

Occurrence and behavior of *Bacillus cereus* in naturally contaminated ricotta salata cheese during refrigerated storage

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1 Occurrence and behavior of *Bacillus cereus* in naturally contaminated Ricotta salata cheese
2 during refrigerated storage

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12

13 Abstract

14 The present study shows the fate of *B. cereus* in refrigerated ricotta salata cheese during shelf-life.

15 144 ricotta salata cheese belonging to nine naturally contaminated batches were stored refrigerated

16 and analyzed at 24 h, 30, 60 and 90 days of storage. Total bacterial count, *B. cereus* spores and

17 vegetative forms, intrinsic properties and composition were determined. The presence of spores was

18 sporadic while the prevalence and the level of *B. cereus* vegetative cells decreased respectively

19 from 83.3% and 4.65 ± 0.74 cfu g⁻¹ at the beginning of the observation period to 33.3% and

20 1.99 ± 0.55 cfu g⁻¹ after 90 days. No information is currently available on the fate of *B. cereus* in

21 ricotta salata. The production process of ricotta salata includes steps such as whey heating followed

22 by slow cooling of clots, which expose to the risk of spore germination and successive growth to

23 levels compatible with toxins production. The prolonged refrigerated storage was not favorable to

24 sporulation, explaining the successive death of vegetative cells. The present study demonstrate the

25 potential risk of food poisoning as consequence of pre-formed emetic toxins in ricotta salata. Food

26safety of ricotta salata relies on the rapid refrigeration of the product during critical phases for
27cerulide production.

28Keywords: *Bacillus cereus*; whey cheese; sheep's milk

29

301. Introduction

31The *Bacillus cereus* group includes Gram-positive rod shaped spore-forming bacteria, which are
32widely distributed in the natural environment. Within the group, *B. cereus sensu stricto* is the most
33important organism causing food spoilage and food-borne illness (Kramer and Gilbert, 1989). *B.*
34*cereus* causes two clinical forms of foodborne illness: the emetic and the diarrheal syndrome
35(Granum and Lund, 1997). A dose of 10^5 - 10^8 cells or spores per gram is generally considered
36necessary to cause illness (ICMSF, 1996; Granum and Lund, 1997). *B. cereus* is frequently isolated
37from raw milk and dairy products thus, representing a serious concern for the dairy industry
38(Svensson et al., 2006). Due to its ubiquitous nature and the extreme resistance of endospores to
39several harsh conditions (Nicholson, et al., 2000), it is difficult to avoid the contamination of dairy
40products. *B. cereus* can enter the dairy chain mainly through raw milk contaminated at farm level
41(Heyndrickx, 2011). However, contamination may also arise from the food-processing environment
42(da Silva Fernandes et al., 2014). Dairy products have been seldom associated with human illness
43despite the frequent contamination with *B. cereus* (EFSA, 2005). Whey products processed at high
44temperatures and successively stored refrigerated are particularly exposed to the risk of *B. cereus*
45(Heyndrickx and Scheldeman, 2002). The endospores are activated by whey heating applied during
46protein denaturation ($>80^\circ\text{C}$) and vegetative cells are then facilitated in their growth by the absence
47of competing microbiota, inactivated by the heat treatment (Scheldeman et al., 2006). *B. cereus*
48psychotropic strains can grow to temperature as low as 4 - 5°C and during the refrigerated storage
49can reach levels potentially harmful for human health (Huck et al., 2007). Ricotta salata is a
50traditional dry and salted sheep's milk whey cheese produced in Sardinia (Italy). Technology and
51microbiological profile of ricotta salata have been previously described (Spanu et al., 2015). The

52attributed shelf-life of ricotta salata is generally up to several months under refrigerated storage
53(Casti et al., 2016). The present study was conducted following a case of large *B. cereus*
54contamination of ricotta salata occurred in one sheep's milk cheese-making plant operating in
55Sardinia. During the period September-October 2014, a local food business operator observed the
56presence of *B. cereus* contamination in ricotta salata samples during routine microbiological testing
57conducted as part of their procedure based on HACCP principles. The mean level of contamination
58was $5.57 \pm 0.15 \log_{10} \text{ cfu g}^{-1}$ in a batch. Although no food safety criteria for *B. cereus* are applicable to
59foodstuffs placed on the market during their shelf-life (EC Regulation No. 2073/2005), the food
60business operator as corrective action withdrew the entire batch of ricotta salata. The subsequent
61production batches positive for the presence of *B. cereus*, were destined to a durability study. The
62few published data existing on *B. cereus* contamination in ricotta salata produced in Sardinia
63reported a prevalence of ca. 15% and a contamination level ranging from 1 to 3 $\log_{10} \text{ cfu g}^{-1}$
64(Cosentino, et al., 1997; De Santis et al., 2008; Fadda et al., 2012). Despite the reported
65contamination levels are below the dose necessary to cause illness, they demonstrate that *B. cereus*
66in ricotta salata represents a potential concern for consumer's health. No published reports are
67currently available on the fate of *B. cereus* in naturally contaminated ricotta salata stored under
68refrigerated conditions. The aim of the present study was to describe the evolution of *B. cereus* in
69naturally contaminated ricotta salata during shelf-life and to assess the potential health risk
70associated with the microorganism survival or growth.

71

72. **Materials and methods**

73

742.1. *Ricotta salata batches and samples*

75Ricotta salata batches used in the study were selected based on the natural occurrence of *B. cereus*.

76With this aim, during the period September-October 2014, ricotta salata production batches were

77tested on a daily basis for the presence of *B. cereus*. From each positive batch were randomly

78selected sixteen ricotta salata wheels. Samples were immediately vacuum packed in plastic bags,

79transported refrigerated to the laboratory and stored in cold room ($4\pm 2^{\circ}\text{C}$) until analyses were
80performed.

81

822.2. *Experimental design*

83Ricotta samples were analyzed at four different times during the shelf-life. Sampling times were:
84within 24 h after the arrival of ricotta salata wheels defined as time zero (T_0), 30, 60 and 90 days
85after the production defined respectively as time 30 (T_{30}), time 60 (T_{60}) and time 90 (T_{90}). From each
86of the nine different batches and at each sampling time, two samples were used for microbiological
87analysis and two samples for physico-chemical determinations.

88

892.4. *Microbiological analysis*

90Ricotta salata samples were analyzed for the determination of total aerobic mesophilic bacteria (ISO
914833, 2013) and enumeration of *B. cereus* (ISO 7932, 2004). The enumeration of both *B. cereus*
92vegetative cells and spores was determined, respectively before and after heating at 80°C for 10 min
93by plating two 0.1 mL aliquot on selective chromogenic culture media such as Mannitol Egg Yolk
94Polymyxin agar (MYP, Biolife, Milan, Italy) and Polymyxin Pyruvate Egg-Yolk Mannitol
95Bromothymol Blue (PEMBA, Oxoid) agar. Samples were incubated at 30°C in aerobic conditions
96for 24 h. From each positive sample were picked five presumptive *B. cereus* colonies, transferred
97onto Trypticase Soy Agar (TSA, Biolife) and incubated at 37°C for 24 h. Each isolate was
98submitted to phenotypic identification and successively confirmed by PCR (Oh et al., 2012).

99

1002.5. *Intrinsic properties and composition*

101PH and a_w were measured using pH meter GLP22 (Crison Instruments SA, Barcelona, Spain) and
102water activity meter Aqualab 4 TE (Decagon, Pullman, WA, USA), respectively. Determination of
103centesimal composition (% of moisture, fat, protein, salt and total solids) was performed using the
104Near Infrared Transmittance (NIT) compositional analyzer (FOSS, Eden Prairie, MN, USA).

105

1063. **Results**

107

1083.1. *Microbiological profile*

109The mean aerobic mesophilic counts (\log_{10} cfu g^{-1} ; $\bar{x} \pm SD$) of ricotta salata analyzed at T_0 , T_{30} , T_{60}
110and T_{90} were 5.17 ± 1.39 , 5.69 ± 0.54 , 5.99 ± 0.67 and 5.62 ± 0.87 , respectively. The prevalence of *B.*
111*cereus* vegetative cells and the mean contamination level decreased during the refrigerated storage
112($P < 0.05$). At T_0 , the prevalence was 83.3% with counts ranging from $3.45 \log_{10}$ cfu g^{-1} to $6.20 \log_{10}$
113cfu g^{-1} , while at T_{90} the observed prevalence was 33.3% with counts ranging from $1.30 \log_{10}$ cfu g^{-1}
114to $2.56 \log_{10}$ cfu g^{-1} (table 1). The mean reductions over time (ΔT) in *B. cereus* vegetative cells
115concentration (\log_{10} cfu g^{-1}) were 0.38, 1.74 and 2.66 at T_{30} , T_{60} and T_{90} , respectively. The detection
116of *B. cereus* spores after heat activation was observed in two samples belonging to two different
117batches, one at T_{30} ($2.30 \log_{10}$ cfu g^{-1}) and one at T_{60} ($2.0 \log_{10}$ cfu g^{-1}), respectively. Out of 49 total
118positive samples (68.0%) were isolated 245 presumptive *B. cereus* strains of which 101 were
119confirmed by molecular identification.

120
1213.2. *Physico-chemical characteristics*
122The pH values ranged between 6.23 and 6.67 at T_0 and between 5.30 and 6.32 at T_{90} , while a_w
123values ranged between 0.964 and 0.986 at T_0 and between 0.976 and 0.983 at T_{90} . The evolution of
124the mean centesimal composition values ($\%$; $\bar{x} \pm SD$) at different sampling times is reported in table

1252.
126

1274. Discussion

128Despite raw milk is the main source of contamination of dairy product with sporeformers, their
129level is generally low, $< 1-10^2$ cfu mL^{-1} (Vissers et al., 2007). Seasonal variation has been reported
130with counts as high as 10^4 cfu mL^{-1} (Slaghuis et al., 1997; TeGiffel et al., 2002; Coorevits et al.,
1312008). The presence of *B. cereus* in ricotta salata is a rare finding, with maximum contamination
132level of ca. $3 \log_{10}$ cfu g^{-1} (Cosentino et al., 1997; De Santis et al., 2008; Fadda et al., 2012; Spanu et
133al., 2012). The high level of contamination, up to $8.33 \log_{10}$ cfu g^{-1} , observed in the present study
134and the large number of positive batches (nine) over a limited period of time (three months), should

135be considered as an event strictly associated with the late summer and early fall production period.
136Microbiological testing of each production batch, conducted on a regular basis in the frame of the
137food business operator's HACCP procedures, showed no occurrence of *B. cereus* contamination
138during the rest of the year. This could be explained with the typical sheep's milk breeding systems
139adopted in Sardinia. Milk production is seasonal, starting from December until July. The peak of
140milk production is concentrated between January and May, with a decrease between June and
141August, when the sheep start entering in the dry period. Cheese-making during the dry period relies
142on the milk available provided by flocks adopting the out-of-season breeding system. Poor pasture
143quality during this season determines a decline in milk yield and microbiological quality (Sitzia et
144al., 2015). Due to economic reasons, raw milk is picked and transformed every three or five days
145instead that daily. In addition, during winter and spring periods sheeps mainly graze on grass
146pasture, while during the summer and fall periods on stubble with concentrate and feedstuff
147supplement, which may increase the risk of transferring spores into raw milk. Total bacterial count
148and yield records of the milk used to make the ricotta salata used in the present study, were obtained
149by the food business operator. Data confirmed differences in milk yield and microbiological quality
150over the milking season. In the period from January to June, the total bacterial count (geometric
151mean) of raw milk was ca. 140,000 cfu mL⁻¹ with an average production of 2,150,000 l, while in the
152out-of-season period the total bacterial count was ca. 1,100,000 cfu mL⁻¹ with an average milk yield
153of 135,000 l. Therefore, the production of ricotta salata during the out-of-season period was
154characterized by risk factors increasing the likelihood of a high initial level of *B. cereus*
155contamination in the product. The greater relative decrease in *B. cereus* vegetative cells was
156observed after 60 days (-1.36 log₁₀ cfu g⁻¹) and after 90 days (-0.92 log₁₀ cfu g⁻¹) of refrigerated
157storage. Since psychrotrophic strains can grow at temperature as low as 4-5°C, contamination was
158likely due to mesophilic strains which minimum growth temperature is 15°C (ICMSF, 1996).
159Growth and survival characteristics of *B. cereus* vary widely between strains and depend upon a

160complex series of interacting factors such as temperature, pH, water activity (NaCl concentration),
161nutrients and presence of competitive microbiota. *B. cereus* is generally a poor competitor in
162unpasteurized products (Andersson, Ronner, & Granum, 1995). The high total bacterial count (ca. 6
163log), combined with the decrease of pH values (from 6.49 to 5.63) observed over time, suggest the
164possible presence of contaminants from the whey or the environment that may have exerted a
165possible competitive action. A previous study, conducted on vacuum packed ricotta salata, showed a
166mean aerobic mesophilic bacteria count (\log_{10} cfu g^{-1}) after 2 and 4 months of refrigerated storage
167ranging from 7.56 ± 0.85 and 4.57 ± 0.62 on the rind and from 3.64 ± 0.71 and 2.95 ± 0.65 on the inner
168paste, respectively (Spanu et al., 2013). At the beginning of the ricotta salata storage *B. cereus* is
169mainly present in its vegetative form, as consequence of heat activation of spores occurred during
170whey heating. The successive phase of pressing of the warm clots, expose the product to
171temperature at risk for the growth of the vegetative forms to levels compatible with the emetic toxin
172(cerulide) production. The reduction in *B. cereus* vegetative cells count over the storage period
173suggests the death of the microorganism rather than sporulation, since no grow was observed in
174samples analyzed after pasteurization. The sporulation is a complex process which occurs as
175response to stress such as starvation, high cell density (10^6 - 10^7 cfu g^{-1}) or DNA damage and it is
176regulated by hundreds of genes (Eichenberger et al., 2003; Piggot and Hilbert, 2004). Borge et al.
177(2001) concluded that vegetative cells are unlikely to develop endospores in refrigerated media. The
178high levels of ricotta salata contamination with *B. cereus* observed in the present study indicates
179that this product could represents a risk for human health due to the potential presence of pre-
180formed emetic toxins in the product. The low contamination level observed in the product after long
181refrigerated storage could lead to the wrong conclusion that the product is safe, while cerulide still
182persists. In fact, cerulide is highly resistant to heat, low pH, and proteolytic activity of pepsin and
183trypsin (Kramer and Gilbert, 1989; Rajkovic et al., 2008). Further investigation is needed in order to

184 assess whether the origin of the contamination is from ingredients, processing environment or from
185 packaging materials and to determine the pathogenicity of the strains.

186 **5. Conclusion**

187 Ricotta salata production process includes critical phases such as heat coagulation and slow cooling
188 of clots, which support the activation of *B. cereus* spores and the successive growth of vegetative
189 cells, in the absence of competing microbiota. The present investigation demonstrates that the level
190 of *B. cereus* vegetative cells in naturally contaminated ricotta salata decreases during refrigerated
191 storage, while the presence of spores is a rare finding. The control of *B. cereus* in ricotta salata
192 relies on one hand on limiting the level of spores in raw milk, and as consequence in the whey, and
193 on the other hand in preventing germination and successive growth of vegetative cells.

194

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201

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302

303 Table 1. Evolution of *B. cereus* (\log_{10} cfu g^{-1} ; mean \pm SD) in nine batches of vacuum-packed Ricotta
304 salata stored refrigerated until 90 days after production.

Batc	T ₀	T ₃₀	T ₆₀	T ₉₀
h				
1	4.51 \pm 0.00 ^a (n = 1/2)	2.79 \pm 0.79 ^a (n = 2/2)	N.D. (n = 2/2)	N.D. (n = 2/2)
2	4.37 \pm 0.37 ^a (n = 2/2)	5.04 \pm 0.00 ^a (n = 1/2)	2.94 \pm 1.00 ^a (n = 2/2)	N.D. (n = 2/2)
3	4.68 \pm 0.24 ^a (n = 2/2)	4.49 \pm 0.20 ^a (n = 2/2)	3.26 \pm 0.93 ^{ab} (n = 2/2)	1.30 \pm 0.00 ^b (n = 1/2)
4	4.50 \pm 0.33 ^a (n = 2/2)	4.68 \pm 0.20 ^a (n = 2/2)	2.15 \pm 0.00 ^b (n = 1/2)	1.78 \pm 0.68 ^b (n = 2/2)
5	4.66 \pm 0.35 ^a (n = 2/2)	3.85 \pm 0.21 ^a (n = 2/2)	N.D. (n = 2/2)	N.D. (n = 2/2)
6	4.88 \pm 0.00 ^a (n = 1/2)	3.81 \pm 0.74 ^a (n = 2/2)	3.45 \pm 0.16 ^a (n = 2/2)	N.D. (n = 2/2)
7	4.0 \pm 0.00 ^a (n = 1/2)	3.62 \pm 0.00 ^b (n = 1/2)	2.38 \pm 0.00 ^a (n = 2/2)	N.D. (n = 2/2)
8	6.19 \pm 0.14 ^a (n = 2/2)	4.83 \pm 0.43 ^b (n = 2/2)	3.78 \pm 0.00 ^{bc} (n = 1/2)	2.56 \pm 0.00 ^c (n = 1/2)
9	3.77 \pm 0.46 ^{ab} (n = 2/2)	5.43 \pm 0.89 ^b (n = 2/2)	2.46 \pm 1.67 ^a (n = 2/2)	2.26 \pm 0.00 ^a (n = 2/2)
total	4.65 \pm 0.74 ^a (n =15/18)	4.27 \pm 0.90 ^a (n =16/18)	2.91 \pm 0.84 ^b (n =12/18)	1.99 \pm 0.55 ^c (n =6/18)

305 The sampling time (T₀, T₃₀, T₆₀ and T₉₀) were respectively the day of packaging and 30, 60 and 90
306 days after the production. Means in the same row with different superscript letters are significantly
307 different (P < 0.05); values within brackets indicate the prevalence of positive samples. N.D = data
308 not definable, below the detection limit of the method.

309Table 2. Intrinsic properties (mean \pm SD) and composition ($\% \pm$ SD) evolution during storage of
310ricotta salata wheels.

parameter	sampling times			
	T ₀	T ₃₀	T ₆₀	T ₉₀
pH	6.49 \pm 0.10 ^a	6.18 \pm 0.10 ^b	5.73 \pm 0.14 ^c	5.63 \pm 0.28 ^c
a _w	0.978 \pm 0.001 ^{ab}	0.976 \pm 0.002 ^a	0.976 \pm 0.002 ^a	0.980 \pm 0.001 ^b
% moisture	58.28 \pm 2.91 ^a	58.56 \pm 3.29 ^a	57.41 \pm 2.99 ^{ab}	56.23 \pm 2.55 ^b
% total solids	41.72 \pm 2.93 ^a	41.44 \pm 3.26 ^a	42.59 \pm 2.99 ^{ab}	43.77 \pm 2.49 ^b
% fat	23.74 \pm 3.92 ^a	23.13 \pm 4.28 ^a	23.32 \pm 3.73 ^a	23.39 \pm 3.86 ^a
% protein	12.65 \pm 1.12 ^a	12.83 \pm 1.16 ^a	13.01 \pm 0.84 ^a	13.36 \pm 1.14 ^a
% salt	2.60 \pm 0.30 ^{ab}	2.73 \pm 0.24 ^b	2.49 \pm 0.28 ^c	2.01 \pm 0.49 ^d

311The sampling time (T₀, T₃₀, T₆₀ and T₉₀) were respectively the day of packaging and 30, 60 and 90
312days after the production. Means in the same row with different superscript letters are significantly
313different (P < 0.05).