Characterisation of the non-starter lactic acid bacteria (NSLAB) of Gruyère PDO cheese

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Abstract – The diversity of non-starter lactic acid bacteria (NSLAB) of Swiss Gruyère cheese was studied in three factories over a six-month period. A total of 1099 isolates of NSLAB from raw milk, skim milk and Gruyère cheese were characterised at both the species and strain level. Over 90% of the isolates were either Lactobacillus casei or L. rhamnosus, and a combined total of 45 genotypes were found. The genotypes from the three factories differed from one another and varied slowly over the six-month period. At one factory, samples from the corresponding raw milk, skim milk and cheese culture were also analysed. No NSLAB were found in the cheese starter culture. On the contrary, raw milk contained the largest number of different genotypes. The genotypes found in the cheese all came from raw milk.

non-starter lactic acid bacteria / NSLAB / L. casei / L. rhamnosus / Gruyère cheese

Résumé – Caractérisation des bactéries lactiques non levain (NSLAB) du Gruyère AOC. La diversité des bactéries lactiques non levain (NSLAB) du Gruyère suisse a été étudiée sur une période de 6 mois dans trois fromageries. Au total, 1099 isolats de NSLAB provenant de lait cru, de lait écrémé et de Gruyère ont été caractérisés au niveau de l’espèce et de la souche. Plus de 90 % des isolats étaient issus soit de Lactobacillus casei, soit de L. rhamnosus. Parmi ces isolats, on a trouvé un total de 45 génotypes différents. Les génotypes identifiés dans les trois fromageries différaient les uns des autres et variaient faiblement au cours des 6 mois. En outre, dans une fromagerie, des échantillons de lait cru, de lait écrémé et de fromage tous issus de la même matière première ont été analysés. Aucun NSLAB n’a été trouvé dans le levain traditionnel de fromagerie. Par contre, le lait cru contenait le nombre le plus important de génotypes différents. Les génotypes trouvés dans le fromage provenaient tous du lait cru.

bactéries lactiques non levain / NSLAB / L. casei / L. rhamnosus / Gruyère suisse

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1. INTRODUCTION

Non-starter lactic acid bacteria (NSLAB) are defined as lactic acid bacteria found in cheese which do not form part of the starter culture and consist of four main groups of bacteria, i.e. mesophilic lactobacilli, pediococci, enterococci and Leuconostoc. All natural cheeses studied to date contain bacteria from at least one of these four groups. Among the mesophilic lactobacilli, members of the facultatively heterofermentative lactobacilli are most often encountered in cheese. There appears to be some variation in NSLAB depending on the cheese variety, processing and duration of ripening [2]. In a range of cheeses such as Cheddar of different origins, Fiore Sardo, Fossa cheese and other European traditional cheeses, *L. casei*, *L. plantarum*, *L. curvatus* and *L. rhamnosus* were the dominant species of the mesophilic lactobacillus flora, but also various other species occurred at lower levels [3, 8, 11, 12, 14, 16, 21]. Raw milk is the main source of the NSLAB, but also other sources such as other cheese-making ingredients and contamination of plant equipment with persistent biofilms contribute to the NSLAB composition. Although pasteurisation reduces the flora of the raw milk to a large extent, small numbers of mesophilic lactobacilli may survive and subsequently grow in cheese made from pasteurised milk. Mesophilic lactobacilli grow in cheese to levels of $10^6$–$10^8$ cfu·g$^{-1}$ over the first 10–20 weeks of ripening and remain at high concentrations until the end of the ripening period [2, 18, 19]. Comparison of cheeses made from raw milk with those made from microfiltered or pasteurised milk has shown that the raw milk flora plays an essential role in the development of the sensory characteristics of hard cheeses. In most cases it was found that elimination of the raw milk flora leads to a less intense cheese flavour [6, 10, 17]. Several studies further indicated that mesophilic adjuncts can exert a positive effect on flavour [5, 8, 19]. However, it remains difficult to quantify the impact of individual strains of NSLAB on cheese quality since it is impossible to achieve complete control of cheese flora. In order to better understand the role played by this microflora in the development of the sensory characteristics of cheese, it is necessary to identify them not only at the species level but eventually at the strain level using electrophoretic techniques such as pulsed-field gel electrophoresis, hybridisation techniques and PCR-based methods.

Until now, the description of the ecosystem of raw milk and Gruyère cheese has been based on classical microbiological methods which are limited to describing bacteria at the species or sub-species level. Analysis of this microflora using modern molecular biology methods should provide us with further knowledge of these strains as well as their origin, diversity and variability.

Three different cheese factories, that consistently produced Gruyère cheese of high quality, were selected and the diversity of NSLAB of six-month-old cheeses produced at monthly intervals over a period of six months was analysed.

2. MATERIALS AND METHODS

2.1. Manufacture of Gruyère PDO cheese

The present investigation on NSLAB in Swiss Gruyère cheese was performed in collaboration with three typical factories located in Pomy (approval number CH-4418), Mezieres (CH-4089) and La Brévine (CH-4214) during the period of August 2002 until January 2003. Traditional Gruyère PDO cheese was produced according to the list of requirements of the Gruyère association. Briefly, the fat content of the cheese milk was adjusted by adding skimmed raw milk to the untreated raw milk. Mesophilic lactobacilli grow in cheese to levels of $10^6$–$10^8$ cfu·g$^{-1}$ over the first 10–20 weeks of ripening and remain at high concentrations until the end of the ripening period [2, 18, 19]. Comparison of cheeses made from raw milk with those made from microfiltered or pasteurised milk has shown that the raw milk flora plays an essential role in the development of the sensory characteristics of hard cheeses. In most cases it was found that elimination of the raw milk flora leads to a less intense cheese flavour [6, 10, 17]. Several studies further indicated that mesophilic adjuncts can exert a positive effect on flavour [5, 8, 19]. However, it remains difficult to quantify the impact of individual strains of NSLAB on cheese quality since it is impossible to achieve complete control of cheese flora. In order to better understand the role played by this microflora in the development of the sensory characteristics of cheese, it is necessary to identify them not only at the species level but eventually at the strain level using electrophoretic techniques such as pulsed-field gel electrophoresis, hybridisation techniques and PCR-based methods.

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ripened at about 14 °C until the age of six months.

2.2. Sampling of the study materials
In order to study the influence of raw milk, processing equipment and starter cultures on NSLAB composition, samples of raw milk, skimmed milk and traditional whey cultures were taken at monthly intervals over a period of six months from each factory and stored at –80 °C until used for analysis. The corresponding cheese samples were collected after a ripening period of six months.

2.3. Isolation of bacteria and growth conditions
Bacteria were isolated from cheese by adding 10 g of grated cheese to 90 mL peptone water, heating for 10 min at 45 °C and then plating out the diluted suspension on facultatively heterofermentative agar (FH-agar) plates, since this media is specific for facultatively heterofermentative lactobacilli [13]. Furthermore, in a preliminary experiment the suspensions were plated out on lithium propionate agar (LP-agar) [15] in order to determine whether other bacterial species grew in the cheese. Since over 90% of the species were facultatively heterofermentative lactobacilli, it was decided to concentrate on this group for the main study. Fifty colonies were selected at random; the isolates were purified by plating out on MRS-agar three times and further cultured in MRS broth. Bacteria from milk and skim milk were isolated after heat-treatment similar to manufacturing conditions by heating 10 mL at 57 °C for 6 min and then directly plating out on FH-agar. Without this heat-treatment other types of bacteria, such as Weisella, Leuconostoc and Pediococcus, grew on the agar plates. One hundred and thirty-four bacterial colonies were isolated from milk and 65 from skim milk.

2.4. Total DNA preparation
Genomic DNA was prepared from 1 mL of a stationary phase culture as described previously [7].

2.5. Species identification
The 16S rRNA gene was amplified by PCR with the primers bak11w (5’-AGTTTGATCMTGGCTCAG-3’) and Univ-518 (5’-CWATTACCGCGGCKGCTG-3’) [9]. The PCR products were purified using the QIAquick PCR Purification Kit and sequenced using the same primers on an ABI Prism® 310 Genetic Analyser using the BigDye Terminator Cycle sequencing kit. Each species was identified by comparing the DNA sequence obtained with those in the NCBI database.

2.6. Strain identification
Isolated strains were grouped into genotypes by repetitive sequence-based polymerase chain reaction (REP-PCR) which was performed on DNA extracts using the primer GTG5 (5’-GTTGTTGGTGGTG-3’) as described previously [20].

3. RESULTS

3.1. Cheese
In a series of preliminary experiments cheese extracts were plated out on LP-agar. Thermophilic cheese starters did not produce colonies on this agar, thus making it suitable for studying the NSLAB content of cheese. One hundred isolates each from the three cheese factories were identified at the species level by DNA sequencing of 16S rRNA. The different groups found are listed in Table I. Over 90% of the bacteria found were facultatively heterofermentative lactobacilli. This confirms the results found with other cheese types [1, 8, 14, 16] and indicates that these mesophilic lactobacilli can contribute to the flavour of Gruyère cheese.

Fifty randomly selected isolates from each cheese sample were identified by DNA sequencing of the 16S rRNA. The results of the identification are given in Table II. The bacteria were found at concentrations between 10^6 and 10^7 cfu·g⁻¹ and the large majority of isolates were Lactobacillus casei.
The different isolates were then analysed by REP-PCR in order to evaluate the different genotypes present. A combined total of 45 different genotypes were found in the cheeses from the three cheese factories over the six-month period. The genotypes from the three cheese factories were all different from one another. The distribution of the genotypes over the six-month period is shown in Tables III–V.

### 3.2. Milk and skimmilk

In order to determine the origin of the facultatively heterofermentative lactobacilli, we examined the raw milk and skimmilk, used for the manufacture of cheese, for the presence of these bacteria. The January milk samples from cheese factory CH-4214 were chosen as an example, since the cheese made by it contained only the three genotypes C32, C44 and C45 at relatively equal concentrations (Tab. V). Fourteen different genotypes of NSLAB were found in raw milk, whereas only 4 genotypes were found in skimmilk (Tab. VI). All three genotypes present in cheese as well as 3 of the genotypes present in skimmilk were also found in the corresponding raw milk samples. The genotypes and frequency of facultatively heterofermentative lactobacilli found in raw milk, skimmilk and cheese are summarised in Table VI.

### 3.3. Starter cultures

The traditional whey culture used for the manufacture of cheese in factory CH-4214

#### Table I. Bacterial groups found in cheese extracts after growth on lithium propionate agar (100 isolates from each factory were analysed).

<table>
<thead>
<tr>
<th>Group / Species</th>
<th>Factory CH-4089</th>
<th>Factory CH-4214</th>
<th>Factory CH-4418</th>
</tr>
</thead>
<tbody>
<tr>
<td>Facultative heterofermentatively lactobacilli</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L. casei</td>
<td>100</td>
<td>32</td>
<td>58</td>
</tr>
<tr>
<td>L. rhamnosus</td>
<td>57</td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>Pediococci</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enterococci</td>
<td>1</td>
<td>2</td>
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</tr>
</tbody>
</table>

#### Table II. Concentrations and percentages of facultatively heterofermentative lactobacilli in Gruyère cheeses after six months of ripening (50 randomly selected isolates from each factory and each month).

<table>
<thead>
<tr>
<th>Month</th>
<th>Cheese Factory CH-4214</th>
<th>Cheese Factory CH-4418</th>
<th>Cheese Factory CH-4089</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10⁶ cfu·g⁻¹</td>
<td>%</td>
<td>10⁶ cfu·g⁻¹</td>
</tr>
<tr>
<td>August</td>
<td>6</td>
<td>88% L. casei</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>12% L. rhamnosus</td>
<td></td>
</tr>
<tr>
<td>September</td>
<td>2</td>
<td>81% L. casei</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>19% L. rhamnosus</td>
<td></td>
</tr>
<tr>
<td>October</td>
<td>10</td>
<td>100% L. casei</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>November</td>
<td>8</td>
<td>58% L. casei</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>42% L. rhamnosus</td>
<td></td>
</tr>
<tr>
<td>December</td>
<td>6</td>
<td>100% L. casei</td>
<td>3</td>
</tr>
<tr>
<td>January</td>
<td>2</td>
<td>100% L. casei</td>
<td>0.7</td>
</tr>
</tbody>
</table>
NSLAB of gruyère PDO cheese

in January did not contain any facultatively heterofermentative lactobacilli detectable by our method.

4. DISCUSSION

In the present study a total of 1099 isolates of NSLAB from cheese milk, skim milk and Gruyère cheese from three different cheese factories were identified at the species and genotype levels. Previously, Berthier et al. [4] described the diversity of mesophilic lactobacilli in Comté cheese, which has often been compared with Gruyère cheese. Whereas the study with Comté cheese concentrated on the development of mesophilic lactobacilli during ripening, our study examined six-month-old cheeses produced at monthly intervals over a period of six months.

### Table III.
Numbers of facultatively heterofermentative lactobacilli isolates of different genotypes found in Gruyère cheeses after six months of ripening in cheese factory CH-4089 over a period of six months (50 randomly selected isolates from one cheese each month identified).

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Aug</th>
<th>Sep</th>
<th>Oct</th>
<th>Nov</th>
<th>Dec</th>
<th>Jan</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>34</td>
<td>34</td>
<td>6</td>
<td>21</td>
<td>42</td>
<td>41</td>
</tr>
<tr>
<td>C2</td>
<td>9</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td>3</td>
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<tr>
<td>C3</td>
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<td>3</td>
<td>2</td>
<td>6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C4</td>
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</tr>
<tr>
<td>C5</td>
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<td></td>
<td></td>
<td>42</td>
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</tr>
<tr>
<td>C6</td>
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<td>C7</td>
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<td>C9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6</td>
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</table>

### Table IV.
Numbers of facultatively heterofermentative lactobacilli isolates of different genotypes found in Gruyère cheeses after six months of ripening in cheese factory CH-4418 over a period of six months (50 randomly selected isolates from one cheese each month identified).

<table>
<thead>
<tr>
<th>Genotype</th>
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<th>Sep</th>
<th>Oct</th>
<th>Nov</th>
<th>Dec</th>
<th>Jan</th>
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<tbody>
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</tr>
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<td>C12</td>
<td>2</td>
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<td>31</td>
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<td>C13</td>
<td>7</td>
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### Table V.
Numbers of facultatively heterofermentative lactobacilli isolates of different genotypes found in Gruyère cheeses after six months of ripening in cheese factory CH-4214 over a period of six months (50 randomly selected isolates from one cheese each month identified).

<table>
<thead>
<tr>
<th>Genotype</th>
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</tbody>
</table>

in January did not contain any facultatively heterofermentative lactobacilli detectable by our method.
Similarly to us, Berthier et al. [4] found that over 90% of the NSLAB isolates from the mature cheeses belonged to the species of *L. casei* or *L. rhamnosus* and were present at concentrations of $10^6$–$10^7$ cfu·g$^{-1}$ cheese after six months of ripening. We found between 1 and 8 genotypes per cheese, which is also similar to Comté. The genotypes from the three Gruyère cheese factories were different from one another, which is not surprising since the factories are situated at least 50 km apart.

The genotypes of the NSLAB in Gruyère cheese within each factory varied over time, but generally there was an overlap in the genotypes from one month to another. However, the shifts in the genotypes in the cheeses of the three factories were different from one another. The dominant genotype in the cheeses from factory CH-4089 was present over the whole six-month period, whereas the dominant genotype in cheeses from factory CH-4418 changed slowly with time and was completely different after six months. The cheeses from factory CH-4214 were in-between the two, since one of the genotypes found at the beginning of the study was still present after six months, even if it was no longer the dominant genotype. Since we reasoned that the cheese milk would be partially contaminated with NSLAB originating from the factory equipment and starter culture, we also analysed the skimmilk and starter culture from the same day as the cheeses were produced. In contrast to Comté cheese, the culture used for the manufacture of Gruyère cheese did not contain any NSLAB at a detectable level. This is possibly because during the preparation of the cultures for Gruyère the whey undergoes a preheating step at 59 °C before incubation at 38 °C. Pretreatment at 59 °C eliminates a lot of bacteria. This procedure is not undertaken during the manufacture of Comté cheese.

Raw milk was found to be far richer than skimmilk in the number of genotypes of NSLAB that it contained (Tab. VI). This is not surprising, since skimmilk was prepared in the factories early in the morning from

| Table VI. Genotypes of facultatively heterofermentative lactobacilli and their frequency found in cheese milk, skimmilk and the corresponding six-month-old cheese of the January production in factory CH-4214 (RM1–RM14 = genotypes isolated from raw milk, SM1–SM4 = genotypes isolated from skimmilk, C1–C45 = genotypes isolated from cheese). |
|---|---|---|---|
| Genotype | Raw milk (n = 134) | Skimmilk (n = 65) | Cheese (n = 50) |
| RM1 = SM1 = C44 | 3 | 42 | 27 |
| RM2 = SM2 | 1 | 16 | |
| RM3 | 112 | | |
| RM4 = C45 | 1 | 13 | |
| RM5 | 2 | | |
| RM6 | 2 | | |
| RM7 | 1 | | |
| RM8 = C32 | 4 | 10 | |
| RM9 | 1 | | |
| RM10 | 1 | | |
| RM11 | 2 | | |
| RM12 = SM3 | 1 | 6 | |
| RM13 | 2 | | |
| RM14 | 1 | | |
| SM4 | | | 1 |

Similarly to us, Berthier et al. [4] found that over 90% of the NSLAB isolates from the mature cheeses belonged to the species of *L. casei* or *L. rhamnosus* and were present at concentrations of $10^6$–$10^7$ cfu·g$^{-1}$ cheese after six months of ripening. We found between 1 and 8 genotypes per cheese, which is also similar to Comté. The genotypes from the three Gruyère cheese factories were different from one another, which is not surprising since the factories are situated at least 50 km apart.

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Raw milk was found to be far richer than skimmilk in the number of genotypes of NSLAB that it contained (Tab. VI). This is not surprising, since skimmilk was prepared in the factories early in the morning from
individual milks of the first delivering farmers and not from the combined raw milk of all suppliers. Of the four genotypes found in skim milk only one was different from those found in raw milk.

The three genotypes found in the cheese were also found in raw milk, thus indicating that it is the raw milk flora and not the factory environment that principally determines the composition of the NSLAB in Gruyère cheese. Similarly to us, Crow et al. [8] observed a seasonal variation in NSLAB in New Zealand Cheddar cheese manufactured in 6 factories at different times, showing that the flora of the factory equipment did not lead to a persistent NSLAB composition in the cheese. It is interesting to note that in the cheese we could not find the dominant genotype in raw milk, but genotypes found at much lower concentrations. It can be assumed that some genotypes did not survive the temperatures applied during the cheese-making process.

5. CONCLUSION

In conclusion, the present study has shown that NSLAB in Swiss Gruyère mainly derive from raw milk and that generally only a few strains survive the processing conditions and grow during ripening. In addition, the study has shown that with a continuous shift in the NSLAB composition in the ripened cheese over time. If individual strains of NSLAB were responsible for specific aroma development in Gruyère cheese this would imply that the aroma of cheeses from factory CH-4089 should not vary much over time, whereas cheeses from factory CH-4418 would vary from month to month. Since the three investigated factories produced at a constant high quality level, it seems therefore unlikely that any individual strains of NSLAB have a dominant impact on flavour development of Swiss Gruyère cheese. However, further studies are necessary to elucidate the impact of NSLAB on the ripening ability and sensorial properties of this well-known PDO cheese.

REFERENCES


