Presence and persistence of *Pseudomonas* sp. during Caspian Sea-style spontaneous milk fermentation highlights the importance of safety and regulatory concerns for traditional and ethnic foods

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**Abstract**

The aim of this study was to evaluate the performance of Caspian Sea-style spontaneous milk fermentation to improve the quality of pasteurized milk containing high levels of *Pseudomonas* contamination, with a focus on microbiological safety and stability of the final product. Bacterial diversity of pasteurized milk, fermentation process, and after 60 days of storage was analyzed by Illumina-based sequencing, and presence of viable taxa was confirmed by culturing on selective media. Low quality pasteurized milk harbored mainly Gram-negative bacteria, markedly dominated by *Pseudomonas*. Following fermentation, lactic acid bacteria rapidly became dominant with maximum population of 10.15 log CFU/mL at 18 h, represented mainly by *Lactococcus*. However, sequences related to *Pseudomonas*, and to a lesser extent for enterobacteria, remained constant throughout the fermentation process. The cultured-dependent approach confirmed the presence of viable *Pseudomonas*, with a final population of 5.60 log CFU/mL. Biochemical transformations were further analyzed, indicating lactic acid as the main end-metabolite produced (maximum concentration of 5.93 g/L at 24 h). In addition, the increase of 2-nonanone can be correlated as a volatile biomarker of *P. aeruginosa* and related species. Altogether, the results demonstrated that natural milk fermentation may often not inhibit the development of pathogens and food spoilage microorganisms.

**Keywords:** fermented milk; food-borne microorganisms; lactic acid fermentation; food safety; probiotic.

**Practical Application:** Food safety authorities in Brazil need to conduct rigorous surveillance of fermented milk products.

1 Introduction

The emergence of dairying was a critical step in early agriculture, with considerable importance in the human diet (Panesar, 2011). As a rich nutritional source for microbial growth, prehistoric farmers used lactic acid fermentation to prolong the shelf life of milk (Carrer et al., 2016). The finding of abundant milk residues in pottery vessels from seventh–millennium sites from north-western Anatolia provided the earliest evidence of milk processing (Salque et al., 2013). Until now, fermented dairy products have been a vital component in the daily diet of ethnic groups all around the world, and play an important nutritional role in modern life (Granato et al., 2010).

The popularity and the availability of fermented dairy products (e.g., kefir, koumiss, curd, lassi, laben, and Suero costeño) have been increased throughout the world due to their functional properties and prolonged shelf-life, given by the dynamics of the microbial community living there (Grandos Conde et al., 2013; Panesar, 2011; Singh & Shah, 2017). The fermentation is based on the indigenous microbiota present in the raw material or using part of a successful fermentation as back-slopping in order to ensure the dominance of the original microbiota (Capozzi et al., 2012; Pereira et al., 2020; Capozzi et al., 2020). Caspian Sea-style spontaneously fermented milk is widespread as an traditional product. It is usually produced by natural fermentation (12-24 h) of raw cow’s milk at ambient temperature (approximately 25 °C) (Kiryu et al., 2009). The finished product has a highly viscous consistency with a pleasant acid taste, due to the presence of *Lactococcus lactis* ssp. cremoris and, to a lesser extent, *Leuconostoc* sp., *Gluconobacter* sp., and *Acetobacter orientalis* (Ishida et al., 2005; Uchida et al., 2009). In Brazil, a similar fermented milk is produced by different families (private households) that believe the “mother inoculum” is originated from Caucasus region, so-called Caspian Sea-style fermented milk. However, there are no studies of this traditional product circulated in Brazil.

Although natural fermented milk products harbored various beneficial microorganisms, they are susceptible to contamination due to the conduct of fermentation in open systems and poor microbiological quality of the raw material (Capozzi et al., 2017). *Pseudomonas* spp. is a relevant contaminant for fermented milk products (Del Olmo et al., 2018; Reichler et al., 2018; Scatamburlo et al., 2015). Due to their high metabolic versatility, *Pseudomonas* are able to survive in different environments, such as food, soil, water, and air (Scatamburlo et al., 2015). In raw milk, *Pseudomonas* are the dominant group due to
their proteolytic activity (Ercolini et al., 2009). In addition, *Pseudomonas* have also been found in pasteurized milks due to post-pasteurization contamination or processing environment (Gennari & Dragotto, 1992; Reichler et al., 2018). Reichler et al. (2018) identified the contaminating bacteria of pasteurized milk samples from 10 facilities across the northeastern United States. The study revealed that 76.5% of the spoiled samples were contaminated with *Pseudomonas* sp., and 8 out of 10 facilities showed repeat isolation one or more *Pseudomonas* strains, suggesting a difficulty on controlling this microorganism in the supplying industries. Thus, the use of low-quality milk can affect the bacterial ecosystem through fermentation processes and compromise both quality and health-assurance of finalized products (Leitner et al., 2008; Motato et al., 2017).

The emergence of high-throughput sequencing (HTS) technologies has revolutionized the way to investigate microbial diversity of traditional fermentations. These recent platforms have enabled the discovery of several uncultivable microorganisms, sub-dominant populations, and late-growing species, overlooked by culture-based approaches. In the field of food microbial ecology, different HTS platforms have been used for community analysis, including 454 pyrosequencing from Roche, SOLID/Ion Torrent PGM from Life Sciences, and Genome Analyzer/HiSeq 2000/ MiSeq from Illumina (Dertli & Çon, 2017; Humblot & Guyot, 2009; Serafini et al., 2014). These platforms have been used to accurately detect, identify, and characterize foodborne pathogens without any culturing step (Jagadeesan et al., 2019; Leonard et al., 2015).

The aim of this study was to evaluate the performance of Caspian-style milk spontaneous fermentation using low quality milk as a raw material, with focus on microbiological safety and stability of the final product. Due to the high incidence and persistence of *Pseudomonas* found in this study, we applied culture-dependent methods to confirm the presence of viable taxa. In addition, biochemical transformation dynamics (sugar consumption and end-metabolite generation) were examined for better insight into microbial activity during the process.

### 2 Materials and methods

#### 2.1 Fermentation and sampling

The sample of domestic fermented milk was obtained from a private household that traditionally produce Caspian Sea-style spontaneously fermented milk through back-slopping in Curitiba city, Paraná State, Brazil. 50 mL of the “mother” culture was inoculated into 450 mL pasteurized whole cow’s milk and renewed daily in the same proportion (10% vol/vol) at 25 °C for 7 days. This was performed for microbial stabilization before experimental fermentation.

100 mL of resulting fermented milk (“mother” culture) was transferred in triplicate into 2-L Erlenmeyer flasks, containing 900 mL of industrially pasteurized milk of low microbiological quality, and incubated under static condition at 25 °C for 24 h. The low-quality milk, purchased at a local Curitiba market, was selected after quantification of *Pseudomonas*, a common contaminant in pasteurized milk, on *Pseudomonas* F agar (PFA; Thermo Fisher Oxoid). The milk samples that reached plate counts above 20,000 CFU/mL were selected for the fermentation assay (Alles et al., 2018).

Samples (20 mL) of fermenting milk in triplicate were collected at intervals of 6 hours (0, 6, 12, 18, and 24 h) to perform microbiological and metabolite target analysis. At each sampling point, the pH was measured using a digital pH meter (LUCA-210 model, Requipal, Curitiba, PR, Brazil). The resulting fermented product was stored at 4 °C for 60 days (storage stability) and submitted to both culture-dependent and-independent microbiological analyses.

#### 2.2 Total DNA extraction and high-throughput sequencing

Samples of pasteurized milk, fermentation times, and storage stability were withdrawn to perform total genomic DNA extraction and metagenetic analysis. The extraction protocol was performed according Junqueira et al. (2019), with slightly modifications. The cell pellets, obtained after centrifugation of each sample at 12,000 × g for 1 min, were resuspended in 500 μL Tris-EDTA (pH 8.0), vortexed with 10 μL of lysozyme solution at 20 mg/mL (Sigma Aldrich, Saint Louis, MO, USA), and incubated at 30 °C for 60 min. Then, 50 μL of sodium dodecyl sulfate (10% w/v in distilled, deionized water) and 10 μL of proteinase K solution (20 mg/mL in deionized water; Sigma Aldrich) were added to the lysis solution, followed by incubation at 60 °C during 60 min. 150 μL of phenol-chloroform (25:24; Sigma Aldrich) was added, homogenized by inversion, and centrifuged at 12,000 × g for 5 min. The supernatant was collected, and the DNA was precipitated with 3x (v/v) absolute ethanol. Pellets were washed with 80% ethanol, dried, and resuspended in ultrapure water. Extracted DNA quality was checked on a 0.8% (w/v) agarose gel and quantified with the Nanodrop 2000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA).

Twenty ng of the extracted DNA, containing complementary adaptors for Illumina platform, was amplified using degenerated primers for the hypervariable V4 region of 16S (515F and 806R) rRNA gene (Caporaso et al., 2012). Bar-coded amplicons were generated by PCR following conditions described by Junqueira et al. (2019). Samples were sequenced in the MiSeq platform using the 500 V2 kit, following standard Illumina protocols. Resulting sequences in FASTQ files were deposited in the NCBI Sequence Read Archive (SRA) repository with accession BioProject ID PRJNA592162.

#### 2.3 Bioinformatic analyses

After sequencing, chimeric sequences detection, removal of noises from pre-cluster, and taxonomic attribution were performed using standard parameters of QIIME software package, version 1.9.0. Applying the UCLUST method (Edgar, 2010), sequences presenting identity above 97% were considered the same operational taxonomic units (OTUs) according to the SILVA database (Quast et al., 2013).

#### 2.4 Microbial counts

Aliquots of 1 mL of each sample (pasteurized milk, fermentation times, and storage stability) were vortexed with 9 mL of 0.1% saline-peptone water (10⁻¹ solution) and diluted serially. Total aerobic bacteria (TAB) was enumerated on Nutrient Agar medium (NA; Thermo Fisher Oxoid, Waltham, MA, USA), lactic acid bacteria (LAB) on De Man, Rogosa, and
Sharpe Agar (MRS, Thermo Fisher Oxoid), and *Pseudomonas* on *Pseudomonas* F agar (PFA; Thermo Fisher Oxoid); all media containing 0.1% (w/v) nystatin (Sigma Aldrich, Saint Louis, MO, USA) for fungal growth inhibition. NA and MRS plates were incubated at 30 °C for 24 h, and PFA plates were incubated at 37 °C for 48 h. Subsequently, the numbers of cell-forming units (CFU) were recorded.

### 2.5 Substrates and metabolites

Lactose consumption and organic acids production were determined at intervals of 6 hours (0, 6, 12, 18, and 24 h) by high-performance liquid chromatography (HPLC) according to Junqueira et al. (2019), with slightly modifications. Aliquots of 2 mL were centrifuged at 6000×g for 15 min and filtered through a 0.22 μm pore size hydrophilic Polyethersulfone (PES) membrane (Millipore Corp., Burlington, MA, USA). 100 μL of filtered samples were injected into the HPLC system, equipped with an Aminex HPX 87 H column (300 × 7.8 mm; Bio-Rad, Richmond, CA, USA), and a refractive index (RI) detector (HPG1362A; Hewlett-Packard Company, Palo Alto, CA, USA). The column was eluted in an isocratic mode with a mobile phase of 5 mM H₂SO₄ at 60 °C, and a flow rate of 0.6 mL/min.

The extraction of volatile compounds was performed using a headspace (HS) vial coupled to a SPME fiber (CAR/PDMS df75 μm partially crosslinked, Supelco., Saint Louis, MO, USA). For each determination, 2 mL of sample was stored in a 20 mL HS vial, in triplicate. The SPME fiber was exposed for 30 min at 60 °C. The compounds were thermally desorbed into the GC injection system gas phase (GC-MS TQ Series 8040 and 2010 Plus GC-MS; Shimadzu, Tokyo, Japan) at 260 °C. The column oven temperature was maintained at 60 °C for 10 min, followed by two heating ramps of 4 and 10 °C/min until reaching the temperatures of 100 and 200 °C, respectively. The compounds were separated on a column 95% PDMS/5% PHENYL (30 m × 0.25 mm × 0.25 mm film thickness). The GC was equipped with an HP 5972 mass selective detector (Hewlett Packard, Palo Alto, CA, USA). The compounds were identified by comparison to the mass spectra from library databases (Nist'98 and Wiley7n). For quantification, standard solutions of ethanol were prepared in different concentration levels (1, 10, 20, 50, 100 and 1000 μmol L⁻¹) and used to construct a calibration curve. The volatile compound concentrations were expressed as μmol L⁻¹ of headspace, as ethanol equivalent.

### 2.6 Statistical analysis

The data obtained of microbial count and target metabolites were analyzed by post-hoc comparison of means by Duncan’s test. Statistical analyses were performed using the SAS program (Statistical Analysis System Cary, NC, USA). The level of significance was established in a two-sided p-value <0.05.

### 3 Results and discussion

#### 3.1 Microbiological analysis

A total of 651,747 Illumina paired-end reads (average of 93,107 reads/sample) were obtained and clustered into 215 OTUs at 97% sequence similarity. Rarefaction curve analysis showed a trend to level-off at the genus level, indicating that the majority of bacterial communities were covered (Supplementary Material, Figure S1). Sequences were classified, using QIIME and SILVA database, to the lowest possible taxonomic rank (i.e., genus level), and the results are represented in Figure 1. Selected low-quality pasteurized milk harbored mainly Gram-negative bacteria, markedly dominated by *Pseudomonas* (83.74%), and subpopulations of *Acinetobacter*, Enterobacteriaceae, *Sphingomonas*, *Staphylococcus*, and Comamonadaceae, while *Lactococcus* (3.40%) was the only Gram-positive bacteria found. Viable cell count of pasteurized milk on *Pseudomonas*-specific culture medium was 6.40 log CFU/mL, while LAB and TAB were not detected (Figure 2A). The milk used in this study was within the shelf life and stored under conditions ideal. Thus, it is possible to assume that failures occurred during the pasteurization process (Elmoslemany et al., 2010; Vidal et al., 2017; Martin et al., 2018; Russo et al., 2020). Recalls of pasteurized milk contaminated with spoilage bacteria are relatively frequent (Kumaresan & Villi, 2008; Quigley et al., 2013; Samet-Bali et al., 2013; Walsh et al., 2016)

Following fermentation, the number of LAB was always higher, showing the maximum value of 10.15 log CFU/mL at 18 h (Figure 2A). Similarly, TAB increased through the fermentation process, reaching 9.60 log CFU/mL at 18 h. Illumina-based amplicon sequencing showed that LAB population was mainly represented by *Lactococcus*, reaching 66% among the common OTUs at 6 h (Figure 1). Previous studies, using culture-dependent approaches, reported the dominance of *Lactococcus lactis ssp. cremoris* in Caspian Sea yogurt circulated in Japan, accompanied by species of *Leuconostoc, Lactobacillus, Gluconobacter* and *Acetobacter* (Kiryu et al., 2009; Uchida et al., 2009). *Lactococcus* dominance genus was confirmed in this study using Illumina-based amplicon sequencing. *Lactococcus* species have also been found as part of LAB members of other naturally fermented dairy products (Pérez Elortondo et al., 1998; Ferchichi et al., 2001; Fortina et al., 2003; López-Díaz et al., 2000). This microbial group has several important implications for fermentative process, including (i) milk acidification and casein proteolysis; (ii) metabolism of amino acids and fatty acids for flavor development; and (iii) action against food-borne pathogens and spoilage bacteria (El-Ghaish et al., 2011; Matamoros et al., 2009).

In addition, a more complex bacterial diversity, uncovered by the previous traditional cultivation studies, was revealed. These includes *Leuconostoc, Pediococcus, Streptococcus, Plesiomonas, Bacillus, Sphingomonas, Acinetobacter, Comamonadaceae, Ruminococcaceae, Serratia, Prevotella, Staphylococcus, Chryseobacterium, Hydrogenophaga, Methylobacterium* (Figure 1), and other 143 minor bacterial groups with relative prevalence ≤0.1% (Table S1). These findings indicate the need for the use of next-generation sequencing technologies for an in-deep knowledge of the microbial ecology of natural milk fermentation. The discovery of these new taxa will promote the best opportunities to isolate novel microorganisms with functional proprieties and, ultimately, their use as improved starters.

Even though *Pseudomonas* load decreased by more than 99% after 24h of fermentation (8.29 to 5.60 log CFU/mL; Figure 2A), the significant final population can compromise the quality of
Presence of *Pseudomonas* after milk fermentation

the fermented product since *Pseudomonas* are known to produce various enzymes (e.g., lipases, proteases, and phospholipases) that lead to odor, flavor, and body defects (Chen et al., 2011). In addition, it may indicate potential health relevance when consumers believe they are ingesting only beneficial microorganisms. Although the incidence of *Pseudomonas* bacteremia from foods is very rare, some studies reported the presence of virulence in *P. aeruginosa* associated with fresh vegetables, water, and meat (Allydice-Francis & Brown, 2012; Xu et al., 2019). Recent evidence suggests that virulence factors found in environmental isolates, such as *pilin* gene, multidrug efflux transport system, *porin oprD* gene, and haemolytic and proteolytic activities, show no difference with clinical *P. aeruginosa* (Allydice-Francis & Brown, 2012). *P. aeruginosa* is considered an opportunistic pathogen, able to cause urinary tract infections, respiratory dermatitis, soft tissue infections, bacteremia, gastrointestinal infections, and a variety of systemic infections (Lucchetti-Miganeh et al., 2014; Sader et al., 2015; Castaldo et al., 2017). In this sense, great efforts are being explored to prevent contamination by *Pseudomonas* in dairy products (Meesilp & Mesil, 2018; Nan et al., 2016; Picoli et al., 2017; Yasin et al., 2017).

Sequences related to Enterobacteriaceae remained constant throughout the fermentation (Figure 1). The presence of Enterobacteriaceae indicates poor hygiene during the manufacturing process, and high numbers of enterobacteria have been linked to the accumulation of undesirable compounds with implications to flavor and texture defects in dairy products (Linares et al., 2012; Morales et al., 2003). Several studies showed the presence of enterobacteria in kefir grains (Dertli & Çon, 2017; Walsh et al., 2016; Wang et al., 2006) and in cheese samples (Saxer et al., 2013). Dertli & Çon (2017) showed that *Enterobacter* species can pass to the kefir grains from the milk, which should be assessed as they might create safety concerns. However, further studies using enterobacteria-selective culture media should be performed to confirm the presence of viable taxa through the

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**Figure 1.** Relative bacterial abundance and dynamics during Caspian Sea-style spontaneous milk fermentation produced with low quality pasteurized milk. Low prevalence: bacterial groups with relative prevalence ≤0.1%. The complete list of bacteria present at a low prevalence is shown in the supplemental material (Table S1).
Caspian Sea-style spontaneous milk fermentation. In addition, some Enterobacteriaceae genera could be not relevant as food-borne pathogens since many of them are plant commensal organisms (Jha et al., 2011).

There was a substantial discrepancy between viable cell count and Illumina-based amplicon sequencing; for example, *Pseudomonas* spp. accounted for more than 20% of the 16S rRNA gene sequences after 24h (Figure 1), while culture-dependent analysis demonstrated that these organisms account for less than 1% of viable bacterial cells (Figure 2A). This discrepancy is frequently observed in DNA-dependent analyses after contaminants that were present in the raw material are killed while their DNA is still present.

**Figure 2.** Viable cell count and biochemical changes during Caspian Sea-style spontaneous milk fermentation produced with low quality pasteurized milk. (A) Enumeration of lactic acid bacteria (LAB), total aerobic bacteria (TAB) and *Pseudomonas*, and pH monitoring. (B) Course of sugar consumption and organic acids production. Asterisk = significantly higher from one another in a two-sided p value < 0.05 according to Duncan’s test.
amplified (Mayo et al., 2014). This overestimation of β-diversity in culture-independent analysis is recurrently observed in several studies (Martínez et al., 2013; Ursell et al., 2012; Wen et al., 2017).

Interestingly, after the storage process at 4 °C for 60 days, LAB was present at 7.41 log CFU/mL and *Lactococcus* represented 90% of total OTUs by sequencing (Figures 1 and 2). On the other hand, *Pseudomonas* was no longer detected by plating and represented less than 5% of total OTUs (Figures 1 and 2), indicating amplification of reminiscent dead cells (Mayo et al., 2014). The metabolites formed during the fermentation process promotes the survivability of LAB, maintaining their viability over storage time. It is widely known that refrigerated storage is a key point in LAB dominance, increasing shelf life of fermented beverages (Łopusiewicz et al., 2019). Even so, the sampled fermented product should still represent a health concern since Caspian Sea-style spontaneously fermented milk consumption is usually performed within a few days after refrigeration.

### 3.2 Substrates and metabolites

Changes in non-volatiles (lactose and organic acids) and volatile compounds (carboxylic acids, aldehydes, and ketones) were monitored during the course of the fermentation (Figure 2B and Table 1, respectively). The initial lactose content (40.43 g/L) was rapidly reduced to 29.39 g/L within 6 h and remained constant until the end of the process. Lactose is the main carbohydrate in milk, with an average concentration of around 5% (w/v) (Barros et al., 2019). Fermentation reduces lactose in dairy products, helping to prevent symptoms in lactose-intolerant individuals (Savaiano, 2014). In the present study, the highest population of LAB was represented by the genus *Lactococcus*, which uses lactose by active transportation into the cytoplasm via phosphotransferase (PTS) system and hydrolyzing it into glucose and galactose (Mayo et al., 2010). Both monosaccharides enter glycolysis at the level of glucose-6P or metabolized via the Leloir pathway (Kandler, 1983; Mayo et al., 2010).

Lactic acid is the primary end-product observed, showing a continuous increase with maximum concentration of 5.93 g/L at 24 h. The accentuated production of lactic acid is in agreement with the strong dominance of LAB found in the present study (Figures 1 and 2), resulting in pH decrease from 5.10 to 4.10 at the end of fermentation. Lactic acid is the major fermentation product of various bacterial families related to dairy products. This compound is responsible for the raw milk acidification and partial casein coagulation, resulting in the formation of desirable sensory notes and rheological modifications. In addition, lactic acid is the main antimicrobial metabolite produced by LAB, which is responsible for the inhibition of various pathogens and food-borne microorganisms (Chahad et al., 2012; Gálvez et al., 2010; Nakajima et al., 2003; Nakai & Siebert, 2004). Specifically, Nakai & Siebert (2004) showed that *P. aeruginosa* was extremely sensitive to lactic acid, having the lowest MIC (minimum inhibitory concentration) among six different bacteria analyzed. Other factors that also contribute to antagonist action of LAB include the production of hydrogen peroxide, bacteriocins, and antibiotic-like substances.

### Table 1. Concentration of volatile aroma compounds (Area *10°) formed during fermentation of Caspian Sea-style spontaneously fermented milk circulated in Brazil.

| Compounds  | Aroma and taste description     | Milk fermentation (h) |
|------------|--------------------------------|-----------------------|
|            |                                | 0  | 6  | 12 | 18 | 24 |
| **GC**     |                                |    |    |    |    |    |
| Carboxylic acids (6) |                      |    |    |    |    |    |
| Hexanoic acid | Sour, fatty, sweet, cheesy    | 53.89 ± 1.86b | 34.19 ± 3.70b | 55.29 ± 12.27b | 65.81 ± 2.95b | 71.11 ± 1.23b |
| Heptanoic acid | Cheesy, waxy, fermented pineapple | 1.29 ± 0.11ab | 0.69 ± 0.19ab | 0.96 ± 0.39ab | 1.59 ± 0.16a | 1.58 ± 0.29a |
| n-Decanoic acid | Rancid, sour, fatty, citrus | 35.25 ± 11.56a | 31.51 ± 7.26a | 30.01 ± 6.84a | 44.73 ± 0.29a | 48.03 ± 14.83a |
| Benzoic acid | Balsamic                      | ND | 2.06 ± 0.34a | 14.81 ± 3.15a | 20.31 ± 3.23a | 21.00 ± 2.53a |
| Octanoic acid | Fatty, oily, cheesy           | 68.10 ± 3.66a | 62.83 ± 9.76a | 69.31 ± 17.14a | 96.61 ± 1.12b | 97.98 ± 12.41b |
| Nonanoic acid | Cheesy, dairy                 | 0.54 ± 0.01a | 0.53 ± 0.01a | 0.72 ± 0.18a | 0.74 ± 0.00a | 0.81 ± 0.22a |
| **Aldehydes (3)** |                      |    |    |    |    |    |
| Nonanal | Green lemon peel like nuance, citrus, melon rindy | ND | ND | ND | ND | 0.39 ± 0.35 |
| Decanal | Sweet, fatty, citrus, orange peel | ND | ND | 0.44 ± 0.04a | ND | 0.30 ± 0.17a |
| Benzaldehyde | Almond, fruity, powdery, nutty | ND | ND | 1.05 ± 0.47 | ND | ND |
| **Ketones (3)** |                      |    |    |    |    |    |
| 2-Heptanone | Cheesy, fruity, ketogenic      | ND | 0.36 ± 0.01a | 0.86 ± 0.61a | ND | 0.43 ± 0.20a |
| 2-Nonanone | Cheesy, fruity                 | ND | ND | 0.31 ± 0.24a | 0.58 ± 0.03a | 0.35 ± 0.01a |
| δ-dodecaketone | Sweet, creamy, coconut, milky, | ND | 0.20 ± 0.09a | 0.42 ± 0.04a | 0.21 ± 0.09a | 0.47 ± 0.05a |

ND, not detected; GC, gas chromatography. Means of triplicate in each row bearing the same letters are not significantly different (p > 0.05) from one another using Duncan’s Test (mean ± standard variation).
(Arqués et al., 2015). Other minor organic acids produced during the fermentative process can be associated with the growth of sub-dominant bacteria reported by the 16S rRNA gene high-throughput sequencing, including succinic acid produced by Leuconostoc or Acinetobacter, propionic acid from hexose metabolism of Enterobacter species, and acetic acid by both acetic acid bacteria and heterofermentative LAB (Andriani et al., 2019; Souza et al., 2019; Kang et al., 2012).

Volatile compound metabolites were, for the first time, measured during Caspian-sea milk fermentation. Twelve volatile compounds were detected by GC/MS during fermentation, including six carboxylic acids, three aldehydes, and three ketones (Table 1). Amongst the carboxylic acids class, benzoic, hexanoic, and octanoic acids showed a significant increase through the fermentation. Hexanoic and octanoic acids are mainly related to Lactococcus metabolism of lipids and have been associated with cheesy aroma in fermented milk beverages (Azizan et al., 2012; Ziadi et al., 2008). In addition, benzoic acid was detected after 6 h and showed a steady increase, reaching a maximum peak at 24 h. This organic acid is commonly produced by species of Lactococcus, Lactobacillus, and Streptococcus through the conversion of hippuric acid, a natural component of milk. Benzoic acid has an inhibitory effect against spoilage microorganisms, such as yeast, mold, Listeria innocua, Listeria ivanovii, P. aeruginosa, and Oenococcus oeni (Garmiene et al., 2010; Horníčková et al., 2014; Nakai & Siebert, 2004).

Ketones were the second class of volatile aroma reported. Both 2-heptanone and δ-dodecanol were produced after 6 h of fermentation, referring to LAB activity of unsaturated fatty acids hydrolysis (Azizan et al., 2012; Wanikawa et al., 2002). These volatile compounds are commonly reported in the literature for conferring a cheesy-like aroma in fermented milk beverages (Braun, 2019; Walsh et al., 2016). On the other hand, 2-nonanone was produced after 18 h, time in which Pseudomonas count showed a significant increase (Figure 1). This molecule has been recurrently used as a volatile biomarker for rapid detection of Pseudomonas aeruginosa and related species in hospital environments (Savelev et al., 2011; Zechman & Labows, 1985). In this sense, this volatile can also be used for monitoring of quality control in dairy production facilities. Finally, as a minority group, the aldehydes nonanal, decanal, and benzaldehyde were produced at specific fermentation times, being mainly associated with lipid oxidation by LAB (Gänzle et al., 2007).

4 Conclusions

The results of this study demonstrated that Caspian Sea-style spontaneous milk fermentation is not an efficient tool to overcome poor microbiological quality of the milk used as raw material. Although Caspian Sea-style fermented milk showed a high load of LAB and lactic acid content, the presence and persistence of Pseudomonas and enterobacteria through fermentation indicate a potential health risk in the final fermented product. Because of the increased focus on consumption of naturally fermented dairy products, this is a major health concern, since the spread of pathogenic organisms can be facilitated among unsuspecting individuals. The use of poor quality pasteurized milk is also recurrent in other natural milk fermentation (e.g., kefir, koumiss, curd, lassi, laben, and Suero costeño) and further studies should be expanded to cover the safety status of these traditional foods. In addition, for a better understanding of Pseudomonas control using natural milk fermentation, it is crucial to evaluate different Pseudomonas inoculum concentration and more time during storage. The establishment of programs emphasizing hygienic manufacturing procedures can have a major effect on improving the microbiological quality of traditional and ethnic foods circulated in Brazil.

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Supplementary Material

Supplementary material accompanies this paper.

Fig. S1. Alpha rarefaction curves of observed bacterial OTUs (operational taxonomic units) from the temporal samples of Caspian Sea-style fermented milk (A). raw milk. and viability tests (B).

Table S1. Relative abundance (%) of low abundance bacteria at family and genus level during Caspian sea-style fermentation process. ND = not determined; VT = viability test.

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