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Characterization of the microbial community in ripened Pecorino Toscano cheese affected by pink discoloration

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CHARACTERIZATION OF THE MICROBIAL COMMUNITY IN RIPENED PECORINO TOSCANO CHEESE AFFECTED BY PINK DISCOLORATION

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23 Abstract 2

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24 Pink discoloration defect can cause economic losses for cheese producers due to the impossibility to 4 $\frac{6}{2}$ 5 sell the defected cheese, but few knowledge is currently available on the causes of this defect. To 26 gain more insight on the causes that lead to the formation of pink discoloration in Pecorino Toscano 9 $\frac{11}{12}$ 7 cheese with the Protected Designation of Origin (PDO) status, the bacterial community in defected 28 and not defected cheese was characterized by high-throughput sequencing of bacterial 16S rRNA 14 $\frac{16}{12}$ 9 gene. The bacterial community in the defected cheese significantly differed compared to the 30 control. The relative abundance of the genera Acidipropionibacterium, Enterococcus, $^{21}_{231}$ Escherichia/Shigella, Lactobacillus, Lentilactobacillus and Propionibacterium was higher in the 23
2 $\frac{2}{3}$ cheese with pink discoloration defect. The concentration of short chain fatty acids and of lactic acid $5 - 5$ \approx 725 Seil the defected cheese, but lew kn 8 and 2010 12^{\prime} check with the Flotence Designa 13 15 17^3 gene. The bacterial community in 18 (a) $\frac{1}{2}$ (b) $\frac{1}{2}$ (c) $\frac{1}{2}$ (c) $\frac{1}{2}$ (c) $\frac{1}{2}$ (c) $\frac{1}{2}$ 1930 control. The relative abundance of 20 $22²$ 242 cneese with pink discoloration def

 $2\,$ §3 in cheese was measured and a shift towards the production of propionate in the cheese with pink $^{28}_{29}$ discoloration defect was observed. Furthermore, the possible involvement of microbially produced 3135 vitamin B_{12} in the formation of pink discoloration was not supported by the data, since a tendency $33/36$ to a lower concentration of vitamin B₁₂ was measured in the defected cheese compared to the 37 control. 36 25 and 26 an 29^4 alsoloration defect was observed 34 ³ to a lower concentration of vitaming

 $^{40}_{4}$ 39 Keywords: microbiota; pink discoloration; propionic bacteria; sheep cheese; vitamin B₁₂ 4¹39 **Keywords:** microbiota; pink disco

$\frac{45}{45}$ d 1. Introduction 4641 1. Introduction

42 Sheep cheese includes several varieties of dairy products widely produced in Italy. Several of them, 48 $^{50}_{5,43}$ as Pecorino Toscano, have the Protected Designation of Origin (PDO, EC regulation 306/2010) 44 status (Buccioni et al., 2012). Pecorino Toscano PDO cheese is ripened from a minimum of 20 days 53 $^{55}_{64}$ 5 (for fresh cheese) to over 4 months (for ripened cheese). The ageing is carried out in conditioned 46 chambers in which temperature and humidity favorite the proliferation of the bacteria responsible $^{60}_{64}$ 7 for aromas. Sometime, undesirable fermentation, due to a microbial contamination, may occur. Pink 49 51^{43} as Fecorino Toscano, have the T 52 54 56 (for from encese) to over Thront 57 5846 chambers in which temperature a 59 61

48 discoloration (PD) of the loaf rind or of the cheese inner part is a very common defect. Pink 49 discoloration leads to economic losses for the producers due to the impossibility to sell the defected 2 $\frac{4}{5}$ 0 cheese (Daly et al., 2012). Despite the negative consequences potentially linked to this defect, few 31 knowledge is available on the causes. 1 3 50 cneese (Dary et al., 2012). Despite $6\overline{6}$

 $\frac{9}{10}$ In cheese with added colorants (e.g., Cheddar) the PD defect has been often associated to the 53 degradation of the colorant itself (Daly et al., 2012). In cheese without colorant the defect has been 12 $\frac{1}{2}$ 54 associated to Maillard browning or to the microbial activity, such as the activity of some lactobacilli 55 or propionic acid bacteria used as starter cultures (Daly et al., 2012). 10^{22} in cheese with added colorants 11 13 15 associated to maintain crowning σ 16 155 or propionic acid bacteria used as

 $^{1.5}_{2.56}$ Thermus thermophilus has been identified as a possible responsible of PD defect in Continental-21 type cheese (Quigley et al., 2016). T. thermophilus is a Gram-negative, extremely thermophilic, ²⁴58 aerobic, nonpathogenic microorganism, able to produce carotenoids (Tian and Hua, 2010), whose $^{26}_{25}$ 9 occurrence can lead to the presence of a PD in cheese (Quigley et al., 2016). In a recent work, the 2 60 formation of PD close the rind of Pecorino Toscano cheese, has been attributed to the presence of $^{31}_{2}$ 51 Serratia liquefaciens, a psychrotrophic and motile organism (Martelli et al., 2020). The authors 62 have hypothesized that the presence of S. liquefaciens on Pecorino Toscano cheese with PD defect $\frac{3}{2}$ was due to an environmental contamination (Martelli et al., 2020). Despite the papers reported 38
3\$64 above, which attempted to understand the mechanisms that lead to the PD defect in cheese, several $^{4.1}$ 65 aspects of this phenomenon remain unclear. Further understanding of the factors involved in the $^{43}_{44}$ 66 development of PD in cheese can give the opportunity to develop strategies to avoid the formation ⁴ 67 of this defect, with potential economic benefits for the producers. 20 22 ² type cneese (Quigley et al., 2016) 25 279 occurrence can read to the present 28 30 $32²$ berrand hyperacters, a psychion 33 3462 have hypothesized that the presen 35 37° \cdots \cdots 3964 above, which attempted to unders 40 44 pb development of PD in cheese can 45

 $^{48}_{40}$ 68 The aim of this study was to describe the microbial community associated to PD defect in Pecorino 5169 Toscano PDO cheese and to provide new hypotheses on the possible role of the bacterial $\frac{53}{6}$ 70 communities on the development of PD defect. 490 THC and Of this study was to desc. 50 52 54° communities on the development

582 2. Materials and methods

 60 73 2.1. Samples

74 Cheese making was realized in an industrial dairy processing plant located in Tuscany (Caseificio 75 Sociale Manciano, Manciano, Grosseto, Italy) and it was carried out in accordance with the 2 $\frac{4}{57}$ 6 Protected Designation of Origin (PDO) disciplinary of Pecorino Toscano cheese: raw milk was 77 pasteurized and inoculated with milk cultures Streptococcus salivarius subsp. thermophilus, $\frac{9}{10}$ 78 Lactobacillus delbrueckii subsp. bulgaricus, Lactococcus lactis subsp. lactis, Lactococcus lactis 1279 subsp. *cremoris*. Veal rennet was added to the milk and the coagulation occurred at a temperature of $\frac{1}{2}$ 80 35 °C within 30 minutes. The curds were broken at the size of a corn kernel. After, the curd was 81 placed into 2 kg molds for the whey drain by manual pressing, located in a thermostatic chamber at $\frac{1}{2}$ 82 35 °C for 2 h and then plunged in a salt solution (NaCl, 19% w/v) at 12 °C for 24 h. Cheese was 21

283 ripened for 270 days at 8-10 °C. 1 $\overline{3}$ 5 σ Protected Designation of Origin ($6\overline{6}$ $8 \tcdot 1$ 10^{10} Laciobacinus aeidencent subsp. 11 13 15 16 1781 placed into 2 kg molds for the wh 18 $20⁻$ and $20⁻$ 2283 ripened for 270 days at 8-10 °C.

²⁴84 Cheese samples were collected from two lots of Pecorino cheese: the first lot was composed of non-26
285 defected cheese (4 samples were collected from 2 cheese units) and was used as control; the second 2 \ 86 lot was composed of non-defected cheeses (4 samples were collected from 2 cheese units), used as $3\frac{1}{2}$ 87 control, and defected cheese (8 samples were collected from 2 cheese units). Since the defected 88 cheeses showed a PD shading from one flat side to the other, the slices were cut in half, the two $\frac{369}{10}$ samples were processed independently and considered in the defected group (Figure S1). The 8 90 samples collected from the non-defected cheese were considered as the control group (4 cheese 38 ⁴¹91 units, 2 samples for each cheese units) while the 8 samples collected from the cheese with PD (2 $^{43}_{4}$, 49 2 cheese units, 4 samples each) were considered as the defected group. 25 and $\frac{1}{2}$ and $\frac{1$ 2^{3} defected cheese (4 samples were d 28 30 $32²$ control, and defected encese (b) 33 3488 cheeses showed a PD shading from 35 37³ compress the processes margins 390 samples collected from the non-40 44 2 cneese units, 4 samples each) were

$^{48}_{4.9}$ 94 2.2. DNA extraction and 16S rRNA gene sequencing 49^4 2.2. DIVA extraction and TOS TIME

5195 Around 25 g of each cheese sample were homogenized in a mixer and fat was removed. DNA was $^{5.3}_{5.9}$ extracted from 200 mg of each homogenized sample with the DNeasy *mericon* Food Kit (Qiagen) 5 θ 7 according to the manufacturer's instructions.
57
5 $\frac{6}{3}$ 8 The V5-V6 hypervariable regions of the 16S rRNA gene were PCR-amplified using the 783F and 52 54° cannot hom zoo mg of each hy 55 5 β 7 according to the manufacturer's in

 $^{60}_{61}$ 99 1046R primers (Huber et al., 2007; Wang and Qian, 2009) as previously described (Daghio et al., 6199 1046K primers (Huber et al., 200

100 2018). Briefly, the PCR was performed in 2×50 µL reactions with GoTaq®Green Master Mix 101 (Promega Corporation, Madison, WI, USA) and 1 µM of each primer. Amplification conditions 2 10^4 were: 94°C for 5 min, 29 cycles with 94°C for 50 s, 47°C for 30 s, 72°C for 30 s and a final 103 elongation step of 72°C for 5 min. 10^{9} PCR Clean-up System (Promega Corporation, Madison, WI, USA) according to the manufacturer's 105 instructions and quantified using Qubit® (Life Technologies, Carlsbad, CA, USA). All DNA $\frac{1}{4}$ 06 samples were tested for amplification inhibition by sample dilution. The libraries were sequenced 107 by MiSeq Illumina (Illumina, Inc., SanDiego, CA, USA) using a 300 bp × 2 paired-end protocol. ¹/₁₀₈ The sequencing produced a total of 1,656,884 reads with an average of 103,555 \pm 10,525 reads per 21
2109 sample (average \pm standard error). 1 $3 \times 2 \times 1$ μ ₂ were: 94 C for 3 min , 29 cycles $6\overline{6}$ 10^{4} T CK Cican-up System (1 folloga 11 1205 instructions and quantified using 15 16 (1990) - 1990 (1991) - 1990 1107 by MiSeq Illumina (Illumina, Inc 18 20 2μ 9 sample (average \pm standard error).

26 ₂41 2.3. Bioinformatic elaboration $2\frac{1}{2}$ 1 2.3. Dioinjormatic etaboration

2012 Bioinformatic elaborations were performed in R 4.0.3 (R Core Team, 2020) using DADA2 package $31/3$ (Callahan et al., 2016), version 1.16.0. According to the quality profiles, forward reads were 114 truncated at 200 bases, and reverse reads were truncated at 180 bases. The first 20 bases were 3615 removed from both the forward and the reverse reads. Low quality reads (i.e., reads with expected 116 errors higher than 2 and with Ns) were discarded. Specific error rates were estimated for the 38 117 forward reads and for the reverse reads. Filtered reads were dereplicated, the estimated error rates 41 $^{43}_{44}$ 18 were used to infer the Amplicon Sequence Variants (ASVs) (Callahan et al., 2017) and the reads 119 pairs were merged with default parameters. Chimeric sequences were removed and taxonomic 46 $^{48}_{44}$ 20 assignment (confidence 80%) for each ASV was performed against the RDP database (Cole et al., 51121 2014) using the assign Taxonomy function. Species assignment was performed by add Species $\frac{53}{2}$ function (100% identity). Only the ASVs with a relative abundance of 0.01% (or higher) in at least 123 one sample were considered for further processing. After filtering, merging, removal of chimeric $\frac{5624}{4}$ sequences and removal of low abundance ASVs a total of 1,190,658 high-quality sequences were 60
 6425 obtained with an average of 74,416 \pm 7,958 sequences per sample (average \pm standard error). 30 32^2 (Cananan ct al., 2010), version 33 3444 truncated at 200 bases, and reve 37^o \cdots \cdots \cdots \cdots \cdots \cdots \cdots \cdots $31\frac{1}{9}$ errors higher than 2 and with N 40 42 44μ 8 were used to meer the Amplicon 45 47 $49²$ assignment (connuence $80/0$) for 50 54 and $(100\%$ dentity). Only the 55 5423 one sample were considered for 57 6425 obtained with an average of $74,41$

127 2.4. Chemical analyses \sim 3

 Short chain volatile fatty acids (SCVFAs) (C2:0, acetic; C3:0, propionic; C4:0, butyric; iso C4:0, 129 isobutyric; C5:0, valeric; iso C5:0, isovaleric) and lactic acid were extracted as follows: 10 g of each sample were added to 50 mL of 0.1 N H₂SO₄ aqueous solution and homogenized for 2 min by 131 UltraTurrax (IKA®-Werke GmbH & Co. KG, Staufen, Germany). After the extraction 15 mL of $^{14}_{12}$ 32 supernatant were centrifuged at 2500 rpm for 15 min. Five mL of the supernatant were microfiltered 133 $(0.22 \text{-} \mu\text{m})$ to remove solid particles. The resulting sample was directly injected in the HPLC ¹ 234 apparatus using an Aminex 85 HPX-87 H ion exclusion column (300 mm × 7.8 mm; 9-µm particle 135 size; Bio-Rad, Milan, Italy); the detection wavelength was 220 nm. The analyses were carried out applying an isocratic elution (flux 0.6 mL/min) with a 0.008 N H₂SO₄ solution as mobile phase; the $^{26}_{24}$ 37 injection loop was 20 µL. Individual SCVFAs and lactic acid were identified and quantified by 21938 means of an external calibration curve using a standard solution of 4.50 mg/mL of lactic acid, 5.40 $31/39$ mg/mL of acetic acid, 5.76 mg/mL of propionic acid, 7.02 mg/mL of butyric acid and isobutyric 1440 acid, 8.28 mg/mL of valeric acid and isovaleric acid in 0.1 N H_2SO_4 (69775, 338826, 402907, $\frac{36}{14}$ 1 B103500, 58360, 75054, 129542, respectively; Sigma- Aldrich, Milano Italy). The molar 142 concentration of each SCVFA and of lactic acid were estimated and the ratios lactic acid/acetic 143 acid, lactic acid/propionic acid and propionic acid/acetic acid were calculated. Ezo Snort chain volatile latty actus (SC $6\overline{6}$ 8 and 2010 10^{10} cath sample were added to be the $15²$ supernature were contrinuous at 2. 16 (a.e. 1955).
16 (a.e. 1956). $11B3$ (0.22- μ m) to remove solid parti **The Country of the Country of the Second Street** size; Bio-Rad, Milan, Italy); the c \cdots injection loop was 20 μ L. murvi $32³$ ing/life of accur acid, 3.70 ing/li acid, 8.28 mg/mL of valeric acid $37⁻$ 2100000, 00000, 10001, 1250 31942 concentration of each SCVFA and

 $^{43}_{44}$ The samples of pecorino cheese were analyzed for the determination of the concentration of vitamin B₁₂. The main natural forms of cobalamin present in food are hydroxocobalamin, 5²- $^{48}_{44}$ 6 deoxyadenosylcobalamin, methylcobalamin and cyanocobalamin (vitamin B₁₂). All the different 147 forms of cobalamin were converted to cyanocobalamin before analysis, because it is more stable $53/48$ than the others. Ultra-performance liquid chromatography coupled to triple-quadrupole mass spectrometry (UPLC-MS/MS) was used for the quantification of cyanocobalamin in Pecorino 150 cheese. Methotrexate was used as internal standard (IS). The equipment employed consisted of a 151 Waters Acquity UPLC® binary pump, coupled with a Waters Xevo TQ-S Micro triple quadrupole The samples of pecorino cheese w acoxyanchosynovalalilii, iliciliyl and the others. Once performance 5149 spectrometry (UPLC-MS/MS) w waters Acquity OPLC® binary p

152 mass spectrometer equipped with an ESCiTM Multi-Mode Ionization Source (Waters Corporation, 153 Milford MA, USA). Mass spectrometer operated in positive electrospray ionization (ESI+) mode 2 154 and analysis were performed in MRM (multiple reaction monitoring) mode, following two specific 155 transitions for the target analytes: $678.43 > 147.15$, $678.43 > 359.15$ for vitamin B₁₂ and $455.25 >$ 156 308.17, 455.25 > 134.17 for methotrexate (IS). The chromatographic separation was achieved on a 157 Waters Acquity BEH C18 UPLC® column (Waters Corporation, Milford MA, USA). The 12 $^{14}_{12}$ 58 chromatographic conditions were set as follow: constant flow of 0.350 mL/min; the mobile phase 159 was 5 mM ammonium formate in water acidified with 0.05% of formic acid (A) and acetonitrile ¹ $\frac{1}{26}$ 60 with 0.3% of formic acid (B). The extraction procedure was performed on 1 g of Pecorino cheese in 21
2161 accordance with the protocol described by Zironi et al., 2013, with some modifications due to the 162 high lipid component that characterizes the matrix. 24 1 $\frac{3}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ 454 and analysis were performed in Mi $6\overline{6}$ 8 10^{10} 300.17 , 733.23×137.17 for finem 11 13 ¹⁵⁰ cmomatographic conditions were 16 1159 was 5 mM ammonium formate in 18 20 $2E51$ accordance with the protocol desi 23 25 and $\overline{1}$ and \overline

2164 2.5. Statistical analysis

 3165 Data from 16S rRNA gene amplicons sequencing were further processed using the vegan package, 166 version 2.5.7 (Oksanen et al., 2020) in R 4.0.3 (R Core Team, 2020). To estimate the alpha- $\frac{36}{2}$ diversity within each samples group a randomly rarefied dataset (28,000 sequences) was generated, 168 then, the Chao1 index and the Shannon diversity index were calculated. A non-metric 38 169 multidimensional scaling (NMDS) and a permutational multivariate analysis of variance 41 $^{43}_{4470}$ (PERMANOVA) based on Hellinger transformed ASV abundance data were performed using the 171 metaMDS and the adonis2 functions, respectively. Both the NMDS and the PERMANOVA were 46 $^{48}_{44}$, performed on the Bray-Curtis dissimilarity index. A Kruskal-Wallis test (function kruskal.test) was 173 performed to identify the ASVs and the genera with a different relative abundance between the 51 $\frac{53}{27}$ 4 defected cheese and the control cheese. 32 Data from TOD fixty generation 33 3466 version 2.5.7 (Oksanen et al., 2 35 $37'$ 31,58 then, the Chaol index and the 40 42 $44/0$ (PERIVIANOVA) based on Heilin 45 47 $49²$ performed on the Dray-Curtis dist 50 52 54 ⁻⁴ defected encese and the control of

175 A Kruskal-Wallis test was performed in R 4.0.3 (R Core Team, 2020) to identify the significant $\frac{56}{276}$ differences in the content of SCVFAs and of lactic acid, and in the content of vitamin B₁₂. 5175 • A Kruskal-Wallis test was perfor 57 59

178 3. Results and discussion

 $1/179$ The taxonomic composition of the microbial community of Pecorino Toscano PDO cheese was 180 investigated by high-throughput sequencing of 16S rRNA gene amplicons to identify bacteria 4 $^{6}_{181}$ potentially associated to PD defect. The rarefaction analysis, performed on the detected ASVs, 182 indicated that the sequencing depth was enough to describe the biodiversity within the samples $^{11}_{14}$ 83 (Figure S2). The presence of 74 ASVs was observed within the whole dataset: 18 ASVs were 184 detected only in the control cheese, 19 ASVs were detected only in the defected cheese and 37 14 $^{16}_{14}$ 85 ASVs were shared between the two groups (Figure S3). No differences in Chao1 index and in the 186 Shannon diversity index were observed between the defected cheese and the control cheese (Figure $\frac{21}{25}$ 87 1). 1²⁷⁹ The taxonomic composition of the 3 5 **F**81 potentially associated to PD defect 8 and 2010 12° (Figure 32). The presence of \prime -13 1493 The vs were shared between the ty 18 1986 Shannon diversity index were obs 20

 $^{23}_{24}$ 8 The NMDS plot clearly showed a separation between the bacterial community enriched in the ²189 defected cheese and the bacterial community enriched in the control cheese (Figure 2). The 28
 $\frac{28}{290}$ difference between the bacterial communities enriched in the two conditions was further confirmed 3191 by the PERMANOVA $(R^2 = 0.37, p = 0.002)$. 2488 The NMDS plot clearly showed 25 and 26 an 290 anterence between the bacterial c

 $\frac{33}{22}$ The starter culture was composed by microorganisms belonging to the genera Streptococcus, 193 Lactobacillus and Lactococcus which, in total, accounted for \sim 91 % of the sequences in the control $\frac{38}{19}$ 4 cheese and for ~88% of the sequences in the defected cheeses (Table 1). 34 $\overline{34}$ $\overline{4}$ $\overline{34}$ $\overline{34}$ $\overline{4}$ $\overline{34}$ $\overline{34}$ $\overline{34}$ $\overline{$ 35 3193 Lactobacillus and Lactococcus wl $39'$ checke and for 60% or the sequence

195 To date the main genera that have been clearly associated with the cheese PD defect were Serratia 40 ⁴¹96 in Pecorino Toscano cheese (Martelli et al., 2020) and *Thermus* in continental cheese (Quigley et $^{45}_{4}$ 497 al., 2016). Sequences classified within the genera *Serratia* and *Thermus* were not detected in the 4998 whole dataset (Table S1). Furthermore, the presence of Serratia was linked to PD close to the rind $50₅₀$ of cheese (Martelli et al., 2020), while in this study the discoloration defect was observed in the 200 inner part of the cheese (Figure S1). It is therefore possible to exclude the involvement of these 53 $\frac{5}{20}$ 1 genera in the formation of PD defect in the present study. Quigley and colleagues also observed that 202 the pink color in presence of T. thermophilus was more intense raising the levels of *Lactobacillus* $\frac{6}{2}$ 03 *helveticus* in the starter culture and maintaining *Streptococcus thermophilus* at the same level, but 4195 10 date the main genera that have 42 4θ al., 2010). Sequences classified v 47 $5\frac{1}{2}$ of check (Marten et al., 2020), 52 54 $\frac{201}{56}$ genera in the formation of TD den 57 5202 the pink color in presence of T. t. 59

204 the biological reason of this difference was not determined (Quigley et al., 2016). A similar pattern 205 in the relative abundance of these two genera was observed also in this work with a higher ($p <$ $^{4}_{206}$ 0.05) ratio *Lactobacillus/Streptococcus* in the cheese with PD (0.024 \pm 0.002) compared to the $\overline{207}$ control cheese (0.017 \pm 0.002). Furthermore, Martelli and colleagues isolated different strains of 208 Enterobacter spp. together with S. liquefaciens from the pink spots on the rind of Pecorino Toscano 209 (Martelli et al., 2020) and despite the PD was not attributed to the genus *Enterobacter* it is $\frac{14}{12}$ 10 interesting to observe that, in our study, the relative abundance of this genus was higher (p < 0.01) 211 in the samples collected by the defected cheese (Table 1). 1 3 40 aux ratio Laciobaculus/Streptoco $6\overline{6}$ 8 and 2012 **1996** 1900 *Enterboucter* spp. together with 5. 11 1209 (Martelli et al., 2020) and desp 15 merebring to observe that, in our 16 1211 in the samples collected by the det

 $\frac{1}{2}$ 212 Another difference possibly linked to PD defect is related to *Propionibacteriaceae* family. Three 21
2213 • ASVs were classified within this family, one ASV (ASV 21) was close to *Propionibacterium* 2414 *freudenreichii* (Table S2), and the other two ASVs (ASV 27 and ASV 64) were close to 26
 $\frac{26}{24}$ 5 *Acidipropionibacterium olivae* and *Acidipropionibacterium damnosum* (Table S2), previously 216 classified as Propionibacterium olivae and Propionibacterium damnosum (Turgay et al., 2020), 29 $\frac{31}{27}$ both isolated from spoiled packaged green olives (Lucena-Padrós et al., 2014). However, ASV_27 218 and ASV_64 show a high similarity also to Acidipropionibacterium jensenii and to $\frac{36}{219}$ Acidipropionibacterium thoenii (Table S2), two propionic bacteria associated to dairy products 38
320 (Turgay et al., 2020). 20 **CONTRACT SERVICE CONTRACT** $2\frac{1}{2}$ 13 ASVS were classified within the 23 25 24 5 Aciaipropionibacierium otivae a 28 30 $32'$ both isolated from sponce package 33 3418 and ASV_64 show a high 35 37° \cdots \cdots \cdots \cdots \cdots \cdots \cdots \cdots 320 (Turgay et al., 2020).

 42 21 In our study, *P. freudenreichii* was more abundant in the defected cheese (Table 1, p < 0.01). *P.* $^{43}_{42}$ 2 *freudenreichii* subsp. *shermanii* was previously described as a possible agent of PD in Swiss 223 cheese, but no mechanism was suggested (Daly et al., 2012; Park et al., 1967). Regarding the genus 46 $^{48}_{24}$ Acidipropionibacterium, the other member of the family *Propionibacteriaceae*, detected in the 225 dataset (Table S1), its presence was observed only in the defected cheese (Table 1, $p < 0.01$). $\frac{52}{2}$ 6 Members of the genus *Acidipropionibacterium* can produce red pigments that can cause 227 discoloration defect in cheese (Turgay et al., 2020), therefore it is possible to hypothesize that 228 members of the family Propionibacteriaceae were the main actors involved in the formation of PD 58 229 defect in the Pecorino Toscano PDO cheese. The higher relative presence of propionic bacteria in 60 42 442 freudenreichti subsp. *shermanti* 45 47 49⁴ *Acturpropromoderum*, the other 50 5225 dataset (Table S1), its presence 54 Memoers of the genus n_{cusp} . 55 5227 discoloration defect in cheese (1 57 6429 detect in the Pecorino Toscano P

230 the defected cheese has been evaluated by sequencing of the 16S rRNA gene, therefore there are no 231 clues on their metabolic activity within the cheese. Propionic bacteria are characterized by 2 $\frac{4}{23}$ propionic fermentation, which leads to the production of acetate and propionate using lactate as the 233 substrate (Turgay et al., 2020). Therefore, to obtain more information on their activity the SCVFAs 7 234 and the lactic acid molar concentrations (Figure S4) and percentages were measured (Figure 3A). 235 Furthermore, the lactic acid/acetic acid, the lactic acid/propionic acid and the propionic acid/acetic 12 $^{14}_{12}$ 36 acid ratios were calculated (Figure 3B). The percentage of lactic acid was higher in the control 237 cheese compared to the defected cheese ($p < 0.01$), while the percentage of propionic acid ($p <$ $\frac{1}{2}$ 38 0.01) and butyric acid (p < 0.05) was higher in the cheese with PD defect (Figure 3A). The ratios 21

2239 lactic acid/acetic acid and lactic acid/propionic acid were considerably higher ($p \le 0.01$) in the 240 control cheese compared to the defected cheese. These data suggest a higher consumption of lactate 24 $\frac{26}{241}$ for the propionic fermentation in the cheese in the presence of PD (Figure 3B). This observation is 242 in accordance with the highest abundance of propionic bacteria in the samples collected from 29 $\frac{31}{24}$ 3 cheese with PD. Furthermore, the higher ratio propionic acid/acetic acid in the defected cheese (p < 244 0.01) suggested the presence of different pathways of propionic fermentation in the two groups of $\frac{3}{24}$ 45 cheese. Indeed, the fermentation seemed to shift towards the production of propionate in the cheese 246 with PD defect. Therefore, a change in the activity of propionic bacteria could be involved in the 38 424 7 production of PD defect in this study. 1 $\overline{3}$ 452 propionic termentation, which lead $6\overline{6}$ 8 and 2. 10^{4} and the factic acid motal concent 11 13 15^o and ratios were calculated (1.5^a) 16 1287 cheese compared to the defected 20 \cdots , 20 2239 actic acid/acetic acid and lactic 25 and 26 an 24μ for the proprome remierration in 28 30 $\frac{24}{32}$ check which D. Furthermore, the 33 3244 0.01) suggested the presence of d 35 37° enverse masses, are communicated 3446 with PD defect. Therefore, a chai 40 42

 $^{43}_{44}$ 8 A possible metabolite responsible for PD could be vitamin B₁₂: vitamin B₁₂ is pink and it could be 4249 produced by P. freudenreichii when it is growing in food-like conditions (Deptula et al., 2017). $^{48}_{42}$ 50 Furthermore, microorganisms within the genera *Escherichia/Shigella* and *Lactobacillus*, which are 251 more abundant in the defected cheese (Table 1), have been shown to possess the genes encoding for 51 $\frac{53}{25}$ the enzymes required in vitamin B₁₂ biosynthetic pathway (Balabanova et al., 2021). There are four forms of vitamin B_{12} : cyanocobalamin, hydroxocobalamin, methylcobalamin and 254 adenosylcobalamin (Prentice et al., 2013). Hydroxycobalamin, adenosylcobalamin, and 58 $\frac{60}{62}$ methylcobalamin are the major forms of vitamin B₁₂ in bovine milk and hard cheese (Gille and 4448 A possible inetabolite responsible 45 47 490 runnermore, interoorganisms with 50 52 54 and $\frac{3}{24}$ and $\$ 55 vitamin 57 59 6455 methylcobalamin are the major t

256 Schmid, 2015). The involvement of vitamin B_{12} in the PD defect in this study was excluded since 257 the content of vitamin B₁₂ did not differ in the defected cheese (24 \pm 1 ng/g) compared to the $\frac{4}{258}$ control cheese (29 ± 2 ng/g), though a tendency (p < 0.1) to a lower concentration in the defected 259 cheese was observed (Figure S5). These values are in accordance to the values reported in literature 7 260 for other dairy products (Gille and Schmid, 2015; Souci et al., 2008) since a vitamin B₁₂ 261 concentration of 3.8 ng/g was reported for curd, but a concentration ranging from 10 ng/g to 31 ng/g 12 $\frac{1}{26}$ 2 was reported for other cheeses (Gille and Schmid, 2015). The work of Prentice et al. (2013), on 263 formation of pink color in therapeutic proteins, suggests a threshold between 300 and 500 ng/g $\frac{1}{26}$ 4 hydroxycobalamin for visible pink. This threshold, even if referred to a different matrix, is well 21 above the average values found in this work. 1 $\overline{3}$ $\overline{4}58$ control cheese (29 ± 2 hg/g), thought $6\overline{6}$ $8 \qquad \qquad 8$ 1900 for other dairy products (Office 11 13 15 $\frac{152}{25}$ $\frac{152}{$ 16 1263 formation of pink color in therap 18 20 225 above the average values found in

$\frac{26}{2}$ 67 4. Conclusions 2^{2p} 4. Conclusions

268 Pink discoloration is an important issue in cheese manufacturing but its causes are still not 29 $\frac{31}{269}$ completely understood. In this study, bacteria belonging to the genera *Thermus* and *Serratia*, which 270 have been associated to PD defect were excluded as their presence was not observed in the samples $\frac{36}{27}$ 1 analyzed. As well, data do not support the involvement of vitamin B₁₂ in the formation of PD. 272 Considering the differences observed in the bacterial community of Pecorino Toscano PDO cheese 38 $\frac{42}{7}$ 3 with PD defect, the involvement of microorganisms belonging to the genera *Propionibacterium* and $^{43}_{42}$ 4 *Lactobacillus*, has been hypothesized. These genera were more abundant in the defected cheese 275 compared to the control and a different propionic fermentation was observed in the two groups of 46 $^{48}_{27}$ 6 cheese. Anyway, further studies could be helpful to elucidate the mechanisms that causes PD in 5277 Pecorino cheese. 30 $\frac{20}{32}$ completely understood. In this sta 33 3470 have been associated to PD defect 35 $37²$ and $37²$ 38/2 Considering the differences obser 40 42 44⁴ *Laciobaciuus*, nas been hypoines 45 47 49° chccsc. Anyway, further studies 50

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352 Tables $1 \qquad \qquad \blacksquare$

353 Table 1 Average relative abundance of the microorganisms detected in the defected cheese and in $\frac{6}{354}$ the control cheese. Only the genera with a relative abundance of 0.1%, or higher, in at least one 355 sample are reported. Values are reported average \pm standard error. N.D. = not detected (i.e., relative ¹¹/₁356 abundance = 0). Significance codes: * p < 0.05; ** p < 0.01. 3_{p4} the control cheese. Only the gener 8 and 2010 **120 and 2010 120 and 2010 120 and 2010 120 and 2010 120 and 2010** 12^{20} abundance -0). Significance code

	Control $(\%)$		Defect $(\%)$			
Genus	Average	Std. Error	Average	Std. Error	p-value	
Acidipropionibacterium	N.D.		0.06	0.02	0.001	$***$
Corynebacterium	0.01	0.01	N.D.		0.317	
Enterococcus	0.01	0.01	0.06	0.02	0.008	$***$
Escherichia/Shigella	0.07	0.02	0.19	0.04	0.046	\ast
Lacticaseibacillus	1.78	0.64	0.75	0.18	0.142	
Lactiplantibacillus	0.41	0.13	0.49	0.08	0.401	
Lactobacillus	1.44	0.18	1.96	0.18	0.046	\ast
Lactococcus	4.49	0.76	4.07	0.42	0.674	
Lentilactobacillus	1.37	0.38	3.58	0.49	0.006	$***$
Leuconostoc	0.67	0.15	0.30	0.19	0.027	$***$
Pediococcus	0.24	0.07	0.04	0.02	0.004	$***$
Propionibacterium	0.06	0.06	0.09	0.02	0.016	$***$
<i>Streptococcus</i>	84.94	1.90	81.87	0.92	0.208	
Weissella	0.00	0.00	0.04	0.02	0.371	
Other genera	0.06	0.01	0.07	0.01		
Unclassified	4.44	0.20	6.44	0.45		

360 Figure captions

361 Figure 1 Alpha diversity indexes calculated for ASV abundance. The alpha diversity is not $\frac{6}{36}$ different in the defected cheese compared to the control cheese. c 1 c 3p2 different in the defected cheese con

 $^{11}_{13}$ 64 Figure 2 - Non Metric Mutidimensional Scaling (NMDS) based on the Bray-Curtis distance, 1365 calculated on the Hellinger transformed ASV relative abundance data. Stress $= 0.061$. The bacterial $\frac{16}{136}$ community enriched in the defected cheese and the bacterial community in the control cheese were 367 different. 12^{24} right $2 -$ iven include maturem $17²$ community emicroe in the defect $18 \quad \ldots$

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 $23\overline{4}$ 69 Figure 3 – Relative abundance of the detected SCVFAs and lactate in the control cheese and in the 370 defected cheese (A). The proportion of lactic acid was lower in the defected cheese, and the 371 proportion of propionic acid was lower in the control cheese. Ratios of fatty acids and of lactic acid 372 in the control cheese and in the defected cheese (B). Lactic acid/acetic acid and lactic acid/propionic $\frac{33}{27}$ 3 acid ratios were higher in the control cheese, while propionic acid/acetic acid ratio was higher in the defected cheese. Significance codes: * $p < 0.05$; ** $p < 0.01$. $24p9$ Pigure 3 – Relative abundance of 25 and 26 an 27 (a) $\sqrt{27}$ $29/1$ proportion of propionic acid was a and ratios were ingred in the com- 3374 defected cheese. Significance code

