence Variants (ASVs) the raw-reads were analyzed with *DADA2* package (Callahan et al., 2016) in R environment (R version 4.1.1; <u>http://www.r-project.org</u>). The pipeline described by Botta et al. (2022) was applied to filter and merge raw-reads: 157,311 paired-end bacterial reads (19,664 reads/sample) and 270,743 paired-end fungal reads (45,124 reads/sample) passed the filtering parameters and were used to construct ASVs tables.

Taxonomy was assigned with a 99 % sequence similarity through Bayesian classifier method (Wang et al., 2007) by matching bacterial ASVs to the 2021 release (version 138.1) of Silva prokaryotic SSU reference database (https://zenodo.org/record/4587955#.YObFvhMzZRE). Fungal ASVs taxonomy was assigned at 99 % against an internal database of 26S rRNA (Mota-Gutierrez et al., 2019). Taxonomy assignment for 16S and 26S was double checked by using BLASTn suite (https://blast.ncbi.nlm.nih.gov).

Fungal and bacterial ASVs were aligned with *DECIPHER* package and two unrooted phylogenetic tree were constructed with *phangorn* package (Schliep, 2011; Wright, 2016). Alpha diversity metrics and weighted UniFrac beta-diversity distance were calculated with *phyloseq* and *picante* packages (Kembel et al., 2010; McMurdie & Holmes, 2013): rarefaction limit was set to the lowest number of sequences/sample.

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227 2.7. GC-MS analysis of volatile components

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Two g of fresh salami were weighed in a 10 mL screw cap septum vial, then the vial was equilibrated at 40 °C for 15 min in a thermostatic bath. The static headspace was sampled by a 65 μm PDMS/DVB SPME fibre (Supelco, Bellefonte, PA) for 15 minutes. Chromatographic separation of volatiles was performed by a fused silica capillary column ZB-5 (30 m L, 0.25 mm ID, 0.25 μm FT; Phenomenex, Torrance, CA) mounted on a Varian 3900 gas chromatograph (Varian Analytical Instruments, Walnut Creek, CA); the injector was operating in splitless mode for 0.1 min at a constant temperature of 250 °C; oven temperature was increased from 40 °C to 220 °C at a rate of 6 °C/min, then held at the final temperature for 5 min; carrier gas (He) was set at constant flow mode (1.0 mL/min). The gas chromatograph was coupled to an ion trap 236 mass detector Saturn 2100T (Varian Analytical Instruments, Walnut Creek, CA): the trap and the transfer line were set at 237 200 °C and 220 °C, respectively; electron impact (70 eV) mass spectra were acquired in the mass range of 31-250 amu. 238 Volatile compounds were identified by matching mass spectral data collected in the NIST/EPA/NIH Mass Spectral 239 Library (Version 2.0a, build July 1 2002; National Institute of Standards and Technology) and Kovats Retention Indexes 240 (RIs) available in the public access database Pubchem (https://pubchem.ncbi.nlm.nih.gov/). A C8-C20 normal alkanes 241 mixture (Sigma-Aldrich, St. Louis, MO) was used to calculate RIs. An automated spreadsheet (Lucero et al., 2009) was 242 used for simplifying the calculation of RIs ok unknown components and speeding up the comparison with published 243 indexes.

- 244
- 245 2.8. Isolation and characterization of L. sakei
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### 247 2.8.1. Isolation and identification

As previously described, colonies of lactic acid bacteria grown on MRS agar added with cycloheximide were randomly selected and subsequently sub-cultured to purity under the same conditions. The obtained isolates were then stored at -80 C.

251 Prior to further analysis, lactic acid bacteria were thawed and sub-cultured twice on MRS agar at 37 °C for 48 h. DNA 252 was extracted from the lactic acid bacteria isolates according to Osimani et al. (2015) and its purity and quantity was 253 verified with a NanoDrop ND 1000 (Thermo Fisher Scientific, Wilmington, DE, USA). DNAs were standardized to a 254 final concentration of 100 ng  $\mu$ L<sup>-1</sup> and subjected to PCR in a My Cycler Thermal Cycler (BioRad Laboratories, Hercules, 255 CA, USA) using the universal prokaryotic primers 27f and 1495r, as described by Osimani et al. (2015). The amplification 256 was verified by electrophoresis in 1.5% (w  $v^{-1}$ ) agarose gel in 0.5X Tris/Borate/EDTA (TBE) buffer containing 0.5  $\mu$ g 257 mL<sup>-1</sup> GelRed® Nucleic Acid Gel Stain, 10,000X in water (Biotium, San Francisco Bay Area, USA). The electrophoretic 258 run included the HyperLadder<sup>™</sup> 1 kb (Meridian Bioscience, Cincinnati, Ohio, USA) as molecular weight standard and 259 was carried out at 75 V for 3.5 h and visualized under UV light. The amplicons were then shipped to Genewiz (Takaley, 260 UK) for their purification and sequencing.

The raw sequences were analysed with UCHIME2 software tool to uncover chimeras (Edgar, 2016) and were trimmed to remove NNNs and misleading data from the terminations. Afterwards, a BLAST search was exploited to compare the obtained sequences with 16S rRNA sequences of type strains from GenBank DNA database (http://www.ncbi.nlm.nih.gov/). The sequences of the lactic acid bacteria cultures were finally submitted to GenBank DNA database to acquire the respective accession numbers.

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#### 267 2.8.2. Acidification in synthetic medium

268 L. sakei isolates were sub-cultured twice in MRS broth incubated at 30 °C for 18 h (Wen et al., 2021). The cultures were 269 centrifuged at 4,000 rpm for 5 min and the pellets washed with sterile physiological solution (0.9 % w v<sup>-1</sup>) prior to 270 resuspension in the same diluent. The concentration of bacterial cells in the suspensions was determined using a 271 spectrophotometer (Shimadzu UV-1800, Shimadzu Corporation, Kyoto, Japan) by measuring the optical density (OD) at 272 600 nm. All the isolates were inoculated to 8 log cfu mL<sup>-1</sup> in 10 mL of: i) MRS broth; ii) MRS broth added with 150 mg 273 Kg<sup>-1</sup> of E250 (sodium nitrite); iii) MRS broth added with 150 mg Kg<sup>-1</sup> of E252 (potassium nitrate); iv) MRS broth added 274 with 3% (w v<sup>-1</sup>) of NaCl; v) MRS broth added with 150 mg Kg<sup>-1</sup> of E250, 150 mg Kg<sup>-1</sup> of E252, and 3% (w v<sup>-1</sup>) of NaCl. 275 The pH values of the media were measured prior to inoculation (t<sub>0</sub>) and after incubation at 30 °C for 0, 4, and 24 h.

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### 277 2.8.3. Semi-quantitative assessment of enzymatic activities

278 The semi-quantitative micromethod API® ZYM (bioMérieux, Marcy-l'Etoile, France) was used for the assessment of key 279 enzymatic activities of the selected pool of lactic bacteria in accordance with manufacturer's instructions. Each API® 280 ZYM strip is composed of 20 cupules containing synthetic substrates that are inoculated with the microorganism to be 281 tested in a water suspension, this latter used to rehydrate the enzymatic substrates. The 20 cupules are designed to evaluate 282 the activity of the following enzymes: 1 - control; 2 - alkaline phosphatase; 3 - esterase (C 4); 4 - esterase lipase (C 8); 5 283 - lipase (C 14); 6 - leucine arylamidase; 7 - valine arylamidase; 8 - cystine arylamidase; 9 - trypsin; 10 - alpha-284 chymotrypsin; 11 - acid phosphatase; 12 - naphthol-AS-BI-phosphohydrolase; 13 - alpha-galactosidase; 14 - beta-285 galactosidase; 15 - beta-glucuronidase; 16 - alpha-glucosidase; 17 - beta-glucosidase; 18 - N-acetyl-ß-glucosaminidase; 286 19 - alpha-mannosidase; 20 - alpha-fucosidase. The metabolic end-products produced during the incubation period were 287 detected through coloured reactions revealed by the addition of reagents.

Briefly, a suspension in API Suspension Medium (2 mL) of each isolate grown on MRS agar was prepared according to a turbidity of 5-6 McFarland. Sixty-five µL of the obtained suspension were used for the inoculation of each cupule of the API® ZYM strips with incubation for 4 hours at 37°C. After incubation, 1 drop of ZYM A reagent and 1 drop of ZYM B reagent were added to each cupule until colour development (at least 5 minutes). For each cupule, a value ranging from "0" to "5" was assigned, corresponding to the colour developed: 0 corresponding to a negative reaction, 5 to a reaction of maximum intensity and 1, 2, 3, or 4 were intermediate reactions depending on the level of intensity (3, 4, or 5 being considered as positive reactions).

295

### 296 2.8.4. Assessment of antimicrobial activity

297 The antimicrobial activity of the selected pool of isolates ascribed to L. sakei was performed by following the agar well 298 diffusion assay, described by Parente et al. (1995). Briefly, molten Brain Heart Infusion (BHI) soft agar (0.75% agar) 299 (VWR Prolabo Chemicals) was inoculated (2%, v v<sup>-1</sup>) with the indicator microorganism *Listeria innocua*. Twenty mL of 300 the inoculated medium were poured into 90 mm Petri dish and left to solidify. A cone of a 200 µL sterile tip was utilized 301 to create wells of  $\sim 50 \,\mu\text{L}$  capacity on BHI soft agar. Previously, lactic acid bacteria were sub-cultured twice in MRS 302 broth at 37 °C for 48 h. The broth cultures were then added with 0.1 N NaOH solution to reach pH 7.0 to neutralise the 303 organic acids produced during the bacterial growth. A filtration step on sterile PES membrane filter of 0.22 µm pore size 304 (Laboindustria S.p.A., Padova, Italy) followed. For each lactic acid bacteria isolate, 4 wells were formed on BHI soft 305 agar, each containing: (i) 50  $\mu$ L of the sub-cultured suspension; (ii) 50  $\mu$ L of the neutralised suspension adjusted to pH 306 7.0; (iii) 50  $\mu$ L of the filtered neutralised suspension; (iv) 50  $\mu$ L of sterilised water as a negative control. Afterwards, the 307 Petri dishes were incubated at 37 °C for 24 h and examined for the presence of zones of inhibition. In the case of positive 308 results (presence of inhibition halo), 3 spots of 5 µL each of pepsin (Fluka<sup>TM</sup>, Honeywell, Morristown, USA), trypsin 309 (Fluka<sup>TM</sup>) or Pronase (Merck KGaA, Darmstadt, Germany) were set along the circumference of the inhibition zone to 310 evaluate the protein nature of the antimicrobial. The Petri dishes were further incubated at the same conditions. The 311 antimicrobial activity of lactic acid bacteria isolates due to the synthesis of bacteriocins was confirmed by the formation 312 of crescents.

313

### 314 2.8.5. In-vitro EPS production

315 The selected pool of isolates ascribed to L. sakei was screened for EPS production, based on the method already reported 316 by Hilbig et al. (2019) with some modifications. In more detail, the isolates were retrieved from cryo-protective 317 suspensions and sub-cultured twice on MRS broth at 37 °C for 48 h. The EPS production was visually observed by adding 318 aliquots (5 µL) of each bacterial culture on the following solid media: (i) MRS agar supplemented with sucrose (80 g L 319 <sup>1</sup>) to promote the synthesis of homopolysaccharides (HoPS); MRS agar supplemented with yeast extract (10 g L<sup>-1</sup>), meat 320 extract (10 g  $L^{-1}$ ), galactose (20 g  $L^{-1}$ ), and lactose (20 g  $L^{-1}$ ) to promote the synthesis of heteropolysaccharides (HePS). 321 After an incubation period of 48 h at 30 °C, the colonies were classified as positive whenever presenting a mucoid aspect 322 (visible shiny and slimy appearance) or a ropy consistence (able to produce detectable filaments by using a sterile 323 toothpick). For each isolate, the analyses were conducted in duplicate.

324

325 2.9. Statistical analysis

- 326 To assess statistical differences within sausage samples, the Tukey-Kramer's Honest Significant Difference (HSD) test
- 327 (level of significance 0.05) was used by one-way analysis of variance (ANOVA). Tests were performed through JMP

328 v11.0.0 software (SAS Institute Inc., Cary, NC).

- 329 Permutational Multivariate Analysis of Variance (PERMANOVA) of the metataxonomic data were performed and
- 330 mycobiota/microbiota compositions were compared between the two batches by Wilcoxon rank sum test (P value
- adjustment method: FDR) in R environment.
- 332 Correlation between taxa (ASVs merged at highest taxonomic level achieved) and volatile organic compounds (VOCs)
- 333 was performed by means of Spearman's rank correlation.
- 334
- **335 3. Results**
- 336
- 337 *3.1. Physico-chemical analyses*
- 338

339 The results of physico-chemical analysis carried out on the analyzed *Ciauscolo* PGI samples are reported in Table 1.

- 340 In more detail, pH values ranged between 5.94 and 5.23, with samples collected at  $t_0$  showing the highest values,
- 341 irrespective of the production batch.

342 As for  $a_w$ , the detected values were comprised between 0.961 and 0.936, with samples collected from  $t_{10}$  to  $t_{20}$  showing

- 343 statistically significant lower values.
- 344 TTA % ranged between 0.594 and 0.990; for this parameter, as expected, the highest % were detected in samples at the
- 345 end of ripening, irrespective of the production batch.
- Regarding organic acids, lactic acid content ranged between 0.368 and 0.901 g 100 g<sup>-1</sup>, with the highest values recorded

in samples at the end of ripening, irrespective of the production batch. Acetic acid content was comprised between 0.014 and 0.069 g  $100g^{-1}$ , with statistically significant higher values recorded at the end of ripening, irrespective of the

- 349 production batch.
- 350
- 351 3.2. Morpho-textural properties of Ciauscolo PGI salami
- 352
- 353 The results of color measurements during ripening of the analyzed sausages are reported in Figure 2.

In more detail, the lightness (L\*) attested between 49.89 and 55.13 for batch 1, and between 54.03 and 62.33 for batch

- 355 2. For each batch, the values for this parameter showed no significant differences during the ripening of *Ciauscolo* PGI
- 356 salami.

- 357 As for redness/greenness (a\*), the detected values were comprised between 9.57 and 12.7 (batch 1), and between 9.87
- and 12.64 (batch 1). In the two batches, the highest  $a^*$  values were observed at  $t_5$ .
- 359 Finally, blueness/yellowness (b\*) values were comprised between 1.44 and 2.69 (batch 1), and 1.64 and 2.75 (batch 2).
- 360 No differences were observed for the b\* values of the samples collected from batch 1, whereas, in samples from batch 2,
- 361 the lowest  $b^*$  values were observed at  $t_{10}$  and  $t_{20}$ .
- 362 The results of texture analysis are reported in Table 2.
- 363 In more detail, values for hardness were comprised between 8.23 and 13.80 N (batch 1), and 7.79 and 13.21 N (batch 2).
- 364 In both the analyzed batches, the highest values for this texture parameter were detected at t<sub>20</sub>.
- 365 As for cohesiveness, the detected values ranged between 0.33 and 0.48 (batch 1), and 0.28 and 0.48 (batch 2). Within the
- 366 same batch, no statistically significant values were observed among samples.
- 367 Finally, springiness ranged between 3.60 and 6.35 mm (batch 1), and 3.10 and 6.25 mm (batch 2). The highest springiness
- 368 values were observed at  $t_0$  for samples form batch 1, and at  $t_{10}$  for samples from batch 2.
- 369
- 370 *3.3. Microbiological analyses*
- 371
- The results of viable counts are reported in Table 3.
- 373 In more detail, the counts of lactic acid bacteria were comprised between 2.87 and 8.95 log cfu g<sup>-1</sup>, with statistically lower
- 374 values recorder at t<sub>0</sub>, irrespective of the production batch. No statistically significant differences of lactic acid bacteria
- 375 counts were highlighted among samples from t<sub>5</sub> to t<sub>20</sub>.
- 376 Regarding coagulase-negative cocci, Enterobacteriaceae, and eumycetes, the counts ranged from 3.45 to 5.94 log cfu g<sup>-1</sup>,
- 377 <1 to 2.87 log cfu g<sup>-1</sup>, and 3.07 to 4.40 log cfu g<sup>-1</sup>, respectively, with the lowest values recorded at t<sub>0</sub>, irrespective of the
- 378 production batch and the microbial group.
- 379 For C. botulinum, the multiplex real-time PCR revealed the absence of the target genes bont/A, bont/B, bont/E, bont/F,
- 380 and *4gyrB (CP)*, encoding botulinic toxins, in all analyzed samples.
- Finally, neither L. *monocytogenes* nor *Salmonella* spp. could be detected in 25 g of product, irrespective of the sampling
  time.
- 383
- 384 *3.4. Metataxonomic analyses*
- 385
- 386 *3.4.1. Microbiota*

387 Microbiota of *Ciauscolo* PGI salami differed significantly (PERMANOVA, P[FDR] < 0.05) if the day of production (to) 388 was compared to the rest of ripening period in both batches (Fig. 3). A certain segregation between batch 1 and 2 was 389 anyway observed in the PCoA plot, especially at to, although with no statistical significance (P[FDR] > 0.05). No 390 differences in the alpha diversity parameters were observed between batches and along time (data not shown).

391 By considering the composition of the microbiota, at to, in both batches, three *Pseudomonas* species s (*P. fragi*, *P.* 

392 psychrophila, P. lundensis) represented the dominant taxa (Fig. 3). From t<sub>5</sub>, L. sakei was dominant in batch 1 until the

- 393 end of ripening (t<sub>20</sub>), whereas in the same timeframe *Dellaglioa algida* represented the most abundant species in batch 2.
- 394 Unassigned species of *Burkholderia-Caballeronia-Paraburkholderia* and *Escherichia-Shigella* groups were significantly
- 395 more abundant in the batch 1 (Wilcoxon's Test, *P*[FDR]<0.001). On the other hand, *Leuconostoc carnosum* was detected
- 396 from t<sub>5</sub> to t<sub>20</sub>, irrespective of the production batch (Supplementary Table 1).
- 397

#### 398 *3.4.2. Mycobiota*

As far as the fungi are concerned, a core mycobiota composed by five species of yeast and one genus of mold that cumulatively represented more than 50% of the relative abundance in all samples was observed (Fig. 4, Supplementary Table 2). In fact, *Cladosporium cladosporioides*, *Debaryomyces hanseni*i, *Kurtzmaniella zeylanoides* and *Malassezia restricta/globosa* were the most abundant yeasts occurring in all samples, followed by less ubiquitous ASVs. Filamentous fungi were mainly represented by *Penicillium* sp., which was found from t<sub>0</sub> until to t<sub>10</sub> in batch 2, whereas, in batch 1, it was not observed at t<sub>0</sub> and t<sub>5</sub>.

- 405 As far as compositional variability of the mycobiota is concerned, PERMANOVA did not highlight any significant effect
  406 of time and batch (data not shown).
- 407
- 408 3.5. Characterization of L. sakei isolates
- 409

410 The BLAST search allowed the unambiguous identification of 44 *L. sakei* isolates to be obtained; the alignment results

411 of the 16S rRNA sequences obtained from these isolates are reported in Table 4.

412 Regarding antimicrobial activity assessed through agar well diffusion assay, no *L. sakei* isolate showed inhibitory activity

413 against *L. innocua* (Table 4).

- 414 As for production of EPS, 21 out of the 44 isolates produced sucrose-dependent EPS, this latter revealed by the formation
- 415 of mucoid colonies on agar plates (Table 4). By contrast, no isolate showed the production of sucrose-independent EPS.

- 416 The result of acidification in the synthetic media assayed are reported in Table 5. In more detail, pH values measured in
- 417 the uninoculated growth media were 6.30, 6.27, 6.24, 6.11, and 6.13 in MRS, MRS + 150 mg Kg<sup>-1</sup> of E250, MRS + 150
- 418 mg Kg<sup>-1</sup> of E252, MRS + 3% NaCl, MRS + 150 mg Kg<sup>-1</sup> of E250, 150 mg Kg<sup>-1</sup> of E252 and 3% NaCl, respectively.
- 419 Regarding acidification in MRS broth, according to ANOVA results (data not shown), isolates C55 and C53 showed the
- 420 highest pH after 24 h (4.9), whereas the isolates C8, C9, C17, C19, C31, C34, C37, and C4 0reached the lowest pH values
- 421 (4.1) after 24 h.
- 422 As for acidification in MRS broth added with 150 mg Kg<sup>-1</sup> of E250, according to ANOVA results (data not shown), the
- 423 isolate L. sakei C55 showed the highest pH after 24 (4.9), whereas the medium inoculated with the isolates C8, C15, C17,
- 424 C21, and C29 reached the lowest pH (4.1) after 24 h.
- 425 Regarding MRS broth added with 150 mg Kg<sup>-1</sup> of E252, according to ANOVA results (data not shown), again the isolates
- 426 C55 and C53 showed the highest pH after 24 h, whereas the medium inoculated with the isolate C37 reached the lowest 427 pH value (3.9) after 24 h.
- 428
  - In MRS broth added with 3% NaCl, according to ANOVA results (data not shown), the isolates C50, C51, C53, C54,
  - 429 C55, C57, C59, C60 showed the highest pH values after 24 h (4.7). By contrast, the growth medium inoculated with the
  - 430 isolates C17, C29, C31, or C37 reached the lowest pH (3.9) after 24 h.
  - 431 As for MRS broth added with 150 mg Kg<sup>-1</sup> of E250, 150 mg Kg<sup>-1</sup> of E252, and 3% NaCl, according to ANOVA results
  - 432 (data not shown), the isolate C2 showed the lowest pH drop after 24 h (5.0), whereas the medium inoculated with the
  - 433 isolate C8 reached the lowest pH (3.9) after 24 h.
  - 434 For all these growth media, according to ANOVA, a wide variability of the results was observed among the isolates (data 435 not shown).
  - 436 The results of the semi-quantitative assessment of selected enzymatic activities are reported in the heat map represented 437 in Figure 5.
  - 438 According to the manufacturer's instructions, only the isolates with color development, scored as 3, 4, or 5 were 439 considered as positive for the tested enzymatic activity.
  - 440 As for alkaline phosphatase, only two isolates (C4 and C15) showed a positive reaction. Thirty-nine out of the 44 L. sakei 441 isolates showed a strong activity for leucine arylamidase, whereas 36 isolates showed a positive reaction for valine 442 arylamidase. Acid phosphatase activity was observed in 27 isolates, whereas only 10 isolates were positive for naphthol-443 AS-BI-phosphohydrolase activity. Six isolates were positive for alpha-galactosidase, whereas 18 showed a positive
  - 444 reaction for beta-galactosidase. Alpha-glucosidase, beta-glucosidase, and N-acetyl-ß-glucosaminidase were observed in
  - 445 3 isolates, being C16, C28, and C39. Finally, no isolate was positive for esterase (C 4), esterase lipase (C 8), lipase (C

446 14), cystine arilamidase, trypsin, alpha-chymotripsin, beta-glucuronidase, alpha-mannosidase, or alpha-fucosidase447 activity.

448

#### 449 3.6. GC-MS analysis of volatile components

450

451 Volatile compounds detected in the static headspace of the two batches of *Ciauscolo* PGI salami samples are reported in
452 Table 6.

453 Monoterpene hydrocarbons (limonene, sabinene,  $\alpha$ -pinene,  $\beta$ -pinene,  $\beta$ -carene,  $\alpha$ -thujene) and sesquiterpene 454 hydrocarbons ( $\beta$ - and  $\alpha$ -copaene) dominated the aroma profiles of samples, throughout the whole ripening time. Allyl 455 methyl sulphide and diallyl disulphide were the major aliphatic sulphur compounds, but lower amounts of diallyl sulphide 456 and allyl methyl disulphide were also detected. Samples from batch 1 were characterized by higher levels of monoterpene 457 hydrocarbons and alcohols (n-hexanol and n-pentanol, mainly) and lower levels of sulphur compounds than those 458 belonging to batch 2. Different dynamics were observed in the aroma composition during the ripening. In more detail, 459 increasing levels of monoterpene hydrocarbons (sabinene,  $\alpha$ -pinene, camphene) and 2-methyl-1-butanol, and decreasing 460 amounts of 1-pentanol, ethyl hexanoate, and octanal were observed in samples from batch 1; whereas samples from batch 461 2 were characterized by very limited changes in the aroma composition during ripening, where only an increase of ethyl 462 ester levels (ethyl isopentanoate) and a decrease of 1-pentanol were observed.

463 Regarding the correlation between bacteria-fungi and VOCs, we observed patterns of positive and negative correlations

464 (*P*-value [FDR adjusted] <0.05) mainly influenced by the batch and then by the sampling time (Fig\_X). L. sakei was

- 465 negatively correlated with cis-sabinene hydrate and positively with to ethyl isopentanoate (ester) and with 3-methyl-1-
- 466 butanol together with *L. curvatus* while *Dellaglioa algida*, with allyl-methyl disulfide and α-thujene. L. plantarum showed
- 467 positive relationship with ethyl pentanoate and 1-pentanol.

468 Regarding fungi we observed a direct relationship between *Debaryomyces hansenii* and *Kurtzmaniella zeylanoides* with

- 469 β-caryophillene and 2-methyl-1-butanol (alcohol), respectively.
- 470 [modify in relation to the discussion]
- 471
- 472 **4. Discussion**
- 473

The microbiota of fermented sausages has widely been investigated along time; however, traditional fermented sausages still represent a niche of undisclosed microbial diversity and a source of pro-technological microorganisms that could successfully be applied by the industry of fermented meat for product improvement. In the present study, the morpho477 textural, physico-chemical (including volatile compounds), and microbiological characteristics of *Ciauscolo* PGI salami
478 during ripening were investigated. *L. sakei* isolates collected from the analyzed samples were also tested for some pro479 technological activities to select potential cultures to be used as starter or adjuncts.

480 As for pH values detected in the analyzed samples, the data were in accordance with those already detected by Belleggia 481 et al. (2020a) and Trani et al. (2010) in Ciauscolo salami produced by other manufacturers. In the analyzed samples, a 482 progressive decrease in pH was observed during time, irrespective of the production batch. Of note, since Ciauscolo is 483 ready for consumption only after 20 days, the pH values herein detected in the end products reflect the short ripening 484 time. According to the production disciplinary, the pH of ready-to-eat Ciauscolo PGI salami should be > 4.8. In 485 accordance with pH reduction, an increase in TTA values as well as in lactic and acetic acid content was also observed in 486 the analyzed samples. The trends of the abovementioned parameters can likely be explained by the activity of the lactic 487 acid bacteria naturally occurring in the raw material which produced organic acids through fermentative metabolism of 488 meat carbohydrates (Fadda, López, & Vignolo). Of note, the presence of both lactic and acetic acid in the analyzed 489 samples attests the presence of homofermentative as well as heterofermentative lactic acid bacteria.

490 In the analyzed samples, a reduction of a<sub>w</sub> was observed during time, thus attesting the progressive drying of the product 491 with further improvement of the safety and stability of *Ciauscolo* PGI salami. The detected values were in accordance 492 with those detected by Trani et al. (2010) and Belleggia et al. (2020a) in the same type of fermented sausage.

Interestingly, pH and a<sub>w</sub> values herein detected were higher than those detected by Aquilanti et al. (2007) in the same type
of fermented sausage, thus suggesting a great variation of these parameters based on the artisan method applied during
production.

496 Based on Commission Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs, ready-to-eat products 497 with pH > 4.4 or  $a_w > 0.92$  or products with pH > 5.0 and  $a_w > 0.94$  are prone to support the growth of *Listeria* 498 *monocytogenes*, hence, such issue should be taken into consideration when producing *Ciauscolo* salami, whose pH and 499  $a_w$  values can sometimes be permissive for the growth of the pathogen.

500 For color measurements, it is noteworthy that Ciauscolo PGI salami is a short-time ripened fermented sausage, hence, the 501 color of the end product (at t<sub>20</sub>) may not substantially differ from that of the stuffed meat batter at t<sub>0</sub>. To the authors 502 knowledge, no data reporting the color of Ciauscolo PGI salami are available in the scientific literature for further 503 comparison of results. Regarding lightness, this parameter can vary from 0 (black) to 100 (white) (Méndez-Cid et al., 504 2017); hence, the high level of lightness detected in the samples herein analyzed can likely be attributed to the high 505 amount of fat that characterizes the meat batter of this fermented sausage (Estévezet al., 2005). As for a\* parameter, this 506 axis represents the green-red opponent colors, with values < 0 toward green and values > 0 toward red (Méndez-Cid et 507 al., 2017). In fermented sausages, the intensity of the red color is enhanced by the presence of nitrates and nitrites as well

508 as by the microbial activity of pro-technological microorganisms (e.g., lactic acid bacteria and coagulase-negative cocci) 509 (Holck et al., 2017). In the samples herein analyzed, the a\* values were in the range of the red color, with slight variations 510 detected only at  $t_5$ . Regarding the b\* parameter, this axis denotes the blue–vellow opponents, with values < 0 toward blue 511 and those > 0 toward yellow. In the present study, samples showed values of yellow around 1-2, with only slight variations 512 occurring at t<sub>10</sub> and t<sub>20</sub> in the samples of batch 2. In general, an increase of b\* value in fermented sausages could denote 513 fat rancidity due to lipid oxidation (Méndez-Cid et al., 2017), hence, the low values detected in the samples herein 514 analyzed represent a positive quality characteristic of this high-containing lard salami and suggest a low level of lipid 515 oxidation.

516 Regarding texture, as expected, the longer the ripening time, the higher the measured hardness. This evidence was also 517 supported by the lower  $a_w$  values measured in the samples a  $t_{20}$  in respect with those at  $t_0$ , thus highlighting the achievement 518 of the required drying of the product. As for the cohesiveness, in fermented sausages, this parameter is affected by the 519 solubility of muscle proteins (Farouk et al., 2002); moreover, as reported by Bis-Souza et al. (2020), the amount of fat 520 strongly influences this parameter. In more detail, Bis-Souza et al. (2020) observed a progressive increase of cohesiveness 521 as well as of springiness in salami produced with a progressive reduction of pork fat. Interestingly, the values of 522 cohesiveness measured in the samples herein analyzed were as low as those reported by Bis-Souza et al. (2020) for Italian 523 salami containing 20% (w/w) of pork fat, attesting at about 0.5.

524 Viable counts carried out on Ciauscolo PGI salami samples highlighted a progressive increase of pro-technological
 525 microbial groups as well as a decrease of indicators of enteric contamination.

526 As it has previously been elucidated, raw meat can be contaminated by pathogenic or spoilage microorganisms during 527 animal slaughtering (Taylor & Aiyegoro, 2022). In more detail, the Enterobacteriaceae family includes important 528 foodborne pathogens such as Salmonella spp., Yersinia enterocolitica, Escherichia coli (including the pathogenic species 529 E. coli O157:H7), Shigella spp., and Cronobacter spp. Enterobactericeae, together with coliforms, are often assessed in 530 foods to highlight poor hygiene or inadequate processing, process failure, and post-process contamination, being therefore 531 considered as indicator microorganisms (Petruzzelli et al., 2016). In the present study, the two analyzed batches were 532 characterized by a constant reduction in Enterobacteriaceae counts that corresponded to a notable increase in pro-533 technological microorganisms as lactic acid bacteria and coagulase-negative cocci. Of note, for Enterobacteriaceae, pH 534 tolerance is often influenced by the acidulant they are exposed to, with lactic acid being more inhibitory than mineral 535 acids. This feature explains the progressive reduction of Enterobacteriaceae in the analyzed samples at  $t_{20}$ , where the 536 amount of lactic acid was maximum.

Lactic acid bacteria represent a key microbial group in meat fermentation. Their metabolic activity improves the safety
and sensory quality of fermented sausages (Fadda et al., 2010). Indeed, lactic acid and other organic acids (e.g., acetic

539 acid) produced by lactic acid bacteria prevent the development of spoilage and pathogenic microorganisms. Of note, some 540 strains of lactic acid bacteria can act as protective cultures in fermented meat products through the production of 541 bacteriocins. These compounds are proteins with potential antimicrobial activity against specific pathogens (e.g., Listeria 542 monocytogenes) (Junges da Costa et al., 2021). Moreover, lactic acid bacteria metabolism leads to the stabilization of 543 meat color and to texture development (e.g., formation of a distinctive gel-like texture due to protein denaturation () 544 (Fadda et al., 2010). Lactic acid bacteria are also responsible for production of volatile compounds through the release of 545 free amino acids (Fadda et al., 2010). In the present study, the counts of lactic acid bacteria were consistent with those 546 already detected by Belleggia et al. (2020a) and Aquilanti et al. (2007) in the same salami at 20 days of ripening, attesting 547 at about 8 log cfu g<sup>-1</sup>.

548 In fermented meat, coagulase-negative cocci enhance the aroma and texture of the product. In more detail, this microbial 549 group encompasses microorganisms that produce nitrate reductase, this latter enzyme converting nitrate into nitrite 550 (Khusro, & Aarti, 2022). Nitrites exert an antimicrobial activity against clostridia, especially C. botulinum, with a 551 consequent reduction of food poisoning risk; they also exert an antioxidative activity and promote the formation of 552 nitrosomyoglobin with the subsequent development and stabilization of a pleasant red color (Khusro, & Aarti, 2022). 553 Furthermore, coagulase-negative cocci improve flavor and aroma of fermented sausages through the formation of volatile 554 compounds deriving from proteolysis and lipolysis (Khusro, & Aarti, 2022). In the present study, an increase in coagulase-555 negative cocci counts was observed during time, with final counts that were similar to those already detected by Aquilanti 556 et al. (2007) in Ciauscolo PGI at 20 days of ripening, but higher than those recently detected by Belleggia et al. (2020a) 557 in the same salami, thus suggesting the occurrence of a certain variability among producers based on production process 558 and the raw materials used.

As for eumycetes (yeasts and molds), their role in fermented sausages is mainly related to the production of proteolytic (exoproteases) and lipolytic (exolipases) enzymes that are responsible for the development of flavor and aroma (Copetti, 2019; Sunesen & Stahnke, 2003). Of note, yeasts are common environmental contaminants of fermented sausages and are well adapted to this high-salt and acidic environment (Osei Abunyewa, Laing, Hugo, & Viljoen, 2000). As reported by Cocolin et al. (2011), molds occurring on the surface of fermented sausages facilitate the dehydration process, since they create micro-pores on the casing. Finally, external mold layer protects lipids from oxidation in the presence of light (Cocolin et al., 2011).

566 Of note, the absence of the major pathogens as *L. monocytogenes*, *Salmonella* spp., and of the target genes *bont/A*, *bont/B*,

567 *bont/E, bont/F,* and *4gyrB (CP)*, encoding botulinic toxins, in all the samples herein analyzed attests the high quality of

the raw materials and the proper application of good manufacturing practices.

569 Metataxonomic analysis highlighted differences in the microbiota of the analyzed samples, including stuffed meat batter,

570 fermented sausages sampled during ripening, and fermented sausages sampled at the end of maturation; in more detail, a 571 progressive substitution of meat spoilage bacteria of with pro-technological microorganisms was seen.

572 Pseudomonas, that encompasses psychotropic bacteria, constituted the core microbiota of stuffed salami at to. P. fragi is 573 a spoilage microorganism that produces slime on meat during storage and it is responsible for off-odors and soften of the 574 flesh (Shao et al. 2021). P. psychrophila is a facultatively psychrophilic bacterium that can be responsible for meat 575 spoilage (Yumoto et al., 2001). Regarding *P. lundensis*, this is a further spoilage species that has already been isolated 576 from beef and lamb fresh meat, where it was responsible for the production of off-odors (Shao et al. 2021). As reported 577 by Fidan et al. (2022), the presence of organic acids (as lactic or acetic acid) can inhibit putrefactive bacteria, including 578 those belonging to the genus *Pseudomonas*, thus explaining the reduction of such microorganisms as the fermentation of 579 the sausages herein analyzed has progressed.

580 As for the most represented pro-technological species, L. sakei and L. carnosum dominated in salami of batch 1 after 5 581 days of ripening, whereas the same two species, together with D. algida, were mostly represented in samples of batch 2. 582 L. sakei represents the emblematic lactic acid bacteria species in fermented meat products for its rapid acid production 583 (Amadoro, Rossi, Piccirilli & Colavita, 2015; Zagorec & Champomier-Vergès, 2017). Indeed, this psychrotrophic and 584 facultative anaerobic species has already been recognized as the key lactic acid bacterium in fermented sausages produced 585 in Western Europe, as well as in fermented meat products manufactured in Asia and South America (Zagorec & 586 Champomier-Vergès, 2017). In meat or fermented sausages, L. sakei takes advantage of meat nutrients by encoding 587 oligopeptide transporters and intracellular peptidases through genes up-regulation. The endo and exo-peptidases produced 588 by L. sakei increase the concentration of free amino acids, thus positively affecting flavor development (Belleggia et al., 589 2022b). Moreover, L. sakei can use the ribose present in raw meat as carbon source through an ATP-dependent system 590 (Zagorec & Champomier-Vergès, 2017). Based on the abovementioned features, in fermented sausages, L. sakei shows a 591 shorter lag phase and a growth rate higher than other lactic acid bacteria; moreover, it is highly salt tolerant (up to 6.5% 592 NaCl) (Amadoro et al., 2015).

As reviewed by Honrada Perez, Zendo, & Sonomoto (2022), *L. sakei* can produce multiple bacteriocins, namely sakacin P, sakacin T, and sakacin X, depending on the strain. At this regard, none of the 44 *L. sakei* cultures herein isolated showed an antimicrobial activity against *Listeria innocua*. The production of bacteriocins represents a highly desirable feature due to the competitive advantage obtained by the producing strain and for the increased safety of the product. Of note, the genes encoding for active bacteriocins are frequently located in operon clusters, harbored in the genome, plasmid, or in other mobile genetic elements; moreover, the expression of these operons is complex and can be induced by the presence of auto-inducer peptides (Kumariya et al., 2019). The abovementioned features might explain the lack of bacteriocins production in the analyzed *L. sakei* isolates and suggest that bacteriocin production by *L. sakei* is less
common than expected as already reported by Belleggia et al. (2022c), who observed no listericidal effect of 22 *L. sakei*cultures isolated from fermented fish sausages.

As reported by Wang, Song, Zhao, Han, & Zhou (2019), strains of *L. sakei* also showed a high-yield EPS production with unique characteristics for their exploitation in large-scale industrial food applications. EPS produced in fermented sausages by lactic acid bacteria can impact on the morpho-textural properties of the end product (Hilbig et al., 2019). At this regard, none of the 44 *L. sakei* cultures herein assayed showed the production of sucrose-independent EPS, whereas 21 isolates showed the formation of mucoid colonies in the presence of sucrose, thus suggesting a good *in vitro* synthesis of EPS to be further investigated.

Regarding the acid production capacity, and the capability to cope with the main environmental stresses encountered during the fermentation process (pH, NaCl, sodium nitrite, and potassium nitrate), the good performances of the tested *L. sakei* isolates inoculated in synthetic media were in accordance with those reported by Wen et al. (2021) for the same lactic acid bacteria species, thus suggesting its potential application in salami production for the formulation of starter or adjunct cultures.

In is noteworthy that the acidification occurring in fermented sausages affects proteolysis and thus flavor formation. At this regard, fermented sausages produced in Northern European countries are characterized by a pH drop below 5.0 from fermentation throughout ripening, whereas in those produced in Southern European countries a moderate drop in pH is usually observed during fermentation, with final pH values comprised between 5.5 and 6.0 (Berardo et al., 2017). Hence, almost all the isolates herein assayed could serve as suitable acidifiers in fermented meat products.

Among the enzymatic activities observed in the selected pool of isolates ascribed to *L. sakei*, the most relevant were leucine arylamidase and valine arylamidase activities showed by most of the isolates. The presence of these two aminopeptidases has already been reported in *L. sakei* strains. Of note, these enzymatic activities contribute to the hydrolysis of the sarcoplasmic proteins with the formation of free amino acids, these latter representing precursors of favor compounds in fermented sausages (Wang et al., 2013). The results on the presence of aminopeptidases obtained in the present study are in accordance with those reported by Ammor et al. (2005) that observed a strong leucine and valine arylamidase but no cystine arylamidase activity, in *L. sakei* isolated from traditional French dry sausages.

626 In the present study, 27 isolates showed a high acid phosphatase activity. Of note, this hydrolase liberates phosphate ions

627 from organic esters at pH values ranging from 4.5 to 6.0. These results are in accordance with those reported by

628 Papamanoli et al. (2003) that observed a high acid phosphatase activity in *L. sakei* cultures isolated from a Greek dry-

629 fermented sausage.

630 As a positive trait, the absence of beta-glucuronidase in all the tested *L. sakei* isolates deserves attention. Indeed, the 631 activity of this enzyme leads to the production of carcinogens and mutagens in the colon of the consumer, thus also 632 increasing the probability of tumor induction (Li et al., 2023).

633 In the analyzed *Ciauscolo* PGI salami, *D. algida* and *L. carnosum* were also detected among the major pro-technolgical
634 taxa.

*D. algida* (basonym *Lactobacillus algidus*) has already been detected by Belleggia et al. (2020a) in *Ciauscolo* PGI salami.
The occurrence of *L. algidus* has also been reported by Greppi et al. (2015) in traditional Piedmontese sausages. This
lactic acid bacterium represents a fastidious microorganism in meat since it can be the causative agent of spoilage with
production of biogenic amines (Säde et al., 2020). Raw meat contaminated be *L. algidus* has a higher sour smell (Schirmer,
Heir, & Langsrud, 2009); notwithstanding, when a sour smell is produced in fermented sausages, this could not necessarily
represent a negative trait.

As for *L. carnosum*, the occurrence of this species has already been reported by Belleggia et al. (2020a) in *Ciauscolo* PGI salami. As reported by many authors, *L. carnosum* can produce anti-listerial bacteriocin (leucocin) with a broad spectrum of activity (Hornbæk et al., et al., 2004; Osmanağaoğlu, 2007; Woraprayote et al., 2021), thus suggesting the use of selected *L. carnosum* isolates as protective adjunct culture in *Ciauscolo* manufacturing.

645 A few eumycetes species characterized the mycobiota of the *Ciauscolo* PGI salami herein analyzed.

646 The presence of *C. cladosporioides* in *Ciauscolo* PGI salami has already been reported by Belleggia et al. (2020a) and by 647 Vila et al. (2019) among the minor fraction of filamentous fungi isolated from dry-cured sausages produced in Argentina. 648 This species of eumycetes, usually occurring on the surface of fermented sausages, can originate from contaminated air 649 and raw materials (Parussolo et al., 2019). As reported by Lozano-Ojalvo (2015), *Cladosporium* species are generally not 650 able to heavily colonize dry-cured meat products, although some xerotolerant and halotolerant strains can produce black 651 spots on the surface of the sausages.

652 D. hansenii is one of the most widely detected yeast species in fermented sausages, and it contributes to the development 653 of the flavour of fermented meat products through the production of endo- and exo-peptidases (Flores & Toldrá, 2011; 654 Medina-Córdova et al., 2018). Indeed, the enzymatic activities carried out by D. hansenii enrich the meat matrix in free 655 amino acids and peptides that characterize the aroma of the end product (Flores & Toldrá, 2011). Furthermore, the enzyme 656 glutaminase produced by *D. hansenii* neutralizes the acidic pH of fermented sausages and generates L-glutamate that can 657 act as flavor enhancer (Flores & Toldrá, 2011). Interestingly, in the last decade, D. hansenii attracted the attention of the 658 fermented meat industry as bioprotective agents against toxigenic penicillia (Núñez et al., 2015). 659 Regarding K. zeylanoides (formerly known to as Candida zeylanoides), the presence of this yeast species has already

been reported by Belleggia et al. (2020a) in *Ciauscolo* PGI salami and in fermented sausages produced in Italy (Giarratana

et al., 2014) and other Southern European countries (Belleggia et al. 2020b; Belleggia et al., 2022a; Belleggia et al.,
2022b; Encinas, López-Diáz, Garciá-López, Otero, & Moreno, 2000).

The SPME-GC/MS analysis performed during ripening of *Ciauscolo* PGI salami allowed the composition of the major and minor volatile components to be disclosed. In more detail, the presence of spices-derived components (mono- and sequiterpene hydrocarbons, sulphides and disulphides), detected throughout the whole ripening time of samples was in accordance with the results reported by other authors for different kind of Italian salami (Bianchi et al., 2007; Jerković et al., 2010; Moretti at al., 2004). Particularly, the presence of the major aliphatic sulphur compounds (allyl methyl sulphide and diallyl disulphide) detected in the analyzed samples can likely been explained by allicin decomposition, that was recognized as main character in fresh garlic smell.

Hexanal was the most abundant lipid oxidation marker, thus confirming the results already observed in other Italian salami
(Bianchi et al., 2007; Moretti et al., 2004). Interestingly, a marked decrease of hexanal in batch 1 samples came with an
increase of hexanol, probably due to the reducing conditions inside the product.

Branched chain alcohols (3-methyl-1-butanol, 2-methyl-1-butanol) were the most represented markers of microbial activity, together with ethyl esters of of isopentanoic, hexanoic (the most abundant), and octanoic acids (Chaves-López et al., 2011; Janssens et al., 2012) contributing to fruity notes of salami aromas. In fact, the esterifying activities of several yeasts, molds, and bacteria usually present in cured meats have previously been reported (Bianchi et at., 2007; Lorenzo et al., 2013).

Only a limited number of aroma components herein detected could be related to microbial activity; among these, branched chain alcohols originated from amino acid catabolism (3-methyl-1-butanol, 2-methyl-1-butanol) were the most represented (Chaves-López et al., 2011; Janssens et al., 2012). Ethyl esters of isopentanoic, hexanoic (the most abundant), and octanoic acids were recognized as typical markers of microbial activity, as well. According to Bianchi et at. (2007) and Lorenzo et al. (2013), several yeasts, molds, and bacteria that are usually present in cured meat could contribute to fruity notes of salami aromas through their esterifying activities.

684 Interestingly, a marked decrease of the main lipid oxidation product (hexanal) came with an increase of hexanol, probably
685 due to the reducing conditions inside the product.

686

#### 687 5. Conclusions

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689 The results of the present study provided an advancement of knowledge on the morpho-textural, microbiological, and 690 volatile features of *Ciauscolo* PGI salami. As for colour and texture characteristics, for the first time this research provides 691 objective parameters to be used as product quality indices. Regarding microbiota characterization, the dominance of *L*. 692 sakei in the analyzed sausages was once again confirmed, together with other co-occurring species, namely Dellaglioa 693 algida and Leuconostoc carnosum. The pool of L. sakei cultures isolated from the analyzed samples showed a suitable 694 acid production capacity, together with the capability of coping with the main environmental stresses occurring in 695 fermented sausages. For most of the isolates, a strong aminopeptidase activity (due to leucine arylamidase and valine 696 arylamidase) was observed. Moreover, most isolates showed the in vitro production of sucrose-dependent EPS. 697 Interestingly, no isolate were positive for beta-glucuronidase activity. Based on these findings, a few candidates as starter 698 or adjuncts cultures, with potential use for product safety and quality improvement, were found, being the isolates L. sakei 699 C5, C7, C11, C31, C45, C48, C53, C55, and C60. Further research is needed to assess their supposed pro-technological 700 traits in trials for salami manufacturing. The absence of pathogenic microorganisms attests the high quality of the raw 701 materials and the production process, thus confirming Ciauscolo PGI salami as an Italian food excellence. To the authors' 702 knowledge only one available study has previously investigated the volatilome of Ciauscolo PGI salami; hence, the 703 similarity of the volatile pattern detected in the present study with that already published allows a kind of volatile 704 fingerprint of the Ciauscolo PGI salami to be drawn.

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### 706 CRediT authorship contribution statement

707

Andrea Osimani: Conceptualization, Writing - Review & Editing, Supervision, Resources. Luca Belleggia:
Investigation, Formal analysis. Ilario Ferrocino: Investigation, Formal analysis, Writing - Original Draft, Resources.
Cristian Botta: Investigation, Formal analysis. Vesna Milanović: Formal analysis. Federica Cardinali: Investigation,
Formal analysis, Writing - Original Draft. M. Naceur Haouet: Investigation. Cristiana Garofalo: Formal analysis.
Massimo Mozzon: Investigation, Formal analysis, Resources. Roberta Foligni: Investigation, Formal analysis. Lucia
Aquilanti: Review & Editing, Resources.

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## 715 Declaration of Competing Interest

716

The authors declare that they have no known competing financial interests or personal relationships that could haveappeared to influence the work reported in this paper.

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- 728
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- 730
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- 732
- 733 References
- 734
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982	FIGURE CAPTIONS
983	
984	Fig. 1. Slice of ready-to-eat Ciauscolo PGI salami.
985	
986	Fig. 2. Images of cross sections of Ciauscolo PGI salami during drying.
987	
988	to, day of production; t5, samples after 5 days of ripening; t10, samples after 10 days of ripening; t20, samples after 20 days
989	of ripening.
990	Means $\pm$ standard deviations of triplicate independent measurements are shown.
991	Within each batch, for the same color parameter, means followed by different letters are significantly different ( $P < 0.05$ ).
992	L* value describes the lightness; a* value describes the redness/greenness; b* describes the blueness/yellowness.
993	
994	Fig. 3. Microbiota composition.
995	
996	PCoA charts (on the left) displaying weighted UniFrac distance matrix ( $\beta$ -diversity). Batches and sampling time are
997	defined by different colours and shapes (legend); dashed ellipses are indicating significant different communities and
998	results of PERMANOVA are reported in the graph. Stacked bar plots (on the right) showing core microbiota composition
999	(relative abundance) at the Species/Genus rank level and relative colour coding key.
000	Samples are grouped by batch and displayed according to time; taxa are sorted in the legend from the most to the least
001	abundant (> 0.2 % of average abundance).
002	to, day of production; t5, samples after 5 days of ripening; t10, samples after 10 days of ripening; t20, samples after 20 days
003	of ripening.
004	
005	Fig. 4. Mycobiota composition.
006	
007	Stacked bar plots showing core mycobiota composition (relative abundance) at the Species/Genus rank level and relative
008	colour coding key.
009	Samples are grouped by batch and displayed according to time; taxa are sorted in the legend from the most to the least
010	abundant (> $0.2$ % of average abundance).
011	to, day of production; t5, samples after 5 days of ripening; t10, samples after 10 days of ripening; t20, samples after 20 days
012	of ripening.
	24

013	
014	Fig. 5. Heat map representing the results of semi-quantitative assessment of enzymatic activities of Latilactobacillus sakei
015	isolated (C1-C60) from Ciauscolo PGI salami.
016	
017	For each enzymatic reaction, a value ranging from 0 to 5 was assigned, corresponding to the colors developed: 0
018	corresponds to a negative reaction (blue dots), 5 to a reaction of maximum intensity (dark red dots), and values 1, 2, 3
019	or 4 are intermediate reactions depending on the level of intensity (3, 4, or 5 being considered as positive reactions).
020	
021	
022	Fig X. Correlation between metataxonomic and volatolomic data.
023	Tile plots showing the existing pairwise correlations between VOCs and bacterial-fungal ASVs (merged at the
024	species/genus level). Taxa are ordered from the more to the less abundant, whereas the VOCs are grouped in relation to
025	their chemical class (refer to the caption for the codes). Colours represents level of Spearman's Rho correlation (from -
026	1 to 1; caption) and significant positive and negative correlations are highlighted with asterisks (P-value [FDR
027	adjusted]: $* = <0.05$ ; $** = <0.01$ ; $*** = <0.001$ ).