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Model cheese aroma perception is not only explained by in vivo aroma release but also by salivary composition and oral processing parameters

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Abstract

The aim of the present paper was to determine, on four model cheeses differing in fat content and firmness consumed by fourteen well characterised subjects, the respective impacts of in vivo aroma release, bolus rheology, chewing activity, mouth coating and saliva composition on dynamic aroma perception. The originality of the approach was to consider all the parameters together and to be able to evaluate their relative contribution using multi-block partial least square (MB-PLS) regression. Fruity aroma perception of the more hydrophilic compound (ethyl propanoate) was related to its dynamic release parameters before swallowing whereas blue cheese aroma perception of the more hydrophobic compound (nonanal) was related to its dynamic release parameters after swallowing and was highly impacted by mouth coating. Moreover MB-PLS approach made it possible to evidence the combined effects of saliva composition and cross-modal interactions to understand why in some cases dynamic aroma perception could not be explained by dynamic in vivo aroma release data. Subjects with low sodium content in saliva perceived fruity aroma which is not congruent with saltiness as less intense and salt-congruent (blue cheese) aroma as more intense, which was explained by their higher sensitivity to salt. Subjects with a high lipolysis activity perceived fruity aroma which is not congruent to fat as less intense and fat-congruent (blue cheese) aroma as more intense, which should be explained by the link between lipolysis activity and fat sensitivity. These results could be considered for the reformulation of foods towards specific populations taking into account nutritional recommendations.

Key words: aroma perception, aroma release, cheese, bolus rheology, saliva composition, chewing behaviour
Introduction

Aroma perception is an important aspect of food acceptability by consumers. However it is often very difficult to directly explain aroma perception during food consumption by the amount of aroma compounds in the food. One explanation is that aroma compounds have first to be released from the food matrix to the vapour phase in the buccal cavity then transferred into the nasal cavity via the velum to reach the olfactory receptors. The partition of aroma compounds between the food matrix and the vapour phase depends on the affinity of aroma compounds for the macromolecules present in the food which can be determined by measuring the vapour/matrix partition coefficients of aroma compounds. These coefficients depend both on the physico-chemical properties of the aroma but also on the food matrix composition. As an example hydrophobic aroma compounds are more soluble in fat than in water and thus are better released in the vapour phase from products with a reduced fat content.

Proteins are able to form reversible or irreversible binding with aroma compounds which will impact aroma release in the vapour phase. In case of irreversible binding such as the formation of Schiff base between amino groups of proteins and aldehydes, the aroma compounds cannot be released in the vapour phase and thus cannot be perceived. In case of reversible binding the aroma perception is lowered. The major effect of hydrocolloids is to increase food viscosity and thus decrease the transfer of aroma compounds from matrix to vapour, however inclusion complexes are formed between amylose and specific aroma compounds which also impacts aroma perception. During consumption, food is mixed with saliva and broken down during the masticatory step to form a swallowable bolus. During this step, aroma compounds are transferred from food to saliva before being released in the nasal cavity. This transfer of aroma compounds is highly dependent on food composition and texture but also on subject’s physiology. Thanks to the development of on line aroma release technologies, such as atmospheric pressure chemical ionisation (APCI-MS) or proton transfer reaction mass spectrometry (PTR-MS), it was possible to better understand the impact of dynamic in vivo aroma release of food matrix composition and structure and of physiological parameters such as chewing, swallowing, saliva flow. Concerning the impact of food matrix, increasing fat content results in a decrease in maximum intensity and overall in vivo aroma release together with an increase in aroma remanence as was observed in gelatin gels, whey protein gels or model cheeses, which was explained by a lower solubility of hydrophobic aroma compounds in fat. Even if food rheology could explain chewing activity during consumption, bolus properties and in-mouth bolus rheology better explain in vivo aroma release. The amount of cheese remaining in the mouth after swallowing has also been found important to explain in vivo aroma release. A higher amount of fat product remaining in the mouth leads to a lower release of the more hydrophobic compounds before swallowing due to their high affinity for the food bolus and to a longer aroma persistence in the breath. The less hydrophobic compounds are less
impacted by mouth coating but are more quickly released from firmer cheeses, due to a higher food breakdown\textsuperscript{16}. When consuming solid foods, differences in chewing activities between subjects are responsible for differences in aroma release, an intense chewing work leads to a higher amount of aroma release\textsuperscript{17}. However despite the development of \textit{in vivo} measurements, results in the literature in the field of dairy products showed that no clear relationship exists between \textit{in vivo} aroma release and aroma perception due to the co-existence of physicochemical and cognitive mechanisms\textsuperscript{23}. Cognitive mechanisms are due to cross-modal interactions which vary according to the type of texture\textsuperscript{24}. In the case of semi-liquid foods such as yoghurts, an increase in viscosity induced a decrease in both aroma release and aroma perception\textsuperscript{25}, whereas for solid cheeses aroma perception decreased with firmness without any noticeable difference in aroma release\textsuperscript{26}. This finding was explained by the attention paid by the subjects to the texture perception which likely led to a decrease in the aroma perception. By delivering simultaneously different kinds of texture and different odorants, Bult et al.\textsuperscript{27} showed that the perceived aroma intensity was reduced with increasing viscosity when the odour was presented either ortho or retronasally. Moreover the perceived thickness was increased only when the odour was presented retronasally simultaneously with swallowing time. An adaptation phenomenon was also evidenced in gel candies as being responsible for the influence of taste and texture on aroma perception\textsuperscript{28}, by comparing the \textit{in vivo} aroma release curves and sensory time intensity curves which were recorded simultaneously. This study showed that time at maximum chewing activity came first followed by time at maximum in nose concentration then time at maximum perceived intensity. The time delays between release and perception increased when the product remained longer in the mouth. The difficulties to relate results from analytic and sensory approaches was also pointed out in brandies\textsuperscript{29} and explained by sensory interactions between fruity and woody aroma\textsuperscript{30}. In coffee, the burnt sensory notes could be associated with pyrazines detected by \textit{in vivo} PTR-MS analysis but the effect of adding sugar produced a change in aroma perception from burnt to caramel which could not be associated to any change in aroma composition\textsuperscript{31}, confirming the existence of sweet-aroma interactions\textsuperscript{23}. Odour-taste interaction basically depends on the capacity of the two stimuli to form an appropriate combination in a food product context (congruency)\textsuperscript{32}. Thus, odour–taste integrated perception highly depends on learned associations, the context in which the food is consumed and the consumer’s previous experience\textsuperscript{33}. In experimental situations, odour–taste interactions were found to be affected by the so-called dumping and halo effects\textsuperscript{34}, which resulted from the tendency for sensory panellists to dump their sensations on several available scales (taste and/or odour). Moreover, in more complex cheese flavour mixtures, NaCl, lactic acid, and aroma were found to be able to enhance cheese flavour intensity and to compensate each other towards cheese flavour intensity\textsuperscript{35}, thus suggesting complex taste-taste and taste-aroma interactions involved in the overall cheese flavour. So far, no study has investigated cross-modal interactions using parameters related to oral physiology.
During the last decade, saliva has gained more and more interest in the field of sensory science. Indeed, due to its role in oral clearance, bolus moisturizing and hydrolytic properties, saliva promotes the disintegration of the matrix and thus the release of palatable active substances during food consumption. For example the salivary flow rate influences aroma release by a dilution effect. Salivary flow has also been positively associated with liking for fat. Moreover, amylase activity can modify the bolus rheological properties and thus modulates the perception of salt in starch-based matrices. Even subject to controversy, lipase might be responsible for hydrolysis of food triglycerides and thus fat perception. Finally the catalytic activity of saliva against some aroma compounds (e.g. esters, aldehydes) has been demonstrated in vivo and in vitro with consequences on aroma release and thus on perception. Saliva is also composed of small and large molecules that contribute to maintain a "salivary homeostasis" whose background level can regulate the dynamics of molecule release and thus their sensory impact (sensory or nutritional). For instance, it has been shown that a human subject with a high salivary sodium concentration is less sensitive to saltiness.

A previous paper highlighted the role of bolus rheology and composition of stimulated saliva on aroma release from cheeses varying in fat content and firmness. The aim of the present paper is to better understand the physicochemical and physiological parameters which drive dynamic aroma perception. For that purpose, a group of fourteen subjects were selected as representative of those participating to our previous study in order to evaluate aroma perception as a function of time and to determine, on these well characterised subjects, the respective impacts of in vivo aroma release, bolus rheology, chewing activity, mouth coating and saliva composition on dynamic aroma perception.

**Experimental**

**Cheese products**

Four processed model cheeses were used. They were composed of a mixture of cheddar, soft cheese, butter, melting salts, protein powder (casein), salt and water. Two levels of texture (S=soft, F=firm) were obtained by varying the water content and two ratio of fat to dry matter, a low level at 25% for low fat cheeses (lfF and lfS) and a high level at 50% for high fat cheeses (hfF and hfS). The pH ranged from 5.27 to 5.55. The rheological properties of the cheeses were measured in a large deformation at a rotation of 0.01 rad.s\(^{-1}\) for 240 s using a Haake Viscotester (VT550 - Thermo electron GmbH, Karlsruhe, Germany), as previously described. The breakdown stress corresponds to the maximum strength necessary to cause cheese breakdown, with the lowest values for the softest cheese (8129 ± 469 Pa for lfS and 8022 ± 1309 Pa for hfS) and the highest values for the firmest cheese (15253 ± 1231 Pa for lfF and 15556 ± 2307 Pa for hfF). The critical strain at breakdown corresponds to the maximum rotation angle required to cause breakdown, with the lowest values for cheeses
with the highest fat content (0.273 ± 0.022 rad for hFS and 0.348 ± 0.061 rad for hFF), and the highest values for cheeses
with the lowest fat content (0.804 ± 0.056 rad for lFS and 0.836 ± 0.036 rad for lFF). Two aroma compounds were added
during cheese production, a hydrophobic ketone, nonan-2-one (NO: logP= 2.9, 6 mg.kg⁻¹) and a hydrophilic ester, ethyl
propanoate (EP: logP= 1.4, 25 mg.kg⁻¹).

Subjects

In a first step, 48 subjects (23 female and 25 males) were selected from a group of 100 volunteers based on their good
dental and oral status, and on the repeatability of physiological parameters (salivary flow rate under resting and stimulated
conditions, respiratory flux, salivary composition⁰¹, ²⁰. They were characterised for their oral volume, saliva composition
and flow, in vivo release measurement, chewing activity, bolus saliva content, bolus rheology and mouth coating. From this
group of 48 subjects, a subgroup of fourteen subjects (6 females and 8 males, average age: 40 years ± 9) was selected for
the sensory analyses, on their ability to detect and recognise the two aroma notes. The subjects were not allowed to
smoke, eat or drink starting at least one hour before the different test sessions. All the subjects were informed of the
observational nature of this study. They gave their signed consent and received a financial compensation for their
participation. The study protocol was submitted to an Ethics Committee and was approved on 17 April 2008 by the Comité
de Protection des Personnes Est-1 (N°2008/15) and on 8 August 2008 by the Direction Générale de la Santé - France (N°
DGS2008-0196).

Subjects were characterised for specific physiological parameters described in the following sections.

Oral volume

Oral volume was measured using an Eccovision® acoustic pharyngometer (Hood Laboratories, USA), as described
previously⁰⁹. This device consists of two microphones and a horn driver mounted on a wave tube and connected to a PC-
compatible computer with signal conversion capabilities. The signal was converted into the surface change (cm²) as a
function of the length of the oral cavity (cm). The subjects held the mouthpiece in their mouth with their teeth against the
flange and their tongue in a low position. To prevent air leaks, which could cause measurement errors, the subjects placed
their lips over the flange, sealing the mouthpiece. The subjects were instructed to breathe with their nose during the
experiment. Values are expressed in cm³ and correspond to the average of 10 measures.

Saliva samples and flow

Since our previous study²² highlighted that resting saliva poorly contributed to explaining aroma release whereas
stimulated saliva significantly impacted in vivo aroma release from soft cheeses, only results obtained with stimulated saliva
are taken into account in the present paper. Stimulated saliva was collected as previously described. The subjects chewed a piece of Parafilm™ (0.5 g ± 0.2 g) for a period of 5 min and spit out the saliva every 30 seconds into a pre weighed cup over a period of 5 minutes. The cups were weighted and the salivary flow rates were expressed in mL.min⁻¹. Immediately after collection, the saliva samples were standardized by a first step of centrifugation for 30 min at 15000-x g to remove bacteria and cellular debris. Thereafter the supernatants were stored at -80°C to stop metabolism until subjected to biochemical analyses.

Biochemical analyses of saliva samples

**Protein concentrations.** Protein concentrations (Prot) were measured with a standard Quick Start Bradford protein assay (Bio-Rad, France) with bovine serum albumin as the calibration standard.

**Enzyme activities.** All enzyme activities were expressed in International Enzyme Activity Units (U) per ml of saliva. One U is defined as the amount of enzyme that catalyses the conversion of 1 micromole of substrate per minute. The lipolytic (Lipolysis), proteolytic (Proteolysis), lysozymal (Lysozyme) and amylolytic (Amylase) activities were determined as previously described.

**Sodium analysis.** The saliva samples were diluted to 1/20 (50 μL saliva in 950 μL filtered 18 mΩ Milli-Q-water (Millipore, Bedford, MA, USA)) and filtered through a membrane (pore size = 0.45 μm, C.I.L., Sainte-Foy-La-Grande, France). The amounts of sodium (Na) in saliva were determined by HPLC ionic chromatography using a Dionex ICS2500 ion chromatographic system (Dionex, Voisins le Bretonneux, France) as previously described. Quantifications were performed using calibration curves realised with sodium standard solutions ranging from 0.1 to 10 mM in 22 mM sulfuric acid (R²=0.999).

Sensory analyses

All sessions took place in an air-conditioned (21 ± 2°C) sensory testing room of the ChemoSens platform (Centre des Sciences du Goût et de l’Alimentation, INRA, Dijon) using standardized booths equipped with computers. Subjects were instructed to place each piece of cheese (6 g) in the mouth, and freely consume it while keeping the lips closed. The products were presented in a random order at 17° C. All measurements were done in triplicate. Bread, apple and water were used as mouth cleansers between two tests.

The subjects were firstly trained during eight sessions to recognise the odour of the two aroma compounds, blue cheese for nonan-2-one (NO) and fruity for ethyl propanoate (EP), using triangular tests and recognition tests. Other training sessions were conducted to familiarize the subjects with the discontinuous time-intensity methodology used. The last training
sessions were dedicated to the use of a continuous scale (0 to 10) to score the intensity of the two aroma compounds. Two

test sessions were then done using Fizz® software to score the intensity of the two aroma notes during the mastication, to

indicate the time of the first swallowing and then to score the intensity of the two aroma notes after each swallowing

event, during a total time of 3 min. During mastication and at each swallowing time, the intensity of each aroma note was

scored on the continuous scale (0-10). Each of the four cheeses was presented three times, in a random order, at 17°C.

From the aroma intensity perception at each swallowing event and the time of each swallowing event, time-intensity curves

were reconstituted for each aroma note and the following parameters were extracted: Imax_S for maximum intensity,

Tmax_S for time to reach maximum intensity, Ideg_S for intensity at the first swallowing and Tend_S for time to reach the

end of perception (Fig. 1). The rate of perception (Vmax_S = Imax_S/Tmax_S) was then calculated as a fifth variable.

Insert Figure 1

In vivo aroma release measurement

The same protocol was applied for cheese consumption than that described for sensory analysis. The release of the two

aroma compounds was followed simultaneously in the nasal cavity as previously described using Atmospheric Pressure

Chemical Ionisation-Mass Spectrometry (APCI-MS) with an ion trap Esquire-LC mass spectrometer (Bruker Daltonique,

Wissembourg, France) according to their protonated molecular ion (MH+), which is the main ion: ethyl propanoate

(m/z=103) and nonan-2-one (m/z=143). Air was sampled from the nose at an average flow rate of 37 mL min−1 via a fused

silica capillary tubing (i.d. = 0.53 mm) heated at 150 °C and to which a 5 kV positive ion corona pin discharge was applied.

Each subject was asked to position the plastic tube in one nostril (the same for all the experiments) and to breathe

normally. This period (breath-blank phase) was used to record the potential residual signal of the previous sample until

return to the baseline and to control the regularity of breathing.

The curves were smoothed using a wavelet decomposition method to eliminate signal fluctuations due to the subjects’
brathing patterns. Two release phases were identified, the chewing phase (1) extended from placing the cheese in the

mouth to the first swallowing, and the post-swallowing phase (2) extended from the first swallowing to the time at which

the signal returned to its baseline level. For both release phases and for each aroma compound, four main parameters were

extracted from each individual release curve: the area under the curve (A1_P and A2_P (a.u.: arbitrary unit)) representing

the quantity of aroma released, the maximum intensity (IMax1_P and IMax2_P (a.u.)), the time to reach maximum intensity

tMax1_P and tMax2_P (min)) and the release rate (Vmax_P= IMax1_P / tMax1_P (a.u./min)). These data, also not

quantitative, allow a direct comparison of the different release curves.

Chewing activity
Chewing activity was monitored during cheese consumption, simultaneously to aroma release. The muscle activity of the superficial masseter and temporis muscles (left and right) during chewing was recorded by electromyography (EMG) using gold surface electrodes (Grass technologies, West Warwick, RI, U.S.A), at 382 Hz, then the signal was amplified and digitalized. The following parameters were extracted from EMG data: number of chewing cycles (Nber_Cycle), chewing duration (Chew_time expressed in s), total muscle work (W_tot expressed in mV.s-1) and mean amplitude of contraction (Ampl expressed in mV) which corresponds to a mean calculated from the amplitude values of each chewing cycle registered in a whole chewing sequence.

**Bolus saliva content**

The percentage of dry matter and water content were determined using an infrared dryer for all cheeses and boluses obtained just before swallowing. The percentage of moistening (Moist %) into the bolus was calculated from the bolus water content (Bwc %), the bolus dry matter (Bdm %), the cheese dry matter (Cdm %) and the cheese water content (Cwc %) as follows:

$$\text{Moist} \, \% = \left( \frac{\text{Bwc}}{\text{Bdm}} \times \text{Cdm} \right) - \text{Cwc}$$

Three replicates per cheese and per subject were performed.

**Bolus rheology**

The subjects were instructed to chew the cheese samples until swallowing and to spit out the bolus into a truncated syringe. Bolus rheological properties were measured using a compression test on an aliquot of 3 mL of bolus. The test was performed using a mobile circle upper plate and a fixed circle lower plate as compression device, with a compression rate of 1 mm.s$^{-1}$. The bolus was subjected to a force $F$ ranging between 0.01 N and 50 N. From the compression curve, particularly two phases were highlighted. A "flow phase" during which the suspension begins to flow and the particles move significantly in relation to one another at a height denoted as $h_{\text{flow}}$ (mm). Yield stress and viscous effects were described respectively by the parameters $s_{\text{flow}}$ (Pa) and $K_{\text{flow}}$ (Pa.s). A "particle phase" during which the mechanical response is governed by the particle size which is represented by a height denoted $h_{\text{part}}$ (mm) and the yield stress component denoted $s_{\text{part}}$ (Pa). At the end of the compression, $h_{\text{end}}$ (mm) denotes the final height and $S_{\text{end}}$ (mm$^2$) the area generated under the maximal force. All measurements were done in triplicate.

**Mouth coating**
The amount of food that sticks to the oral surface after food ingestion (QR8) was quantified by the “mouth rinse” method\(^4\).

Curcumin (Naturex, France) was added during cheese production (30 mg.kg\(^{-1}\)). Each subject was asked to place a piece of cheese (6 g, at 17°C) in the mouth and to chew normally until swallowing. The subjects swallowed without cleaning movement and then rinsed their mouth (with cleaning movements) with 4 mL of warm water at 50°C for 30 s, and spat it into a vial. This rinsing procedure was applied two times consecutively and the spittle was cumulated in the same vials. The fluorescence intensity of curcumin was quantified using a Perkin Elmer 1420 Multilabel Counter Victor 3V at an excitation wavelength of 450 nm and an emission wavelength of 510 nm. All measurements were done in triplicate.

### Statistical analyses

Analyses of variance (ANOVA) and Principal Component Analysis (PCA) were performed using XLSTAT\(^7\) Software (Excel 97, version 8.0, Paris, France). When a significant effect (\(p<0.05\)) was revealed by applying ANOVA, the Student-Newman-Keuls test was used to compare the differences in least-squares (LS) means.

Statistical treatments for Partial lest Square (PLS) analysis were performed using the free software R 3.3.0 (http://cran.r-project.org/), as already described for the treatment of \textit{in vivo} release data obtained with the same cheeses and 48 subjects\(^2\). The main R package used for multivariate data analyses was «pls 2.5-0»\(^5\). In a preliminary stage, the statistical treatment consisted in a pre-processing step\(^6\). More precisely, all the variables are mean centered; then the blocks of variables are set to the same total variance. Finally, in order to explore the systematic variation patterns in the X blocks which are likely to predict the systematic variation patterns in Y, Multiblock -PLSR (MB-PLSR) is applied.

The different variables used in the MB-PLS approach are presented in Table 1. They have been divided in six blocks. The Y block corresponds to the variables to be explained that is, the sensory variables. The five other blocks correspond to the explanatory variables (X1-X5). X1 is related to aroma release variables before (1) and after swallowing (2) as previously described. The variables in the subsequent blocks were selected as highly impacting the \textit{in vivo} aroma release\(^2\). X2 is related to bolus rheological variables and bolus moistening. X3 is related to mouth coating and oral volume. X4 is related to masticatory variables extracted from the EMG signals. X5 is related to the properties of stimulated saliva.

### Results

#### Subjects’ physiology

The 14 subjects were selected from the 48 subjects participating to the \textit{in vivo} release measurements\(^1\). Their physiological characteristics (Table 2) cover the range of variability observed for the 48 subjects as presented in the PCA representation.
The principal plan represents 47% of the information. Axis 1 (26.76%) is explained by the salivary flow (Flux_S) and the amount of sodium (Na_S) on its positive part, which are higher for subjects S027 and S094. Axis 2 (20.14%) is explained by the amount of amylase (Amylase_S), the amount of salivary proteins (Prot_S) and the oral volume (Oral_vol) on its positive part, in relation with subjects S001, S052 and S086. At the opposite, S004 has a very low amount of amylase.

Time intensity perception

From the time intensity curves obtained for the four cheeses, 14 subjects and two odour notes, four variables were extracted namely, maximum intensity (Imax_S), time to reach the maximum intensity (Tmax_S), intensity at the first swallowing (Ideg_S) and time at the end of perception (Tend_S). The fifth variable, rate of perception (Vmax_S = Imax_S/Tmax_S) was calculated. Due to the high inter-individual differences between subjects only two variables show significant differences between the cheeses. Tmax_S is significantly higher (p = 0.002) for the blue cheese aroma (NO) detected in low fat cheeses (lfS and lfF) and Vmax_S is significantly higher (p = 0.07) for the fruity aroma (EP) detected in high fat soft cheese (hfS). Means and standard deviations for the different cheeses are reported in supplementary Table S1.

The high inter-individual variability was also observed on the in vivo release variables for the 48 subjects, which was explained by differences in physiological parameters. Results of in vivo aroma release have already been published\cite{16}, However the data obtained for the 14 subjects selected for the sensory analysis are available in supplementary Table S2. In order to better highlight the subject effect, a multiblock PLS analysis was performed on these 14 subjects to explain sensory perception for each cheese and each aroma compound by the in vivo release variables together with the physiological variables (Table 1).

Relating sensory perception to in vivo aroma release and physiological parameters using MB-PLS

MB-PLS is an extension of the PLS method, a class of regression models attempting to find relationships between explanatory and response variables. In MB-PLS, the predictor variables are separated into subsets or blocks that are standardized in order to balance for the size effect due to the measurement scale. It is a statistical approach particularly relevant when different data sets reflecting different dimensions (physiology, physic, chemistry ..) and with a different number of variables in each set are considered.

MB-PLS analyses were conducted on the different data sets to assess the extent to which the X blocks of variables explain sensory perception during cheese matrix consumption (Fig. 3 and Fig. 5). The four cheeses and the two
aroma compounds nonan-2-one (NO) and ethyl propanoate (EP) were considered separately for the statistical treatment.

Two different results are presented from MB-PLS analysis in the following sections, the importance of the block and the projection of the variables in the correlation circle. At first, the importance of the blocks is calculated from the sum of the beta-weight of the different variables constitutive of the block related to the corresponding component and is expressed in %. It thus reflects the contribution of the block of variables to the determination of the component. Higher is the percentage, higher is the contribution of the block for explaining sensory data.

Complementary to the importance of the blocks, the projections correspond to the depiction in a correlation circle of the variables belonging to the different blocks. They thus represent the importance of the variables to the components. Higher a variable is correlated to the components, higher it contributes to the model and thus to explain perception.

The choice of the number of MB-PLS components to be retained for the importance of the block and the projections was based on the total variance of block Y recovered by these components. For more details regarding this aspect, we refer to a previous paper. We restrict ourselves to the first two components because they explain between 45.3% and 56% of the total variance of Y for the high fat cheeses and the two aroma compounds and between 36.5 and 50.4% for the low fat cheeses and the two aroma compounds.

In order to avoid cumbersome graphical displays on the projections, only the variables with a correlation coefficient with one of the two first components above 0.5 are depicted. The font size for each variable on the projection (Fig. 4 and Fig. 6) reflects the importance of the correlation coefficient with the first two components; with large font size indicating large correlations.

The results from the MB-PLS are presented and discussed successively for high fat cheeses and for low fat cheeses.

**Relative Importance of the blocks in the projection for high fat cheeses**

The importance of the blocks of variables for the two first MB-PLS components is shown in figure 3 for the two high fat cheeses and the two aroma compounds. The release block is important for the high fat firm cheese (hfF) and both molecules and this on the two components (31.7% for NO on component 1 and 37% for EP on component 2) and less important for high fat soft cheeses (with a maximum of contribution of 18.5%). The bolus rheology block is mainly reflected by the second component for EP (25.9% for hfF-EP and 31.5% for hfS-EP) and less reflected for NO. The importance of coating and oral volume is higher for EP than for NO, on the first component. The masticatory variables take an important part in the explanation for both cheeses and both aroma compounds. Notice also the high contribution of stimulated saliva, mainly for hfS-NO (48.9% on component 1 and
24.8% on component 2) for which the masticatory variables are less important (12.5% on component 1 and 21.3% on component 2).

Insert Figure 3

Insert Figure 4

Projection of the main variables for high fat cheeses

Figure 4 shows the projection of the variables from each block for the two high fat cheeses (hfF and hfS) and the two aroma compounds (EP and NO). The sensory variables from the Y block (variables to be explained) are negatively correlated with component 1 and for the high fat soft cheese, the rate of perception is negatively correlated with component 2. The other variables (explanatory variables) are projected differently according to both the cheese and the aroma compound. For the high fat firm cheese and ethyl propanoate (hfF-EP), the masticatory parameters are correlated with component 1, with the chewing time (Chew_time) and number of chewing cycles (Nber_Cycle) on the positive part and the amplitude (Ampl) on the negative part, the release parameters are correlated with component 2, with a higher correlation coefficient for Vmax_P and Imax1_P, the amount of product remaining in the mouth (QRB) is projected at the opposite of the maximum intensity perceived (Imax_S). Among the salivary parameters, the amount of proteins (Prot_S) is positively correlated with component 1, opposite to the amount of sodium (Na_S) and of amylase (Amylase_S). The amount of lipolysis (Lipolysis_S) is projected close to the QRB. For the high fat soft cheese and ethyl propanoate (hfF-EP), the most correlated explanatory variables are the bolus rheology properties, with Send and Moist negatively correlated with component 1 together with the sensory variables, opposite to hend and hpart. The masticatory parameters are also correlated with component 1, with Chew_time and Nbe_Cycle on the positive part and Ampl on the negative part. Among the release parameters, only Imax1_P and A1_P are positively correlated with component 1 and Vmax_P with a lower correlation. QRB is also projected opposite to Imax_S. Concerning the high fat firm cheese and nonan-2-one (hfF-NO), the masticatory parameters (Chew_time and Nbe_Cycle) are also positively correlated with component 1 and negatively with component 2, the release parameters before swallowing (Imax1_P and A1_P) are positive correlated with component 1 whereas the release parameters after swallowing (Imax2_P, A2_P) are negatively correlated with component 2. Concerning the high fat soft cheese and nonan-2-one (hfS-NO), the salivary parameters are well represented on the projection, mainly Flux_S and Na_S, positively correlated with component 1 and component 2 and opposite to the maximum intensity perceived. The release parameters have only low correlations with these two components as represented by their small font size, but the time to reach maximum intensities both before and after swallowing (tmax1_P and tmax2_P) are positively correlated with component 2, opposite to the rate of perception. The masticatory parameters and QRB are negatively correlated with component 1.

Relative Importance of the blocks in the projection for low fat cheeses
The importance of the blocks of variables for the two first MB-PLS components is shown in figure 5 for the two low fat cheeses and the two aroma compounds. The relative importance of the release block is always below 28%, it is higher for nonan-2-one and the firm cheese (lfF-NO, 27.5% on component 1 and 14.9% on component 2) and for ethyl propanoate and the soft cheese (lfS-EP, 20.7% on component 1 and 19.8% on component 2). The bolus rheology block is reflected on the two components with a contribution of 19.2% on component 1 and 6.8% on component 2 for lfF-NO, 17.4% on component 1 and 29% on component 2 for lfS-NO and intermediates for EP. Coating and oral volume are highly important for lfF-NO (25.2% on component 1 and 29.8% on component 2) and lfS-EP (48.7% on component 2). The masticatory properties have a high impact on fruity aroma (EP) in the low fat firm cheese (46.5% on component 1 and 47% on component 2), and in the low fat soft cheese (41.3% on component 1) but they seem to impact less blue cheese aroma (NO). For nonan-2-one the salivary parameters are more relevant to explain sensory perception with 22.2% of contribution on component 1 and 28.5% on component 2 for low fat firm cheese and 14.8% on component 1 and 46.5% on component 2 for low fat soft cheese.

Projection of the main variables for low fat cheeses

Figure 6 shows the projection of the variables from each block for the two low fat cheeses and the two aroma compounds on the two first MB-PLS components. For the low fat firm cheeses (lfF), the sensory variables related to aroma persistence (Tmax_S and Tend_S) are negatively correlated with component 1, whereas the rate of perception (Vmax_S) and the maximum intensity of perception (Imax_S) are negatively correlated with component 2 for ethyl propanoate and positively for nonan-2-one. For the low fat soft cheeses, the rate of perception (Vmax_S) is positively correlated with component 1, the maximum intensity (Imax_S) negatively with component 2, whereas the times (Tmax_S and Tend_S) are negative correlated with component 1 for ethyl propanoate and negatively correlated with component 2 for nonan-2-one.

Concerning the explanatory variables, the masticatory parameters (Chew_time and Nber_Cycle) are negatively correlated with component 1 for lfS cheese and the 2 aroma compounds and for lfF cheese and ethyl propanoate but negatively correlated with component 2 for lfF cheese and nonan-2-one, whereas the amplitude is negatively correlated with component 2 only for lfF cheese and ethyl propanoate. Concerning the bolus rheology parameters Hend, hpart and Kflow are always positive correlated with component 1 and component 2 whereas Moist and Send are negatively correlated with component 1 and component 2. Concerning the release parameters, tmax2_P is always negative correlated with component 1, the other parameters have lower correlations on this plan, except Vmax_P on the positive part of component 1 for lfF cheese and ethyl propanoate, Imax1_P on the positive part of component 1 and negative part of component 2 for lfF cheese and nonan-2-one, A2_P and A_P for both cheese
and nonan-2-one. The salivary parameters have only small correlations on this plane for the low fat soft cheese but for the low fat firm cheese, the salivary flow (Flux_S) and the amount of sodium (Na_S) are negatively correlated with component 2.

**Discussion**

Considering the results of MB-PLS on high fat cheeses (Fig. 4), different trends are observed for firm and soft cheeses. For the firm cheeses, the perception of fruity aroma is more related to the release of ethyl propanoate (EP) before swallowing whereas the perception of blue cheese aroma is more related to the release of nonan-2-one (NO) after swallowing. Ethyl propanoate which is less hydrophobic is released faster from the fat cheese matrix, and thus perceived at the beginning of the oral processing for the firm cheese whereas nonan-2-one which is hydrophobic is released later and mainly after swallowing, which explains the longer time to reach maximum intensity and time to reach the end of perception. For the soft cheeses the same correlations are found for nonan-2-one whereas no such correlation is found between sensory parameters and release parameters for ethyl propanoate. Regarding the masticatory variables, it appears that the mean amplitude of contraction (Ampl) is well correlated with the rate of fruity perception (EP) on component 1 for firm cheese and component 2 for soft cheeses. This variable was already highlighted in our previous paper as responsible for a higher rate of EP release. Thus, we confirm that subjects with high amplitude per burst will release more rapidly the hydrophilic compound (EP) and we demonstrate that this higher release rate induce a more rapid perception. The other masticatory variables better explain the blue cheese perception (NO) and more specifically aroma persistence. Subjects with a longer chewing time (Chew_time) and larger number of bursts (Nber_Cycle) have a longer duration of blue cheese aroma (Tmax_S and Tend_S) for both cheeses. This could also be explained by a longer time to reach the maximum NO release intensity after swallowing (tMax2_P). The bolus rheology properties impact fruity perception (EP) more, with a positive relationship between bolus moistening (Moist) and time to reach the end of perception (Tend-S), which could be explained by an important bolus spreadability (Send). Bolus moistening (Moist) impacts EP release intensity (Imax1_P), a high bolus moistening decreases the amount of release before swallowing. This could be explained by dilution with saliva which decreases the rate of release of hydrophilic aroma compounds as already proposed using *in vitro* and *in silico* models and thus will delay the perception by increasing the persistence. The amount of product remaining in the mouth (QRB) explains the perception of both aroma notes. QRB corresponds to the fat coating at the surface of the tongue and oral mucosa after swallowing. For blue cheese perception (NO), it could be thus due to the retention of NO, hydrophobic aroma compounds, in the fat, which delays its release and then enhances its perception. Conversely, the fruity perception (EP) is projected opposite to fat coating in the mouth, i.e. higher is the coating lower is the perception. Many salivary parameters impact aroma perception. Sodium concentration in saliva (Na_S) positively influences fruity perception (EP) and negatively blue cheese perception (NO), whereas it was not found to have a strong impact on *in vivo* aroma release.
Salivary lipolysis (Lipolysis_S) was depicted in the same direction of QRB in three projections (hfF-EP, hfF-NO and hfS-NO) witnessing a different impact on sensory perception depending on the fruity (EP) or blue cheese (NO) note. Salivary flow (Flux_S), protein content (Prot_S), amylase activity (Amylase_S) also impact sensory perception but differently according to the cheese and the aroma notes.

As for high fat cheeses, different trends are observed between firm and soft low fat cheeses (Fig. 6). For the low fat firm cheese (lfF) aroma persistence (Tmax_S and Tend_S) can be explained by the time to reach maximum aroma release after swallowing (tMax2_P), the amount of product remaining in the oral cavity (QRB) and the oral volume (Oral_Vol) for the two aroma compounds whereas the chewing time (Chew_time) and the number of chewing cycles (Nber_Cycle) mainly explain fruity perception (EP). These correlations are logical because a higher amount of product remaining in the mouth increases the amount of aroma released after swallowing which should increase aroma persistence. Moreover a longer chewing time increases the total amount of aroma released and thus aroma perception. The mean amplitude of contraction (Ampl) is depicted very similarly to the rate of perception (Vmax_S) of the fruity note as it was the case for high fat cheeses. Similar to high fat cheeses, stimulated saliva composition and in particular the sodium content (Na_S) impact aroma perception. High sodium content is related to a high fruity aroma (EP) and low blue cheese aroma (NO) intensity.

For the low fat soft cheeses (lfS), similar trends are observed for the fruity aroma (EP) but not for blue cheese aroma (NO). The blue cheese aroma perception (NO) cannot be logically explained by NO release but is related to saliva composition and more precisely to lipolysis (Lypolysis_S). Bolus moistening and bolus rheology highly contribute to the perception of the two aroma notes. A higher aroma perception is related to a higher moistening (Moist) and higher bolus spreadability which is explained by the area at the end of compression (Send). These results are also in line with the higher salivary flow (Flux_S).

Thanks to MB-PLS statistical approach, some general trends can be highlighted concerning the explanation of sensory perception by release and physiological properties. When consuming solid foods containing fat such as cheeses, masticatory behaviour highly impacts the dynamic of aroma release and thus aroma perception. A large mean amplitude of contraction increases rate of release and rate of perception. A longer chewing time increases the time to reach maximum intensity after swallowing and the time to reach maximum perceived intensity. Concerning the variables from the other blocks, the relationships are not always so simple to interpret, which may be due to interactions between these variables. A high salivary flow leads to a high bolus moistening, a large bolus area at the end of compression indicating a high bolus spreadability, which results in a lower rate of aroma release but the direct consequence on aroma perception is not clearly evidenced.
However some variables were found to be related to sensory perception and not to aroma release. This is the case for specific variables of saliva composition. The most striking finding is that lipolysis which is depicted in the same direction to fat coating for NO and EP, is also depicted in the NO projection in the same direction to sensory variables, while the opposite is observed for EP. This was particularly emphasized for high-fat cheeses. Salivary lipolysis has been described as a marker of fat sensitivity and liking in human\textsuperscript{41}. For example, a positive correlation was found between the level of lipolysis and perceived fat intensity while the opposite was observed for liking in subjects\textsuperscript{51} with a normal weight. On obese subjects, \textit{in vivo} inhibition of lingual lipase led to a significant enhancement of their sensory threshold for triolein\textsuperscript{59}. Moreover, compared to normal-weight subjects, obese individuals exhibited a significant low level of salivary lipolysis thus reflecting the higher liking for fat as reported in this population\textsuperscript{48, 60}. Concerning coating, fatty perception of oil in water emulsions has been previously related to the level of fat retention at the surface of the tongue after consumption\textsuperscript{61, 62}. Altogether, these findings tend to substantiate a positive correlation between the level of salivary lipolysis, fat coating and fat perception. These findings support our hypothesis of cross-modal interactions between aroma perception and fattiness that differs depending on the aroma. For NO (blue cheese aroma), the similar depiction of lipolysis, fat coating and sensory variables should indicate a congruency between aroma perception and fat perception. Contrariwise, for EP (fruity aroma), the depiction between lipolysis, fat coating and sensory variables should indicate a negative interaction because of the non-congruency between this aroma and fat perception.

A similar hypothesis can be proposed for the effect of sodium content in stimulated saliva on aroma perception. This property is depicted in the same direction as sensory variables for EP whereas it is in the opposite direction for NO. The amount of sodium in saliva has already been described as an important factor which contributes in the sensitivity for salt, i.e. the higher the concentration, the higher the detection threshold\textsuperscript{63-66}. As for lipolysis, it is likely that subjects exhibiting a high sodium concentration had a lower saltiness perception. Following our hypothesis regarding the existence of salty-aroma interactions and considering that blue cheese aroma, unlike fruity aroma, could be congruent to saltiness, subjects with a low salivary sodium content may perceive the cheeses as being saltier. Since the subjects were instructed to rate the aroma intensity only and not the salty taste in the cheese products during the sessions, they should have reported their salty perception on the blue cheese aroma which is congruent with salty perception, due to a dumping effect\textsuperscript{34}. Contrariwise, the perception of a fruity note may be increased for the subjects with high sodium content who perceived the cheeses as less salty, because this aroma is not congruent with salty.

\textbf{Conclusions}
As a conclusion, this study allowed us to highlight the respective impacts of *in vivo* aroma release and physiological properties on sensory perception. The originality of the approach is to consider all the variables together and evaluate their relative contribution. Our study confirms the important role of masticatory variables for firm cheeses and that of salivary properties for soft cheeses. The perception of the fruity aroma of the more hydrophilic compound (ethyl propanoate) is related to its dynamic release parameters before swallowing whereas the blue cheese aroma of the more hydrophobic compound (nonan-2-one) is related to its dynamic release parameters after swallowing and is highly impacted by mouth coating. Moreover, it was evidenced that dynamic *in vivo* aroma release does not always explain sensory perception. MB-PLS approach made it possible to evidence the combined effects of saliva composition and cross-modal interactions to understand why in some cases dynamic aroma perception could not be explained by dynamic *in vivo* aroma release data. Our study confirms the key role of salivary sodium content and lipolysis activity in sensory perception. The main finding is that subjects with a high sodium content in saliva perceive aromas which are not congruent with saltiness as more intense and salt-congruent aromas as less intense and that subjects with a high lipolysis activity perceive aromas which are not congruent to fat as less intense and fat-congruent aromas as more intense. These findings could help to better understand the inter-individual differences in aroma perception and could be considered for the reformulation of foods targeted for specific populations taking into account nutritional recommendations.

**Acknowledgements**

The acknowledgements come at the end of an article after the conclusions and before the notes and references.

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References


Table 1: presentation of the different blocks of variables used in the –MB-PLS Analyses. Y: variables to be explained, X: explanatory variables. a.u.: arbitrary units

<table>
<thead>
<tr>
<th>Block</th>
<th>Abbreviation</th>
<th>Definition of the variable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y: sensory parameters</td>
<td>Imax_S</td>
<td>Maximum intensity perceived (a.u.)</td>
</tr>
<tr>
<td></td>
<td>Tmax_S</td>
<td>Time to reach the maximum intensity (s)</td>
</tr>
<tr>
<td></td>
<td>Vmax_S</td>
<td>Rate of perception (a.u./s)</td>
</tr>
<tr>
<td></td>
<td>Ideg_S</td>
<td>Maximum intensity at swallowing (a.u.)</td>
</tr>
<tr>
<td></td>
<td>Tend_S</td>
<td>Time to reach the end of perception (s)</td>
</tr>
<tr>
<td>X1: aroma release parameters</td>
<td>A1_P</td>
<td>Area under the curve before 1\textsuperscript{st} swallowing (a.u.)</td>
</tr>
<tr>
<td></td>
<td>A2_P</td>
<td>Area under the curve after 1\textsuperscript{st} swallowing (a.u.)</td>
</tr>
<tr>
<td></td>
<td>A_P</td>
<td>Total area under the curve (A = A1_P + A2_P)</td>
</tr>
<tr>
<td></td>
<td>IMax1_P</td>
<td>Maximum intensity before 1\textsuperscript{st} swallowing (a.u.)</td>
</tr>
<tr>
<td></td>
<td>tMax1_P</td>
<td>Time to reach maximum intensity before 1\textsuperscript{st} swallowing (s)</td>
</tr>
<tr>
<td></td>
<td>IMax2_P</td>
<td>Maximum intensity after 1\textsuperscript{st} swallowing (a.u.)</td>
</tr>
<tr>
<td></td>
<td>tMax2_P</td>
<td>Time to reach maximum intensity after 1\textsuperscript{st} swallowing (s)</td>
</tr>
<tr>
<td></td>
<td>Vmax_P</td>
<td>Rate of release (a.u./s)</td>
</tr>
<tr>
<td>X2: Bolus rheology</td>
<td>Moist</td>
<td>Moistening of the bolus just before the swallowing (%)</td>
</tr>
<tr>
<td></td>
<td>sflow</td>
<td>Yield stress at flow phase of compression curve (Pa)</td>
</tr>
<tr>
<td></td>
<td>spart</td>
<td>Yield stress at particle phase of compression curve (Pa)</td>
</tr>
<tr>
<td></td>
<td>hpart</td>
<td>Bolus height at the beginning of the particle phase of compression curve (mm)</td>
</tr>
<tr>
<td></td>
<td>Kflow</td>
<td>Consistency at the flow phase, which reflects bolus consistency (Pa.s)</td>
</tr>
<tr>
<td></td>
<td>hflow</td>
<td>Bolus height at the beginning of the flow phase of compression curve (mm)</td>
</tr>
<tr>
<td></td>
<td>hend</td>
<td>Bolus height at the end of compression (mm)</td>
</tr>
<tr>
<td></td>
<td>Send</td>
<td>Area at the end of compression (mm\textsuperscript{2})</td>
</tr>
<tr>
<td>X3: Coating- oral vol</td>
<td>QRB</td>
<td>Amount of product remaining in the oral cavity after swallowing (%)</td>
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<tr>
<td></td>
<td>Oral_Vol</td>
<td>Volume of the oral cavity (cm\textsuperscript{3})</td>
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<td>X4: Masticatory parameters</td>
<td>Nber_Cycle</td>
<td>Number of chewing cycle</td>
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<td></td>
<td>Chew_time</td>
<td>Chewing duration (s)</td>
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<tr>
<td></td>
<td>Ampl</td>
<td>Mean amplitude of contraction (mV)</td>
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<tr>
<td></td>
<td>W_tot</td>
<td>Energy expended in chewing (mV/s)</td>
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<td>X5: Properties of</td>
<td>Flux_S</td>
<td>Salivary flow stimulated saliva (ml/min)</td>
</tr>
<tr>
<td>stimulated saliva</td>
<td>Prot_S</td>
<td>Amount of salivary proteins stimulated saliva (mg/ml)</td>
</tr>
<tr>
<td></td>
<td>Lipolysis_S</td>
<td>Amount of Lipolysis in stimulated saliva (mU/ml)</td>
</tr>
<tr>
<td></td>
<td>Amylase_S</td>
<td>Amount of Amylase in stimulated saliva (U/ml)</td>
</tr>
<tr>
<td></td>
<td>Proteolysis_S</td>
<td>Amount of Proteolysis in stimulated saliva (U/ml)</td>
</tr>
<tr>
<td></td>
<td>Na_S</td>
<td>Amount of sodium in stimulated saliva (mM)</td>
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</tbody>
</table>
Table 2: oral physiological characteristics of the 14 subjects included in the study, mean, standard deviation, minimum and maximum values.

<table>
<thead>
<tr>
<th></th>
<th>minimum</th>
<th>maximum</th>
<th>mean</th>
<th>Standard deviation</th>
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<tr>
<td>Oral volume (cm$^3$)</td>
<td>Oral_Vol</td>
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<td>39.03</td>
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<td>Stimulated salivary flow (ml/min)</td>
<td>Flux_S</td>
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<td>Protein (mg/ml)</td>
<td>Prot_S</td>
<td>0.57</td>
<td>1.31</td>
<td>1.04</td>
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<td>Lipolysis (mU/ml)</td>
<td>Lipolysis_S</td>
<td>&lt;0.01</td>
<td>1.32</td>
<td>0.90</td>
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<td>Amylase (U/ml)</td>
<td>Amylase_S</td>
<td>1.27</td>
<td>30.6</td>
<td>20.86</td>
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<tr>
<td>Proteolysis (U/ml)</td>
<td>Proteolysis_S</td>
<td>0.04</td>
<td>0.51</td>
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<tr>
<td>sodium content (mM)</td>
<td>Na_S</td>
<td>2.09</td>
<td>37.82</td>
<td>15.82</td>
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Figure captions:

Figure 1. Time intensity curves obtained for one subject for blue cheese aroma (NO: nonan-2-one) for the four cheeses (hfS: high fat soft, hfF: high fat firm, lfS: low fat soft, lfF: low fat firm) showing the extracted parameters (Imax_S: maximum intensity, Tmax_S: time to reach maximum intensity, Ideg_S: intensity at swallowing and Tend_S: time to reach the end of perception.

Figure 2. Bibplot representation from the PCA performed on the 14 subjects and their physiological parameters.

Figure 3. Bar charts representing the relative importance (%) of the different blocks of variables (X1-X5) for the different components obtained by means of MB-PLS analysis performed on fruity aroma (EP) and blue cheese aroma (NO) sensory data set and for the high fat (hfF & hfS) cheese products. Blue chart: X1-Aroma release, Green chart: X2-rheology, Orange chart: X3-coating and oral volume, Red chart: X4-EM data, Violet chart: X5-stimulated saliva composition.

Figure 4. MB-PLS results on dim1/dim2 for high fat (hf) cheeses: relationships between the X-blocks of explanatory variables (Blue arrows: X1-aroma release, Green arrows: X2-rheology, Orange arrows: X3-coating and oral volume, Red arrows: X4-EMG data, Violet arrows: X5-stimulated saliva composition) and the Y block of variables to be explained (Black arrows: sensory data for fruity aroma (EP) and blue cheese aroma (NO). Top: firm high fat cheeses (hfF), bottom: soft high fat cheeses (hfS). The font size for each variable reflects the importance of the correlation coefficient with the two components.

Figure 5. Bar charts representing the relative importance (%) of the different blocks of variables (X1-X5) for the different components obtained by means of MB-PLS analysis performed on fruity aroma (EP) and blue cheese aroma (NO) sensory data set and for the low fat (lfF & lfS) cheese products. Blue chart: X1-Aroma release, Green chart: X2-rheology, Orange chart: X3-coating and oral volume, Red chart: X4-EM data, Violet chart: X5-stimulated saliva composition.

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254x190mm (96 x 96 DPI)
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705x448mm (72 x 72 DPI)
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