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Impact of thistle rennet from Carlina acanthifolia All. subsp. acanthifolia on bacterial diversity and dynamics of a specialty Italian raw ewes' milk cheese

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Impact of thistle rennet from Carlina acanthifolia All. subsp. acanthifolia on bacterial diversity and dynamics of a specialty Italian raw ewes' milk cheese / Cardinali, Federica; Osimani, Andrea; Taccari, Manuela; Milanovic, Vesna; Garofalo, Cristiana; Clementi, Francesca; Polverigiani, Serena; Zitti, Silvia; Raffaelli, Nadia; Mozzon, Massimo; Foligni, Roberta; Franciosi, Elena; Tuohy, Kieran; Aquilanti, Lucia. - In: INTERNATIONAL JOURNAL OF FOOD MICROBIOLOGY. - ISSN 0168-1605. - ELETTRONICO. - 255:(2017), pp. 7-16. [10,1016/j.ijfoodmicro.2017.05.018] *Availability:* 

This version is available at: 11566/248306 since: 2020-04-12T23:04:51Z

Publisher:

Published DOI:10.1016/j.ijfoodmicro.2017.05.018

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note finali coverpage

(Article begins on next page)

Elsevier Editorial System(tm) for International Journal of Food Microbiology Manuscript Draft

Manuscript Number:

Title: Impact of thistle rennet from Carlina acanthifolia All. subsp. acanthifolia on bacterial diversity and dynamics of a specialty Italian raw ewes' milk cheese

Article Type: Short Communication

Keywords: Sopravissana raw ewes' milk cheese; vegetable milk coagulant; bacterial biota; culture; PCR-DGGE; high-throughput sequencing.

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Ancona 01-02-2017

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Dear Editor,

would you please consider the manuscript titled "Impact of thistle rennet from *Carlina acanthifolia* All. subsp. *acanthifolia* on bacterial diversity and dynamics of a specialty Italian raw ewes' milk cheese" for publication as short communication in International Journal of Food Microbiology.

Caciofiore della Sibilla is an Italian specialty soft cheese manufactured with Sopravissana raw ewes' milk and thistle rennet prepared with young fresh leaves and stems of Carlina acanthifolia All. subsp. acanthifolia, according to an ancient tradition deeply rooted in the territory of origin (mountainous hinterland of the Marche region, Central Italy). In this study, the impact of thistle rennet on the bacterial dynamics and diversity of Caciofiore della Sibilla cheese was investigated by applying a polyphasic approach based on culture and DNA-based techniques (Illumina sequencing and PCR-DGGE). A control cheese manufactured with the same batch of ewes' raw milk and commercial animal rennet was analyzed in parallel. Overall, a large number of bacterial taxa were identified, including spoilage, environmental and pro-technological bacteria, primarily ascribed to Lactobacillales. Thistle rennet was observed clearly to affect the early bacterial dynamics of Caciofiore della Sibilla cheese with several bacterial groups (Lactobacillus alimentarius/paralimentarius and Lactobacillus plantarum/paraplantarum/pentosus) being detected in the phyllosphere of C. acanthifolia All, thistle rennet and curd obtained with thistle rennet. Other bacterial taxa, hypothetically originating from the vegetable coagulant (Enterococcus faecium, Lactobacillus brevis, Lactobacillus delbrueckii, Leuconostoc mesenteroides/pseudomesenteroides), were exclusively found in Caciofiore della Sibilla cheese by PCR-DGGE. At the end of the maturation period, Illumina sequencing demonstrated that both cheeses were dominated by Lactobacillales; however curd and cheese produced with thistle rennet were co-dominated by Lactobacillus and Leuconostoc, whereas Lactoccous prevailed in curd and cheese produced with commercial animal rennet followed by Lactobacillus. Differences in the bacterial composition between the two cheeses at the end of their maturation period were confirmed by PCR-DGGE analysis.

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Thank you very much for your kind consideration. Best regards Lucia Aquilanti



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# To whom it may concern

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# Highlights

- Caciofiore della Sibilla cheese and a control cheese were analyzed in parallel
- A polyphasic approach based on culture and DNA-based techniques was applied
- Thistle rennet affected early bacterial dynamics of *Caciofiore della Sibilla* cheese
- The two cheeses differed in their bacterial composition at the end of ripening

1	Impact of thistle rennet from Carlina acanthifolia All. subsp. acanthifolia on bacterial diversity and dynamics of
2	a specialty Italian raw ewes' milk cheese
3	
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32 Abstract

33

Caciofiore della Sibilla is an Italian specialty soft cheese manufactured with Sopravissana raw ewes' milk and thistle 34 35 rennet prepared with young fresh leaves and stems of Carlina acanthifolia All. subsp. acanthifolia, according to an 36 ancient tradition deeply rooted in the territory of origin (mountainous hinterland of the Marche region, Central Italy). In 37 this study, the impact of thistle rennet on the bacterial dynamics and diversity of Caciofiore della Sibilla cheese was 38 investigated by applying a polyphasic approach based on culture and DNA-based techniques (Illumina sequencing and 39 PCR-DGGE). A control cheese manufactured with the same batch of ewes' raw milk and commercial animal rennet 40 was analyzed in parallel. Overall, a large number of bacterial taxa were identified, including spoilage, environmental 41 and pro-technological bacteria, primarily ascribed to Lactobacillales. Thistle rennet was observed clearly to affect the 42 early bacterial dynamics of Caciofiore della Sibilla cheese with several bacterial groups (Lactobacillus 43 alimentarius/paralimentarius and Lactobacillus plantarum/paraplantarum/pentosus) being detected in the phyllosphere of C. acanthifolia All, thistle rennet and curd obtained with thistle rennet. Other bacterial taxa, hypothetically 44 45 originating from the vegetable coagulant (Enterococcus faecium, Lactobacillus brevis, Lactobacillus delbrueckii, 46 Leuconostoc mesenteroides/pseudomesenteroides), were exclusively found in Caciofiore della Sibilla cheese by PCR-47 DGGE. At the end of the maturation period, Illumina sequencing demonstrated that both cheeses were dominated by 48 Lactobacillales; however curd and cheese produced with thistle rennet were co-dominated by Lactobacillus and 49 Leuconostoc, whereas Lactoccous prevailed in curd and cheese produced with commercial animal rennet followed by 50 Lactobacillus. Differences in the bacterial composition between the two cheeses at the end of their maturation period 51 were confirmed by PCR-DGGE analysis.

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Keywords: *Sopravissana* raw ewes' milk cheese, vegetable milk coagulant, bacterial biota, culture, PCR-DGGE, high throughput sequencing

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63 **1. Introduction** 

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*Caciofiore della Sibilla* is a specialty soft cheese manufactured in a restricted central Italian geographical area using
 *Sopravissana* raw ewes' milk and thistle rennet obtained from young fresh leaves and stems of *Carlina acanthifolia* All.
 subsp. *acanthifolia* according to an ancient local tradition.

68 The term "thistle" refers to plants belonging to the tribe Cynareae (synonym: Cardueae) especially ascribed to the 69 genera Carduus, Cirsium, and Onopordum. However, plants outside this tribe are sometimes considered thistles, 70 including those within the genera Cynara, Scolymus, Silybum, Onopordum, and Carlina. In the Mediterranean area, the 71 exploitation of thistle rennet, especially that obtained from Cynara spp., is particularly widespread in western Africa 72 (García et al., 2012; Cardinali et al., 2016), Italy, and the Iberian peninsula, the latter boasting a large number of 73 Protected Designation of Origin (PDO) cheeses manufactured with this peculiar coagulant (Aquilanti et al., 2011; 74 Cardinali et al., 2016). Cheeses coagulated with thistle rennet are generally manufactured at family-run or artisan dairy 75 farms, most often located in marginal areas (e.g., high altitude pastures, dry lands or islands) using raw ewes' or goats' 76 milk, or a mixture of both. Though these cheeses are greatly appreciated by consumers for their unique, distinctive 77 flavor, their manufacturing is generally seasonal, mainly due to limitations on the availability of young leaves or 78 flowers from spontaneously growing thistles.

*Caciofiore della Sibilla* cheese undoubtedly falls within this cheese category. The manufacturing technology of this cheese, which had been lost for more than 50 years in the original area of production, has very recently been revived by two local family-run dairies located in Pieve Torina and Belforte del Chienti (Macerata district), respectively. The cheese produced by these local dairies from late spring to early summer is 3-4 cm tall, has an average weight of 0.2 to 0.8 kg and is characterized by a very thin straw-white outer rind and a cream-white soft core, with a sweetish buttery smell, a delicate but incisive flavor with a scent of wild herbs, and a slightly acidulous, pleasant taste.

To date, few studies have been conducted to identify the bacterial biota harboured by thistle-rennet cheeses (Sousa and Malcata, 1997; Vioque et al., 2000; Gómez et al., 2001; Roseiro et al., 2003; Fernández-Salguero et al., 2002; Tejada and Fernández-Salguero, 2003; Aquilanti et al., 2011; Galán et al., 2012; Ordiales et al., 2013) and almost all of these investigations have been focused on Protected Designation of Origin (PDO) or specialty cheeses manufactured with thistle coagulants obtained from *Cynara* spp.

Based on these premises, this study uses a polyphasic molecular approach based on culture and DNA-based techniques to assess the impact of an unexplored milk coagulant obtained from *Carlina acanthifolia* All. subsp. *acanthifolia* on the bacterial dynamics and diversity of *Caciofiore della Sibilla* cheese. To this end, two cheese manufactures produced with 93 the same batch of *Sopravissana* raw ewes' milk and coagulated with either thistle rennet or commercial powdered 94 animal rennet were analyzed in parallel and the results comparatively evaluated.

95

#### 96 2. Materials and Methods

97

## 98 2.1. Cheese-making process and sampling

99 *Caciofiore della Sibilla* cheese was made in a family-run dairy farm located in Pieve Torina (Italy) following an ancient
 100 local manufacturing method without any addition of starter cultures.

101 The sole Caciofiore della Sibilla cheese manufacture produced by the dairy farm during spring and summer 2015 was 102 sampled and analysed. Raw milk obtained from Sopravissana ewes during one milking day was filtered and separated 103 into two batches; three cheese wheels were produced from each batch. The first batch (labelled "C") was coagulated 104 with commercial powdered calf rennet (Caglificio Clerici, Cadorago, CO, Italy; 1:10,000), whereas the second batch 105 (labelled "Cf") was coagulated with a crude aqueous extract prepared from C. acanthifolia All. subsp. acanthifolia. The 106 traditional preparation of the crude aqueous extract based on the use of fresh young leaves and stems - containing a 107 considerable quantity of latex - of plants spontaneously growing in the high altitude pastures (>1000 m a.s.l.) of Monti 108 Sibillini National Park has previously been detailed by Cardinali et al. (2016). Milk coagulants were added to raw milk 109 pre-heated at 35 °C and gently stirred. After clotting (ca. 30'), the curds were manually broken into rice-sized grains, 110 transferred into plastic perforated molds (8 x 8 cm) and manually pressed to remove the whey. Molded cheeses were 111 held at 12-13 °C for 4 h, dry salted and ripened for 20 days under controlled conditions (12-13 °C and 70 % relative 112 humidity).

Ewes' raw milk, fresh young leaves and stems of *C. acanthifolia* All. subsp. *acanthifolia*, thistle rennet, curds and cheeses (after 1, 3, 6, 10 and 20 days of ripening) were collected in triplicate. Samples were transported to the laboratory under controlled temperature (4 °C) and processed within 24 h. Triplicate samples were pooled before viable counting and molecular analysis.

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The pH of samples collected during cheese-making was measured using the pH meter 300 (Hanna Instruments, Padova, Italy), equipped with a solid electrode (HI2031, Hanna Instruments). Three independent measurements were performed for each sample, and the mean values  $\pm$  standard deviations were calculated.

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<sup>118 2.2.</sup> pH measurements

125

126	For the microbial counts, 10 grams of each solid sample were accurately homogenized in 90 mL of sterile aqueous
127	citrate (2 % w/v) for 2 min at 260 rpm using a Stomacher apparatus (400 Circulator, VWR International PBI, Milan,
128	Italy). Aliquots of decimal dilutions of raw milk, thistle rennet and the homogenates were inoculated in duplicate on
129	opportune solid media to determine the load of (i) presumptive lactococci, thermophilic cocci and lactobacilli; (ii)
130	coagulase-negative cocci; (iii) total mesophilic aerobes; (iv) enterococci; (v) and finally Enterobacteriaceae (Garofalo
131	et al., 2017). Viable counts were expressed as log colony forming units (log cfu) per gram or mL of sample $\pm$ standard
132	deviations. For all samples but cheeses, bulk cells were prepared by harvesting confluent colonies from both the lowest
133	MRSA and M17 dilution plates and the countable plates with a number of colonies ranging from 30 to 300 (Garofalo et
134	al. 2015). For cheeses, the sole colonies grown on the countable plates were harvested. Harvested cells were suspended
135	in 2 mL of sterile saline solution added with glycerol, and stored at -20 °C prior to DNA extraction (Garofalo et al.,
136	2015).

137

### 138 2.4. DNA extraction from cheese samples and bulks

139

For raw milk, thistle rennet, curd and cheese homogenates, total microbial DNA was extracted using a commercial kit (PowerFood<sup>TM</sup> Microbial DNA Isolation Kit, Mo Bio Laboratories, Carlsbad, USA), as previously elucidated (Cardinali et al. 2016). For bulk cells, DNA extraction was performed using the method proposed by Hynes et al. (1992) with some modifications as reported by Osimani et al. (2015). Assessment of quantity and purity of DNA extract was carried out as described by Osimani et al. (2016).

145

#### 146 2.5. PCR amplification and DGGE analysis

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The universal primers 338F (5'-ACT CCT ACG GGA GGC AGC AGC AG-3'), added with the GC clamp at the 5' end (Ampe et al. 1999), and 518R (5'-ATTACC GCG GCT GCT GG-3') were used to amplify the V3 region of the 16S rRNA gene (Alessandria et al., 2010).

Approximately 100 ng of bacterial DNA were amplified as previously reported by Osimani et al. (2015) using the thermal cycler My Cycler (Bio-Rad Laboratories, Segrate, Italy). DGGE runs were performed as reported by the same authors. Sequencing of DNA eluted from selected DGGE bands was performed in accordance to Taccari et al. (2016). Sequences were compared with those deposited in the GenBank DNA database (http://www.ncbi.nlm.nih.gov/) using

155	the Basic Local Alignment Search Tool (BLAST). A sequence identity $\geq 97$ % was chosen as a threshold for
156	unambiguous assignment into species.
157	
158	2.6. DNA amplification and Illumina sequencing
159	
160	The extracted DNA was used to study the bacterial diversity of the samples (raw milk, thistle rennet, curds and cheeses)
161	by 16S rRNA amplicon Illumina sequencing. A 464-nucleotide sequence of the V3-V4 region of the 16S rRNA gene
162	was analyzed as previously described by Alfonzo et al. (2017).
163	
164	2.7. Illumina data analysis and sequence identification by QIIME
165	
166	Sequences obtained from Illumina sequencing were processed using Quantitative Insights Into Microbial Ecology
167	(QIIME) software package version 1.9 (Caporaso et al., 2010) as previously detailed by Alfonzo et al. (2017). The data
168	generated by Illumina sequencing were deposited in the NCBI Sequence Read Archive (SRA) and are available under
169	Ac. No. PRJNA340351.
170	
171	2.8. Statistical analysis
172	
173	For each cheese batch, viable counts recorded after 1, 3, 6, 10 and 20 days of maturation were analyzed by an ANOVA
174	and a Tukey's test ( $\alpha$ = 0.05) for means separation. At each sampling time, data from the two batches "C" and Cf" were
175	compared using the Student's t-test ( $\alpha$ =0.05). A principal component analysis (PCA) with a correlation matrix was
176	carried out on standardized data to visually identify bacterial dynamics over time. All the statistical analyses were
177	performed using JMP statistical software v. 11.0 (SAS Institute, Cary, NC, USA).
178	
179	3. RESULTS
180	
181	3.1. pH measurement and viable counting
182	
183	Raw ewes' milk and thistle rennet were characterized by pH mean values of 6.39±0.10 and 3.79±0.05, respectively; the
184	results of pH measurements carried out in curds and cheeses sampled at different time points are reported in Fig. 1. As

revealed by ANOVA, curds and cheeses coagulated with thistle and animal rennet significantly differed in their pH values up until the 10<sup>th</sup> day of ripening, whereas after 20 days of maturation, both cheeses reached similar pH values.

The results of viable counting are shown in Table 1. The phyllosphere of *C. acanthifolia* All. was characterized by high levels of total mesophilic aerobes, lactococci, *Enterobacteriaceae* and coagulase-negative cocci and, conversely, by low levels of presumptive thermophilic cocci, enterococci and lactobacilli. Low bacterial loads were seen in thistle rennet.

No significant differences were observed between raw ewes' milk and curds in the viable counts of presumptive lactobacilli, total mesophilic aerobes and enterococci, whereas significant differences were observe in the loads of presumptive lactococci, termophilic cocci and coagulase-negative cocci.

Viable counts of lactobacilli, total mesophilic aerobes, enterococci and *Enterobacteriaceae* increased progressively during the ripening of *Caciofiore della Sibilla* cheese; moreover, at the end of the maturation period (20 days), the two cheeses differed significantly in the load of all the bacterial groups assayed, except for lactobacilli and *Enterobacteriaceae*.

When PCA was computed, the first component accounted for 85.7 % of total variability whereas the second component accounted for an additional 5.3 % (Fig. 2). As clearly evidenced by the plot, the *Caciofiore della Sibilla* cheese had a greater range of variation in the bacterial loads than the control cheese, especially up until the third day of maturation.

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### 201 3.2. PCR-DGGE analyses

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203 The DGGE profiles obtained by analyzing the DNA extracted from both the samples and the bulk cells harvested from 204 selected dilution plates are shown in Fig. S1 and S2, and the results of multiple alignments of DNA sequences from 205 selected DGGE bands are shown in Table S1. These results are summarized in Tables 2 and 3. In the phyllosphere of C. 206 acanthifolia All., the closest relatives to Pantoea agglomerans, Pseudomonas alcaliphila Acinetobacter baumannii, the Lactobacillus alimentarius/paralimentarius group, the Lactobacillus graminis/curvatus group, Enterobacter spp., 207 208 Pseudomonas spp., and Bacillus spp. were found. Members of the latter genus were also detected in thistle rennet, along 209 members of the Lactobacillus plantarum/paraplantarum/pentosus with group and the Lactobacillus 210 alimentarius/paralimentarius group.

In raw ewes' milk, the following taxa were detected: Acinetobacter johnsonii, Enterobacter cloacae, Enterobacter
 hormaechei, Pseudomonas spp. and Lactococcus lactis. The latter species was detected at both the lowest and highest
 dilution.

214 Completely different bacterial compositions were seen in the two curds: curd obtained with commercial animal rennet 215 was dominated by *Enterobacteriaceae* bacteria, while curd obtained from thistle rennet was dominated by lactic acid 216 bacteria. In general, the bacterial diversity of cheeses during ripening was higher than that of raw milk, thistle rennet 217 and curds. Furthermore, if the two cheeses were comparatively evaluated, a few taxa, such as Enterococcus faecium, Lactobacillus brevis, Leuconostoc mesenteroides and Weisella spp., were exclusively found in Caciofiore della Sibilla 218 219 cheese, whereas other taxa, such as Bifidobacterium dentium, Citrobacter spp., Enterobacter hormaechei, Erwinia 220 chrysanthemi and the Lact. alimentarius/paralimentarius group, were found only in the control cheese. By contrast, the 221 following bacteria were identified in both cheeses, namely, Enterococcus feacalis, L. lactis, the Lactobacillus 222 casei/paracasei/rhamnosus group, the Lact. plantarum/paraplantarum/pentosus group and Staphylococcus sciuri. With the exception of Bacillus subtilis, Acinetobacter johnsonii and Enterobacter cloacae, all of the remaining taxa were 223 detected in the highest dilution plates, thereby suggesting a load of at least  $10^5$ - $10^8$  cfu/g. 224

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226 *3.3. Illumina sequencing* 

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The DNA extracted from ewes' raw milk, curds and cheeses was successfully amplified in the bacterial V3-V4 16S rRNA gene region. After splitting and quality trimming of raw data, 62,772 reads remained for subsequent analysis. After alignment, OTUs were clustered at a 3 % distance and the doubles and singletons (OTUs counting only two or one reads, respectively) were discarded by a filter script implemented in QIIME. Chao1 estimator predicted an average of 688 and 231  $\pm$  43 OTUs in milk and cheese samples, respectively, and the average observed counts were 661 and 176  $\pm$  40 OTUs, respectively, suggesting that we were able to capture approximately 96 % and 76 % of the OTUs estimated as present in the milk and cheese bacterial populations, respectively.

235 In most samples, a number of OTUs belonged to chloroplasts. These OTUs were not very abundant in raw milk, curd produced with animal rennet, and Caciofiore della Sibilla cheese at 3 and 20 days of ripening, accounting for less than 236 237 1.2 % of total reads number, with the exception of Caciofiore della Sibilla cheese at 3 days of ripening, where they 238 constituted 10.6 % of the total reads number. Conversely, chloroplast OTUs constituted the majority of the reads 239 number in curd obtained with thistle rennet and Caciofiore della Sibilla cheese collected after 1 day of ripening, 240 accounting for 81.9 % and 78.5 % of the total reads number, respectively. The relative abundance (%) of the different bacterial groups identified in raw milk, curds and cheeses are reported in Fig. 3. Only the groups with an incidence of 241 242 0.1 % were considered. The majority of milk OTUs belonged to three different groups: Lactobacillales, 243 Enterobacteriaceae and Clostridiales, which together accounted for 60 % of the total number of bacterial OTUs in raw 244 milk. Various minority taxa were also identified, including Staphylococcus, Planococcaceae and Bacteroidales.

As the bacterial dynamics were comparatively evaluated along with the two cheese-making processes, the following evidence emerged. In curd obtained with commercial animal rennet, *Planococcaceae* and *Enterobacteraceae* prevailed, followed by *Lactobacillales*, *Staphylococcus* and *Macrococcus*. *Enterobacteraceae*, *Lactobacillales* and *Staphylococcus* were also found in curd obtained with thistle rennet. At the end of the maturation period, both cheeses were dominated by *Lactobacillales*, which accounted for approximately 65 % of the total number of bacterial OTUs; however, different minority taxa were found, namely, *Enterobacteriaceae* in control ewes' milk cheese, and both *Macrococcus* and *Acidobacteria* in *Caciofiore della Sibilla* cheese.

In Fig. 4, a finer taxonomic composition within the order *Lactobacillales* is shown. In raw ewes' milk, OTUs corresponding to *Lactobacillus*, *Enterococcus*, *Lactococcus*, *Staphylococcus*, *Carnobacterium*, *Leuconostoc* and *Weissella* were found, though for the latter two taxa, a relative abundance lower than 0.1 % was seen. In curd and cheese obtained with thistle rennet, *Lactobacillus* and *Leuconostoc* co-dominated, whereas *Lactococcus* prevailed in curd and cheese produced with commercial animal rennet, followed by *Lactobacillus*. The two cheese manufactures also differed in the composition of minority taxa, with *Leuconostocs* and *Pediococci* occurring at low levels in both the control and *Caciofiore della Sibilla* cheese, respectively.

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#### 260 **4. Discussion**

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To the best of the authors' knowledge, this report is the first to describe the contribution of thistle rennet obtained from *C. acanthifolia* All. subsp. *acanthifolia* to cheese bacterial composition and dynamics. Moreover, no previous investigations have been conducted on *Caciofiore della Sibilla* cheese, a Mediterranean soft ewes' raw milk cheese manufactured according to a revived local ancient technology.

As far as raw *Sopravissana* ewes' milk was analysed, viable counts of presumptive lactobacilli, lactococci, thermophilic cocci, total mesophilic aerobes, *Enterobacteriaceae* and coagulase-negative cocci were within the range reported by other authors for raw ewes' milk (Quigley et al. 2013).

Regarding thistle rennet, viable counts of total mesophilic aerobes and presumptive lactobacilli were comparable to those found by other authors (Aquilanti et al. 2011; Barbosa et al., 1981) in thistle rennets obtained from *Cynara* spp. dried flowers; by contrast, slightly lower counts of presumptive lactococci and thermophilic cocci were found compared to those reported by the same authors (Aquilanti et al. 2011; Barbosa et al. 1981).

At the end of ripening (20 days) *Caciofiore della Sibilla* cheese showed viable counts of total mesophilic aerobes and presumptive lactobacilli comparable with those reported by Aquilanti et al. (2013) in a raw cow's milk *Caciotta* cheese manufactured with an aqueous extract of *C. cardunculus*, whereas the counts of coagulase-negative cocci and *Enterobacteriaceae* were higher. Higher counts of *Enterobacteriaceae* in cheeses at the end of their ripening might suggest a lack of hygiene during milking and cheese-making. However, the metabolic activity of members of this bacterial family has been positively correlated to high proteolysis and lipolysis in artisanal cheeses, resulting in the
production of volatile aroma compounds (Tabla et al., 2016).

When DNA-based techniques were applied to the profiling of the bacterial biota, numerous taxa were identified, 280281 including starter lactic acid bacteria (LAB), namely Lactococcus lactis, which is known to lower the pH and control 282 fermentation rapidly) and a wide variety of adventitious microorganisms (such as environmental, spoilage and non-283 starter lactic acid bacteria (NSLAB), which gain access to the cheese at any stage of the manufacturing process). 284 NSLAB are mainly responsible for key physico-chemical transformations, e.g., proteolysis and lipolysis, which greatly 285 impact basic cheese characteristics, such as flavor, appearance and texture (Gobbetti et al., 2015). Illumina sequencing allowed both the majority and minority taxa to be identified, whereas single bacterial species were unambiguously 286 287 recognized by PCR-DGGE. When the latter technique was applied, discrepancies were seen in the fingerprints obtained 288 by the bulk and the direct approach, in terms of both the number of taxa identified and their relative abundance; this 289 finding was in agreement with a recent finding by the same authors on semi-hard cheese models (Aquilanti et al., 2016) 290 In Sopravissana raw ewes' milk, PCR-DGGE allowed the sole species L. lactis to be identified among LAB. Illumina 291 sequencing confirmed the occurrence of Lactococcus spp. in raw ewes' milk, though as a minority taxon with respect to 292 the members of the genus Lactobacillus. The presence of both lactococci and lactobacilli in raw ewes' milk has 293 previously been documented, as reviewed by Quigley et al. (2013). The presence of L. lactis was also evidenced in two 294 previous studies (Aquilanti et al., 2011; 2013) carried out onto raw cow's milk Caciotta cheeses manufactured with 295 aqueous extract of Cynara cardunculus dried flowers. The occurrence of additional genera, including Aerococcus 296 Leuconostoc, Streptococcus and Enterococcus, has previously been found in raw ewes' milk by next-generation 297 sequencing (Quigley et al., 2013).

298 In both the phyllosphere of C. acanthifolia All. and thistle rennet, culture-dependent PCR-DGGE identified various 299 LAB species, as well as both spoilage and environmental microorganisms. To date, there is a paucity of data on the 300 bacterial biota colonizing the aerial surfaces of thistles and thistle rennets. In 1968, Mundt and Hammer (1968) first 301 enumerated and identified lactobacilli on plant surfaces; in such a pioneering systematic study, L. plantarum, L. 302 fermenti and L. brevis were the species most frequently isolated, whereas L. casei, L. viridescens, L. cellobiosus, L. salivarius, and L. buchneri occurred at lower frequencies. A widespread but sporadic distribution of low counts of 303 304 lactobacilli in the phyllosphere of higher plants was also highlighted by the same authors. In accordance with data 305 collected in the present study, Widnyana and Javandira (2016) recently reported the presence of Bacillus spp. and 306 Pseudomonas spp. on plant surfaces, where these microorganisms exert a plant-growth stimulating activity as well as an 307 inhibitory activity towards plant pathogens.

Lactobacilli ascribed to the *Lact. alimentarius/paralimentarius* group and the *Lact, plantarum/paraplantarum/pentosus* found by PCR-DGGE in both the phyllosphere of *C. acanthifolia* All. and thistle rennet were also detected in curd obtained by milk coagulation with thistle rennet, thus strongly suggesting a role of the coagulating agent in the early bacterial dynamics of *Caciofiore della Sibilla* cheese. By contrast, the occurrence of *P. acidilactici* in the sole curd obtained by milk coagulation with thistle rennet is more likely related to an environmental contamination, since this species was apparently absent in ewes' raw milk, thistle phyllosphere and thistle rennet.

A different picture emerged in curd obtained by milk coagulation with commercial animal rennet. In this case, LAB were found to be the minority by Illumina sequencing; among these microorganisms, *Lactococcus*, likely derived from raw ewes' milk, was predominant. PCR-DGGE confirmed the occurrence of *L. lactis* as the sole pro-technological species, along with various spoilage and pathogenic bacteria.

318 As far as the bacterial dynamics are considered, the stable presence of L. lactis in both the Caciofiore della Sibilla and 319 control cheeses suggested by the results of PCR-DGGE might be, at least in part, ascribed to the contribution of raw 320 ewes' milk, where lactococci and L. lactis have been revealed by llumina sequencing and culture-dependent PCR-321 DGGE, respectively. It is worth noting that when this latter technique was applied, closest relatives to L. lactis were 322 found in the bulk cells harvested from both the lowest and the highest dilution plates, thus suggesting a load of L. lactis  $\geq 10^5$  ufc/mL. L. lactis has a wide ecological distribution and is mostly associated with the milk environment and dairy 323 324 products (Cavanagh et al., 2015). As recently elucidated by Ruggirello et al. (2016), this species can persist in late 325 ripening of cheese in both the viable and the viable but nonculturable (VNC) states, shifting its catabolism to peptide 326 and amino acid consumption.

Besides *Lactococcus*, bacteria ascribed to *Leuconostoc* and *Lactobacillus* dominated during ripening of *Caciofiore della Sibilla* cheese; both genera have been detected from t1 to t20 by both Illumina sequencing and PCR-DGGE analysis.

329 The dominance of Leuconostoc during the maturation of Caciofiore della Sibilla cheese might be explained by the 330 higher initial pH of the curd obtained with thistle rennet compared to that produced with commercial animal rennet. 331 Indeed, as reported by Hemme and Foucaud-Scheunemann (2004), Leuconostoc can be affected by the acidification of 332 the growth medium. Interestingly, Leuconostoc was indicated by Illumina sequencing as a minority bacterial taxon in 333 control ewes' milk cheese; however, no closest relatives to this genus were identified by PCR-DGGE in this cheese 334 manufacture in either of the two approaches, thus strongly emphasizing the usefulness of a combined analytical 335 strategy. Moreover, since apparently neither raw ewes' milk nor thistle and thistle rennet harboured leuconostocs, as 336 revealed by PCR-DGGE, the occurrence of these microorganisms in *Caciofiore della Sibilla* cheese might be tentatively 337 ascribed to a contamination from the dairy environment. However, members of the genus *Leuconostoc* have previously 338 been detected in a raw cow's milk Caciotta cheese manufactured with thistle rennet from Cynara cardunculus dried

flowers (Aquilanti et al., 2011), as well as in other cheeses produced with vegetable coagulants, such as *Serra da Estrela* PDO cheese and *La Serena* PDO cheese (Macedo et al., 2004). Based on these latter evidences, it might be also hypothesized that in thistle, thistle rennet and curd obtained with thistle rennet, leuconostcocs occurred at levels below the limit of detection of the PCR-DGGE technique.

343 In addition, Lactobacillus delbrueckii, Lactobacillus brevis, and Enterococcus faecium were retrieved by PCR-DGGE 344 exclusively in Caciofiore della Sibilla cheese throughout its maturation. Since these species could not be detected by PCR-DGGE before t1, again it might be hypothesized that they originate from the vegetable coagulant, where they 345 346 occur at very low levels. Lactobacillus delbrueckii includes a subspecies that can be positively affected by the addition 347 of vegetable extracts to the dairy matrix (Joung et al., 2016); even Lactobacillus brevis has been retrieved in vegetable-348 based matrices (Ruiz Rodriguez et al., 2016). Finally, enterococci are NSLAB that are naturally present in the 349 gastrointestinal tract of humans and animals; they can enter the dairy environment via cross-contaminations, where they 350 play a crucial role during cheese ripening, due to their high adaptation to the cheese environment (e.g., high salt concentration and low pH) (Giraffa 2003) and to acknowledged technological traits, such as the production of 351 352 bacteriocins (İspirli et al., 2017).

Finally, members of two additional bacterial groups, namely, *Lactobacillus casei/paracasei/rhamnosus* and *Lactobacillus plantarum/pentosus/paraplantarum*, were identified by PCR-DGGE in both cheese-manufactures; since these microorganisms could not be detected in either raw ewes' milk or thistle rennet, once again, a contamination from the dairy environment might be hypothesized.

An important piece of evidence addresses the occurrence of *Enterobacteriaceae* in the two cheese manufactures: a higher occurrence of this microbial group was seen in control ewes' milk cheese by applying both DNA-based techniques. However, this finding was not supported by viable counts; indeed, at the end of the ripening period, comparable loads of *Enterobacteriaceae* were found in the two cheeses by culturing in selective VRBA medium, thus suggesting the need for an integration between culture-dependent and culture-independent techniques. It is worth remembering that Illumina sequencing provides a relative but not absolute abundances of single microbial taxa, whereas PCR-DGGE allows a qualitative or, at most semi-quantitative, overview of heterogeneous microbial communities.

364

#### 365 **5. Conclusions**

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367 Overall, the combined analytical approach enabled differences and similarities in the bacterial biota and dynamics of the 368 two cheeses under study to be defined. The integration of data from viable counting, Illumina sequencing and PCR-369 DGGE provided the detection of major and minor bacterial components (Illumina sequencing and viable counting), as 370 well as the identification of single bacterial species (PCR-DGGE) that could not be revealed with the first two 371 techniques. The primary result confirmed the association of lactobacilli, lactococci and leuconostocs with cheese and 372 the dairy environment; moreover, the vegetable coagulant was demonstrated to affect the early bacterial dynamics of 373 Caciofiore della Sibilla cheese, and at the end of its maturation, the composition of the bacterial biota significantly 374 differed from the control ewes' milk cheese. 375 376 Acknowledgments 377 The authors thank "Caseificio degli Angeli" (Pievetorina, MC, Italy) for supplying raw milk, curds and cheeses. 378 379 Funding: This study was financially supported by the Regione Marche within the project "Valorizzazione di prodotti 380 alimentari della tradizione marchigiana". 381 382 References 383 384 Alessandria, V., Dolci, P., Rantsiou, K., Pattono, D., Dalmasso, A., Civera, T., Cocolin, L., 2010. Microbiota of the 385 Planalto de Bolona: an artisanal cheese produced in uncommon environmental conditions in the Cape Verde Islands. World J. Microbiol. Biotechnol. 26, 2211-2221. 386 Alfonzo, A., Miceli, C., Nasca, A., Franciosi, E., Ventimiglia, G., Di Gerlando, R., Tuohy, K., Francesca, N., 387 388 Moschetti, G., Settanni, L., 2017. Monitoring of wheat lactic acid bacteria from the field until the first step of 389 dough fermentation. Food Microbiol. 62, 256-269. Ampe, F., Ben Omar, N., Moizan, C., Wacher, C., Guyot, J.P., 1999. Polyphasic study of the spatial distribution of 390 391 microorganisms in Mexican pozol, a fermented maize dough, demonstrates the need for cultivation-392 independent methods to investigate traditional fermentations. Appl. Environ. Microbiol. 65, 5464-5473. 393 Aquilanti, L., Babini, V., Santarelli, S., Osimani, A., Petruzzelli, A., Clementi, F., 2011. Bacterial dynamics in a raw 394 cow's milk Caciotta cheese manufactured with aqueous extract of Cynara cardunculus dried flowers. Lett. 395 Appl. Microbiol. 52, 651-659. 396 Aquilanti, L., Santarelli, S., Babini, V., Osimani, A., Clementi, F., 2013. Quality evaluation and discrimination of semi-397 hard and hard cheeses from the Marche region (Central Italy) using chemometric tools. Int. Dairy J. 29, 42-52 398 Aquilanti, L., Santarelli, S., Babini, V., Osimani, A., Garofalo, C., Polverigiani, S., Clementi, F., 2016. PCR-DGGE for 399 the profiling of cheese bacterial communities: strengths and weaknesses of a poorly explored combined 400 approach. Dairy Sci. Technol. 96, 747-761.

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491 Figure legends

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493 Fig. 1. Results of pH measurements of curd obtained from milk coagulation with commercial rennet, curd obtained
494 from milk coagulation with thistle rennet, *Caciofiore della Sibilla* and control ewes' milk cheese during cheese-making
495 and ripening.

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- 497 \* pH values of curd obtained from milk coagulation with commercial rennet ( $\Diamond$ ) and from milk coagulation with thistle 498 rennet (Δ).
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Fig. 2. Loading and score plot of principal components analysis (PCA) based on viable counts evolution over time of
 *Caciofiore della Sibilla* (dots) and control ewes' milk cheese (triangles).

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- **Fig. 3.** Relative abundances (%) of bacterial groups and vegetable genome identified by MySeq Illumina in raw ewes' milk (rm); curd obtained by milk coagulation with commercial animal rennet (cc); curd obtained by milk coagulation with carline thistle rennet (ct); control raw ewe's milk cheese and *Caciofiore della Sibilla* cheese collected at 1, 3, and 20 days of ripening ( $C_1$ ,  $C_3$ ,  $C_{20}$  and  $Cf_1$ ,  $Cf_3$ ,  $Cf_{20}$  respectively).
- 507 "Other" represents all genera that were unidentified, belonged to uncultured or whose abundance was less than 0.1 %.508
- **Fig. 4.** Relative abundances (%) of the different LAB genera identified by MySeq Illumina in raw ewes' milk (rm); curd obtained by milk coagulation with commercial animal rennet (cc); curd obtained by milk coagulation with carline thistle rennet (ct); control raw ewe's milk cheese and *Caciofiore della Sibilla* cheese collected at 1, 3, and 20 days of ripening ( $C_1$ ,  $C_3$ ,  $C_{20}$  and  $Cf_1$ ,  $Cf_3$ ,  $Cf_{20}$  respectively).
- 513 Each percentage is related to the total amount of LAB found in each sample.
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**Fig. S1.** Bacterial DGGE profiles of the DNA extracted from the bulk of colonies harvested from selected plates and the DNA extracted directly from the samples of raw ewes' milk (rm); curd obtained by milk coagulation with carline thistle rennet (ct) (panel **a**); fresh young leaves of *Carlina acanthifolia* All. (ca); carline thistle rennet (tr) (panel **b**); curd obtained by milk coagulation with commercial animal rennet (cc) (panel **c**). Lanes  $I_H$  and  $I_L$  indicate DNA extracted from the bulk of colonies harvested from the MRS Agar plates spiked with the highest and lowest dilutions, respectively. Lanes  $I_H$  and  $I_L$  indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 22 °C spiked with the highest and lowest dilutions, respectively. Lanes  $II_H$  and  $II_L$  indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 45 °C spiked with the highest and lowest dilutions, respectively. Lane m indicates DNA extracted from the samples. Mix' refers to the mixture of species: a -*Lactobacillus plantarum* DSMZ 2601; b - *Pediococcus pentosaceus* DSMZ 20336<sup>T</sup>; c - *Lactobacillus parabuchneri* DSMZ 5708; d - *Lactobacillus buchneri* DSMZ 20057; e - *Lactobacillus paracasei* NRRL 4560<sup>T</sup>; f - *Lactobacillus casei* NCIMB 4114.

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**Fig. S2**. Bacterial DGGE profiles of the DNA extracted directly from the samples of control raw ewes' milk cheese (batch C) and *Caciofiore della Sibilla* cheese (batch Cf) (panel **a**) and the DNA extracted from the bulk of colonies harvested from selected plates of control raw ewes' milk cheese (panel **b**) and *Caciofiore della Sibilla* cheese (panel **c**) collected at 1, 3, 6, 10 and 20 days of ripening (C<sub>1</sub>, C<sub>3</sub>, C<sub>6</sub>, C<sub>10</sub>, C<sub>20</sub> and Cf<sub>1</sub>, Cf<sub>3</sub>, Cf<sub>6</sub>, Cf<sub>10</sub>, Cf20, respectively).

Lane I indicates DNA extracted from the bulk of colonies harvested from the MRS Agar plates spiked with the highest

dilutions. Lane II indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 22

<sup>534</sup> °C spiked with the highest dilutions. Lane III indicate DNA extracted from the bulk of colonies harvested from the M17

535 Agar plates incubated at 45 °C spiked with the highest dilutions. Mix' refers to the mixture of species: a - Lactobacillus

- 536 plantarum DSMZ 2601; b Pediococcus pentosaceus DSMZ 20336<sup>T</sup>; c Lactobacillus parabuchneri DSMZ 5708; d -
- 537 Lactobacillus buchneri DSMZ 20057; e Lactobacillus paracasei NRRL 4560<sup>T</sup>; f Lactobacillus casei NCIMB 4114.

Table 1 Results of viable counting of bacteria in *Caciofiore della Sibilla* and control ewes' milk cheese.

	<i>Caciofiore della</i> <i>Sibilla</i> cheese	Control cheese		Caciofiore della Sibilla cheese	Control cheese		<i>Caciofiore della</i> <i>Sibilla</i> cheese	Control cheese				
6 1		Lactococci	_	Т	hermophilic cocci		Lactobacilli					
Sample		(M17 22°C)			(M17 45°C)			(MRSA)				
Raw milk	5.00 ±	0.06		4.08	3 ± 0.06		4.78	± 0.02				
Carlina acanthifolia All.	$6.68\pm0.03$			$4.88 \pm 0.04$			$3.54\pm0.10$					
Thistle rennet	$3.13\pm0.14$			$2.10\pm0.28$			$4.01\pm0.00$					
Curd	$4.11 \pm 0.00$	$4.82 \pm 0.12$		$4.67\pm0.00$	$4.28\pm0.02$		$4.90\pm0.06$	$4.74 \pm 0.11$				
1 day of ripening	$5.56 \pm 0.02^{d}$	$8.57\pm0.01^{\rm c}$	<.0001*	$4.68 \pm 0.00^{e}$	$7.50\pm0.06^{\text{b}}$	0.0002*	$5.06\pm0.05^{\rm d}$	$8.57 \pm 0.00^{b}$	0.0001			
3 days of ripening	$7.30 \pm 0.00^{\circ}$	$7.98 \pm 0.04^{d}$	0.0016*	$6.32 \pm 0.01^{d}$	$7.37\pm0.08^{bc}$	0.0028*	$8.08\pm0.00^{\rm c}$	$8.89 \pm 0.01^{ab}$	<.0001			
6 days of ripening	$8.62 \pm 0.13^{b}$	$9.00 \pm 0.14^{b}$	0.1098	$8.26\pm0.00^{\rm b}$	$8.18\pm0.08^{\rm a}$	0.3251	$8.66\pm0.09^{\rm b}$	$9.03\pm0.01^{\rm a}$	0.0312*			
10 days of ripening	$9.12\pm0.18^{\rm a}$	$9.14\pm0.06^{\text{b}}$	0.9113	$8.31\pm0.00^{\rm a}$	$8.00\pm0.02^{\rm a}$	0.0012*	$8.68\pm0.01^{\rm b}$	$8.96\pm0.01^{\rm a}$	0.0009*			
20 days of ripening	$8.90\pm0.05^{ab}$	$9.52\pm0.06^{\rm a}$	0.0079*	$7.42 \pm 0.03^{\circ}$	$7.24 \pm 0.05^{\circ}$	0.0450*	$9.05\pm0.03^{\rm a}$	$8.87 \pm 0.22^{ab}$	0.3557			

	Caciofiore della Sibilla cheese	Control cheese		Caciofiore della Sibilla cheese	Control cheese		Caciofiore della Sibilla cheese			Caciofiore della Sibilla cheese	Control cheese	
Sample	Coag	ulase-negative coc (MSA)	ci	Total	mesophilic aerobe (PCA)	es	(Sla	Enterococci anetz-Bartley Agar)		Ente	erobacteriaceae (VRBGA)	
Raw milk	4.01	± 0.03		5.14	± 0.16		4.30	$\pm 0.06$		3.93 ±	0.06	
Carlina acanthifolia All.	$5.24\pm0.01$			$7.27\pm0.04$			$3.97 \pm 0.16$			$5.34\pm0.03$		
Thistle rennet	$3.06\pm0.03$			$4.07\pm0.04$			$3.60\pm0.01$			$1.00\pm0.00$		
Curd	$5.06\pm0.08$	$4.52\pm0.04$		$4.99\pm0.01$	$5.13 \pm 0.07$		$4.37\pm0.01$	$4.23\pm0.05$		$1.10\pm0.00$	$3.39\pm0.16$	
1 day of ripening	$5.16 \pm 0.02^{d}$	$6.75 \pm 0.10^{\rm e}$	0.0022*	$5.37 \pm 0.02^{e}$	$8.45\pm0.03^{a}$	<.0001*	$4.60\pm0.05^{e}$	$6.09 \pm 0.05^{e}$	0.0011*	$1.74 \pm 0.06^{\rm e}$	$7.55\pm0.05^{\rm a}$	<.0001*
3 days of ripening	$6.34\pm0.02^{\rm c}$	$7.97\pm0.04^{\rm d}$	0.0003*	$8.15\pm0.08^{\rm d}$	$8.87\pm0.02^{\rm a}$	0.0056*	$5.14\pm0.08^{\rm d}$	$7.12 \pm 0.02^{\circ}$	0.0099*	$2.19\pm0.02^{\rm d}$	$7.30 \pm 0.01^{b}$	<.0001*
6 days of ripening	$7.47\pm0.04^{\rm b}$	$8.91\pm0.19^{\rm a}$	0.0086*	$8.58\pm0.00^{\rm c}$	$8.71\pm0.73^{a}$	0.8181	$7.78\pm0.00^{\rm c}$	$7.55 \pm 0.01^{b}$	0.0007*	$5.41\pm0.02^{\rm c}$	$7.13 \pm 0.07$ <sup>b</sup>	0.0009*
10 days of ripening	$8.72\pm0.02^{\rm a}$	$8.40\pm0.03^{\text{b}}$	0.0059*	$8.82\pm0.01^{\text{b}}$	$9.10\pm0.03^{\rm a}$	0.0073*	$8.26\pm0.01^{\text{b}}$	$7.90\pm0.01^{\rm a}$	0.0004*	$6.17\pm0.05^{\rm b}$	$7.07 \pm 0.09^{b}$	0.0073*
20 days of ripening	$7.44\pm0.01^{\text{b}}$	$7.34\pm0.01^{\text{d}}$	0.0074*	$9.20\pm0.02^{\rm a}$	$8.87\pm0.00^{a}$	0.0017*	$8.51\pm0.07^{\rm a}$	$6.94\pm0.04^{d}$	0.0012*	$6.40\pm0.00^{a}$	$6.32\pm0.03^{\:c}$	0.0597

Viable counts of lactococci, thermophilic cocci, lactobacilli, coagulase-negative cocci, total mesophilic aerobes, enterococci and *Enterobacteriaceae* expressed as mean values  $\pm$  st. dev. of samples collected during ripening of *Caciofiore della Sibilla* and control ewes' milk cheese. Mean separation test throughout ripening: different letters on the same column indicate significant differences over time according to Tukey's test ( $\alpha = 0.05$ ). Least significant difference (LSD) is reported. \*Occurrence of significant differences due to cheese batch (C or Cf) according to Student's t test ( $\alpha = 0.05$ ).

Bacterial species	Raw ewes' milk			Carlina acanthifolia All.				Thistle rennet				Curd obtained from milk coagulation with thistle rennet				Curd obtained from milk coagulation with commercial rennet							
	MRS	M17 22°C		М	MR	S	M17 22°C	M17 45°C	M	MR	S	M17 22°C	M17 45°C	М	MRS	M17 22°C	M17 45°(		MRS	5	M17 22°C	M17 45°C	М
	ни			,	н	L	H L			Н	L	H L			H L				H	L	H L		
Acinetobacter baumannii								•	•														
Acinetobacter johnsonii			• •	•																•	• •		
Bacillus cereus												•	•										
Bacillus sp.									• •				• •			•		•					
Bacillus subtilis												•						•					
Enterobacter cloacae			• •																			•	•
Enterobacter hormaechei			• •																			•	•
Enterobacter sp.							• •	•														•	•
Escherichia coli																							•
Kurthia gibsonii																						•	• •
Lactobacillus																							
alimentarius/paralimentarius					•	•									•	• •	•						
group																							
Lactobacillus graminis/curvatus					•	•																	
group					-	-																	
Lactobacillus																							
plantarum/paraplantarum/										•	•				• •	•		•					
pentosus group																							
Lactococcus lactis	• •																		•	•	• •		
Pantoea agglomerans							•																
Pediococcus acidilactici															•	•							
Pseudomonas alcaliphila							•																
Pseudomonas sp.		•	• • •				• •	•	•														
Staphylococcus chromogenes																						•	• •

**Table 2** Bacterial species identified by PCR-DGGE analysis in raw ewe's milk, Carlina acanthifolia All. subsp. acanthifolia, thistle rennet, curd obtained from milk coagulation with thistle rennet and curd obtained from milk coagulation with commercial rennet.

• DGGE bands resulting from the analysis of the DNA extracted from the bulk of colonies harvested from high (H) and low (L) dilution agar plates used for viable counting from selected plates or the DNA extracted directly from samples showing  $\geq$ 97% of similarity with the sequences of the closest relatives found by a BLAST search in the GenBank. M indicate the DNA extracted directly from a cheese samples matrix.

Bacterial species		t <sub>1</sub>				t <sub>3</sub>			t <sub>6</sub>			t <sub>10</sub>				t <sub>20</sub>				
	MRS	M17 22°C	M17 45°C	Μ	MRS	M17 22°C	M17 45°C	М	MRS	M17 22°C	M17 45°C	М	MRS	M17 22°C	M17 45°C	М	MRS	M17 22°C	M17 45°C	М
Bifidobacterium dentium																				-
Citrobacter sp.							-				•				•				•	
Enterobacter hormaechei				•				•												
Enterobacter sp.			-				-													
Enterococcus faecalis	•		•=						•	• =	• =		•	•	-				• =	
Enterococcus faecium	٠	•	•				•			•	•		•	•	•	٠	•		•	٠
Erwinia chrysanthemi																				
Lactobacillus alimentarius/paralimentarius group				•																•
Lactobacillus brevis	٠				٠	٠											•			
Lactobacillus casei/ paracasei/rhamnosus group				٠				٠									•			٠
Lactobacillus delbrueckii	٠	•	•			•			•	•			•	•				•		
Lactobacillus plantarum/ pentosus/paraplantarum																				
group																	• •	•		
Lactococcus lactis	• =	• =	• =			• =			• =	• =	-	• =	• =	• =		• =		• =	-	
Leuconostoc mesenteroides/pseudomesenteroides								٠										•		٠
Staphylococcus sciuri		٠	• =				• =		•=		-			-						
Weissella sp.					•															

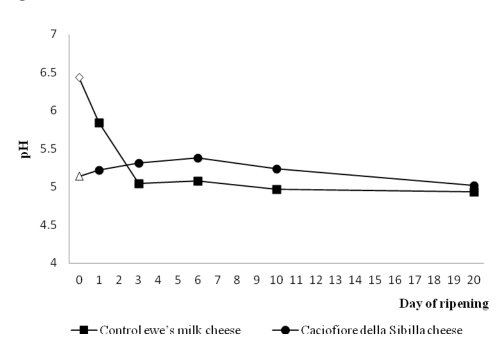
Table 3 Bacterial species identified during the manufacture and ripening of control ewe's milk cheeses and Caciofiore della Sibilla by PCR-DGGE analysis.

• DGGE bands resulting from the analysis of the DNA extracted from the bulk of colonies harvested from selected plates or the DNA extracted directly from *Caciofiore della* Sibilla cheese samples showing  $\geq$ 97% of similarity with the sequences of the closest relatives found by a BLAST search in the GenBank; • DGGE bands resulting from the analysis of the DNA extracted from the bulk of colonies harvested from selected plates or the DNA extracted directly from control ewe's milk cheese samples showing  $\geq$ 97% of similarity with the sequences of the closest relatives found by a BLAST search in the GenBank; • DGGE bands resulting from the analysis of the DNA extracted from the bulk of colonies harvested from selected plates or the DNA extracted directly from control ewe's milk cheese samples showing  $\leq$ 97% of similarity with the sequences of the closest relatives found by a BLAST search in the GenBank;

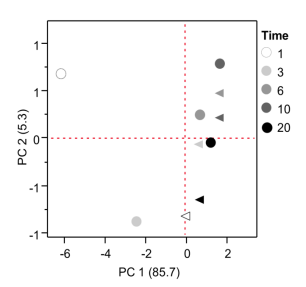
t1: cheese sampled after 1 day of maturation; t3: cheese sampled after 3 days of maturation; t6: cheese sampled after 6 days of maturation; t10: cheese sampled after 10 days of maturation; t20: cheese sampled after 20 days of maturation.

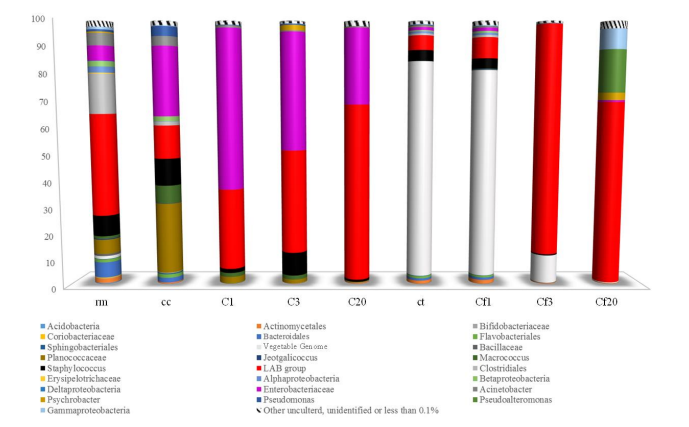
M: DNA extracted directly from a cheese samples matrix.





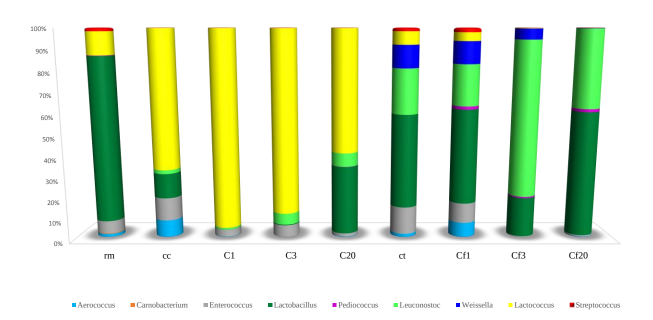






### Fig. 3.

Fig.4.

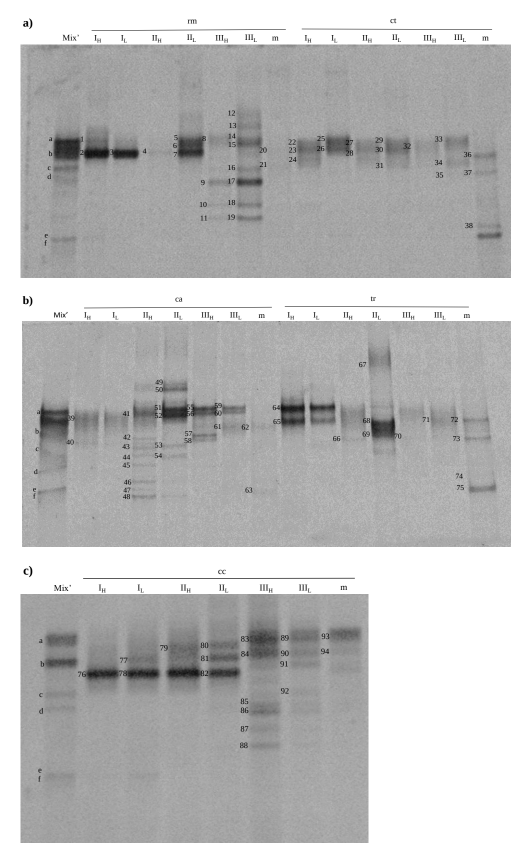


Band(s)	Closest relative	% Ident. <sup>a</sup>	Acc.no. <sup>b</sup>
1, 2, 3, 76, 78, 82, 98, 103, 106, 107, 115, 123, 124,	Lactococcus lactis	100%	KT429894
125, 126, 129, 134, 135, 136, 140, 153, 155, 164,			
168, 178, <b>180, 181</b> , 183, 184, 194, <b>199</b> , 218, 226, 237, 241			
4, 5, 7, 8, 14, 41, 49, 50, 51, 52, 53, 55, 59	Pseudomonas sp.	100%	HF546529
<b>6</b> , 15, <b>21</b> , <b>77</b> , <b>79</b> , <b>80</b> , <b>81</b>	Acinetobacter johnsonii	99%	LN774358
43, 44, 46, 47, 48, 57, 58, 86, 127, 131, 132, 133, 142, 146, 149, 150	Enterobacter sp.	100%	HQ439419
11, 12, 19, 85	Enterobacter cloacae	99%	KP226566
<b>13</b> , <b>20</b> , <b>36</b> , <b>37</b> , <b>38</b> , <b>63</b> , <b>66</b> , <b>72</b> , <b>73</b> , <b>74</b> , <b>75</b> , <b>91</b> , <b>100</b> , <b>102</b> , <b>105</b> , <b>108</b> , <b>114</b> , <b>116</b> , <b>161</b> , 165, 169, <b>137</b> , <b>138</b> , <b>139</b> , <b>182</b> , <b>186</b> , <b>201</b> , <b>253</b>	Failed		
9, 10, 16, 17, 18, 87, 88, 95, 99	Enterobacter hormaechei	98%	HM584024
22, 25, 26, 29, 33, 64, 65, 175, 176, 246, 247, 249,	Lactobacillus plantarum	99%	KJ775808
252	/pentosus		KT634227
	/paraplantarum		KU315098
<b>23, 27,</b> 30, <b>32, 40, 101</b>	Lactobacillus alimentarius	98%	M58804
	/paralimentarius		AB626063
<b>24,</b> 28	Pediococcus acidilactici	99%	KM921945
<b>31, 34, 61, 62, 70,</b> 200	Bacillus sp.	99%	HM755812
35, 69	Bacillus subtilis	98%	GQ392055
39	Lactobacillus graminis	96%	LC063167
	/ curvatus		KP117256
42, 45	Pantoea agglomerans	99%	AB681812
54	Pseudomonas alcaliphila	97%	EU144361
56, 60	Acinetobacter baumannii	100%	AY269241
67, 68,71	Bacillus cereus	100%	KF782833
83, 89, 93	Kurthia gibsonii	99%	KT165384
84, 90, 94	Staphylococcus chromogenes	99%	KR028439
92	Escherichia coli	99%	KC539468
96, 104	Bifidobacterium dentium	100%	AP012326
97	Erwinia chrysanthemi	96%	DQ123809
109, 113, 122, 179	Lactobacillus casei	97%	KM921936
	/paracasei		KT626389
	/rhamnosus		KT626387
110, 111, 112, <b>118</b> , 119, 120, <b>250</b>	Leuconostoc mesenteroides	96%	KR137536
	/pseudomesenteroides		KJ186948
117, 121, 191, 195, <b>196</b> , 202, 206, <b>213</b> , 215, 219, <b>220</b> , 221, <b>222</b> , <b>223</b> , <b>227</b> , <b>228</b> , 229, 230, 231, 233, 234, 235, 238, 239, 242, 243, 244, 245, 254, 255, <b>256</b>	Enterococcus faecium	98%	KM921922
130, 152, <b>157</b> , 163, 167, 171, 187, 192, 204, 207, <b>224</b> , 232, 257	Enterococcus faecalis	98%	KP298396
<b>128</b> , <b>141</b> , <b>143</b> , 144, <b>145</b> , 151, <b>154</b> , 156, 162, 166, <b>170</b> , 177, 197, <b>203</b> , <b>205</b> , 214, 216	Staphylococcus sciuri	99%	JQ511682
147, 148, 158, 159, 160, <b>172,173,174, 185, 188,</b> <b>189</b>	Citrobacter sp.	99%	KP212094
190, 210, 211, <b>248</b>	Lactobacillus brevis	99%	KP221640
<b>193</b> , 198, 208, 212, 217, 225, 236, 240, 251	Lactobacillus delbrueckii	97%	JN969331
209	Weissella sp.	99%	KF598906

**Table S1** Results from the sequencing of the bands cut from the DGGE gels.

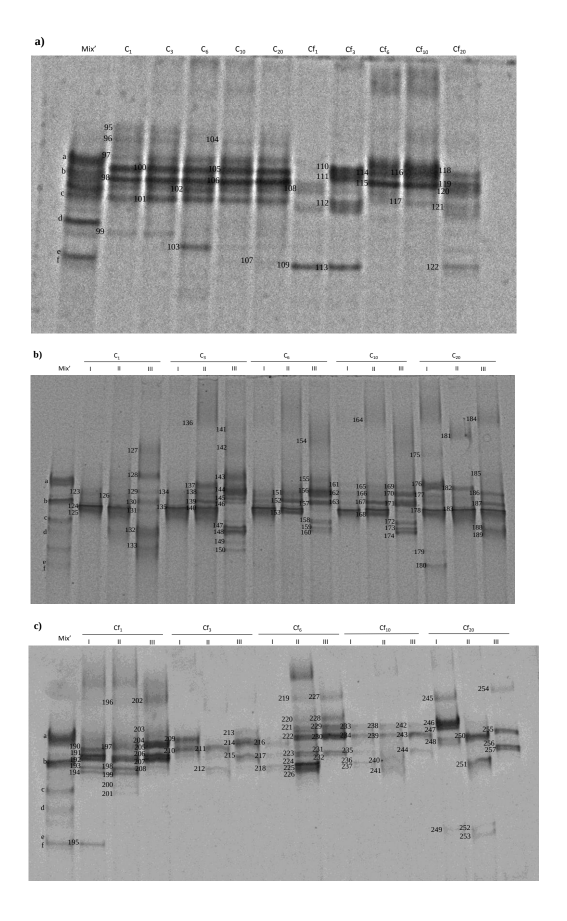
<sup>a</sup> Percentage of identical nucleotides in the sequence obtained from the DGGE band and the sequence of the closest relative found in the GenBank DNA database.
 <sup>b</sup> Accession number of the sequence of the closest relative found by BLAST search.

**Fig. S1**. Bacterial DGGE profiles of the DNA extracted from the bulk of colonies harvested from selected plates and the DNA extracted directly from the samples of raw ewe's milk (rm); curd obtained by milk coagulation with carline thistle rennet (ct) (panel **a**); fresh young leaves of *Carlina acanthifolia* All. (ca); carline thistle rennet (tr) (panel **b**); curd obtained by milk coagulation with commercial animal rennet (cc) (panel **c**).



Lanes  $I_H$  and  $I_L$  indicate DNA extracted from the bulk of colonies harvested from the MRS Agar plates spiked with the highest and lowest dilutions respectively. Lanes  $II_H$  and  $II_L$  indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 22°C spiked with the highest and lowest dilutions respectively. Lanes  $III_H$  and  $III_L$  indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 45°C spiked with the highest and lowest dilutions respectively. Lanes III\_H and III\_L indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 45°C spiked with the highest and lowest dilutions respectively. Lane m indicate DNA extracted from the samples. Mix' refers to the mixture of species: a - *Lactobacillus plantarum* DSMZ 2601; b - *Pediococcus pentosaceus* DSMZ 20336<sup>T</sup>; c - *Lactobacillus parabuchneri* DSMZ 5708; d - *Lactobacillus buchneri* DSMZ 20057; e - *Lactobacillus paracasei* NRRL 4560<sup>T</sup>; f - *Lactobacillus casei* NCIMB 4114.

**Fig. S2.** Bacterial DGGE profiles of the DNA extracted directly from the samples of control raw ewe's milk cheese (batch C) and *Caciofiore della Sibilla* cheese (batch Cf) (panel **a**) and the DNA extracted from the bulk of colonies harvested from selected plates of control raw ewe's milk cheese (panel **b**) and *Caciofiore della Sibilla* cheese (panel **c**) collected at 1, 3, 6, 10 and 20 days of ripening (C<sub>1</sub>, C<sub>3</sub>, C<sub>6</sub>, C<sub>10</sub>, C<sub>20</sub> and Cf<sub>1</sub>, Cf<sub>3</sub>, Cf<sub>6</sub>, Cf<sub>10</sub>, Cf<sub>20</sub> respectively).



Lane I indicate DNA extracted from the bulk of colonies harvested from the MRS Agar plates spiked with the highest dilutions. Lane II indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 22°C spiked with the highest dilutions. Lane III indicate DNA extracted from the bulk of colonies harvested from the M17 Agar plates incubated at 45°C spiked with the highest dilutions. Mix' refers to the mixture of species: a - *Lactobacillus plantarum* DSMZ 2601; b - *Pediococcus pentosaceus* DSMZ 20336<sup>T</sup>; c - *Lactobacillus parabuchneri* DSMZ 5708; d - *Lactobacillus buchneri* DSMZ 20057; e - *Lactobacillus paracasei* NRRL 4560<sup>T</sup>; f - *Lactobacillus casei* NCIMB 4114.