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Microbial Communities of Raw Milk Cheeses, A Review

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Microbial communities play a fundamental role in shaping the taste, aroma, and texture of cheeses. They consist of starter and secondary microorganisms. Starters contribute to acid development during cheesemaking, while secondary microbiota play a crucial role in the ripening process. Their diversity is a subject of significant importance, shaped by various factors such as the cheesemaking environment, employed starters, physicochemical conditions, and manufacturing procedures. In this review, we attempted to provide an accurate picture of the microbial communities commonly found in raw milk cheeses and tried to list their origins, factors influencing their existence, and the approaches used for their screening. The research employed information retrieval methods, mainly focusing on specific keywords. We systematically searched various databases for relevant articles and reviews, prioritizing the retrieval of the most recent publications and those deemed most relevant to the objectives of this review. This review disclosed the frequently identified bacterial genera in cheeses, such as *Lactococcus* and *Lactobacillus*. In terms of fungi, regularly isolated species included *Candida*, *Kluyveromyces*, *Saccharomyces*, *Yarrowia*, *Goetrichum*, among others. Our investigation enabled the unveiling of both the core microbiota shared across diverse cheeses, crucial for cheese fermentation and ripening, and the variable microbiota contributing to the diversity in cheese characteristics.

4 ABSTRACT

1 - Introduction

The story says that curd formation was accidentally discovered when milk was kept in bags made from animal stomachs. Hence, the idea of milk preservation through curdling emerged [1]. What initially began as milk conservation has transformed into an impressive variety of cheeses, each possessing its unique cheesemaking process, texture, flavor, and shape. Consequently, the classification systems developed for categorizing cheeses are diverse and sometimes confusing. To date, no classification scheme has been universally recognized as complete and satisfactory [2].

 Today, there are over 4,000 varieties of cheese produced worldwide [3], with the majority manufactured in accordance with strict guidelines to ensure the quality of the final product. In 2015, global cheese production reached approximately 20 million tonnes, marking a 23% increase from 2005 to 2015 [4].

 Obtaining various types of cheese, even when similar ingredients and manufacturing procedures are employed, raises questions about this enigmatic product. Scientific evidence supports the notion that cheese is a living system, and its variability stems from fermentation and ripening steps guided by a multitude of cheese microorganisms [5]. Understanding these primary actors in the cheese production process is crucial for a more precise appraisal and the enhancement of manufacturing conditions. The primary objective of this study was to compile the most recent, albeit not exhaustive, findings on the microbiota of raw milk cheese.

 This review is divided into five main sections. We started by giving an overview of raw milk cheeses. Then, we listed the main sources of their indigenous microorganisms; we discussed some of the factors that influence their presence as well as the approaches that researchers use to screen them, including culture -dependent and culture -independent tools. Finally, we used research examples to summarize recent insights into cheese microbial populations. This review could provide valuable insights for novice researchers interested in cheese microbiology.

 The information retrieval methods utilized in this research predominantly involved the use of specific keywords, including "raw milk," "artisanal cheese," "raw milk cheese," "metagenomics," "high throughput sequencing," and "cheese biodiversity." Various databases, such as the Web of Science, Science Direct, Pubmed and ResearchGate were systematically searched for relevant articles and reviews. The selection process prioritized the retrieval of the most recent publications and those deemed most relevant to the objectives of this review. The extensive reading of numerous articles and reviews aimed to extract the most pertinent information for a comprehensive understanding of the subject matter.

2- LITERATURE

2.1. Raw versus pasteurized milk cheeses

Raw milk, as defined by the code of hygienic practice for milk and milk products (CAC/RCP 57 -2004), is "*milk which has not been heated beyond 40°C or undergone any treatment that has an equivalent effect*." In contrast, pasteurized milk is defined as milk subjected to a heat treatment to reduce the number of any potential pathogenic microorganisms to a level at which they do not pose a significant health hazard. Pasteurization is commonly conducted at 72°C for 15 seconds [6].

 Recently, cheeses crafted from raw milk have garnered renewed interest, primarily owing to their authentic taste, aromas, and pronounced flavors. The distinctiveness of raw milk cheeses arises from the rich and diverse microbial populations naturally present in the raw milk, including primary and secondary cultures [5].

 Nevertheless, the consumption of raw milk products is not without consequences and may pose serious impacts on public health. This is why public health authorities mandate a ripening period of at least 60 days for raw milk or raw milk cheese that has not undergone pasteurization [7]. It is widely acknowledged that pasteurization eliminates bacteria from raw milk, but it is also recognized that it eradicates microbial entities responsible for flavor and health benefits. For this reason, cheese connoisseurs often refer to pasteurized cheeses as "*dead cheeses* " [8] **.**

 Ensuring the production of a cheese that is both safe and flavorful is among the primary challenges faced by cheesemakers. To address this, some producers use selected starters isolated from high -quality raw milk cheeses to "*restitute*" the natural taste.

2.2 Microbial origin of raw milk cheeses

 While a growing body of research explores the intricacies of cheese microbiota, few studies have delved into its origin. Cheese harbors several types of microbiota, including bacteria, yeasts, and molds, coexisting at different stages of production —both during manufacturing and ripening—forming dynamic and complex ecosystem. These microorganisms predominantly originate from milk (a key ingredient in cheesemaking) and the cheese -making environment.

2.2.1. Origins of milk microbiota

 Milk provides an ideal environment for a diverse microbiota, owing to its rich composition of fats, proteins, sugars, vitamins, and minerals. Several factors contribute to this microbial diversity, including the microbiota found on animals' teats, the overall farm environment, interactions with dairy workers, and the dairy equipment. For example, the teat skin microbiota demonstrates significant biodiversity, consisting of 66 different species from genera such as *Enterococcus* , *Pediococcus* , *Enterobacter* , *Pantoea* , *Aerococcus* and *Staphylococcus* —commonly

found in raw milk. Many of these taxa have been identified as key contributors to flavor development during the cheese ripening process [9]. Stable air is also recognized as a significant source of contamination. Vacheyrou et al. [10] demonstrated this by examining the microbial communities in stable air alongside those in raw milk, revealing a notable transfer of microbiota. Additionally, various environmental factors, including soil and silage, can influence the microbial composition of milk. This was demonstrated in a study focusing on the mesophilic lactic acid bacteria (LAB) of milk. Indeed, *Staphylococcus warneri* and *Staphylococcus* sp. were identified in soil, silage, and milk suggesting a potential transfer of these strains from the farm ecosystem to the milk [11]. Moreover, research has shown that hygienic practices wield significant influence over the microbial diversity of raw milk. Under stringent hygiene conditions, milk predominantly features Corynebacteriaceae and Micrococcaceae. Conversely, when produced under less stringent hygienic conditions, milk reveals the presence of diverse taxa, including gram -negative bacteria, *Lactococcus lactis*, *Brevibacterium linens*, and *Leuconostoc mesenteroides* species. This illustrates the direct impact of farming practices on the microbial communities present in raw milk [12]. Milk equipment also plays a significant role. In a study conducted by Didienne et al. [13], it was found that wooden vats significantly enriched the microbial populations in stored milk. Interestingly, deliberate inoculation of milk with pathogens such as *Listeria monocytogenes*, *Salmonella*, or *Staphylococcus aureus* within the same research had no apparent effect on the biodiversity of the vats. The vats remained uncontaminated and uncolonized.

2.2.3. Origins of cheese microbiota

 The microbial composition of cheese is primarily influenced by the indigenous microbiota of milk. However, the cheesemaking environment, including surfaces, curd tanks, molding machines, and ripening Safae Azzouz et al. **Microbial Communities of Raw** ...

rooms, also exerts a significant impact. Furthermore, the inclusion of starter cultures, such as whey, plays a crucial role in shaping the microbiota of cheese. Kamimura et al. [14] investigated microbial diversity in samples from the cheesemaking environment, cheese ingredients, and ripened Serra da Canastra cheese (21 days) across three different farms in Minas Gerais, Brazil. Among its key findings, the study indicates a strong link between the microbiota found in 'pingo' (a natural fermented whey) used as a starter culture and the ripened cheese. In a separate study, Quijada et al. [15] examined the bacterial communities of the Vorarlberger Bergkäse (VB) cheese rind and different processing surfaces in the ripening cellars. Certain bacterial groups, such as coryneforms, *Staphylococcus equorum*, and *Halomonas*, were found both in the cheese and on most environmental surfaces, suggesting their transfer to the cheese surface during ripening. Additionally, **Calasso et al.** [16] conducted another study revealing the presence of specific house microbiota, such as *Lactobacillus plantarum*, in the rind of Caciocavallo Pugliese cheese, the curd tank, and molding machines. Furthermore, other bacteria were detected in both the ripening rooms and the rinds of Caciotta and Caciocavallo Pugliese cheeses. In a comparative study exploring different production methods, primarily distinguished by the type of equipment used during cheese making (stainless steel and wooden equipment), it was revealed that particular strains within the *Enterococcus* spp. group including *Enterococcus faecalis*, *Enterococcus casseliflavus*, and *Enterococcus gallinarum* originated from wooden vats and persisted during the ripening of Caciocavallo Palermitano cheese. In contrast, these strains were not present in the cheese made by the standard method [17].

2.3. Factors influencing cheese microbiota

The growth of microbial communities can be either positively or negatively influenced throughout the cheesemaking process. Several

factors come into play, including the pasteurization and refrigeration of milk, the temperature of curd cooking, pH levels, salt content, the season of cheesemaking, and various other parameters. While investigating the impact of milk refrigeration on cheese microbiota, it was found that towards the end of the ripening process for Serra de Estrela cheese, non -refrigerated milk cheeses exhibited elevated counts of Enterobacteriaceae and diminished counts of LAB in comparison to refrigerated milk cheeses [18]. In a separate study examining the effect of salt on starter viability, strains of *L*. *lactis* subsp. *cremoris*, *L*. *lactis* subsp. *lactis* biovar diacetylactis, and *L*. *lactis* subsp. *lactis* isolated from DL -starter commercial cultures underwent viability tests using varying NaCl concentrations. The results revealed strain-dependent outcomes but indicated a negative correlation between viability and elevated NaCl concentrations for all strains [19]. The growth of certain strains can also be influenced by the cooking temperature. In a study involving a semi -hard cheese with *Streptococcus thermophilus* as a starter, the levels of this strain were observed to be impacted by the cooking temperature during production. Notably, elevating the cooking temperature from 47° C or 50° C to 53° C resulted in a reduction in the mean viable cell numbers of *S*. *thermophilus* strains during various manufacturing stages, particularly during the transition from maximum scald temperature to pre -press and brining steps. Conversely, it was observed that the cooking temperature had no discernible effect on the viability of *Lactobacillus helveticus* and other non -starter lactic acid bacteria (NSLAB) strains [20]. The pH of whey during drainage also plays a pivotal role in influencing the microbial composition of cheese. In a specific study, the reduction in whey pH during the drainage process of mozzarella cheese was linked to higher counts of mesophilic and thermophilic lactobacilli during the subsequent refrigeration phase of the cheese [21]. The season of cheese manufacturing may also exert an influence on cheese microbiota. Artisanal cheeses collected

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during the dry and rainy seasons exhibited distinct microbial patterns. In the dry season, cheeses were predominantly characterized by the presence of *Enterococcus* and *Lactobacillus*, while in the rainy season, there was a prevalence of *Lactococcus* and *Weissella* genera [22]. The composition of the starter culture also plays a crucial role in shaping the bacterial growth of naturally occurring bacteria in cheese. The investigation into the microbiology of a 90 -day ripened Graviera cheese delved into the impact of natural starter culture (NSC) and commercial starter culture (CSC). The study revealed that nonstarter bacteria, specifically *Lactobacillus paracasei* and *Lactobacillus plantarum*, prevailed in cheeses produced with commercial starters. In contrast, cheeses crafted with natural starters predominantly featured indigenous strains of *Enterococcus faecium* and *Enterococcus durans* [23].

2.4. Investigation of the cheese microbiota by Culture -dependent and culture -independent techniques

 With the advent of molecular methods, researchers have attained a more comprehensive understanding of microbial communities in cheese. Numerous molecular tools are currently at their disposal, and these can be categorized into two types: culture dependent and culture -independent methods. The former necessitates an isolation and culturing step, while the latter omits this requirement, enabling the analysis of extracted DNA/RNA directly from the matrix of interest.

2.4.1 Culture -dependent methods

 Randomly Amplified Polymorphic DNA (RAPD) is a PCR -based method that employs single small primers and low annealing temperatures. This generates multiple bands, usually 3 –12 bands of varying sizes, providing distinct profiles for each strain. The method is quick and easy to operate, but its major drawbacks include poor reproducibility

and a lack of standardization among laboratories [24].

Repetitive Sequence-Based PCR (rep -PCR) technique amplifies genomic regions between repetitive sequences using primers like REP, ERIC, or BOX. Each species produces unique patterns with bands of varying sizes, enabling differentiation from others [25].

 16S rRNA gene sequencing, a widely used technique, allows species identification by providing information about the partial or complete nucleotide sequence of the 16S rRNA gene [26]. The obtained sequence is compared with a database, such as NCBI, containing numerous available sequences to determine closely related species.

2.4.2. Culture -Independent methods

 Among PCR -based culture -independent methods, examples include **Temporal Temperature Gel Electrophoresis (TTGE)** and **Denaturant Gradient Gel Electrophoresis (DGGE)**. TTGE employs a constant temperature and denaturant, while DGGE utilizes an increasing gradient of chemical denaturant (urea and formamide) along with a constant temperature. This allows the separation of amplicons of similar length but with dissimilarities in nucleotide sequence. For species identification, the bands obtained are either compared with co -migrated reference strain amplicons or excised from the gel for sequencing [27].

 Culture -independent approaches also include the single conformation polymorphism (SSCP) method. In this technique, the sequence signature governs the single -stranded DNA conformation [28].

 High -throughput sequencing (HTS) is increasingly utilized and gaining popularity for analyzing microbes in food samples. This technique can be used in three ways: by targeting a specific sequence, mainly the 16S rRNA gene (amplicon sequencing), analyzing non -targeted sequences (shotgun metagenomics), or by investigating the total expressed genes in a sample (metatranscriptomics, RNASeq) [29].

2.5. Current state -of-knowledge on cheese microbial communities

2.5.1. Fresh unripened cheeses

Table 1

The microbial composition of fresh unripened cheeses is detailed in Table 1. In the variety of freshly made unripened cheeses, Ranchero cheese underwent a thorough examination of its LAB, revealing a total of 172 isolated species. Within this diverse spectrum, 42.5% were identified as *Lactococcus* spp., 41.8% as *Enterococcus* spp., and 8.1% as *Lactobacillus* spp. When categorizing *Lactococcus* spp. at the species level, *L*. *lactis* subsp. *lactis*, succeeded by *L*. *garvieae*, emerged as the most prevalent species [30]**.** El Galiou et al. [31] explored the microbiota of Jben cheese made with goat milk by using different cheesemaking procedures. In the cheese produced using the traditional cheesemaking method involving calf rennet and uncooled milk, the following genera were identified: *Enterococcus*, *Lactococcus*, *Lactobacillus* and *Listeria*. An investigation into yeast communities in artisanal cheeses from various regions in Morocco revealed the presence of 18 species distributed in a cheese dependent manner. The most common species included *Kluyveromyces lactis, Saccharomyces cerevisiae, Yarrowia lipolytica, Candida parapsilosis, Kazachstania unispora, Kluyveromyces marxianus and Pichia fermentans* [32] **.** In another study, LAB were isolated from Oaxaca cheese at various stages of processing and sequenced using the 16S rDNA gene. The main species found in ready made Oaxaca cheese were *E. faecalis,* followed by *L. lactis* subsp. *lactis*, *E. faecium*, *Lb. plantarum*, *Lb. paracasei* subsp. *paracasei and* finally *Lactobacillus rhamnosus*. Additionally, Rep -PCR analysis showed high molecular diversity within *L. lactis* species [33] **.** In a comprehensive examination of Buryatian

cottage cheese, PacBio sequencing identified seven distinct phyla in samples from various regions. Notably, Firmicutes (73.57%) and Proteobacteria (26.27%) emerged as the predominant phyla. Both culture -dependent and culture -independent methods revealed the presence of diverse genera, including *Lactobacillus, Lactococcus, Leuconostoc, Pediococcus, Streptococcus, Acetobacter, Klebsiella, Acinetobacter, Pseudomonas, Raoutella* among others. Remarkably, the most abundant genera observed were *Lactococcus* spp. and *Streptococcus* spp. [34] **.**

Mendez Romero et al. [35], investigated the bacterial composition of 36 samples of fresco cheese from Sonora, Mexico. According to HTS results, the most representative phylum was Firmicutes, followed by Proteobacteria. At the family level, Streptococcaceae dominated with 52.87%, followed by Lactobacillaceae (9.61%) and Leuconostocaceae (7.19%). The representative genera included *Lactococcus* with a relative abundance of 28.22%, followed by *Streptococcus* (16.08%), *Lactobacillus* (8.03%), and *Leuconostoc* with a percentage of 6.82%. The Venn diagram facilitated the identification of shared genera, namely *Lactococcus* spp., *Streptococcus* spp., *Lactobacillus* spp., and *Leuconostoc* spp. PCA analysis indicated that the dominant genera are *Lactococcus* spp., *Streptococcus* spp., and *Leuconostoc* spp.

2.5.2. Soft cheeses

Table 2

 The microorganisms found in soft cheeses are presented in table 2. Minas cheese, sourced from distinct geographical regions (Serro, Canastra, Araxá, Serra do Salitre, and Campo das Vertentes), underwent analysis using both culture -dependent methods (sequencing of LAB isolates) and independent methods (LH -PCR, Rep -PCR). Ten genera were discerned, with *Lactobacillus* emerging as the most prevalent. Additional taxa included *Lactococcus*, *Enterococcus*, *Weissella*,

Pediococcus , *Leuconostoc*, *Streptococcus*, *Escherichia*, *Kocuria*, and *Staphylococcus*. Distinct microbial patterns were observed in the cheeses, suggesting a dependence on the cheese farm and geographical region. Furthermore, four species were exclusively found in specific cheeses; *Lactobacillus delbrueckii* in Canastra cheese, *Kocuria kristinae* in Serra do Salitre cheese, and (*L*. *garvieae*, *Lactobacillus acidipiscis*) in Campo das Vertentes cheese [36]. Three white pickled cheeses collected at various ripening stages from two distinct locations (L1) and (L2), underwent yeast and mold content analysis. All of the cheeses were dominated by *Debaryomyces hansenni* and *Candida zeylanoides.* Only cheese from location two was found to have *Y. lipolytica* and *Galactomyces goetrichum. K. lactis* was detected in the three cheeses. No mold was observed in any of the cheese samples [37].Tomme d'Orchies cheese was evaluated for its surface and core microbiota. For ripened cheese (21 days), *Streptococcus* spp*.* and *Lactococcus* spp*.* were the most abundant genera in the cheese's core while *Corynebacterium* spp*.* and *Brevibacterium* spp*.* were the least abundant. Notably, bacterial diversity was more pronounced on the cheese's surface compared to the core. Major taxa identified in the ripened cheese included *Corynebacterium* spp. and *Psychrobacter* spp. [38].

 Metagenomic analysis uncovered a notable fungal diversity, particularly yeast, in Tomme d'Orchies cheese. Similar to a prior study (84), the surface microbiota exhibited greater diversity compared to the core microbiota. The prevalent operational taxonomic units (OTUs) in the cheese were *Y. lipolytica* and *G. goetrichum.* Despite its use as a starter, *D. hansenii* exhibited low percentages in both the core (0.02% -0.08%) and on the surface (0.1%). Mold genera identified included *Aspergillus, Lewia, Nectria, Myrothecium, Cladosporium and Mucor.* The core was dominated by *A. niger* and *Lewia infectoria,* while the surface was dominated by

A. niger and C. cladosporioides [39] *.* Exploration of the bacterial and fungal populations of Portuguese Queijo de Azeitao PDO cheese was conducted through 16S and 26S rRNA gene amplicon sequencing, respectively. Nine 20 -day ripened samples were collected from three different producers. Results revealed tha*t L. mesenteroides* and *L. lactis* were the most representative bacteria, followed by other species such as *L. zeae*, *L. kefiri*. Regarding fungal population, the cheeses were characterized by the dominance of *Y. lipolytica* followed by *C. ethanolica*, *K. zeylanoides*, *G. candidum*, *G. goetrichum* and others [40] .

2.5.3. Semi -hard cheeses

Table 3

The microbial composition of semi -hard cheeses is depicted in Table 3. A metagenomic analysis of Serra da Canastra cheese involved sequencing the V3 -V4 region of the 16S rRNA gene. The microbial content of the cheese remained consistent across farms, despite variations in whey and milk. Major bacterial players included Firmicutes, Proteobacteria, Actinobacteria, and Bacteroidetes. The relative abundance of Firmicutes exceeded 86% regardless of farm origin. It was also observed that the bacterial composition of the final product was influenced by the Non -Starter Cultures (NSC) used, as identical microorganisms (*Streptococcus* spp. and *Lactococcus* spp.) were found in both [14]. The bacterial composition of Paipa cheese from different producers was examined through 16S rRNA gene sequencing, revealing producer dependent variations. The predominant phyla included Firmicutes with the highest percentages ranging from 59.2% to 82%, followed by Proteobacteria, Actinobacteria, and Bacteroidetes. Lactic Acid Bacteria constituted a significant component, primarily characterized by the *Lactococcus* genus. The *Enterococcus* genus was consistently identified in all samples. Apart from LAB, spoilage bacteria, pathogens, and potentially toxin -

producing bacteria were also detected [41]. Higher bacterial differences were identified in a study comparing Serpa cheese from PDO and non industries. Microflora was investigated using both culture -dependent and culture -independent methods. *Lb*. *brevis* was prominently present in non -PDO cheeses, while *Lb*. *paracasei* /*casei* dominated in PDO cheeses. High -throughput sequencing confirmed the presence of LAB, with the dominant genera being *Lactococcus* spp., *Lactobacillus* spp., and *Leuconostoc* spp. Enterobacteria were detected at the end of the ripening process [42]. High throughput sequencing was employed to examine the microbiota of Gouda cheese. The primary genera identified were *Lactococcus* spp. and *Leuconostoc* spp., maintaining prevalence throughout the ripening period from the 6th to the 24th week. In fact, these genera constituted the starter composition added to the milk. Species belonging to the coliform group exhibited a gradual decrease during ripening, reaching negligible levels in matured cheese [43].

 The bacterial communities of 120 -day Raclette du Valais PDO cheeses were examined through 16S rRNA gene amplicon sequencing. All 21 analyzed samples contained *L. paracasei* , *L. lactis* , *L. helveticus* and *S. thermophilus*. More than 80% of the samples exhibited the presence of ten distinct species, all belonging to the Streptococcaceae or Lactobacillaceae families [44]. Thirteen Van Herby cheese samples from diverse Turkish regions showed varied results based on enrichment (OP) or non -enrichment (KOP). In KOP samples, *Lactobacillus*, *Lactococcus*, and *Streptococcus* prevailed, while OP samples displayed dominance by *Enterococcus*, *Streptococcus*, and *Bacillus*. At the species level, KOP had *L*. *raffinolactis* and *Streptococcus salivarius* prevalence, while enrichment led to *E*. *faecalis* and *S*. *salivarius* dominance. Despite differences, Firmicutes was the most abundant phylum in both KOP and OP samples [45]. Table 3 lists the most prevalent genera.

2.5.4. Hard cheeses

Table 4

 Table 4 provides a comprehensive list of microbiota in various hard cheese varieties. A study comparing Silter cheese with and without an autochthonous starter culture unveiled that predominant species in the 200 day cheese belonged to the *Streptococcus* spp. and *Lactococcus* spp. genera. Some strains from these genera were employed as autochthonous starter cultures. Additionally, other genera were identified, including *Lactobacillus* spp., *Leuconostoc* spp., and *Enterococcus* spp. [46].

De Pasquale et al. [47] investigated metabolically active bacteria in Pecorino Siciliano cheese using pyrosequencing. The most abundant genera were *Lactobacillus* spp. and *Streptococcus* spp. Additionally, sub dominant genera, each constituting less than 0.4%, were identified, including *Planococcus, Alkalibacterium, Enterococcus, L. and* $Peptostreptococcus.$ Culture-independent methods were employed to analyze Istrian cheese from two farms at various ripening stages. Microbial populations from both farms are summarized in Table 4. In the 90 -day cheese from farm 1, *L. lactis* (86.47%) predominated, followed by the *Enterococcus* spp. (7.05%) genus. Conversely, in cheese from farm 2, *Enterococcus* spp. (42.65%) and *Streptococcus parauberis* (40.33%) were the most prevalent. Detection of *E. coli* / *S*. *flexneri* and *Salmonella* spp. exclusively occurred in farm 2 cheeses, with high percentages in fresh cheeses: 58.98% for *E*. *coli* / *S*. *flexneri* and 20.44% for *Salmonella* spp. [48] **.** The microbiota of Alberquilla cheese was investigated using both culture -dependent (RAPD -PCR, species -specific or 16S rRNA gene sequencing) and culture -independent methods (PCR -TTGE and 16S gene sequencing for TTGE bands). Various isolates were categorized into different species, with the majority belonging to the *Lactobacillus* genus, particularly *Lb*. *paracasei*. Gram -negative

bacteria, primarily *H*. *alvei*, were also well represented. Direct DNA analysis via the TTGE approach revealed *L*. *lactis* and *Enterococcus* spp. as the most common species. Additionally, culture-dependent approaches exclusively detected *L*. *pseudomesenteroides* and *P*. *urinaequi* [49] **.** A metagenomic study examined four samples of Bulgarian green cheese for their microbial content. The dominant phylum was Firmicutes (50.61%), followed by Actinobacteria (42.96%) and Proteobacteria (6.43%). Representative genera included *Streptococcus* spp., *Lactobacillus* spp., *Lactococcus* spp., *Staphylococcus* spp., *Brevibacterium* spp., and *Corynebacterium* spp. In the fungal population, species belonging to Ascomycota were prevalent, including *P* . *roqueforti*, *S* . *flava*, *D*. *hansenii*, *P*. *membranifaciens*, and others [50] .

3 - CONCLUSIONS

 While not exhaustive, the data and examples presented in this review provide a comprehensive overview of the microbial communities present in raw milk cheeses, allowing for an assessment of their origins and the factors influencing their growth and presence. Based on the current knowledge, it can be inferred that raw milk cheeses serve as significant reservoirs of microbial populations, primarily originating from raw milk and Natural Whey Starters (NSW) used in the cheesemaking process. These ingredients themselves undergo inoculation from the environment, equipment, and farmers. Additionally, contamination can occur during various stages of cheesemaking and ripening. Procedures like pasteurization, milk refrigeration, curd cooking, salting, and ripening exert direct influences on microbial growth during cheesemaking.

 It is crucial to note that the outcomes derived from microbial screening heavily rely on the methodology employed; culture dependent methods identify viable and dominant cells, while culture -independent techniques such as HTS offer an overview of the existing microbiota, typically at the genus level and without distinguishing between dead and live cells. Therefore, the combination of both approaches is of utmost importance.

Based on the aforementioned findings, it is evident that microbial communities vary depending on the type of cheese under study. Nevertheless, the most commonly identified taxa in cheeses include Firmicutes, *Lactococcus* spp., and *Lactobacillus* spp. In terms of fungi, the frequently isolated species belong to *Candida*, *Kluyveromyces*, *Saccharomyces*, *Yarrowia*, *Goetrichum*, and others.

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Table 1. Microbiota of fresh unripened cheeses

*: the starter culture was not added

**: The authors did not mention whether they used the starter or not.

*: the starter culture was not added

**: culture independent

Table 3. The microbial composition of semi-hard cheeses

*: the starter culture was not added

**: Culture independent

Table 4. Detailed list of microbiota belonging to different varieties of hard cheeses

