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To cite this version:
Anne Thierry, Marie-Bernadette Maillard, Romain Richoux, Jean-René Kerjean, Sylvie Lortal. Propionibacterium freudenreichii strains quantitatively affect production of volatile compounds in Swiss cheese. Le Lait, INRA Editions, 2005, 85 (1-2), pp.57-74. hal-00895594

HAL Id: hal-00895594
https://hal.archives-ouvertes.fr/hal-00895594
Submitted on 1 Jan 2005

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Propionibacterium freudenreichii strains quantitatively affect production of volatile compounds in Swiss cheese

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Abstract – Cheese flavour is the result of a complex mixture of volatile compounds, originating mainly from the enzymatic degradation of curd components by cheese microflora during cheese ripening. Directing cheese flavour development requires knowledge on inter- and intra-species contributions to flavour development, i.e. identification of the volatile (flavour) compounds produced by each strain. The aim of this study was to identify the volatile compounds produced in Swiss cheese by Propionibacterium freudenreichii, one of the species essential for the development of the characteristic flavour of this type of cheese. The volatile profile of compounds obtained from small-scale (1/100) Swiss cheeses, with or without P. freudenreichii, were compared (three strains tested, in association with three thermophilic lactic starters, i.e. twelve cheeses, manufactured in duplicate). Neutral volatile compounds, extracted by dynamic headspace, and free fatty acids were identified using gas chromatography-mass spectrometry. The concentrations of all carboxylic acids and 14 of 58 neutral compounds were significantly higher in the presence of propionibacteria (PAB). The three PAB strains tested produced the same volatile compounds, but observed quantitative differences were strain-dependent. Propionic acid and four propionate esters were detected only in the presence of PAB. Moreover, cheeses with PAB contained two- to three- fold higher levels of free fatty acids derived from lipolysis and five- to fifty- fold higher levels of branched-chain compounds derived from isoleucine catabolism (2-methylbutanal, 2-methylbutanol and 2-methylbutanoic acid) and from leucine catabolism (3-methylbutanoic acid). Lactic starters induced significant variations in the concentrations of some of the compounds produced by PAB, such as methylbutanoic acids and free fatty acids, which varied by 2.0 and 1.4, respectively, as a function of the lactobacilli strains. PAB strains affect the concentration of varied volatile compounds and could therefore have distinct contributions to the formation of Swiss cheese flavour.

Propionibacterium freudenreichii / volatile compound / Swiss cheese

Résumé – Les souches de Propionibacterium freudenreichii influencent la concentration de composés volatils variés dans l’emmental. Pour maîtriser la formation de la flaveur du fromage, qui est due à une multitude de composés volatils essentiellement produits par la microflore au cours de l’affinage, il est nécessaire de déterminer comment chaque espèce et souche contribue à la formation de composés d’arôme. L’objectif de cette étude était d’identifier les composés volatils produits dans l’emmental par Propionibacterium freudenreichii, une espèce clé du développement de la flaveur caractéristique de ce fromage. Douze mini-emmentals (échelle 1/100) ont été fabriqués en double, en associant P. freudenreichii (trois souches) et un témoin sans propionibactéries, à trois

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levains lactiques. Les composés volatils neutres de ces fromages et les acides gras libres ont été analysés par chromatographie en phase gazeuse. Seuls les fromages ensemencés en propionibactéries contenaient de l’acide propionique et quatre esters de propionate. Ces fromages contenaient également 2 à 3 fois plus d’acides gras libres et 5 à 50 fois plus de composés ramifiés volatils issus du catabolisme de l’isoleucine (2-méthylbutanal, 2-méthylbutanol et acide 2-méthylbutanoïque) et de la leucine (acide 3-méthylbutanoïque), avec des différences très significatives selon les souches. Les lactobacilles associés modulaient les concentrations de certains composés produits par *P. freudenreichii* (acides méthylbutanoïques et acides gras libres). Les souches de *P. freudenreichii*, en faisant varier la concentration de divers composés volatils, pourraient avoir des contributions distinctes à la formation de la flaveur de l’emmental.

**Propionibacterium freudenreichii** / composé volatil / emmental

1. INTRODUCTION

Mature cheeses contain a wide range of flavour compounds, including volatile and non-volatile compounds [13, 49]. The flavour-active compounds have been identified in several cheese varieties, including Emmental, Gruyere and Camembert cheese [24, 33, 34, 38]. Most of these compounds originate from the action of the cheese microflora and their enzymes on lactose, lactate, lipids and proteins of cheese curd [30]. Cheese is a microbial ecosystem that evolves throughout manufacture and ripening, where complex interactions between microflora generally occur [5]. Several species and strains may be involved in the pathway of formation of flavour compounds, as shown for the formation of volatile compounds derived from amino acid catabolism by lactic acid bacteria [1, 21]. To control or modify cheese flavour development, an understanding of the role of cheese microflora and of their interactions in the formation of flavour compounds is necessary.

The flavour of Swiss cheese is comprised of various compounds, such as furanones, short- to medium-chain acids, esters, diacetyl, and branched-chain (BC) aldehydes [33, 34, 39]. Two main flora are known to be essential for the development of the characteristic flavour of Swiss cheese: thermophilic lactic acid bacteria and propionibacteria (PAB) [26, 32]. Thermophilic lactic acid bacteria (streptococci and lactobacilli) are specifically involved in the fermentation of lactose into lactic acid which occurs during curd acidification, within the first day of manufacture. They also have a major contribution to cheese proteolysis, and the activity of their enzymes results in the release of free amino acids, which are the precursors of many flavour compounds. Moreover, lactic acid bacteria are also capable of forming various volatile compounds, and in particular, significant amounts of carbonyl compounds [17, 19, 35, 48, 51]. PAB have been known for a long time to contribute to the development of Swiss cheese flavour and cheese opening, achieved through the fermentation of lactic acid into propionic acid, acetic acid and CO$_2$ [6, 18, 40]. More recently, PAB were shown to play a prominent role in the formation of free fatty acids (FFA) and of short BC acids that result from triglyceride hydrolysis and from leucine/isoleucine catabolism, respectively [10, 41, 45]. The production in cheese of both FFA and short BC acids has been shown to be strain-dependent [10, 46]. However, the possible contribution of PAB to the formation of other volatile compounds has not been fully investigated [31]. Thierry et al. first studied the ability of one strain of *P. freudenreichii* to produce volatile compounds in cheese, by comparing the profiles of volatiles of experimental Swiss-type cheeses manufactured with or without addition of PAB, using headspace gas chromatography-mass spectrometry [45]. The authors showed the contribution of this strain to the formation of a variety of volatiles, such as alcohols, esters and ketones, which were present at significantly greater concentrations in the
presence of PAB than in the control cheeses without PAB. No data are available regarding the possible influence of LAB on PAB activity, although the existence of lactobacilli-PAB interactions in cheese has been reported [2, 9, 16, 31].

The aim of the present study was to determine the contribution of PAB to the formation of Swiss cheese volatile compounds and to evaluate the effect of PAB strains and of thermophilic lactobacilli-PAB interactions on the volatiles produced. To this end, three strains of *P. freudenreichii* and one control without PAB were each associated with three different thermophilic lactic starters in controlled-flora mini-Swiss cheeses, and the acidic and neutral volatile compounds quantified at two ripening stages.

2. MATERIALS AND METHODS

2.1. Cheese starters

All the lactic and propionic strains used in this study were obtained from the collection of the Institut Technique Français du Fromage (Rennes, France), except the starter LH100 (a mixture of strains of *Lactobacillus helveticus* and *Lactobacillus delbrueckii* subsp. *lactis*), which was obtained from Rhodia-Food (Dangé Saint-Romain, France). The other lactic starters used were *Lactobacillus helveticus* ITGLH56 and ITGLH77, *Lactobacillus delbrueckii* subsp. *lactis* ITGLL57 and *Streptococcus thermophilus* ITGST82 and ITGST87. Three *Propionibacterium freudenreichii* strains (ITGP14, ITGP17 and ITGP23) were chosen from previous screening experiments in mini-Swiss cheeses as they showed similar kinetics of propionic fermentation [36, 37] and different abilities to produce 2-methylbutanoic/3-methylbutanoic acids [47].

2.2. Cheese manufacture

Small-scale (1/100) experimental Swiss cheeses were manufactured from thermised and microfiltered milk according to a standardised cheese-making process previously described [36, 37, 46]. Cheeses were manufactured according to a factorial experimental design where two factors, lactic starter and propionic starter, were studied. Three lactic starters (LH100, ITGLH56 + ITGLL57 and ITGLH77, each associated with ITGST82 + ITGST87) were combined with three PAB strains (ITGP14, ITGP17 and ITGP23) and one control cheese without PAB, resulting in twelve starter associations. Two batches of each of the twelve cheeses (800 g) were manufactured on two different days, over a period of five weeks in December and January. Each cheese was brined on day one and then divided into one cheese block of 400 g and 8 sectors of 50 g before being wrapped under vacuum in a Cryovac BK1L film (Cryovac-Europe, Épernon, France). The cheese ripening conditions used were similar to those of industrial cheese-making, ripening at 12 °C for 21 d, then transferred to 24 °C (warm room) until 80% of the initial lactic acid had been utilised. To achieve this, the increase in cheese volume, determined by immersion of the cheese in water as previously described [37], was followed for the 400-g cheeses. As soon as their volume had increased by 2–3%, i.e., after 2–11 d depending on the PAB strains, two sectors were taken from the warm room, one for control of propionic fermentation and the other one placed at 4 °C for 8 weeks of cold ripening. The same operation was renewed three times at 2- to 3-d intervals for the 3 × 2 resting sectors. For each of the 24 cheeses, the four sectors sampled during the ripening in the warm room were analysed for their content in organic acids by HPLC as described hereafter. The samples containing approximately 20% of the initial content of lactic acid were chosen to be further analysed for FFA and neutral volatile compounds. The corresponding samples were taken from cold ripening after 8 weeks. All the samples were kept frozen at –80 °C until analysis.

2.3. Compositional analysis

Samples of ripened cheeses were analysed for moisture (oven drying at 103 °C),
for protein (Kjeldahl), fat (Heiss butyro-
metric method) and pH by classical methods
as previously described [37]. Carboxylic
acids (lactic, propionic and acetic acids)
determined in cheese homogenate, prepared
as described in 2.5 for neutral volatile
compound analysis, by HPLC on an Aminex
A-6 ion exchange column (Bio-
Rad, Hercules, CA, USA) at 55 °C with 0.01
NH$_2$SO$_4$ as eluent, at a flow rate of
1.0 mL·min$^{-1}$. Both UV (210 nm) and refrac-
tometric detectors were used. Acetic and
propionic acids were also determined by gas
chromatography as previously described [46].

2.4. Microbiological analyses

For microbiological analyses, performed
after 2 weeks of ripening in the warm room,
samples of cheeses (20 g) were dispersed in
180 g of a 2% sodium citrate solution, homog-
enised, diluted with peptone saline solution
and plated on specific media. Propionibacte-
ria were enumerated on lithium-glycerol
agar [42] incubated at 30 °C anaerobically
for 6 d. Non-starter lactic acid bacteria, which
can grow in cheeses during the ripening,
were enumerated on FH agar [20] incubated
at 30 °C anaerobically for 3 d and on MRS
agar incubated at 15 °C anaerobically for
14 d [46].

2.5. Determination of volatile
compounds

FFA (C$_{4:0}$ to C$_{20:1}$, including iC$_{5:0}$ +
aC$_{5:0}$ and conjugated linoleic acid) analyses
were performed by gas chromatography
(GC) by ITERG (Pessac, France) according
to De Jong and Badings [14]. Briefly, FFA
were extracted from the cheese with ether-
heptane after grinding with sodium sulphate
and addition of sulphuric acid, isolated from
lipids using an aminopropyl column and
analysed by GC under the following condi-
tions: cooler on-column injector, column:
QUADREX – FFAP capillary column,
30 m × 0.32 mm × 0.25 µm film thickness;
carrier gas: hydrogen, 1.1 bar; temperature
programme, heating rate: 10 °C·min$^{-1}$ from
50 °C up to 240 °C, maintained for 15 min,
flame-ionisation detector operated at 260 °C.
As the chromatographic conditions did not
efficiently separate 2-methylbutanoic acid
from 3-methylbutanoic acid, the proportion
of these two acids was obtained by deter-
mining the proportion of their respective
methyl ester. Methyl esters were synthe-
sised according to the method of Beck et al.
[3], with the following modifications: 1 mL of
cheese homogenate, 1 mL of a saturated solu-
tion of NaHSO$_4$ and 1 mL of a 205 mg·L$^{-1}$
methanol solution were mixed in a 9-mL
pyrex tube. The esterification reaction was
performed by placing the closed tube in a
water bath at 90 °C for 2 h. The two methyl
esters of methylbutanoic acids, i.e. methyl
2-methylbutanoate and methyl 3-methyl-
butanoate, were extracted by dynamic head-
space and analysed by gas chromatography
as described in the next section, and quanti-
tified by the ions 88 and 74, respectively.
Samples of control cheese homogenates
were spiked with 50 mg·kg$^{-1}$ of methyl-
butanoic acids containing varying propor-
tions of the two isomers (0:100, 25:75, 50:50,
75:25 and 100:0) and were subjected to
esterification as described above. Data were
used to determine the regression curve of
the proportion of acid isomers consisted of
2-methylbutanoic acid (y) and the propor-
tion of ester isomers consisted of methyl 2-
methylbutanoate (x). The equation obtained
(y = 1.047 x, R$^2 = 0.9928$) shows that the
proportion of the two ester isomers synthe-
sised fitted with the proportion of the two
acid isomers that were added.

Neutral volatile compounds were iden-
tified and quantified by dynamic headspace
GC-MS. Before analysis, cheese samples
were thawed and cut into cubes (2.5 mm ×
2.5 mm × 2.5 mm) which were mixed. A 15 g
sample was homogenised with 60 g of
0.5 mol·L$^{-1}$ sodium citrate solution by mixing
for 4 min at 20 500 rpm using an Ultraturrax
blender (Janke & Kunkel, Staufen, Germany).
A 7-g sample of this cheese homogenate
(± 0.05 g) was used for each headspace GC-
MS analysis. Each sample was analysed in
duplicate. Analyses were performed as
described in detail previously [43]. Briefly,
Cheese volatiles produced by propionibacteria

Volatiles were trapped on a Vocarb 3000 trap (Supelco, Bella Fonte, PA, USA), thermally desorbed at 250 °C and cryofocused at –100 °C, before being injected into a HP5890 (Agilent Technologies, Palo Alto, CA, USA) gas chromatograph – HP5972A quadrupole mass spectrometer (GC-MS). Volatiles were separated on a HP5 capillary column (60 m × 0.32 mm × 1.0 µm film thickness) under the following conditions: carrier gas: helium, 29 cm·s⁻¹ at 35 °C; temperature programme: 35 °C for 5 min, heating rate: 5 °C·min⁻¹ up to 140 °C then 15 °C·min⁻¹ up to 250 °C. MS was operated in the scan mode within a mass range of m/z 25–173 at 4.83 scan·s⁻¹, after ionisation by electronic impact at 70 eV. Thirty-three compounds were identified by comparison of mass spectra and retention times with those of authentic standards. Twenty-three other compounds, for which standards were not available, were tentatively identified on the basis of mass spectral data from the Hewlett Packard Chemstation NIST 75K mass spectral Database. Twenty-three neutral volatile compounds were quantified as follows. Peaks were quantified by the areas of either the total ion current (TIC) or selected fragments (m/z). To avoid the approximations related to the commonly used internal standard calibration [15], eight high purity chemicals (3-methylbutanal, 3-methylbutanol, ethyl acetate, ethyl propionate, ethyl butanoate, 2,3-butanedione, 2-heptanone and dimethyl disulphide), purchased from Sigma-Aldrich (St-Quentin-Fallavier, France) were used as external standards. Stock solutions of standard compounds were prepared in high purity (99.8%) methanol at concentrations of 4–7 mg·g⁻¹ and stored at –20 °C. Amounts of each stock solution were used to prepare a standard mix containing 50 to 2500 µg·g⁻¹ of each standard compound. An aliquot (~15 mg) of standard mix was accurately weighed and used to spike a 35-g sample of control cheese homogenate, resulting in final concentrations of 100 to 5000 µg·g⁻¹ cheese. Seven additional calibration standard solutions were prepared by further dilution (weight to weight) of the spiked cheese homogenate in blank cheese homogenate, in order to obtain eight different concentrations covering the following ranges: 0.8–100 ng·g⁻¹ cheese (dimethyl disulphide), 1.4–180 ng·g⁻¹ (3-methylbutanol), 8–1000 ng·g⁻¹ (2-heptanone), 17–2200 ng·g⁻¹ (ethyl acetate, ethyl propionate, ethyl butanoate, 3-methylbutanol), 40–5000 ng·g⁻¹ (2,3-butanedione). Five compounds (3-methylbutanol, 3-methylbutanal, ethyl propionate, 2,3-butanedione and 2-heptanone) were quantified from the regression curve of the corresponding standard. Eighteen other compounds were quantified from the regression curves of closely related standards (same chemical function, close molecular mass), as follows: 2-methylbutanol, 2-butanol and 2-pentanol from 3-methylbutanol; 2-methylpropanal, 2-methylbutanal and pentanal from 3-methylbutanal; methyl propionate from ethyl acetate; n-propyl acetate from ethyl propionate; isopropyl propionate and propyl propionate from ethyl butanoate; 2-butanol and 2,3-pentanedione from 2,3-butanedione; 4-methyl-2-pentanone, 2-hexanone, 6-methyl-2-heptanone, 5-methyl-2-heptanone and 2-octanone from 2-heptanone and dimethyl trisulphide from dimethyl disulphide.

2.6. Statistical analyses

Concentration data of each compound of duplicate cheeses were used for statistical analysis. Analyses of variance (ANOVA) were performed using the General Linear Model procedure of Statgraphics Plus (Statistical Graphic Corp., Englewood Cliffs, NJ, USA) to determine the effect of PAB strain, the effect of thermophilic lactic acid bacteria strains, and the effect of the interaction between PAB and lactic acid bacteria strains on PAB growth and on the concentration of each volatile compound. Differences between the treatment means were compared at the 5% level of significance using the Fisher’s least significance difference (LSD) test.

The concentrations of compounds which did not significantly discriminate between
cheeses ($P > 0.01$) were removed from the subsequent analyses. The data were then standardised (1/standard deviation) and Principal Component Analysis (PCA) was performed using Statbox (GrimmerSoft, Paris, France).

3. RESULTS AND DISCUSSION

3.1. Cheese rough composition and microbiology

The rough composition of the cheeses (62.4 ± 0.5% total solids, 45.6 ± 0.5% fat in dry matter and 52.5 ± 0.5% moisture in the non-fat substance) was consistent with the expected values for this type of mini-cheese [36, 46], and was not influenced by the lactic and propionic starters used (data not shown).

Non-starter lactic acid bacteria, enumerated on two different media in cheeses after 2 weeks of ripening in the warm room, were under detection level in half of the cheeses, irrespective of the starter added. Low numbers of non-starter lactic acid bacteria, ranging from $5 \times 10^1$ to $7 \times 10^4$ colony-forming units (cfu) per g cheese, were found in the remainder of the cheeses. Indigenous PAB were not detected in cheeses that were not inoculated with PAB (control cheeses).

3.2. Growth of propionibacteria and propionic fermentation

In cheeses inoculated with *P. freudenreichii*, PAB numbers ranged from $4 \times 10^9$ to $7 \times 10^9$ cfu·g$^{-1}$ after 14 d in the warm room. ANOVA showed that PAB numbers were significantly ($P < 0.001$) influenced by the lactobacilli strains used, but not by the strain of PAB and by PAB-lactobacilli interactions. The absence of the effect of the strain of PAB on their levels was expected as the three PAB strains used in the present study were chosen from strains showing similar kinetics of propionic fermentation in this type of mini-Swiss cheese [36, 37]. Levels of PAB detected were, on average, 50% higher in the presence of lactobacilli LH100 than in the presence of lactobacilli LH56-LL57; the third lactobacilli strain exhibited an intermediate behaviour in respect to its effect on PAB. The influence of lactobacilli was the same regardless of the strain of PAB used, which is in accordance with the absence of statistical PAB-lactobacilli interactions. The influence of thermophilic lactobacilli on PAB in cheese has been previously reported [2, 9, 16]. As a consequence of the influence of lactobacilli strains on propionic fermentation, using the same ripening period for cheeses containing different lactic strains can result in a bias of the results if the abilities of PAB strains to produce flavour compounds are to be compared.

The sampling procedures used in the present study were designed to counteract this bias, and cheeses were taken out of the warm room after a similar degree of propionic acid fermentation had been achieved, fixed at a level of ~80% of the initial lactic acid utilised. This level of propionic fermentation, which corresponded to the production of around 550–600 mg propionic acid per 100-g cheese, was achieved following a wide range of time periods in the warm room (11.8 ± 3.7 d). These time periods were not affected by PAB strain but significantly ($P < 0.05$) depended on the lactic starter, with a mean time in the warm room of 9.3, 11.3, and 14.8 d in the presence of starters LH100, LH77, and LH56-LL57, respectively. This result is in agreement with the observed effect of lactic starters on PAB growth in the present study, and with previous works showing that lactobacilli can induce marked differences in kinetics of propionate formation in Swiss cheese [2, 9, 16]. As expected from the sampling procedure, the concentrations of propionic and acetic acids did not differ significantly at the end of the warm room ripening (Tab. I). The average values of acetic and propionic acids in cheeses containing PAB were 211 ± 23.5 and 574 ± 45.7 mg·100 g$^{-1}$, respectively, versus 50 ± 15.6 and 1 ± 1.1 mg·100 g$^{-1}$, respectively, in the control cheeses.
Table I. Effect of three *Propionibacterium freudenreichii* strains (P17, P23 and P14) on the formation of acids in mini-Swiss cheeses.

<table>
<thead>
<tr>
<th>Acids (mg·100 g⁻¹)</th>
<th>At the end of warm room period</th>
<th>After two months’ storage at 4 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ctrl</td>
<td>P17</td>
</tr>
<tr>
<td>Acetic acid (C₂:0)  #</td>
<td>49.7 a</td>
<td>219.2 b</td>
</tr>
<tr>
<td>Propionic acid (C₃:0)</td>
<td>1.1 a</td>
<td>608.5 b</td>
</tr>
<tr>
<td>3-Methylbutanoic acid (iC₅:0)†</td>
<td>0.2 a</td>
<td>0.2 a</td>
</tr>
<tr>
<td>2-Methylbutanoic acid (αC₅:0)†</td>
<td>0.0 a</td>
<td>0.4 b</td>
</tr>
<tr>
<td>Butanoic acid (C₄:0)</td>
<td>1.5 a</td>
<td>3.2 b</td>
</tr>
<tr>
<td>Hexanoic acid (C₆:0)</td>
<td>0.6 a</td>
<td>2.3 b</td>
</tr>
<tr>
<td>Octanoic acid (C₈:0)</td>
<td>0.8 a</td>
<td>2.2 b</td>
</tr>
<tr>
<td>Capric acid (C₁₀:0)</td>
<td>1.8 a</td>
<td>4.3 b</td>
</tr>
<tr>
<td>Lauric acid (C₁₂:0)</td>
<td>2.6 a</td>
<td>6.5 b</td>
</tr>
<tr>
<td>Myristic acid (C₁₄:0)</td>
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<td>18.5 b</td>
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<td>Myristoleic acid (C₁₄:1)</td>
<td>0.7 a</td>
<td>1.7 b</td>
</tr>
<tr>
<td>Pentadecanoic acid (C₁₅:0)</td>
<td>1.0 a</td>
<td>3.3 b</td>
</tr>
<tr>
<td>Palmitic acid (C₁₆:0)</td>
<td>19.9 a</td>
<td>64.0 b</td>
</tr>
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<td>Palmitoleic acid (C₁₆:1)</td>
<td>2.4 a</td>
<td>5.4 b</td>
</tr>
<tr>
<td>Stearic acid (C₁₈:0)</td>
<td>6.8 a</td>
<td>15.6 b</td>
</tr>
<tr>
<td>Oleic acid (C₁₈:1)</td>
<td>18.3 a</td>
<td>47.5 b</td>
</tr>
<tr>
<td>Linoleic acid (C₁₈:2)</td>
<td>2.5 a</td>
<td>6.4 b</td>
</tr>
<tr>
<td>Linolenic acid (C₁₈:3)</td>
<td>0.7 a</td>
<td>1.1 ab</td>
</tr>
<tr>
<td>Conjugated linoleic acid ‡</td>
<td>0.4 a</td>
<td>1.1 b</td>
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<tr>
<td>Arachidic acid (C₂₀:0)</td>
<td>0.2 a</td>
<td>0.2 a</td>
</tr>
<tr>
<td>Gadoleic acid (C₂₀:1)</td>
<td>0.3 a</td>
<td>0.6 b</td>
</tr>
<tr>
<td>Unsaturated FFA #</td>
<td>25.2 a</td>
<td>63.8 b</td>
</tr>
<tr>
<td>Saturated C₄:0-C₈:0</td>
<td>3.0 a</td>
<td>7.7 b</td>
</tr>
<tr>
<td>Saturated C₁₀:0-C₂₀:0</td>
<td>38.5 a</td>
<td>112.3 b</td>
</tr>
<tr>
<td>Total free fatty acids</td>
<td>66.7 a</td>
<td>183.8 b</td>
</tr>
</tbody>
</table>

*a All acids were analysed by GC, but acetic and propionic acids were analysed separately from the other acids, as described in the Materials and Methods section.
†: 2- and 3-methylbutanoic acids were calculated as follows: concentration of (2-methylbutanoic + 3-methylbutanoic acids) × proportion of each isomer of methylbutanoic acid. The proportions of acids were determined from the proportion of the corresponding methyl esters, which were synthesised and analysed as described in the Materials and Methods section.
‡: CLA constituted a mixture positional and geometric isomers of conjugated linoleic acid.
#: Sum of C₁₄:1, C₁₆:1, C₁₈:1, C₁₈:2 including CLA, C₁₈:3, and C₂₀:1.
*p Results are means of six experiments (averaged values across three lactic starters and duplicates). Values in the same row at the same ripening stage with the same superscript were not significantly different according to the LSD test (α<0.05).
© Ctrl: control cheese manufactured under the same conditions without *P. freudenreichii*.

Probability of F-test: *** P < 0.001; ** 0.001 < P < 0.01; * 0.01 < P < 0.05; NS, P > 0.05.
Both acids were further produced during the cold storage, with values of 272 ± 21.7 and 721 ± 27.8 mg·100 g−1, respectively, in the presence of PAB and 66 ± 16.0 and 1 ± 0.5 mg·100 g−1, respectively, in the control cheeses, at the end of ripening. These values are in the range of concentrations reported in Swiss cheese but the molar ratio of propionate and acetate that can be calculated from these concentrations was higher in the present study (2.2 in cheeses containing PAB) than in previous reports (0.5 to 2.1) [4, 7, 23, 25, 34, 39]. From the concentrations observed in the present study in the control cheeses, manufactured in the absence of PAB, it can be estimated that the total propionic acid and most (75%) of the acetic acid produced resulted from PAB metabolism. Taking these data into account, the molar ratio of propionate and acetate produced by propionibacteria could be evaluated as 2.8.

### 3.3. Production of short branched-chain fatty acids

Both 2-methylbutanoic acid and 3-methylbutanoic acid (also referred to as isovaleric acid) were produced in the cheeses. Using a simple esterification method, we showed that the peak formed by the co-elution of both isomers of methylbutanoic acids actually constituted 58 to 88% of 2-methylbutyric acid in cheeses containing PAB, and ~35% in the control cheeses. 2-Methylbutyric acid was essentially produced by PAB, with concentrations 12 to 50 times higher in cheeses containing PAB than in the control cheeses. Significant differences ($P < 0.001$) of 2-methylbutanoic acid production were observed between PAB strains, at both ripening stages (Tab. I), with strains ITGP14 and ITGP17 producing the highest and the lowest levels, respectively. The same classification of PAB strains was observed for the production of 3-methylbutanoic acid (Tab. I). These results confirm, using the same PAB strains but different lactic starters, previous observations in mini-Swiss cheeses which showed that 2-methyl/3-methylbutanoic acids were mainly produced by PAB [46]. From the concentrations of methylbutanoic acids detected in the control cheeses, it could be estimated that thermophilic lactic starters directly contribute to the production of only ~3% and 20% of the total levels of 2-methylbutanoic and 3-methylbutanoic acids produced, respectively, in cheeses. Thermophilic lactic starters also had an indirect contribution to the formation of BC acids, by modulating the production of these acids by PAB. Hence, the concentrations observed in the presence of starter LH77 were significantly ($P < 0.05$, 2-methylbutanoic acid and $P < 0.001$, 3-methylbutanoic acid) higher than the ones observed for starter LH56-LL57 (Fig. 1). The main production of short BC acids (59 ± 14% and 61 ± 13% of 2-methyl- and 3-methylbutanoic acid, respectively) occurred during the cold storage in all cheeses, as was previously observed [47]. Methylbutanoic acids impart a cheesy/sweaty note to cheese and are thought to play a role in cheese flavour [49, 51]. In Swiss cheese, 3-methylbutanoic acid was found to contribute to cheese taste, whereas 2-methyl-butanoic acid was not cited [34].

### 3.4. Production of free fatty acids

The concentrations of all the FFA derived from lipolysis were 2 to 8 times higher in the presence of *P. freudenreichii*, depending on the strain and the fatty acid considered (Tab. I). The strain *P. freudenreichii* ITGP23 produced the highest concentration of FFA throughout ripening (Tab. I, Fig. 1). Palmitic (C$_{16:0}$), oleic (C$_{18:1}$), myristic (C$_{14:0}$) and stearic (C$_{18:0}$) acids were the most abundant acids in all the cheeses, constituting 45–50% of FFA, as in milk fat [12]. This result indicates that FFA essentially arise from a non-specific hydrolysis of milk triglycerides, in agreement with previous results [26]. It should be noted, however, that the three PAB strains induced a greater release of medium-chain and long-chain saturated FFA (C$_{14:0}$–C$_{15:0}$, C$_{16:0}$ and C$_{18:0}$). They were five- to eight-fold higher in the presence of PAB than in the control cheeses,
Figure 1. Production of volatile compounds in mini-Swiss cheeses made using different combinations of lactobacilli starters (L1, LH56-LL57; L2, LH77; L3, LH100) and propionibacteria (three strains of *P. freudenreichii*, ITGP17, ITGP23 and ITGP14, and a control cheese without PAB). Production of compounds is shown at the end of ripening in the warm room (open bars) and during a subsequent cold storage period for 8 weeks at 4 °C (hatched bars). Values are means of duplicate batches of cheeses. A, 2-methylbutanoic acid; B, 3-methylbutanoic acid; C, sum of short-chain (C4:0 - C8:0) and unsaturated (C14:1 - C20:1) FFA; D, sum of medium- and long-chain saturated FFA (C10:0 - C20:0); E, ethyl propionate; F, ethyl butanoate.
compared with the release of short-chain (C₄:0, C₆:0 and C₈:0), and unsaturated FFA (C₁₆:1, C₁₈:1, C₁₈:2 and C₁₈:3), which showed a two- to three-fold increase in the presence of PAB. From values observed in the control cheeses, it could be assumed that lactic starters produced 16% of saturated FFA, 32% of unsaturated FFA, and 41% of short-chain FFA of the total amounts found in the cheeses. This study in controlled-flora cheeses confirms the initial observations of Chamba et al. studying full-size Emmental cheeses, who showed that PAB lipolytic activity was the main factor of lipolysis in Swiss-type cheeses [10]. However, lactobacilli starters also had an indirect contribution to the formation of FFA, by modulating the effect of PAB on lipolysis. The concentrations of long-chain saturated FFA released, in particular, were significantly \((P < 0.05)\) higher (+31%) in the presence of lactobacilli strains LH56-LL57 than in the presence of lactobacilli strain LH100, as shown in Figure 1. This effect could be related to the time period in the warm room, which was 60% longer, on average, for cheeses made using LH56-LL57 than for cheeses made using LH100, as noted above. The greatest production of FFA was during the warm room ripening stage, but significant levels were still produced at 4 °C (41 ± 13%, 25 ± 4%, and 25 ± 15% of long-chain saturated FFA, short-chain FFA, and unsaturated FFA, respectively). Short- and medium-chain FFA are considered as important contributors to cheese flavour, even though levels of lipolysis of less than 1% have been reported in good quality Swiss cheeses [10, 34].

3.5. Neutral volatile compounds

Fifty-six neutral volatile compounds were identified in the cheeses by headspace GC-MS (Tabs. II and III). All compounds detected have previously been reported in Swiss-type cheeses [8, 29, 43, 45] except 3-heptanone, which has been reported in butter and in other cheese varieties such as Parmesan [29]. The concentrations of twenty-three volatiles were significantly influenced by the presence of PAB. Nine volatile compounds were detected at greater concentrations in the presence of each of the three PAB strains, and six compounds in the presence of one or two PAB strains (Tab. II). The main groups of neutral volatile compounds detected at a higher concentration in the presence of PAB than in the control cheeses were esters of propionate and some BC compounds. In contrast, eight compounds were detected at lower concentrations in the presence of PAB: two alcohols (2-butanol and 2-pentanol), three aldehydes (2-methylpropanal, 3-methylbutanal and pentanal), two ketones (2,3-butandione and 2-heptanone) and dimethyl trisulphide (Tab. II).

Five of the sixteen esters detected in the cheese were more abundant in the presence of PAB: four esters of propionate, and propyl acetate (Tab. II). The presence of propionate esters, as expected, was strictly dependent on the presence in the cheese of propionic acid, i.e. of PAB. Ethyl propionate was by far the most abundant ester in mini-Swiss cheese, as previously reported in this type of mini-cheese [45]. The concentrations of ethyl propionate were similar with the three PAB strains, but were significantly \((P < 0.05)\) influenced by the strains of lactobacilli, with the highest concentrations detected in cheeses manufactured using starter LH56-LL57 (Fig. 1). Similarly, two ethyl esters, ethyl hexanoate and ethyl heptanoate, were detected at significantly \((P < 0.05)\) higher levels in the presence of starter LH56-LL57 than in the presence of starter LH77 (data not shown). All the identified esters were synthesised mainly during the cold storage of the cheese (70–81% of the total amount produced), as shown for ethyl propionate and ethyl butanoate in Figure 1. The observation of similar kinetics of ester synthesis in the absence and in the presence of PAB, as observed for ethyl butanoate synthesis, suggests that enzymes of lactic acid starters are involved in their synthesis. PAB are therefore not necessarily directly involved in the
### Table II. Effect of three strains of *Propionibacterium freudenreichii* (P17, P23 and P14) on the concentration of neutral volatile compounds in mini-Swiss cheeses.

<table>
<thead>
<tr>
<th>RT (min)</th>
<th>Compound (ng·g⁻¹)</th>
<th>QI</th>
<th>At the end of warm room period</th>
<th>After two month’s storage at 4 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ctrl e</td>
<td>P17</td>
<td>P23</td>
<td>P14</td>
</tr>
<tr>
<td><strong>alcohols</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.92</td>
<td>2-Butanol</td>
<td>45</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>11.90</td>
<td>2-Pentanol</td>
<td>45</td>
<td>3.9 b</td>
<td>0.6 a</td>
</tr>
<tr>
<td>13.63</td>
<td>3-Methylbutanol †</td>
<td>70</td>
<td>26.5 a</td>
<td>31.5 a</td>
</tr>
<tr>
<td>13.75</td>
<td>2-Methylbutanol</td>
<td>57</td>
<td>3.4 a</td>
<td>70.2 b</td>
</tr>
<tr>
<td><strong>aldehydes</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.29</td>
<td>2-Methylpropanal</td>
<td>72</td>
<td>1.6</td>
<td>0.9</td>
</tr>
<tr>
<td>10.00</td>
<td>3-Methylbutanal †</td>
<td>TIC</td>
<td>8.5 b</td>
<td>2.2 a</td>
</tr>
<tr>
<td>10.42</td>
<td>2-Methylbutanal</td>
<td>TIC</td>
<td>2.7 a</td>
<td>17.3 b</td>
</tr>
<tr>
<td>11.84</td>
<td>Pentanal   #</td>
<td>44</td>
<td>2.1 a</td>
<td>1.2 a</td>
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<td></td>
<td></td>
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<tr>
<td>9.02</td>
<td>Methyl propionate #</td>
<td>88</td>
<td>0.0 a</td>
<td>3.7 b</td>
</tr>
<tr>
<td>12.51</td>
<td>Ethyl propionate †</td>
<td>TIC</td>
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<td>27.6 a</td>
</tr>
<tr>
<td>12.64</td>
<td>n-Propyl acetate</td>
<td>43</td>
<td>0.2</td>
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<tr>
<td>14.40</td>
<td>Isopropyl propionate #</td>
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<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>16.79</td>
<td>Propyl propionate</td>
<td>57</td>
<td>0.02</td>
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<tr>
<td><strong>ketones</strong></td>
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<tr>
<td>7.29</td>
<td>2,3-Butanedione †</td>
<td>86</td>
<td>387</td>
<td>197</td>
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<tr>
<td>7.60</td>
<td>2-Butanone</td>
<td>72</td>
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<td>278 a</td>
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<td>11.74</td>
<td>2,3-Pentanedione #</td>
<td>100</td>
<td>1.9 a</td>
<td>12.1 b</td>
</tr>
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<td>13.77</td>
<td>4-Methyl-2-pentanone</td>
<td>43</td>
<td>1.5 a</td>
<td>1.8 a</td>
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<tr>
<td>15.95</td>
<td>2-Hexanone</td>
<td>TIC</td>
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<td>9.9 ab</td>
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<td>TIC</td>
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<td>104.6</td>
</tr>
<tr>
<td>22.53</td>
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<td>0.1</td>
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<tr>
<td>23.01</td>
<td>5-Methyl-2-heptanone #</td>
<td>43,58</td>
<td>0.0 a</td>
<td>1.0 bc</td>
</tr>
<tr>
<td>23.95</td>
<td>2-Octanone #</td>
<td>58</td>
<td>1.7</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>23.65</td>
<td>Dimethyl trisulphide</td>
<td>126</td>
<td>1.1 b</td>
<td>0.3 a</td>
</tr>
</tbody>
</table>

a RT, retention time in GC-MS.
b Values are concentrations expressed in ng·g⁻¹. Five compounds, marked with †, were quantified by a regression curve obtained using authentic standards, and the other compounds quantified using the regression curve of a closely related standard, as described in Materials and Methods. Compounds marked with # were tentatively identified by comparison of mass spectral data with those of the NIST 75K database.
c QI, quantification ions: TIC, total ion current peak areas were used or selected fragment(s) as indicated. Results are means of six experiments (averaged values across three lactic starters and duplicates). nd, not detected. Values in the same row at the same ripening stage with the same superscript were not significantly different according to the LSD test (α < 0.05).
d Ctrl: control cheese manufactured under the same conditions without *P. freudenreichii*.

### Notes
- *P < 0.001; ** 0.001 < P < 0.01; * 0.01 < P < 0.05; NS, P > 0.05.*
The synthesis of esters of propionate, but may only act as providers of the acid moiety (as propionic acid or propionyl CoA). Esters are common volatile compounds in cheese, but their mechanism of synthesis in cheese is still largely unknown [28]. Esters are responsible for fruity flavours that can be considered as a flavour defect or contribute positively to cheese flavour [28]. In Swiss cheese, esters are thought to contribute to the fruity note [33, 34], and their presence has also been associated with the sweet odour of this cheese [27].

Numerous BC compounds were detected at a higher concentration in cheeses made using PAB than in the controls (Tab. II). They include products from the catabolism of isoleucine (2-methylbutanol and 2-methylbutanal) and three BC ketones. PAB were the main producers of isoleucine-derived products. Control cheeses without PAB contained only 10–20% and 4–15%, respectively,
of the amounts of 2-methylbutanol and 2-methylbutanal found in cheeses containing PAB, irrespective of the lactic starter used. It should be noted that products of leucine catabolism (3-methylbutanol and 3-methylbutanal) reflected different patterns from products of isoleucine catabolism. Concentrations of 3-methylbutanol in cheeses containing PAB did not significantly differ by the end of ripening from the controls, whereas levels of 3-methylbutanal detected were lower in cheeses containing PAB.

Significant differences ($P < 0.01$) in the production of BC compounds were observed between PAB strains, at both ripening stages (Tab. II). Strain ITGP14 produced the highest concentrations of 3-methylbutanol and 2-methylbutanal, as noted above for the production of BC acids. This strain also produced two BC ketones, 4-methyl-2-pentanone and 5-methyl-2-heptanone. The origin of BC ketones has not, to the authors’ knowledge, been investigated, but these ketones have already been related to the presence of PAB [45]. The fact that their levels have been correlated with those of some other BC compounds suggests that BC ketones could also originate from BC amino acid catabolism.

Thermophilic lactobacilli contributed only slightly to the direct production of products originating from isoleucine catabolism. However, they significantly ($P < 0.05$) influenced the production of 2-methylbutanol by PAB, which was ~20% higher in cheeses made using starter LH77 than in cheeses made using both other lactic starters (data not shown). This stimulation of the production by this strain of lactobacilli had been observed above for the production of BC acids (Fig. 1). The mechanisms by which lactobacilli influenced the production of BC compounds by PAB have not been elucidated. Thermophilic lactobacilli are known to play a main role in the release of peptides and free amino acids in Swiss cheese, through the activity of their numerous aminopeptidases [11], and this activity is enhanced by cell lysis [50]. They are also capable of converting amino acids into various BC volatile compounds in vitro, and in particular catalyse transamination reaction, which is the first step of amino acid catabolism in cheese [17, 22, 51]. Therefore, the stimulatory effect that LH77 exerted on the synthesis of BC volatile compounds by PAB may have been due to its contribution to the release of BC precursors of volatile compounds, which may be amino acids or keto acids resulting from transamination. However, as neither proteolysis, nor lactobacilli lysis, nor enzymic activities were determined in this study, no conclusion can be drawn on this point.

BC alcohols and aldehydes constituted only minute amounts of the total amounts of BC compounds detected (~1% and 0.1%, respectively, the remainder being BC acids). As both BC alcohols and BC aldehydes can be converted to their corresponding BC acids by cheese bacteria [44, 51], they probably constitute only transitory compounds in Swiss cheese. BC aldehydes are characterised by their malty flavour, and 3-methylbutanal is considered as an odourant in Swiss cheese [33, 34]. BC alcohols, like other alcohols identified in cheese, are believed not to have a direct influence on cheese flavour [26]. They can, however, be indirectly involved as precursors in ester formation. Only trace amounts of esters derived from BC alcohols were detected in the present study, but BC esters have been previously reported in Swiss-type cheeses [8, 29, 45]. In particular, the presence of BC esters of 2-methylbutanol and 3-methylbutanol (2-methyl and 3-methyl acetate, 2-methyl and 3-methyl propionate) was previously reported in mini-cheeses made using PAB strain ITGP22, although the concentrations of acids and BC alcohols were very similar in this study and the present study [45]. This result suggests that some PAB could have original abilities to synthesise BC compounds, and that these abilities are strain-dependent.

Thirty-three other neutral volatile compounds were identified in the cheeses, but
their concentrations were not significantly influenced by the presence of PAB or by the lactic starter used (except for the two ethyl esters mentioned above, ethyl hexanoate and ethyl heptanoate). These compounds include most alcohols, most aldehydes (except the BC ones), several ketones and the esters, not including those of propionate (Tab. III).

**3.6. Global fingerprint of propionibacteria**

The factors responsible for the similarities and the differences between cheese profiles of volatile compounds were identified by Principal Component Analysis (PCA). PCA was performed on the concentrations of the compounds listed in Tables I and II that ANOVA found significantly ($P < 0.01$) discriminated between the cheeses. As the data for individual FFA were highly correlated, only the sum of FFA was included in the data set for the first two PCA performed (Figs. 2 and 3). In the case of the third PCA, the sum of FFA, the sum of short-chain FFA (C4:0 to C12:0), and the sum of long-chain FFA (from C14:0 to C20:1) were included in the data set (Fig. 4). PCA was performed on data obtained at the end of the warm period room and at the end of cold storage, and the first two components (PC1 and PC2) are represented graphically in Figure 2 and Figure 3, respectively. A cumulative variation of 74% and 61%, respectively, were explained by the first two PCs at each of these two stages. In both PCAs performed, PC1 separated the control cheeses from those containing PAB, and accounted for 54% and 49% of the variation, respectively.
Cheese volatiles produced by propionibacteria

in PCA performed at each of the two stages of ripening (Figs. 2 and 3). Cheeses containing PAB were characterised by high levels of propionic acid, acetic acid, esters of propionic acid, FFA, and most BC compounds, in particular those arising from isoleucine catabolism (2-methylbutanoic acid, 2-methylbutanol and 2-methylbutanal) (Fig. 2). At the end of ripening, additional compounds, such as BC ketones (4-methyl-2-pentanone, 6-methyl-2-heptanone), 2,3-pentanedione, and propyl propionate also associated positively with PC1 (Fig. 2). PC2 differentiated cheeses made using strain ITGP14 from the cheeses made using the other two PAB strains, and accounted for 20% and 12% of the variation, in the PCA performed on data at the end of the warm room period and at the end of cold storage, respectively. Cheeses made using ITGP14 were characterised on PC2 by a high content of most BC compounds, in particular products derived from leucine catabolism (3-methylbutanoic acid and 3-methylbutanol). To emphasise the differences between cheeses containing PAB, a third PCA was performed by excluding the data obtained for control cheeses. The first two PCs accounted for 49% of the experimental variance obtained. PC1, representing 27% of the variation, separated the cheeses made using starter LH56-LL57 from the cheeses made using the other two lactic starters. Cheeses manufactured using starter LH56-LL57 were characterised on PC1 by a high content of FFA and ethyl propionate. PC2, representing 22% of the variation, separated the cheeses made using ITGP14 from the cheeses made using the other two PAB strains on the basis of their content in BC compounds originating from isoleucine catabolism (Fig. 4).

4. CONCLUSIONS

The presence of PAB in cheese resulted in an increase in a variety of products which originate from various sources, such as fermentation of lactic acid, lipolysis, BC

![Figure 3. Results of principal component analysis on the volatile profile of mini-cheeses at the end of cold storage. Biplot of PCA loading and scores for the first two principal components (PCs). Cheeses codes: see Figure 2. Variables: see Figure 2 legend and: mpropanal, 2-methylpropanal; prop-propp, propyl propionate; 6mheptanone, 6-methyl-2-heptanone; 2-butanol; acetone (2-propanone); diacetyl, 2,3-butanedione; pdione, 2,3-pentanedione.](image-url)
amino acid catabolism and other pathways. These results confirmed previous results obtained in mini-cheeses made using another PAB strain [45]. BC compounds originating from isoleucine catabolism (2-methylbutanoic acid, 2-methylbutanol and 2-methylbutanal) were detected in higher concentrations than the corresponding products originating from leucine catabolism in the presence of PAB, whereas the opposite was observed in the control cheeses. The presence of high levels of 2-methyl-branched-chain compounds appears as a characteristic feature of Swiss-type cheeses, in comparison with other internal bacterially ripened cheese varieties, as previously suggested [25, 41].

PAB produced volatile compounds in the cheeses both during their growth at 24 °C and during the subsequent cold storage period at 4 °C. The proportions of volatile compounds produced during the warm and cold storage periods, respectively, greatly differed depending on their origin. Hence, ~20% of propionic and acetic acids, ~25% of short-chain and ~41% of long-chain FFA derived from lipolysis, ~60% of BC acids originating from isoleucine/leucine catabolism, and ~80% of esters were produced during the cold storage period. These results are in agreement with previous studies in full-size Emmental cheese, which showed that esters were the compounds showing the largest increase (×5–7) from 3 to 12 months of ripening, followed by 2-heptanone (five-fold increase) and short-chain FFA (two- to three-fold increase) [39].

This study clearly shows that the concentrations of volatile compounds produced by P. freudenreichii originating from lipolysis and BC amino acid catabolism are strain-dependent. Several of these volatiles could play a role in the formation of cheese flavour. Sensory evaluation now has to be performed to find out if the flavour profiles of the cheeses are perceived as different, and in which attributes they differ.

Figure 4. Results of principal component analysis on the volatile profile of mini-cheeses containing propionibacteria, at the end of cold storage. Biplot of PCA loading and scores for the first two principal components (PCs). Cheeses codes: see Figure 2 (control cheeses without PAB not included). Variables: see Figure 2 and 3 legends and SCFFA, short-chain FFA; LCFFA, long-chain FFA.
Acknowledgements: This work was supported by funds provided by the Brittany Region. We are grateful to Dr. J.A. Hannon for critical reviewing of the manuscript, and we thank G. Roset and L. Aubert for skilled cheese manufacture.

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