Fermented Dairy Product for a Low-Fat Bakery Product Application: Chemical and Sensory Analysis

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Abstract


Whey was fermented using different lactic acid bacteria in order to obtain an ingredient as a substitute for butter in pastry products. Lactococcus lactis subsp. lactis biovar diacetylactis was selected based on its capacity to produce buttery flavours like diacetyl. Fermentation parameters such as temperature, aeration, citrate addition, and the duration of each fermentation step were selected in order to boost diacetyl production. Fermented whey was then added as an ingredient to white sandwich bread which did not initially contain any butter in the recipe. Sensory analysis brought to light that the buttery aroma was under the perception threshold in the bread containing the mixture even though the fermented whey contained diacetyl.

Keywords: Lactococcus lactis subsp. lactis biovar diacetylactis; buttery flavour; diacetyl, butter; pastry product

Over recent decades, campaigns in European and the other Western countries have underlined the nutritional impacts of modern food on consumer health. Indeed, world organisations, such as WHO or FAO, and scientific groups, have recommended the setting up of nutritional strategies to improve the overall health of a country’s population. One important programme set up in France in 2001 is the Programme National Nutrition Santé – PNNS (national nutrition and health programme)(Hercberg et al. 2008). In this context, several food companies signed an agreement with the government to certify, for instance, their engagement to reduce the energy content of their foodstuffs.

In pastry products, butter is one of the most important ingredients, generating pleasant organoleptic characteristics. Besides having the high lipid content, butter also contains a lot of aroma compounds (Bennion & Bamford 1997). Consumers may be put off choosing alternative low-fat products without butter because of the alteration this implies to their organoleptic characteristics: they simply do not recognise the product they have purchased. In order to prevent the deterioration of organoleptic characteristics as a consequence of butter reduction in the recipe, food producers supplement their formulas with flavourings such as diacetyl, a molecule known to be a key compound in the butter aroma. But too frequently, this artificial diacetyl flavouring fails to restore a pleasant nutty-creamy buttery aroma (Buhler et al. 1972). In addition, negative publicity over the years has given flavourings a bad name, but they must nonetheless appear on all foodstuff packaging in line with the European Regulation EU/1334/2008. To overcome these difficulties, the addition of fermented ingredients to the recipe could be an alternative to reducing butter. These fermented products are thought to have the capacity to improve the buttery aroma thanks to the action of microorganisms. Diacetyl is produced for instance by Lactococcus lactis subsp. lactis biovar. diacetylactis (but also by many Leuconostoc strains, yeast, and even many Gram-negative bacteria) and results from the citrate metabolism. Besides diacetyl,
other aroma compounds originating from the bacterial metabolism contribute to the pleasant buttery aroma (LONGO & SANROMÁN 2006). This solution could prevent aroma degradation during the different steps of the baking process, especially cooking, thanks to the formation of interactions between the components of the matrix and the buttery aroma molecules – namely diacetyl.

The aim of this work was to study the effect on the aroma characteristics of white sandwich bread (also called in France pain de mie) of adding a buttery aroma mixture – resulting from the citrate fermentation of a food matrix by two Lactococcus strains – to the recipe.

**MATERIAL AND METHODS**

**Bacterial strains.** Two Lactococcus lactis subsp. lactis biovar. diacetylactis strains were used. One of the two lactococci was selected from a pool of citrate positive Lactococcus strains picked up in traditional fermented dairy products and the other was a citrate positive selected starter. The first strain, named F2.3, was isolated from raw goat whey, phenotypically characterised and identified by 16S rRNA gene sequencing. The second strain, named Dx1, was a commercial strain used for producing buttery aroma in dairy fermented products. This strain was used as a control. The strains were kept frozen at –80°C (Thermo Fisher Scientific, Waltham, USA) in a mixture of culture medium (M17 broth; Biokar, Beauvais, France) and glycerol 30% (Sigma-Aldrich, St-Quentin Fallavier, France).

**Lactococcus growth conditions, pH analysis, and diacetyl production.** The food matrix, composed of a sweet whey solution (10%, w/v; Eurosérum, St Martin Belle Roche, France), was prepared and aliquoted in 1 l flasks (500 ml/flask). This volume enabled to optimise the surface-to-volume ratio, thus maximising the oxygenation of the whey solution during the later step of fermentation. This solution was sterilised at 110°C during 10 minutes. Two flasks were each seeded with one strain of Lactococcus lactis subsp. lactis biovar. diacetylactis, F2.3 or Dx1, at the level of 6 log (CFU)/ml (Figure 1). Prelfermentations were performed aerobically during 16 h at 25°C by shaking the whey solution vigorously with a magnet bar rotating at 1100 rpm (RCT Basic magnetic stirrer; IKA, Staufen, Germany). At the end of the pre-fermentation step, the pH was adjusted to 4.5 (PHM210 standard, MeterLab®, Villeurbanne, France) and anhydrous trisodium citrate (1%, w/v; Quimdis, Levallois-Perret, France) was added to the pre-fermented whey solution. During the second step of citrate fermentation, the whey mixture was again vigorously shaken for 24 h at 20°C. During this second step, it was assumed that large amounts of diacetyl and other flavourings were produced via the citrate metabolism. In this article, the fermented whey mixtures are referred to as diacetyl.F2.3 and diacetyl.Dx1 for the products obtained with the strains F2.3 and Dx1, respectively. These cultures were then quickly cooled down in frozen water and the pH was recorded. Each culture was split into two parts: one used for the extraction of volatile components, the other for preparing loaves of white sandwich bread. Each fermentation was performed in triplicate.

**General procedure of diacetyl measurement.** The extraction of diacetyl was performed using acetone. One millilitre of the diacetyl mixture sample was mixed with 1 ml of acetone and shaken vigorously for 30 s at 3000 rpm (Vortex agitator; IKA, Staufen, Germany). After centrifugation at 4000 g for 5 min (5417R microcentrifuge; Eppendorf, Montesson, France), the supernatant was removed for specific diacetyl quantification. This fraction was kept at –80°C for less than 7 days. After filtration on a cellulose acetate filter (0.2 µm), the supernatant was directly injected into the gas chromatography apparatus.

Gas chromatography analysis of diacetyl was performed using a gas chromatograph equipped with a flame ionisation detector (GC-FID) (Perkin Elmer Autosystem GC, Waltham, USA) (MACCIOLA et al. 2008). A capillary column (30 m x 0.32 mm, 0.25 µm film thickness) with a 100% polyethylene glycol phase (ZB-Wax column; Phenomenex, Torrance, USA) was used. Gas chromatography parameters were as follows: helium carrier gas (1.5 ml/min), injected amount 5 µl, split mode injection at 1 : 10 splitting ratio, injector and detector temperatures 240 and 255°C, respectively; the oven temperature ran from 50°C to 240°C at 7°C/min with three isotherms: 2 min at 91°C, 3 min at 107°C, and 3 min at 186°C. The amount of detected molecules was expressed in surface area units (mV/min) calculated with the Azur Version 5.0.10.0 software (Launaguet, France). The means and the standard deviations were calculated with the data obtained from three replications with the same strain culture.

**White sandwich bread preparation.** The control white sandwich bread (named thereafter bread.Ctrl)
was composed of 1000 g of white flour (T450), 250 g of water, 50 g of sugar, 30 g of fresh yeast, 30 g of neutral colza oil, and 10 g of salt. Test breads were made following the same recipe except that 100 g of water were replaced by the equivalent volume of the diacetyl F.2.3 and diacetyl Dx1 preparations. These breads will be named bread F.2.3 and bread Dx1 hereafter.

The ingredients were mixed prior to kneading (SPI 11; VMI Vendée Mécanique Industrie, Montaigu, France) for 15 min at 120 rpm. The dough was left to rest in bulk for 20 min, scaled into 450 g portions, moulded in a small Exoglass® mould (base dimensions: 250 × 80 × 85 mm, lid dimensions: 290 × 95 mm) (Machinefabriek Holtkamp BV, Almelo, Netherlands), and placed in the proofer (BFA; Bongard, Strasbourg, France) set to 28 ± 1°C and 75% relative humidity for 105 minutes. Baking was carried out at 170°C for 30 min in a deck oven (2000; Jolivet, Gommeignies, France). The core temperature of the loaf at the end of the baking stood at 99°C (digital probe thermometer, Kat. Nr. 30.1018; TFA, Düsseldorf, Germany). The loaves were removed from the pan and allowed to cool down for 120 min on cooling racks at room temperature. Individual loaves were stored at 25°C for 24 hours.

Sensory analysis. The loaves were cut with a bread slicer (Industrade, Strasbourg, France) into 10 mm thick pieces. Triangle tests were carried out to assess differences in the characteristic aroma present in the loaves made by the two different recipes. Sessions were carried out in a specific sensory room 1 h after the sample preparation. For each triangle test, samples were given to 30 untrained judges, spanning an age range from 20 to 50, the two genders being present in equal numbers. During the session, judges were asked to taste 3 samples. Panellists were forced to answer the question (forced-choice method). Samples were presented to the judges in a random order. A significant taste difference for a risk of 5% between two breads was obtained when it was perceived by more than 15 judges according to the binomial distribution B (n, 1/3), n being the group size (NF EN ISO 4120:2007 – Sensory analysis Methodology – Triangular test).

RESULTS AND DISCUSSION

pH analysis and diacetyl quantification. During incubation, the two strains F.2.3 and Dx1 raised the pH of their medium from 4.5 at the end of the pre-fermentation step to 6.1 and 5.7, respectively, at the end of the fermentation step (Figure 1). The pH measurement was used as an indicator of citrate metabolism efficiency and the correlative production of flavourings. Indeed, the citrate metabolism is generally assumed to lead to the re-alkalisation of the medium (Gemelas et al. 2014). The pH increase of the whey mixture therefore suggests that diacetyl production has occurred. The analysis of diacetyl by a chromatographic method (GC-FID) confirmed this assumption: diacetyl was present in the two bacterial cultures, F.2.3 and Dx1. The two samples released a pleasant creamy-buttery odour, slightly more pronounced in the case of the Dx1 starter. According to the chromatographic analysis, the areas corresponding to the diacetyl peak were equal to 1.5 ± 0.5 and 23.8 ± 8.3 mV/min for strains F.2.3 and Dx1, respectively, i.e. a 1 : 16 ratio. This latter strain is habitually used to improve the buttery aroma of dairy fermented products. This great difference in a diacetyl amount between the two strains was expected: in an industrial fermentation process, the starter strains are selected for their technological properties, bringing optimum acidity and/or flavour to the product (Wouters et al. 2002). The perception threshold of diacetyl in butter is low. Concentrations ranging from 0.0045 ppm to 0.01 ppm and between

Figure 1. Monitoring of the pre-fermentation and the citrate fermentation steps for the diacetyl production by citrate positive Lactococcus strains (F.2.3 and Dx1); the characters in a frame represent the actions led for the pilotage of the fermentation.
The absence of the butterm aroma in the bread made with the citrate positive cultures used in this experiment could result from the evaporation of aroma compounds during the bread-making process, and especially during cooking. The relative volatility ‒ characterising the capacity of a compound to reach a gaseous phase ‒ influences the rate at which the volatile compound dissipates. This parameter is influenced by three factors. Firstly, compounds with low molecular weight have a greater ability to diffuse throughout the matrix during cooking, i.e. small molecules are retained less than the largest ones (De Roos et al. 2003). Secondly, polar molecules are retained less than nonpolar compounds. Indeed, polar molecules ‒ soluble in water ‒ can spread easily throughout the matrix by diffusion (Rosenberg et al. 1990). Thirdly, the lower the boiling point, the greater the relative volatility of a compound is. If we consider diacetyl, it matches all three of these characteristics: it is a polar diketone with four carbon atoms (and so a small molecule) and low boiling point (88°C). In light of the core temperature of the bread at the end of baking ‒ 99°C, it would seem that all the diacetyl had been evaporated during the bread-making process. In the same way, Gassenmeier and Schieberle (1994) showed that 90% of the diacetyl present in puff-pastry evaporated during the cooking step (12 min at 180°C). Therefore, our postulate concerning the preservation of diacetyl produced by a fermentation process turned out to be inaccurate. Contrary to a fresh milk product, dough is baked and the butterm aroma could have evaporated during the baking step. Therefore, the way to give bread a natural butterm aroma without using butter is still to be found. Relative volatility also depends on the interaction between the aroma compounds and the matrix (Bangs & Reineccius 1982). This way could be explored to preserve diacetyl from evaporation during the baking process. The possible entrapment of diacetyl in a macromolecular network of exopolysaccharides produced by another lactic acid bacterium, Leuconostoc, will be investigated in a future article.

CONCLUSION

In white sandwich bread, the diacetyl produced by a citrate positive lactic acid bacterium and added to the recipe, was not perceptible by the panellists of a triangle test. We can wonder whether panellists are ‘really’ able to perceive butter in a pastry product or
whether butter contributes in other ways to the quality of the final product – texture for instance – but not only to aroma. But we can also postulate that the diacetyl evaporated during the bread-making process and that the residual concentration was below its threshold value of perception. To circumvent this latter problem, the next steps of our research will aim at the use of exopolysaccharides produced by Leuconostoc strains for the entrapment of diacetyl. This work will be the subject of an upcoming article.

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