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INSIGHTS IN AROMA COMPOUND RETENTION BY MUCOSA DURING CONSUMPTION THROUGH MATHEMATICAL MODELLING

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Highlights

- Aroma compound persistence after gaseous sample inhalation was modelled.

- Both physiological and physicochemical parameters were included in the model.

- The respective contributions of wetted mucosa and saliva on release kinetics were assessed.

- The major role of mucosa partition coefficients of molecules was confirmed.
Abstract

A multidisciplinary approach combining physiology and physical chemistry and associating experimental measurements with in silico modelling was applied to explain the release of aroma compounds during food consumption. Experimental release kinetics obtained by inhaling gaseous samples through controlled protocols highlighted different release behaviours, depending on aroma compound properties. The associated mathematical model described mass transfer mechanisms between the different compartments of the naso-oro-pharyngeal cavities and included both physicochemical and physiological parameters. One of the main developments was notably to consider the possible retention of aroma compounds by wetted mucosa. Model sensitivity analysis confirmed the key role of interaction between aroma compounds and mucosa (air/mucosa partition coefficient) and of individual breath parameters (current breath volume and respiratory frequency) on the persistence of aroma compound in exhaled air. These achievements show that the association of an experimental approach and mechanistic modelling constitutes a powerful tool to improve the understanding of aroma release and persistence.

Keywords: aroma release, wetted mucosa, saliva, persistence, interaction, dynamic modelling

Chemical compounds studied in this article:

Short running title: Modelling of in vivo aroma retention

Highlights

- Aroma compound persistence after gaseous sample inhalation was modelled.
- Both physiological and physicochemical parameters were included in the model.
- The respective contributions of wetted mucosa and saliva on release kinetics were assessed.
- The major role of mucosa partition coefficients of molecules was confirmed.
1. Introduction

Olfactory perception is known to largely contribute to overall perception of foods and, consequently, to consumer choice and preferences. A better understanding of this specific perception is therefore of great importance and requires the identification of the main mechanisms at the origin of aroma compound release during food consumption. Several studies have notably focused on orthonasal and retronasal perceptions to highlight the origin of the main differences between these two perception pathways (Espinosa Diaz, 2004; Halpern, 2004; Heilman and Hummel, 2004; Hummel, 2008; Hummel et al., 2006; Sun and Halpern, 2005; Visschers et al., 2006; Welge-Lüssen et al., 2009). The large number and the variety of mechanisms (physical, chemical, physiological, neurobiological, cognitive, etc.) that can be involved at different space and time scales largely contribute to the complexity of perception. Among them, the release dynamics of aroma compounds have long been known to be among key factors to explain aromatic perceptions (Barron et al., 2012; Biasioli et al., 2006; Déléris et al., 2011; Gierczynski et al., 2011; Heenan et al., 2009). Numerous studies in the literature have focused on the identification of the main factors that can impact release kinetics, either related to the physicochemical properties of the molecules, to product characteristics (composition, structure), to individual physiology (saliva composition and flow rate, breath flow rate) or to oral processing (chewing efficiency, product coating, etc.) (Benjamin et al., 2012; Buettner and Beauchamp, 2010; Foster et al., 2011; Frank et al., 2012; Gierczynski et al., 2011; Heath, 2002; Heenan et al., 2011). The existence of aroma compound retention by wetted mucosa has often been proposed to explain specific release behaviours, but little is known about the origin of this phenomenon. However, several mechanisms have been suggested in the literature: the interaction of aroma compounds with the constituents of the mucus layer (mucins, enzymes, antioxidants, ionic compounds), with saliva and/or with the mucosa tissues themselves (Buettner and Beauchamp, 2010); the role of
the contact area between nasal mucus and air (Keyhani et al., 1997); the role of the
physicochemical properties of aroma compounds (Ferreira et al., 2006; Tromelin et al., 2010);
and the role of breath and/or salivary flow rates (Buettner and Mestres, 2005; Hodgson et al.,
2004).
Performing \textit{in vivo} experiments and developing appropriate experimental set-ups constitute
the main difficulties involved in exploring this topic and in validating or not the assumptions.
A previous study proposed various simple protocols to explore and quantify \textit{in vivo} aroma
release and persistence from gaseous samples, depending on the exposed physiological
cavities (nose, mouth or pharynx) (Déléris et al., 2015). Results confirmed the main role of
aroma compound properties and highlighted the possible occurrence of different types of
mechanisms, either physical or biochemical, to explain release behaviours. The global nature
of the approach and the complexity of the phenomena involved did not allow the authors to
clearly identify the relative contribution of each mechanism.
The difficulty of dissociating all of the phenomena that occur during \textit{in vivo} experiments
generally prevents from determining the respective contribution of product properties or of
consumer characteristics to aroma release. Due to these experimental issues, the modelling
approach (\textit{in silico}) can be a useful tool to improve the understanding. It has been largely used
in the fields of pharmacokinetics and toxicology: Quantitative Structure-Activity
Relationships (QSAR) (Geerts and Heyden, 2011), Physiologically-Based Pharmaco-Kinetic
(PB-PK) (Corley et al., 2012; Medinsky et al., 1993; Morris, 2012) and Theoretical Passive
Absorption (TPAM) (Obata et al., 2005; Takano et al., 2006) models have helped to better
understand drug and toxic vapour absorption. In the field of olfaction, the QSAR approach
has also been largely used to identify the main interactions between aroma compounds and
olfactory receptors at the origin of perception (Anker et al., 1990; Chastrette and Rallet, 1998;
Kraft et al., 2000; Rognon and Chastrette, 1994; Sanz et al., 2008). Some of these models
clearly highlighted the need to consider absorption/solubilisation phenomena in tissues of the respiratory tract and/or in the mucus layer to correctly represent the availability of aroma compounds for olfactory receptors. It was demonstrated that the transport of odorant molecules in nasal mucosa clearly differs from the one within an aqueous layer (Kurtz et al., 2004). The main limitation of modelling approaches remains the lack of experimental data, notably concerning the air/mucosa partition or diffusion properties of aroma compounds within the mucus layer, or mucosa characteristics depending on its location (nasal, oral or pharyngeal cavity).

In food science, some mechanistic models describing volatile release have been proposed and sometimes compared to experimental in vivo data (Buffo et al., 2005; Harrison, 2000; Harrison and Hills, 1997; Hodgson et al., 2005; Normand et al., 2004; Wright and Hills, 2003). These models, based on physical, chemical and physiological parameters, led to more or less good predictions of the release kinetics of aroma compounds, but only for liquid food products. Only the models of (Wright and Hills, 2003) and (Normand et al., 2004) included a term representing possible interactions between aroma compounds and mucosa and/or salivary constituents. Even though many publications exist on molecular mechanisms that explain interactions between aroma compounds and proteins in the mucus of the nasal cavity of rats (Odorant Binding Proteins, OBPs), results cannot be directly used to explain in vivo release kinetics in humans (Borysik et al., 2010; Yabuki et al., 2011). All of these studies constitute a first step in describing the phenomena involved but do not yet provide a clear understanding. In a previous publication, a mathematical model was proposed to predict in vivo aroma release from masticated food products that considered food properties and the physiological characteristics of the individuals (Doyennette et al., 2014). Comparison between experimental and predicted kinetics highlighted the possible specific retention of one hydrophobic aroma compound by wetted mucosa and mucus in the naso-oropharyngeal
cavities. This model thus needs to be further developed to propose a satisfactory quantitative description of the retention phenomenon at the origin of aroma persistence.

In this context, the main goal of the present study is to better understand the mechanisms underlying aroma release and persistence. The originality of the proposed approach is to combine: (i) in vivo aroma release measurements (using controlled protocols to ensure aroma supply by flavoured air inhalation, without the interference of any food product); with (ii) the detailed mechanistic modelling of mass transfer to investigate the key mechanisms responsible for the release profiles and/or retention of aroma compounds.

2. Material and methods

Even if this study was not performed in the field of medical research, a detailed research protocol containing the relevant information in agreement with the World Medical Association Declaration of Helsinki was done. Only single-use materials were used with panellists. Aroma compounds were all food grade and their liquid concentrations were adjusted to limit gaseous concentration and ensure panellist comfort and avoid sensory saturation. Only one session (45 minutes) per week was planned for each panellist and the number of samples during one session was limited to five. Samples were coded to protect the privacy of panellist and the confidentiality of their personal information. Subjects were clearly informed of the observational nature of this study, gave their free and informed consent and received compensation for their participation.

2.1. Aroma compounds

Food grade quality aroma compounds (ethyl propanoate, 2-nonanone and (Z)-3-hexen-1-ol) were purchased from Sigma Aldrich (France) (Table 1).

They were selected since they belong to several chemical classes and present different physicochemical properties and different release behaviours in terms of persistence (Déléris et al., 2015). Concentrated stock solutions were prepared in polypropylene glycol (Sigma
Aldrich, France) and used throughout the study. Diluted solutions were prepared extemporaneously.

2.2. Gaseous sample preparation

An aroma compound mixture was used to reduce the number of experimental sessions. Gaseous samples were prepared as previously described (Déléris et al., 2015). The concentrations of aroma compounds in the liquid phase were high enough to be detected during PTR-MS measurements, while being acceptable from a sensory point of view for the panellists: 1000 mg/kg for (Z)-3-hexen-1-ol, 150 mg/kg for ethyl propanoate and 100 mg/kg for 2-nonanone.

Twenty-five mL of flavoured aqueous solution were stored at ambient temperature for 4 hours before measurements in 250-mL flasks (Schott, France), closed by caps equipped with valves (equilibrium establishment). To control the inhaled volume of gaseous sample (and, therefore, the amount of inhaled aroma compounds) between the different assays, a specific set-up was developed to prepare gaseous samples (Figure 1): a manual pump was connected to one of the cap valves and used to push some fresh air into the flasks. By way of this procedure, flavoured air was introduced into a balloon positioned on the other valve of the flask cap. Three pump strokes were needed to prepare 200 mL of gaseous sample, which was considered as appropriate to be inhaled by panellists in one breath. Once inflated, balloons were closed with plastic pliers.

The study of the variation of aroma concentration within balloons during storage highlighted the fact that this preparation had to be done less than 30 s before measurement to avoid any loss of aroma compounds (not shown). Even if some interactions between aroma compounds and balloon material could occur, they were assumed to always be the same and to not influence the results since all conditions were controlled.

2.3. Panellists
Eight panellists (four men and four women, 22-45 years old) were recruited for the study. They were instructed not to smoke, eat, drink or use any persistent-flavoured product for at least one hour before Proton Transfer Reaction-Mass Spectrometry (PTR-MS) or saliva collection sessions.

When dealing with aroma release and food oral processing, lots of studies in literature largely highlighted the key role that anatomy and physiology can have on the dynamics of phenomena (Buettner and Beauchamp, 2010; Féron et al., 2014; Foster et al., 2011; Repoux et al., 2012b). Some physiological measurements were thus performed on panellists who participated to this study to define the range of variation of these parameters for the panel.

The volumes of the oral, nasal and pharyngeal cavities of the subjects were measured with the Eccovision Acoustic Rhinopharyngometer (Sleep Group Solutions, North Miami Beach, FL, USA). Software was developed to automatically calculate the air/product areas of the oral and pharyngeal cavities for each individual (Doyennette et al., 2011). The tidal volume of each individual was measured with a spirometer (Pulmo System II, MSR, Rungis, France) (Repoux et al., 2012a).

Non-stimulated saliva was collected by asking volunteers to swallow the saliva in their mouth before starting and to then spit each 30 s for 5 min into ice-chilled vessels. The final saliva weight was measured and the flow rate was calculated as g/min. Whole saliva samples were centrifuged at 13400×g for 5 min at 4°C to remove cellular debris (Eppendorf, model 5415 R, Germany). The supernatants were frozen and stored at -80°C before analysis. Protein concentration (expressed in mg/mL) was obtained by standard Bradford protein assay Quick Start (Bio-Rad, France) using bovine serum albumin (Sigma-Aldrich, France) as the standard calibration. The lipolytic (lipolysis), proteolytic (proteolysis), lysozymal (lysozyme) and amylolytic (amylase) activities of individual salivas (expressed in U/mL) were determined as previously described (Neyraud et al., 2012).
Three replicates per physiological parameter and per panellist were performed. The minimal, median and maximal values of physiological characteristics and associated quartiles are summarized in Table 2.

2.4. Determination of aroma in vivo release kinetics using PTR-MS measurements

Release kinetics were obtained using the reference protocol previously defined (referred to as the Nose, Mouth, Swallowing protocol, or N.M.S.) (Déléris et al., 2015): sample inhalation was performed through the panellist's mouth in one short breath and the measurement of aroma release was made within the panellist's nasal cavity.

During a session, subjects started with the analysis of a blank sample to get used to the protocol and then tested five samples. The measurement procedure was similar to the one previously described (Déléris et al., 2015): room and breath analyses for 10 s and 30 s, respectively, followed by sample inhalation and release measurement. During the assay, panellists were allowed to swallow. Between each sample, panellists cleaned their mouth with mineral water (Evian, Danone) and their breath was retested before each new measurement.

Some differences with the previous study should be mentioned. First, swallowing events were imposed: 20 s after sample inhalation for the first swallow, and then every 30 s until the end of measurement. Secondly, the sample volume was standardised, allowing the quantitative comparison between protocols. Three sessions of 45 min were planned to obtain three replicates of the five samples for each subject. All the measurements were performed within a 21-day period.

The High-Sensitivity Proton Transfer Reaction-Mass Spectrometer (PTR-MS) (Ionicon Analytik, Innsbruck, Austria) was operated at a drift tube temperature, voltage and pressure of 60°C, 600.1 (±0.4) V and 2.0 (±0.01) mbar, respectively \( (E/N=151.4 \ (±1.4) \ \text{Td}) \). Nose-space was sampled via two inlets of a stainless nosepiece placed in both nostrils of the assessors. The inlet of the PTR-MS instrument was connected to the sampling device via a 1/16”
PEEK™ tube maintained at 110°C. Measurements were performed using the Multiple Ion Detection (MID) mode. For a mass/charge ratio (m/z) of 21 (H$_3$O$^+$) and 37 (H$_2$O-H$_3$O$^+$), the dwell time per mass was fixed to 0.05 s. The mean signal for H$_3$O$^+$ was 7.8×10^6 ± 0.8×10^6 counts per second (cps) and its day-to-day variation along the measurement period was 10%. The signal for H$_2$O-H$_3$O$^+$ did not exceed 4% of the one of m/z 21 (in agreement with equipment specifications).

Using the fragmentation patterns of individual compounds (Table 1), the molecules studied were monitored at m/z 83 ((Z)-3-hexen-1-ol), m/z 75 and 103 (ethyl propanoate) and m/z 143 (2-nonanone). For these four specific masses, a dwell time per mass of 0.1 s was selected as a compromise between sensitivity for aroma compound detection and appropriate sampling frequency with regard to phenomena to be measured. In addition, m/z 59 and 93 were monitored with a dwell time per mass of 0.05 s as markers of panellists’ breath (Weel et al., 2002) and of balloon material, respectively. With these settings, exhaled air was sampled every 0.6 s, which was assumed to be appropriate regarding the mean duration of the breathing cycle (Doyennette et al., 2011; Sherwood, 2006; Tortora and Anagnostakos, 1990).

Mean signal-to-noise ratios varied between 2.0 and 38.0 (depending on the ions), meaning that responses during sample analysis sufficiently exceeded the baseline. These measurements led to the determination of molecule release kinetics, *i.e.*, intensity I$_t$ = f(time t), for each panellist. Since solution composition was precisely known, aroma compounds were unambiguously detected at the stated m/z ratio. For this reason and to facilitate text readability, compound names rather than their m/z ratio are used hereafter.

For data handling, experimental release curves were divided into three main periods: (i) the phase before the product was inhaled (phase 0); (ii) the phase before the first swallow (phase 1); and (iii) the phase after the first swallow (phase 2). For each sample, the mean PTR-MS signal measured during phase 0 was subtracted from the PTR-MS signals obtained during
phases 1 and 2. Some quantitative release parameters were extracted from each individual release curve and for each phase of product consumption: maximal intensities ($I_{\text{max}1}$ and $I_{\text{max}2}$), which indicate the maximum concentration reached by a compound; and areas under the curve ($\text{AUC}_1$ and $\text{AUC}_2$), which are related to the total amount of molecule that is released). The ratios between areas under the curve before and after swallowing were also calculated ($\text{AUC}_1/\text{AUC}_2$). Some temporal release parameters such as times at which $I_{\text{max}}$ occurred ($t_{\text{max}1}$ and $t_{\text{max}2}$) and initial release rates (Rate$_1$ and Rate$_2$, calculated by dividing the $I_{\text{max}}$ values with the $t_{\text{max}}$ times) were also determined. Peak widths for each phase were obtained as the difference between the two times at which the intensity was 20% of $I_{\text{max}}$ (after and before release peak) ($\Delta t_{20\%_1}$ and $\Delta t_{20\%_2}$). The difference between the time at which the intensity was 50% of $I_{\text{max}}$ after the peak and $t_{\text{max}}$ ($t_{50\%}-t_{\text{max}}$) was also extracted. In addition, standardised release kinetics were obtained by dividing each intensity value of the curve by the corresponding $I_{\text{max}}$ ($I_{\text{t, stand}}=I_t/I_{\text{max}}$). Standardised areas under the curve ($\text{AUC}_{\text{stand}}$), determined from these standardised kinetics, were used as an indication of persistence behaviour. Because the objective was to compare the persistence of aroma release between molecules, the use of arbitrary units for aroma concentration data was sufficient for data analysis. Since the two ions related to ethyl propanoate behaved in the same way, only the result of $m/z$ 75 is presented in the text.

Non-parametric descriptive analysis was carried out on datasets and comparative analysis was performed using Kruskal-Wallis tests and the Conover-Iman procedure (multiple paired comparisons) to highlight differences in in vivo release kinetics between molecules. The level of significance was set at $p<0.05$.

3. Modelling

3.1. Principles of the model
The aroma release model presented in this study was developed to describe aroma release after one inhalation of a gaseous sample through the mouth. It is based on equations that describe mass transfers that occur between the different physiological cavities (mouth, nose, pharynx), considered as interconnected reactors that vary in volume and that exchange matter.

A schematic representation of the four physiological cavities involved in the model design, as well as their connections and the mechanisms responsible for aroma release, are given in Figure 2. A schematic representation of mass transfer between air, saliva and mucosa within a physiological cavity is presented in Figure 3. All variables and parameters required for the model simulation are specified in these figures and described in Tables 3 and 4, respectively.

The model describes the steps of the experimental protocol: sample inhalation through the mouth, exposure period to the sample until the first swallow, and post-swallow release. Except for the first inhalation, breathing occurs through the nose. Each swallowing step is known to be very short (Martin-Harris, 2006) compared to persistence phenomena (Hodgson et al., 2004; Normand et al., 2004). Swallowing events are thus described as quick simultaneous contractions of the oral cavity and of the pharynx, leading to air expulsion, followed by relaxation and filling with fresh air (Doyennette et al., 2014).

Two compartments in the mouth and in the pharynx (mucosa and saliva) and one compartment in the nose (mucosa) were included in the model to introduce a reservoir effect (Figure 2). In each cavity, the air phase was assumed to be in contact with mucosa and/or saliva layers. The proportions of these contact areas can be changed in the model to evaluate the respective contributions of saliva and mucosa. The volumes of the layers involved in the interaction with aroma compounds were expressed as products between compartment areas and layer thicknesses (Eq. (A.12) to Eq. (A.16) in the Appendix A). Similarly to Doyennette et al. (2014), transfer resistances on the air side (1/k_{Oa}, 1/k_{Fa} and 1/k_{Na}) were assumed to be
negligible when compared to the transfer resistance on the wetted mucosa or saliva sides
(1/k_{Oms}, 1/k_{Fm}, 1/k_{Nm} and 1/k_{Os}, 1/k_{Fs}, respectively).

To improve the readability of the paper, only generic equations describing phenomena are
given in the text. They were written specifically for each compartment to obtain the complete
description of aroma release. The details of all model equations are given in the Appendix A.

3.2. Air flow rates

By convention, the air flow rates indicated in Figure 2 are positive if they follow the direction
of the arrow. With this convention, air flow rates in the different cavities $Q_{Oa}$, $Q_{OFa}$, $Q_{Na}$ and
$Q_{NFa}$ are positive (or null, depending on whether breathing occurs through the nose or the
mouth) during inhalation, and negative (or null) during exhalation. Conversely, air flow rate
from the trachea $Q_{Ta}$ is negative during inhalation and positive during exhalation.

Air flow rate in the trachea due to breathing was assumed sinusoidal (Eq. (A.33)). According
to Figure 2 and to the experimental protocol, the air flow rate in the mouth is given by the
breathing flow rate during the first inhalation and is null afterwards (Eq. (A.34)). The opposite
was considered for the air flow rate in the nasal cavity (Eq. (A.35)). According to Figure 2,
the air balance in the pharynx at any time is given by the equality of inlet and outlet fluxes
(Eq. (A.36)).

3.3. Saliva in oral cavity

The volume of saliva in the oral cavity gradually increases due to the salivary flow rate $Q_{Os}$
and abruptly decreases after swallowing. A minimal residual volume of saliva $V_{Osmin}$ was
assumed to remain in the mouth after swallowing (Doyennette et al., 2014).

3.4. Mathematical description of interfacial conditions and fluxes

3.4.1. Air/mucosa and air/saliva interfaces

The interfacial aroma compound concentrations at air/mucosa or air/saliva interfaces were
obtained from the partition conditions at the interfaces, using the following generic equations:
Recall that transfer resistances on the air side were assumed negligible, hence, in the air, bulk and interfacial concentrations are identical. Specifically, the air/mucosa interfacial aroma compound concentrations are described by Eq. (A.17), Eq. (A.18) and Eq. (A.19) in oral, nasal or pharyngeal cavities, respectively. The air/saliva interfacial aroma compound concentrations are given by Eq. (A.20) and Eq. (A.21) in the oral or pharyngeal cavities, respectively.

Volatile mass fluxes \( \phi_{am} \) and \( \phi_{as} \) between the air and the other compartments (mucosa or saliva) are determined by the resistances located on the mucosa and saliva sides, respectively. They are given by the difference between the mucosa \( (C_m) \) or saliva \( (C_s) \) concentrations and the interfacial concentrations \( (C_{am}^\ast) \) or \( (C_{as}^\ast) \) and are calculated using the following generic equations:

\[
\phi_{am} = k_m \times A_{am} \times \left( C_m(t) - C_{am}^\ast(t) \right), \quad \phi_{as} = k_s \times A_{as} \times \left( C_s(t) - C_{as}^\ast(t) \right)
\]

Specifically, in the oral cavity these fluxes are given by Eq. (A.26) and Eq. (A.29), in the pharynx by Eq. (A.28) and Eq. (A.30) and in the nose by Eq. (A.27).

### 3.4.2. Mucosa/saliva interface

Since the mucosa was assumed to be partially wetted by the saliva, aroma partition and flux between mucosa and saliva compartments was also considered in the oral and pharyngeal cavities. Transfer resistances were considered in both saliva and mucosa layers since they are expected to be of comparable magnitude. Interfacial aroma compound concentrations at mucosa/saliva interface were obtained from the partition conditions at the interfaces, using the following generic equation (Figure 3):

\[
C_{sm}^\ast(t) = \frac{C_{ms}^\ast(t)}{K_{ms}}
\]

The mucosa/saliva interfacial aroma compound concentrations are described by Eq. (A.22) and Eq. (A.24) in oral or pharyngeal cavities, respectively. The partition coefficients between...
mucosa and saliva were calculated based on partition coefficients with the air phase (Eq. (A.23) and Eq. (A.25)):

\[ K_{ms} = \frac{k_{as}}{k_{am}} \quad \text{Eq. 4} \]

Since the mucosa was assumed to be partially wetted by the saliva, a volatile flux between saliva and mucosa \( \phi_{ms} \) was also considered (Eq. (A.31) in the oral cavity and Eq. (A.32) in the pharyngeal cavity):

\[ \phi_{ms}(t) = k_{eq} \times A_{ms} \times \left( C_s(t) - \frac{c_{m(t)}}{K_{ms}} \right) \quad \text{Eq. 5} \]

with \( k_{eq} \) being the equivalent mass transfer coefficients between saliva and wetted mucosa, with saliva taken as reference. It includes resistances in both phases in contact and the partition between them (Marin et al., 1999):

\[ \frac{1}{k_{eq}} = \frac{1}{k_s} + \frac{K_{ms}}{k_m} \quad \text{Eq. 6} \]

3.5. Volatile mass balances

According to Figure 2, in each considered cavity the air may exchange aroma compounds with mucosa, saliva and air of connected cavities, as applicable. The generic volatile mass balance for the air in a cavity has the following form, where the term “source” denotes cavities which supply air to the cavity under consideration:

\[ V_a \frac{dc_a}{dt} = \phi_{am} + \phi_{as} + \sum_{\text{source}} Q_{a \text{source}} (C_{a \text{source}} - C_a) \quad \text{Eq. 7} \]

For the first inhalation, the variation of aroma concentration in the air in the oral cavity \( C_{oa} \) is due to the volatile flux from the inhaled air sample and also to contact with the mucosa and saliva layers (Eq. (A.37)). For the following breathing cycles (through the nose), the mouth is closed (Eq. (A.38)). The mass balance in the air of the nasal cavity is given by Eq. (A.39), with no saliva layer present and the possible source of volatile compounds being the pharynx, during expiration. The pharynx has both saliva and mucosa layers and can exchange air with
the mouth (first inhalation) nose (subsequent breathing) and trachea, as described by Eq. (A.40).

Saliva layers in mouth (Eq. (A.41)) and pharynx (Eq. (A.42)) exchange aroma compounds with air and mucosa, according to the generic mass balance:

\[ V_s \frac{dC_{ps}(t)}{dt} = -\phi_{as}(t) - \phi_{ms}(t) \]  
Eq. 8

with the saliva in the oral cavity being additionally diluted by the fresh saliva flow rate.

Mucosa layers are in contact with air and saliva (except in the nose) and the mass balance for these layers was written on the basis of Eq. 9 (developed as (Eq. A.43), (Eq. A.44) and (Eq. A.45) in the oral, nasal and pharyngeal cavities, respectively):

\[ V_m \frac{dC_{m}(t)}{dt} = \phi_{ms}(t) - \phi_{am}(t) \]  
Eq. 9

3.6. Reference values of the parameters

The reference values of parameters used for simulations are given in Table 4. Some were taken from experimental data or from the literature, and were estimated when little information was available. For instance, the contact area between air and mucosa in the nose \( A_{N_{am}} \) was set to 150 cm² (Levitzky, 2003). Concerning air/mucosa partition properties, values comprised between \( 5.6 \times 10^{-5} \) and \( 4.8 \times 10^{-1} \) were found for the air/mucus partition coefficient of butanol and octanol in bullfrog (Hornung et al., 1987). Thus, a typical reference value of \( 1 \times 10^{-3} \) was selected in this case. Concerning the mucosa layer thickness, values between 500 and 800 µm in the mouth and 100 to 200 µm for the gingival mucosa were reported (Patel et al., 2012; Shojaei, 1998). It is expected, however, that aroma compounds will not necessarily have time to diffuse in the whole epithelium thickness, so these values were considered as upper limits for the mucosa layer thickness involved in aroma retention. That is why the reference values for mucosa layer thicknesses in the present case in the different compartments were fixed at 50 µm. On the basis of previous studies (Doyennette et al., 2014), the respiratory frequency \( F_R \) was set to 0.24 cycles per second. An analysis of model
sensitivity was done to determine the respective influence of each parameter on aroma release kinetics.

3.7. Numerical methods

3.7.1. Solution of model equations

The dynamic model developed in this study consisted in nine coupled nonlinear differential equations: eight for aroma compound concentrations in air, mucosa (oral, pharynx and nasal cavities) and saliva (oral cavity and pharynx) plus one differential equation for saliva accumulation in the oral cavity between swallowing events. Numeric calculations were performed with Matlab® 8 software (The MathWorks Inc., Natick, MA). The variable step, stiff ODE solver “ode15s” in the Matlab ODE suite (Shampine and Reichelt, 1997) was used with both absolute and relative tolerances for all equations set to $10^{-8}$. Integration step was adjusted internally to meet the specified tolerances while results were provided at the required (e.g. measurement) times. Integration was halted and restarted at each swallowing event to allow abrupt changes in state variables, e.g. saliva volume decrease and air mixing in pharynx related to the quick deglutition process (Doyennette et al., 2014).

3.7.2. Sensitivity analysis

The model contains a total of 27 independent parameters (Table 4), i.e. which cannot be calculated based on other ones (such as a volume being the product of an area and a layer thickness). To assess the importance of these parameters for the prediction of the volatile compound concentration in the nasal cavity (model output) a global Monte Carlo sensitivity analysis was performed as follows. The model output was the relative volatile concentration in the nasal cavity ($C_{Na}^r$), i.e. the predicted concentration ($C_{Na}$) of the aroma compound divided by its maximum value. For a given value $p_j$ of each parameter $p$, the relative local sensitivity was defined as:

$$L(p_j) = \frac{1}{n_t} \sum_{i=1}^{n_t} \left[ \frac{C_{Na}^r(t_i \cdot p_j) - C_{Na}^r(t_i \cdot p_j + \delta p)}{\delta p} \right] \times \frac{p_{\text{ref}}}{\max_{0 \leq t \leq 110} C_{Na}^r(t \cdot p_{\text{ref}}) - \min_{0 \leq t \leq 110} C_{Na}^r(t \cdot p_{\text{ref}})}$$

Eq. (10)
where $C_{Na}^*(t, p_j)$ is the output of the model calculated for time $t_i \in [0, 110]$s with the considered parameter value set to $p_j$ and $\delta p$ is a perturbation of the parameter, taken as 10% of its minimum value indicated in Table 4. Thus $L(p_j)$ represents an approximation of the partial derivative of the nasal concentration with respect to the considered parameter, taken in absolute value and averaged over time. To make sensitivities dimensionless, of order of unity and hence comparable among various parameters, the right-hand side scaling factor was introduced, based on the reference value of the parameter given in Table 4.

The global sensitivity of a parameter was calculated as an average of local sensitivities across a large number of samples taken in the parametric space:

$$G(p) = \frac{1}{1000} \sum_{j=1}^{1000} L(p_j) \quad \text{Eq. (11)}$$

The parametric space was defined by the range of variation of each parameter indicated in Table 4 and sampled according to a multidimensional “Latin hypercube” method to ensure a uniform representation of all parameter values. Parameters whose range of variation spanned more than one order of magnitude were evenly sampled on a logarithmic rather than linear scale.

Automatic sensitivity analysis gives global information on the relative importance of various parameters for the nasal concentration prediction, but provides little insight in the involved phenomena. A manual sensitivity analysis was also performed by varying some of the parameters (specified in the results section) one by one while keeping the others at their reference values. Parameters whose possible variation range was large (sometimes several orders of magnitude) were included in the manual sensitivity analysis, while those relatively well known from the experimental protocol and physiological measurements were kept constant.

3.7.3. Model fitting
To test the ability of the model to reproduce release curves observed for different molecules, some of the least well known parameters were estimated based on release data. The number of estimated parameters was kept at a minimum, however. Volumes, contact areas, respiratory frequency and deglutition times were either known from physiological measurements or imposed by the experimental protocol; these parameters were kept fixed to their reference values indicated in Table 4. Partition coefficients between air and saliva ($K_{Oas} = K_{Fas}$) were experimentally determined for the three aroma compounds; these values were used without change. The estimated parameters, assumed to be the same in all cavities, were the transfer coefficient in the saliva ($k_{Os} = k_{Fs}$, the same for all molecules) because in vivo hydrodynamic conditions are poorly known, and parameters related to the mucosa layer: thickness ($e_{Om} = e_{Nm}$, common to all molecules), transfer coefficient ($k_{Om} = k_{Nm}$, also common to all molecules) and air/mucosa partition coefficients ($K_{Oam} = K_{Fam} = K_{Nam}$), expected to vary strongly with the physico-chemical properties of the studied compounds and hence specific to each molecule.

Model fitting was thus performed simultaneously using data from release experiments with the three molecules, the partition coefficients being specific to each molecule but common to all cavities and the other abovementioned parameters common to all molecules and all cavities. The fitting criterion was the sum of the absolute values of the errors between model predictions and experimental measurements of the compound concentration in the nasal cavity, both normalised by their respective maximum values ($C_{Na}$), because measured data was only available in arbitrary units. Since possible ranges of some of the parameters span up to 4 orders of magnitude (Table 4) a “global” optimisation procedure based on genetic programming was used, namely the “ga” implementation in the Matlab® Global Optimization Toolbox, with default settings (e.g. population of 40 individuals, convergence tolerance $10^{-6}$) and the maximum number of generations increased to 200. Parameters whose search range
(Table 4) spanned more than one order of magnitude were sampled on a logarithmic scale, i.e. the logarithm of the parameter was actually searched for by the optimisation algorithm. Since the considered optimisation algorithm is stochastic, several (~10) optimisation runs were performed and the one with the best fit was selected. Consistent convergence to similar values of the parameters (usually within ±15%) was observed in most runs.

4. Results and discussion

4.1. Molecule specific effects on aroma release kinetics

As previously highlighted (Déléris et al., 2015), significant differences between aroma compounds in terms of release descriptors were observed (Figure 4).

Differences in $I_{\text{max}1}$ and $I_{\text{max}2}$ and in AUC$_1$ and AUC$_2$ were explained by differences in inhaled gaseous concentrations of molecules due to different aqueous concentrations and air/water partition properties. They were thus not shown nor discussed here.

Before swallowing (Figure 4-a), no difference between molecules was observed concerning $t_{\text{max}}$. Yet, the peak width $\Delta t_{20\%_1}$ of 2-nonanone was the largest and those of ethyl propanoate the narrowest. After swallowing, differences in release behaviours were more pronounced since some temporal parameters were significantly different between molecules: ethyl propanoate was released the most rapidly and with a quite narrow peak, whereas (Z)-3-hexen-1-ol had the most delayed release, with the largest peak at 50% of $I_{\text{max}}$ ($t_{50\%_1}-t_{\text{max}}$), and 2-nonanone had the largest peak at 20% of $I_{\text{max}}$ ($\Delta t_{20\%_2}$) (Figure 4-b). Initial release rates were molecule-dependent both before and after swallowing (Figure 4-c). Ethyl propanoate was released faster than (Z)-3-hexen-1-ol (before and after swallowing) and 2-nonanone (only before swallowing).

AUC$_{\text{stand}}$, which can be associated with molecule persistence, also reflected differences in release behaviour: the highest value was obtained for 2-nonanone, highlighting quite persistent release behaviour for this molecule (Figure 4-d). In contrast, ethyl propanoate
presented the lowest values of AUC\textsubscript{stand}. The AUC\textsubscript{1}/AUC\textsubscript{2} ratios were higher than 1 for all molecules, meaning that a greater amount was released before swallowing than after. However, a 30-fold increase in AUC\textsubscript{1}/AUC\textsubscript{2} ratio was observed between 2-nonanone and ethyl propanoate, meaning that the latter was mainly released before swallowing, whereas 2-nonanone was released during both phases.

All these results were in agreement with previous observations and confirmed the probable existence of retention phenomena for some aroma compounds. To get insight in the exact nature of these interactions, notably in the respective roles of the physicochemical properties of the molecules, of saliva and/or of mucosa characteristics, model simulations and sensitivity analysis were used. We could also mention that from these results, no clear relationship between aroma release and anatomical or physiological parameters was observed.

4.2. Simulations of aroma compound release kinetics

One of the advantages of using a modelling approach is the possible determination of the time variation of variables that could not be experimentally determined, providing insight in the involved mechanisms. Examples of simulated release kinetics obtained with the model in the different compartments of the naso-oropharyngeal cavities are presented in Figure 5. As a starting point, model parameters were fixed to their reference values, determined either from data in the literature or experimentally (Table 4). Air/mucosa contact areas in the mouth A\textsubscript{Oam} and in the pharynx A\textsubscript{Fam} were fixed at 10% of the total area of the mouth A\textsubscript{O} and the pharynx A\textsubscript{F}, respectively, meaning that 90% of the mouth and pharynx surfaces were wetted by saliva. The resulting kinetics were considered as a reference. The inset on each figure illustrates the first 3 seconds of the release to better understand initial phenomena. Just after sample inhalation through the mouth, the gaseous concentrations of aroma compounds in the pharynx C\textsubscript{Fa} and in the mouth C\textsubscript{Oa} increased (Figures 5-a and 5-d, respectively). In parallel,
molecules accumulated within mucosa and saliva layers both in the mouth (Figures 5-e and 5-f, respectively) and in the pharynx (Figures 5-b and 5-c, respectively). When the mouth was closed (at 2 s), aroma compound gaseous concentration in the air in the mouth started to decrease, as well as in the pharynx. In the mouth, the slow decrease in the gaseous concentration of aroma compounds after 2s was due to adsorption on mucosa and saliva layers. In the pharynx, the expiration flow rate accounts for the much faster decrease since it was responsible for the transport of aroma compounds from the pharynx to the nose (increase in aroma compound concentration in the nose $C_{Na}$ (Figure 5-g) and, therefore, a decrease in aroma compound concentration in the pharynx $C_{Fa}$ (Figure 5-a). The adsorption of aroma compounds on mucosa is also probably involved in this decrease. The pulse in aroma concentration in the nasal cavity led to an increase in aroma concentration within the nasal mucosa layer $C_{Nm}$ (Figure 5-h). Yet, aroma compound concentration in the nose air rapidly decreased to zero since air from the lungs was aroma-free and the aroma transfer from the nasal mucosa to the air in the nose (which occurred as aroma concentration in the nasal mucosa progressively decreased; Figure 5-h) was not rapid enough to compensate for the dilution by the breath flow rate. Similar conclusions can be drawn concerning the variation of concentrations in pharynx compartments.

The first swallow occurred at 20 s. It led to a sudden decrease in saliva volume in the mouth $V_{Os}$ (Figure 5-i), as well as decreases in aroma compound concentrations in the air within the mouth ($C_{Oa}$, Figure 5-d) and in saliva in the mouth $C_{Os}$ (Figure 5-f) and in the pharynx $C_{Fs}$ (Figure 5-c). In all mucosa compartments, concentrations $C_{Fm}$, $C_{Om}$ and $C_{Nm}$ progressively decreased, highlighting the unloading of these compartments (Figures 5-b, 5-e and 5-h, respectively). These mass transfers were probably the limiting steps since aroma concentrations in the different air phases $C_{Fa}$, $C_{Oa}$ and $C_{Na}$ did not increase (Figures 5-a, 5-d and 5-g, respectively). The saliva volume in the mouth followed a cyclic variation, with a
linear increase due to the saliva flow rate between each swallowing event and a sudden
decrease when swallowing occurred.

4.3. Sensitivity analysis of the model to physicochemical and physiological parameters

Modelling makes it possible to easily test the effect of parameters that govern mass transfers,
notably those that cannot be modified when performing *in vivo* studies. Such a sensitivity
analysis can contribute to the determination of the nature of the key factors underlying aroma
release and persistence.

4.3.1. Global sensitivity analysis

Based on the results of the global sensitivity analysis performed as described in section
3.7.2, model parameters were arbitrarily divided in three groups, according to their influence
on the shape of the nasal concentration release curve (recall that nasal concentration was
always normalized by its maximum value). The first group included 3 most influential
parameters, with global relative sensitivities comprised between 0.4 and 0.2: the air/mucosa
partition coefficients in the nose (\(K_{Nam}\)) and in the pharynx (\(K_{Fam}\)) and the respiratory
frequency (\(F_R\)). The second group included 5 moderately influential parameters, with
sensitivities between 0.1 and 0.025: air/mucosa partition coefficient in the mouth (\(K_{Oam}\)),
air/saliva partition coefficient in the mouth (\(K_{Oas}\)) and pharynx (\(K_{Fas}\)), mass transfer
coefficient in the mouth (\(k_{Os}\)) and the tidal volume (\(V_C\)). The other parameters listed in Table
4 had sensitivities less than 0.025.

Overall, these results support the central assumption underlying this work, namely that in
absence of any food product, volatile persistence in consumer’s exhaled air is mainly related
to the interaction between the aroma compound and subject’s mucosa, quantified in the model
via the air/mucosa partition coefficients. A second important factor, already pointed out in
presence of non masticated (Tréléa *et al.*, 2008) and masticated (Doyennette *et al.*, 2014) food
products, is the consumer’s breath via the respiratory frequency and current breath volume.
Mass transfer and geometry (volumes, contact areas) appear to play a smaller role in volatile persistence than in release from food products.

4.3.2. Manual sensitivity analysis

When dealing with mass transfer, the main factors governing molecule transports are the contact area, the driving force (dependent, in particular, on the partition properties of the molecules) and the mass transfer coefficient of the molecules. Simulations were thus performed to evaluate their influence on release kinetics. The ranges of variation of model parameter values were chosen to be in agreement with physicochemical or physiological values (Table 4). Only the effects of these modifications on the simulated release kinetics in the nasal cavity are explained since they correspond to what can be experimentally determined. The variation of aroma concentrations in other compartments were simulated but are not discussed here.

First, to better understand the respective roles of the saliva and mucosa compartments, some simulations were performed without any saliva compartment (air/mucosa contact areas in the mouth and in the pharynx were equal to the total surfaces of the mouth and the pharynx, respectively, so contact areas with saliva were equal to zero, Table 4).

The absence of saliva as well as the modification of mucosa thickness (10-fold variation factor) did not have an impact on the shape of release kinetics of aroma compounds within the nasal cavity. Differences were mainly observed on mucosa concentration and unloading rates (not shown). For example, mucosa concentrations in the mouth and in the pharynx were 5 to 9-fold higher, respectively, without saliva than with saliva, and 10-fold higher or lower when the thickness decreased or increased, respectively. Mucosa compartments also unloaded more rapidly when saliva was not considered or when mucosa thicknesses were lower.

The mass transfer coefficient of aroma compounds in mucosa had a greater impact on the release kinetics in the nasal cavity when it increased than when it decreased (not shown).
With an increased mass transfer coefficient in mucosa, mucosa loading and unloading rates were higher. Since air flow rate remained constant between simulations, mucosa compartments were more rapidly unloaded and the gaseous concentration in the nose was thus lower and decreased more rapidly. The effects with decreased values of mass transfer coefficients were less obvious: below a value of $10^{-6}$ m/s, very slow aroma transport prevents any significant amount of aroma compound to be loaded into the mucosa, so there is almost no impact on aroma concentration in the nose.

Decreasing the air/mucosa partition properties had a big impact on the nasal concentration of aroma compounds and the shape of release kinetics. A low air/mucosa partition means high affinity of the aroma compound for the mucosa compartment. Most of the aroma compound was thus retained within the mucosa and released very slowly in tiny amounts according to breathing cycles. Persistence was thus long but the actual concentration in the nasal cavity was low. A 100-fold increase in the air/mucosa partition from $10^{-3}$ had less effect on the aroma concentration in the nose than a 100-fold decrease, due to the fact that the amounts of aroma compound loaded into the mucosa became negligible.

These results revealed which parameters related to mucosa had an impact on release kinetics. However, except in the nasal cavity, mucosa is always wetted by a saliva film. Mucosa parameters were thus fixed to their reference values and saliva parameters were modified to evaluate their influence on aroma release kinetics. As a reference, it was assumed that saliva covered 90% of the mouth and pharynx surfaces.

Simulations were performed by increasing this percentage up to 100% without any significant effect (not shown). When saliva is considered, mucosa concentrations are lower due to: (i) the reduced contact area between air and mucosa and, consequently, lower loading rates of mucosa compartments in the mouth and in the pharynx; and (ii) the fact that part of the aroma
compound was captured by saliva instead of mucosa. Nevertheless, there was no impact on nasal concentration and release kinetics.

100-fold variations in the mass transfer coefficients of molecules in saliva did not affect release kinetics in the nasal cavity (not shown). The only consequences were modifications of aroma concentrations in saliva.

Concerning the modifications of air/saliva partition properties, only a 100-fold decrease significantly modified release kinetics as a consequence of a higher affinity of aroma compounds for saliva. Saliva becomes the main aroma reservoir, supplying aroma compounds to the air in the nasal cavity until the first swallow at 20 s. Aroma concentrations increased in saliva (50-fold and 60-fold factors in the pharynx and the mouth, respectively) and decreased in mucosa (6-fold and 3-fold factors in the pharynx and the mouth, respectively) (not shown).

Simulations were also performed by varying mouth and pharynx volumes and the salivary flow rate (in the range of experimental values, determined on panellists), but no significant difference was obtained between simulations (not shown), confirming the absence of effect of these parameters in the range of variation that was tested.

On the basis of these results, it can be concluded that both saliva and mucosa had an impact on aroma release when the affinity of aroma compounds for these compartments was sufficiently high. Mechanistic modelling provided insight into the relative time distributions of the aroma compound among these compartments.

4.4. Adjustment of simulated release kinetics to experimental data

To validate model assumptions, model simulations were fitted to experimental data. Three experimental kinetics were used, each one representing specific release behaviour (related to specific aroma compounds). Several parameters were used as degrees of freedom: the effective mucosa thickness participating to aroma retention, the mass transfer coefficients of aroma compounds in saliva and in mucosa and the air/mucosa partition coefficients of aroma.
compounds. For these parameters, values were assumed to be the same in all physiological cavities (mouth, pharynx and nose).

Figures 6-a, 6-b and 6-c indicate the good fit that was obtained between experimental (1 panellist, 1 replicate) and simulated release kinetics of ethyl propanoate, (Z)-3-hexen-1-ol and 2-nonanone, respectively.

Concerning fitted parameters, the final value of mucosa thickness was modified (26 µm) but remained in a realistic order of magnitude. In a first approach, it was set at a similar value for all physiological cavities (nose, pharynx and mouth) since sensitivity analysis did not show a significant influence of this parameter on release kinetic shape. For further studies, it could perhaps be interesting to more accurately study the impact of this parameter since the nose, mouth and pharynx mucosa are clearly physiologically different.

The values of the mass transfer coefficients of aroma compounds reached $6.0 \times 10^{-5}$ m/s in saliva and $4.9 \times 10^{-5}$ m/s in mucosa. They were on the order of magnitude of the one used as a reference for simulations and were the same for the three molecules. This result is in agreement with the fact that these coefficients mainly depend on the hydrodynamics in the system and little on molecule properties (Marin et al., 1999). The fact that the mass transfer coefficient in saliva was higher than the one in mucosa was quite expected and can be explained by the difference in viscosity between saliva and mucus. Yet, the absolute values of these mass transfer coefficients in mucosa and saliva were quite high in comparison with the data in the literature ($3 \times 10^{-6}$ m/s) (Marin et al., 1999). The high degree of mixing that probably exists in the naso-oro-pharyngeal cavities can account for this discrepancy (the data in the literature were related to in vitro studies).

The air/mucosa partition properties of aroma compounds ranged between $4.7 \times 10^{-4}$ (2-nonanone) and $6.9 \times 10^{-3}$ (ethyl propanoate), the one of (Z)-3-hexen-1-ol being intermediate. In the case of ethyl propanoate, the air/mucosa partition coefficient increased by a factor of 7
compared to reference values, so that the model correctly predicted the experimental release kinetics. It thus appeared that this molecule has limited interaction with saliva and mucosa. The shape of release kinetics in this case is therefore mainly explained by the pulse of the aroma compound due to sample inhalation and air renewal in the mouth at swallowing (Figure 6-a). Concerning 2-nonanone, the 2-fold decrease in the air/mucosa partition value suggests that 2-nonanone interacts more with mucosa than the other two compounds. For (Z)-3-hexen-1-ol, the value of air/mucosa partition properties after model fitting was only slightly modified in comparison to the reference values and remained thus in the same order of magnitude than the air/saliva partition coefficient.

On the basis of these results, we could argue that both air/mucosa and air/saliva partition properties are, at least partly, at the origin of the release behaviours that were observed for the molecules: the ones with the lower affinity for saliva and/or mucosa (highest partition coefficients) had the least persistent behaviour.

All these results confirmed that *in vivo* release behaviours were strongly molecule-dependent and highlighted the fact that different types of interactions with mucosa and/or saliva were involved, depending on the molecule's properties. These simulations were in agreement with previous experimental data, which notably showed the limited retention of ethyl propanoate and the specific retention of 2-nonanone in the oral and pharyngeal cavities (Déléris et al., 2015). To improve simulations, further development of the model could be foreseen, notably concerning assumptions on mucosa. This will require experimental determination of mucosa properties.

5. Conclusions

In conclusion, it appears that the proposed model adequately simulated aroma release and retention after the inhalation of a gaseous flavoured sample by panellists. The simulation of the time variation of concentrations that cannot be determined experimentally helped to better
understand the involved phenomena. The sensitivity analysis contributed to distinguish the respective roles of saliva and mucosa on aroma retention phenomena. No clear effect of cavity volumes or saliva flow rate on aroma release kinetics was highlighted in the range of variation of parameters that was tested. But results confirmed the particular role of wetted mucosa can play, depending on aroma compound properties. This study constitutes a first step in understanding aroma persistence and further work is needed to clarify the relationships between the properties of molecules and the type of interactions that are involved.
### 6. Appendix A: model equations

<table>
<thead>
<tr>
<th><strong>Volumes</strong></th>
<th><strong>Oral cavity</strong></th>
<th><strong>Nasal cavity</strong></th>
<th><strong>Pharynx</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>wetted mucosa</td>
<td>Eq. (A.12) $V_{OM} = \varepsilon_{OM} \times A_{OM}$</td>
<td>Eq. (A.13) $V_{NM} = \varepsilon_{NM} \times A_{NM}$</td>
<td>Eq. (A.14) $V_{FM} = \varepsilon_{FM} \times A_{FM}$</td>
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<tr>
<td>saliva</td>
<td>Eq. (A.15) $V_{S} = \varepsilon_{S} \times A_{S}$</td>
<td>n. a.</td>
<td>Eq. (A.16) $V_{F} = \varepsilon_{F} \times A_{F}$</td>
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</table>

<table>
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<tr>
<th><strong>Interfacial concentrations</strong></th>
<th><strong>Air/mucosa</strong></th>
<th><strong>Air/saliva</strong></th>
<th><strong>Mucosa/saliva</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Wetted mucosa</strong></td>
<td>Eq. (A.17) $C_{OM}(t) = \frac{\varepsilon_{OM}(t)}{\varepsilon_{NM}}$</td>
<td>n. a.</td>
<td>Eq. (A.18) $C_{NM}(t) = \frac{\varepsilon_{NM}(t)}{\varepsilon_{NM}}$</td>
</tr>
<tr>
<td><strong>Saliva</strong></td>
<td>Eq. (A.20) $C_{OM}(t) = \frac{\varepsilon_{OM}(t)}{\varepsilon_{NM}}$</td>
<td>n. a.</td>
<td>Eq. (A.21) $C_{FM}(t) = \frac{\varepsilon_{FM}(t)}{\varepsilon_{FM}}$</td>
</tr>
<tr>
<td><strong>Mucosa/saliva</strong></td>
<td>Eq. (A.23) $K_{OM} = \frac{\varepsilon_{OM}}{\varepsilon_{NM}}$</td>
<td>n. a.</td>
<td>Eq. (A.24) $C_{FM}(t) = \frac{\varepsilon_{FM}(t)}{\varepsilon_{FM}}$</td>
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<tr>
<th><strong>Volatile mass fluxes</strong></th>
<th><strong>Air/mucosa</strong></th>
<th><strong>Air/saliva</strong></th>
<th><strong>Mucosa/saliva</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Wetted mucosa</strong></td>
<td>Eq. (A.26) $\phi_{OM}(t) = k_{OM} \times A_{OM} \times (C_{OM}(t) - C_{NM}(t))$</td>
<td>n. a.</td>
<td>Eq. (A.27) $\phi_{NM}(t) = k_{NM} \times A_{NM} \times (C_{NM}(t) - C_{FM}(t))$</td>
</tr>
<tr>
<td><strong>Saliva</strong></td>
<td>Eq. (A.29) $\phi_{OM}(t) = k_{OM} \times A_{OM} \times (C_{OM}(t) - C_{NM}(t))$</td>
<td>n. a.</td>
<td>Eq. (A.30) $\phi_{FM}(t) = k_{FM} \times A_{FM} \times (C_{FM}(t) - C_{FM}(t))$</td>
</tr>
<tr>
<td><strong>Mucosa/saliva</strong></td>
<td>Eq. (A.32) $\phi_{FM}(t) = k_{FM} \times A_{FM} \times (C_{FM}(t) - \frac{\varepsilon_{FM}(t)}{\varepsilon_{FM}})$</td>
<td>n. a.</td>
<td>Eq. (A.33) $Q_{FM}(t) = -\pi \times F_{FM} \times V_{F} \times \sin \theta$</td>
</tr>
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<table>
<thead>
<tr>
<th><strong>Air flow rates</strong></th>
<th><strong>Breathing</strong></th>
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<tr>
<td>Eq. (A.34)</td>
<td>$Q_{OM}(t) = Q_{FM}(t)$</td>
<td>Eq. (A.35)</td>
<td>$Q_{NM}(t) = Q_{FM}(t)$</td>
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<tr>
<td>Eq. (A.36)</td>
<td>$Q_{FM}(t) = Q_{FM}(t)$</td>
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<td>Eq. (A.36)</td>
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<tr>
<th><strong>Volatiles mass balances</strong></th>
<th><strong>Air</strong></th>
<th><strong>Saliva</strong></th>
<th><strong>Mucosa</strong></th>
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<tr>
<td>Eq. (A.37)</td>
<td>$\phi_{OM}(t) = \phi_{OM}(t) + \phi_{OM}(t)$</td>
<td>n. a.</td>
<td>Eq. (A.41)</td>
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<tr>
<td>Eq. (A.38)</td>
<td>$\phi_{OM}(t) = \phi_{OM}(t)$</td>
<td>n. a.</td>
<td>Eq. (A.42)</td>
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n. a.: not applicable
7. Acknowledgments

We gratefully acknowledge the panellists for their contribution to in vivo and sensory measurements. We also thank Gilles Feron from UMR 1324 INRA/AgroSup Dijon/CNRS/Université de Bourgogne Centre des Sciences du Goût et de l’Alimentation (CSGA) for the characterisation of saliva samples, and Gail Wagman for revising the English version of the manuscript.

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equilibrium of aroma compounds in polysaccharide gels: Study by quantitative structure-


Table 1: Physicochemical properties of aroma compounds used in the present study.

<table>
<thead>
<tr>
<th>Aroma compounds</th>
<th>Chemical formulae</th>
<th>Chemical structures</th>
<th>Molecular weights (g/mol)</th>
<th>Log P*</th>
<th>PTR-MS² fragmentation: main m/z peaks (relative abundance)</th>
<th>Air/water partition coefficient (25°C)</th>
<th>Experimental air/water partition coefficient $K_{aw}$ ($\times 10^{-3}$) (37°C)</th>
<th>Experimental air/saliva partition coefficient $K_{as}$ ($\times 10^{-3}$) (37°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Z)-3-Hexen-1-ol</td>
<td>$C_6H_{12}O$</td>
<td><img src="image" alt="Chemical structure" /></td>
<td>100.16</td>
<td>1.61</td>
<td>55 (100); 83 (39)</td>
<td>0.63 $10^{-3}$</td>
<td>0.78 ± 0.07</td>
<td>0.76 ± 0.12</td>
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<tr>
<td>Ethyl propanoate</td>
<td>$C_5H_{10}O_2$</td>
<td><img src="image" alt="Chemical structure" /></td>
<td>102.13</td>
<td>1.21</td>
<td>75 (100); 103 (20)</td>
<td>15.9 $10^{-3}$</td>
<td>14.9 ± 1.4</td>
<td>12.9 ± 5.5</td>
</tr>
<tr>
<td>2-Nonanone</td>
<td>$C_9H_{18}O$</td>
<td><img src="image" alt="Chemical structure" /></td>
<td>142.24</td>
<td>3.14</td>
<td>143 (100); 41 (20)</td>
<td>11.1 $10^{-3}$</td>
<td>19.6 ± 8.4</td>
<td>9.7 ± 1.39</td>
</tr>
</tbody>
</table>

1*: estimation with EPI Suite™ programme

2*: PTR-MS: Proton Transfer Reaction-Mass Spectrometry

3*: from (Déléris et al., 2015)
Table 2: Minimal, median and maximal values of the physiological characteristics of panellists and associated quartiles.

<table>
<thead>
<tr>
<th>Physiological parameters</th>
<th>min</th>
<th>Q1</th>
<th>median</th>
<th>Q3</th>
<th>max</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_C$ (L)</td>
<td>0.49</td>
<td>0.54</td>
<td>0.80</td>
<td>0.86</td>
<td>0.93</td>
</tr>
<tr>
<td>$V_{\text{noe}}$ (cm$^3$)</td>
<td>5.8</td>
<td>9.5</td>
<td>11.1</td>
<td>13.1</td>
<td>16.8</td>
</tr>
<tr>
<td>$V_{\text{mouth}}$ (cm$^3$)</td>
<td>33.1</td>
<td>35.5</td>
<td>39.3</td>
<td>56.7</td>
<td>69.8</td>
</tr>
<tr>
<td>$V_{\text{pharynx}}$ (cm$^3$)</td>
<td>16.8</td>
<td>24.3</td>
<td>27.4</td>
<td>30.2</td>
<td>34.8</td>
</tr>
<tr>
<td>Salivary flux (g/min)</td>
<td>0.32</td>
<td>0.44</td>
<td>0.54</td>
<td>0.66</td>
<td>0.87</td>
</tr>
<tr>
<td>Antioxidant (eq mM Trolox)</td>
<td>75.4</td>
<td>90.7</td>
<td>105.7</td>
<td>127.5</td>
<td>137.3</td>
</tr>
<tr>
<td>Lipolysis (mU/mL)</td>
<td>0.04</td>
<td>0.08</td>
<td>0.15</td>
<td>0.19</td>
<td>0.27</td>
</tr>
<tr>
<td>Amylase (U/mL)</td>
<td>67.4</td>
<td>88.7</td>
<td>124.8</td>
<td>148.6</td>
<td>196.0</td>
</tr>
<tr>
<td>Proteolysis (U/mL)</td>
<td>1.6</td>
<td>1.9</td>
<td>4.1</td>
<td>4.8</td>
<td>11.2</td>
</tr>
<tr>
<td>Lysozyme (U/mL)</td>
<td>313.2</td>
<td>332.2</td>
<td>413.9</td>
<td>444.1</td>
<td>462.1</td>
</tr>
<tr>
<td>Proteins (mg/mL)</td>
<td>0.46</td>
<td>0.55</td>
<td>0.81</td>
<td>0.94</td>
<td>1.15</td>
</tr>
<tr>
<td>Symbol</td>
<td>Unit</td>
<td>Definition</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>--------</td>
<td>------</td>
<td>------------</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$C_{Fa}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the air in the pharynx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{Fm}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the wetted mucosa in the pharynx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{Fs}$</td>
<td>g/cm³</td>
<td>Aroma concentration in saliva in the pharynx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C^*_{Fam}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the air/wetted mucosa interface in the pharynx</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$C^*_{Fas}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the air/saliva interface in the pharynx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C^*_{Fms}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the wetted mucosa/saliva interface in the pharynx, on the mucosa side</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C^*_{Fs}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the wetted mucosa/saliva interface in the pharynx, on the saliva side</td>
<td></td>
<td></td>
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<tr>
<td>$C_{Oa}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the air in the oral cavity</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>$C_{Om}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the wetted mucosa in the oral cavity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{Os}$</td>
<td>g/cm³</td>
<td>Aroma concentration in saliva in the oral cavity</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$C^*_{Oam}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the air/wetted mucosa interface in the oral cavity</td>
<td></td>
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</tr>
<tr>
<td>$C^*_{Oas}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the air/saliva interface in the oral cavity</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$C^*_{Oms}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the wetted mucosa/saliva interface in the oral cavity, on the mucosa side</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C^*_{Os}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the wetted mucosa/saliva interface in the oral cavity, on the saliva side</td>
<td></td>
<td></td>
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<tr>
<td>$C_{Na}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the air in the nose</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>$C_{Nm}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the mucosa in the nose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C^*_{Nam}$</td>
<td>g/cm³</td>
<td>Aroma concentration at the air/mucosa interface in the nose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{Ta}$</td>
<td>g/cm³</td>
<td>Aroma concentration in the trachea</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$Q_{Na}$</td>
<td>cm³/s</td>
<td>Air flow rate into the nasal cavity</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$Q_{NFa}$</td>
<td>cm³/s</td>
<td>Air flow rate from the nasal cavity to the pharynx</td>
<td></td>
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<tr>
<td>$Q_{Oa}$</td>
<td>cm³/s</td>
<td>Air flow rate into the oral cavity (1st inhalation)</td>
<td></td>
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<tr>
<td>$Q_{Ofa}$</td>
<td>cm³/s</td>
<td>Air flow rate from the oral cavity to the pharynx</td>
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<tr>
<td>$Q_{Ta}$</td>
<td>cm³/s</td>
<td>Air flow rate from the trachea</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$t$</td>
<td>s</td>
<td>Time</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$t_{swg}$</td>
<td>s</td>
<td>Swallowing moment</td>
<td></td>
<td></td>
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<tr>
<td>$\phi_{am}$</td>
<td>g/s</td>
<td>Volatile mass flux between the air and the wetted mucosa in the pharynx</td>
<td></td>
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<tr>
<td>$\phi_{as}$</td>
<td>g/s</td>
<td>Volatile mass flux between the air and the saliva in the pharynx</td>
<td></td>
<td></td>
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<tr>
<td>$\phi_{Fms}$</td>
<td>g/s</td>
<td>Volatile mass flux between the wetted mucosa and the saliva in the pharynx</td>
<td></td>
<td></td>
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<tr>
<td>$\phi_{Nam}$</td>
<td>g/s</td>
<td>Volatile mass flux between the air and mucosa in the nasal cavity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\phi_{Oam}$</td>
<td>g/s</td>
<td>Volatile mass flux between the air and the wetted mucosa in the oral cavity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\phi_{Oas}$</td>
<td>g/s</td>
<td>Volatile mass flux between the air and the saliva in the oral cavity</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>$\phi_{Oms}$</td>
<td>g/s</td>
<td>Volatile mass flux between the wetted mucosa and the saliva in the oral cavity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Symbol</td>
<td>Unit</td>
<td>Definition</td>
<td>Reference value</td>
<td>Range of variation</td>
<td>Global sensitivity index (from Monte-Carlo analysis)</td>
</tr>
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<td>--------</td>
<td>------</td>
<td>------------</td>
<td>----------------</td>
<td>-------------------</td>
<td>---------------------------------------------------</td>
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<tr>
<td>$A_F$</td>
<td>cm$^2$</td>
<td>Total area of the pharynx</td>
<td>65</td>
<td>32.5 ... 130</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Fam}$</td>
<td>cm$^2$</td>
<td>Air/mucosa contact area in pharynx</td>
<td>$= 0.1 \times A_F$</td>
<td>$0.1 \times A_F - A_F$</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Fam}$</td>
<td>cm$^2$</td>
<td>Air/saliva contact area in pharynx</td>
<td>$= A_F - A_{Fam}$</td>
<td>/</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Nam}$</td>
<td>cm$^2$</td>
<td>Mucosa/saliva contact area in pharynx</td>
<td>$= A_F - A_{Fam}$</td>
<td>/</td>
<td>+</td>
</tr>
<tr>
<td>$A_O$</td>
<td>cm$^2$</td>
<td>Air/mucosa contact area in nose</td>
<td>150</td>
<td>75 ... 300</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Oam}$</td>
<td>cm$^2$</td>
<td>Total area in oral cavity</td>
<td>116</td>
<td>58 ... 232</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Oam}$</td>
<td>cm$^2$</td>
<td>Air/mucosa contact area in oral cavity</td>
<td>$= 0.1 \times A_O$</td>
<td>$0.1 \times A_O - A_O$</td>
<td>+</td>
</tr>
<tr>
<td>$A_{Oam}$</td>
<td>cm$^2$</td>
<td>Mucosa/saliva contact area in oral cavity</td>
<td>$= A_O - A_{Oam}$</td>
<td>/</td>
<td>+</td>
</tr>
<tr>
<td>$C_{ext}$</td>
<td>µg/cm$^3$</td>
<td>Aroma concentration in air</td>
<td>1</td>
<td>0.5 ... 2</td>
<td>+</td>
</tr>
<tr>
<td>$e_{Fm}$</td>
<td>cm</td>
<td>Thickness of wetted mucosa in pharynx</td>
<td>$5 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$ ... $5 \times 10^{-2}$</td>
<td>+</td>
</tr>
<tr>
<td>$e_{Fm}$</td>
<td>cm</td>
<td>Thickness of saliva layer in pharynx</td>
<td>$= V_F/A_{Fam}$</td>
<td>-</td>
<td>/</td>
</tr>
<tr>
<td>$e_{Om}$</td>
<td>cm</td>
<td>Thickness of wetted mucosa in oral cavity</td>
<td>$5 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$ ... $5 \times 10^{-2}$</td>
<td>+</td>
</tr>
<tr>
<td>$e_{Os}$</td>
<td>cm</td>
<td>Thickness of saliva layer in oral cavity</td>
<td>$= V_O/A_{Oam}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$e_{Nm}$</td>
<td>cm</td>
<td>Thickness of mucosa in nasal cavity</td>
<td>$5 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$ ... $5 \times 10^{-2}$</td>
<td>+</td>
</tr>
<tr>
<td>$F_R$</td>
<td>Cycle/s</td>
<td>Respiratory frequency</td>
<td>0.24</td>
<td>0.12 ... 0.48</td>
<td>+++</td>
</tr>
<tr>
<td>$K_{Fam}$</td>
<td></td>
<td>Air/wetted mucosa partition coefficient in pharynx</td>
<td>$10^{-3}$</td>
<td>$10^{-4}$ ... $10^{-1}$</td>
<td>+++</td>
</tr>
<tr>
<td>$K_{Fam}$</td>
<td></td>
<td>Wetted mucosa/saliva partition coefficient in pharynx</td>
<td>$= K_{Fam} / K_{Fam}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$K_{Nam}$</td>
<td></td>
<td>Air/mucosa partition coefficient in nasal cavity</td>
<td>$10^{-3}$</td>
<td>$10^{-4}$ ... $10^{-1}$</td>
<td>+++</td>
</tr>
<tr>
<td>$K_{Oam}$</td>
<td></td>
<td>Air/wetted mucosa partition coefficient in oral cavity</td>
<td>$10^{-5}$</td>
<td>$10^{-6}$ ... $10^{-1}$</td>
<td>++</td>
</tr>
<tr>
<td>$K_{Oam}$</td>
<td></td>
<td>Air/saliva partition coefficient in oral cavity</td>
<td>$5 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$ ... $5 \times 10^{-2}$</td>
<td>++</td>
</tr>
<tr>
<td>$K_{Oam}$</td>
<td></td>
<td>Wetted mucosa/saliva partition coefficient in oral cavity</td>
<td>$= K_{Fam} / K_{Oam}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$k_{Fm}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in air in pharynx</td>
<td>$10^{3}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$k_{Fm}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in wetted mucosa in pharynx</td>
<td>$10^{5}$</td>
<td>$10^{4}$ ... $10^{4}$</td>
<td>+</td>
</tr>
<tr>
<td>$k_{Fs}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in saliva in pharynx</td>
<td>$10^{6}$</td>
<td>$10^{5}$ ... $10^{4}$</td>
<td>+</td>
</tr>
<tr>
<td>$k_{Feq}$</td>
<td>m/s</td>
<td>Equivalent mass transfer coefficient between saliva and wetted mucosa in pharynx</td>
<td>$1/k_{Feq} = 1/k_{Fs} + K_{Fm}/K_{Fm}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$k_{Na}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in air in nasal cavity</td>
<td>$10^{3}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$k_{Na}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in mucosa in nasal cavity</td>
<td>$10^{6}$</td>
<td>$10^{5}$ ... $10^{4}$</td>
<td>+</td>
</tr>
<tr>
<td>$k_{Na}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in air in the oral cavity</td>
<td>$10^{6}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$k_{Nn}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in wetted mucosa in the oral cavity</td>
<td>$10^{6}$</td>
<td>$10^{5}$ ... $10^{4}$</td>
<td>+</td>
</tr>
<tr>
<td>$k_{Dm}$</td>
<td>m/s</td>
<td>Mass transfer coefficient in saliva in oral cavity</td>
<td>$10^{6}$</td>
<td>$10^{5}$ ... $10^{4}$</td>
<td>++</td>
</tr>
<tr>
<td>$k_{Oeq}$</td>
<td>m/s</td>
<td>Equivalent mass transfer coefficient between saliva and wetted mucosa in oral cavity</td>
<td>$1/k_{Oeq} = 1/k_{Oa} + K_{Oma}/k_{Oam}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$Q_{Dm}$</td>
<td>cm$^3$/s</td>
<td>Average rate of saliva flow rate</td>
<td>0.6</td>
<td>0.15 ... 2.4</td>
<td>+</td>
</tr>
<tr>
<td>$t_{deg}$</td>
<td>s</td>
<td>Swallowing moment</td>
<td>20, 50, 80, 110</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$V_c$</td>
<td>cm$^3$</td>
<td>Current breath volume</td>
<td>800</td>
<td>400 ... 1600</td>
<td>++</td>
</tr>
<tr>
<td>$V_{Fm}$</td>
<td>cm$^3$</td>
<td>Volume of air in the pharynx</td>
<td>30</td>
<td>15 ... 60</td>
<td>+</td>
</tr>
<tr>
<td>$V_{Fm}$</td>
<td>cm$^3$</td>
<td>Volume of wetted mucosa in pharynx</td>
<td>$= C_{Fm} \times A_{Fam}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$V_{Fs}$</td>
<td>cm$^3$</td>
<td>Volume of saliva in pharynx</td>
<td>0.2</td>
<td>0.1 ... 0.4</td>
<td>+</td>
</tr>
<tr>
<td>$V_{Na}$</td>
<td>cm$^3$</td>
<td>Volume of air in nasal cavity</td>
<td>20</td>
<td>10 ... 40</td>
<td>+</td>
</tr>
<tr>
<td>$V_{Nm}$</td>
<td>cm$^3$</td>
<td>Volume of mucosa in nasal cavity</td>
<td>$= e_{Nm} \times A_{Nm}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>----------</td>
<td>-------</td>
<td>--------------------------------</td>
<td>--------------------------</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>$V_{Oa}$</td>
<td>cm$^3$</td>
<td>Volume of air in oral cavity</td>
<td>40</td>
<td>20 … 80</td>
<td>+</td>
</tr>
<tr>
<td>$V_{Om}$</td>
<td>cm$^3$</td>
<td>Volume of wetted mucosa in oral cavity</td>
<td>$= e_{Om} \times A_{Om}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$V_{Os}$</td>
<td>cm$^3$</td>
<td>Volume of saliva in oral cavity</td>
<td>$= e_{Os} \times A_{Os}$</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>$V_{Omin}$</td>
<td>cm$^3$</td>
<td>Minimal volume of saliva in oral cavity after swallowing</td>
<td>0.2</td>
<td>0.1 … 0.4</td>
<td>+</td>
</tr>
</tbody>
</table>

$V_{Omin}$

\[ V_{Omin} = 0.2 \text{ cm}^3 \]

\[ V_{Omin} = 0.1 \text{ cm}^3 \text{ to } 0.4 \text{ cm}^3 \]

\[ V_{Omin} = + \]

\[ V_{Omin} = (Doyennette et al., 2014) \]

\[ a \text{ Global sensitivity index: } +++ \text{ means a highly influential parameter (sensitivity index between 0.4 and 0.2); ++ means a moderately influential parameter (sensitivity index between 0.1 and 0.025); + was used for parameters with sensitivity index below 0.025.} \]
Figure captions

Fig. 1: Schematic representation of the experimental set-up for the preparation of gaseous samples with controlled volume.

Fig. 2: Schematic representation of the interconnected compartments of the naso-oro-pharyngeal cavities and the mechanisms involved in aroma release during the inhalation of gaseous samples.

Fig. 3: Schematic representation of the balances between the different compartments. Bold red lines represent concentration profiles and horizontal dotted lines represent the limits of boundary layers where mass transfer resistance was considered.

Fig. 4: Comparison of parameters extracted from release kinetics that significantly differ between ions (means and associated standard deviations). (a) $t_{\text{max}1}$ and $\Delta t_{20\%_1}$, before swallowing; (b) $t_{\text{max}2}$, $\Delta t_{20\%_2}$ and $t_{50\%-t_{\text{max}2}}$, after swallowing; (c) initial release rates Rate$_1$ and Rate$_2$; (d) AUC$_1$/AUC$_2$ ratio and AUC$_{\text{stand}}$. Significant differences were determined using Kruskal-Wallis tests and the Conover-Iman procedure ($p<0.05$) and highlighted with letters a to c.

Fig. 5: Simulated release kinetics using the model with the reference values of the parameters (Table 4): variation over time of aroma compound concentrations in (a) the air phase within the pharynx; (b) the mucosa layer in the pharynx; (c) saliva in the pharynx; (d) the air phase in the mouth; (e) the mucosa layer in the mouth; (f) saliva in the mouth; (g) the air phase in the nose; (h) the mucosa layer in the nose; and (i) the variation over time of saliva volume in the mouth. For each figure (except i), the insets focus on the first 3 seconds.

Fig. 6: Comparison between individual experimental and simulated release kinetics of: (a) ethyl propanoate ($m/z$ 75); (b) $\ (Z)$-3-hexenol ($m/z$ 83); and (c) 2-nonanone ($m/z$ 143) in the nasal cavity. Experimental release kinetics were obtained by PTR-MS measurements during
the inhalation of gaseous sample by one panellist. d) Values of the parameters that were changed for model fitting (with respect to reference values).
Figure 1. Schematic representation of the experimental set-up for the preparation of gaseous samples with controlled volume.
Figure 2. Schematic representation of the interconnected compartments of the naso-oropharyngeal cavities and the mechanisms involved in aroma release during the inhalation of gaseous samples.
Figure 3. Schematic representation of the balances between the different compartments. Bold red lines represent concentration profiles and horizontal dotted lines represent the limits of boundary layers where mass transfer resistance was considered.
Figure 4. Comparison of parameters extracted from release kinetics that significantly differ between ions (means and associated standard deviations). (a) $t_{max1}$ and $\Delta t_{20\%_1}$, before swallowing; (b) $t_{max2}$, $\Delta t_{20\%_2}$ and $t_{50\%}-t_{max2}$, after swallowing; (c) initial release rates $Rate_1$ and $Rate_2$; (d) $AUC_1/AUC_2$ ratio and $AUC_{stand}$. Significant differences were determined using Kruskal-Wallis tests and the Conover-Iman procedure ($p<0.05$) and highlighted with letters a to c.
Figure 5. Simulated release kinetics using the model with the reference values of the parameters (Table 4): variation over time of aroma compound concentrations in (a) the air phase within the pharynx; (b) the mucosa layer in the pharynx; (c) saliva in the pharynx; (d) the air phase in the mouth; (e) the mucosa layer in the mouth; (f) saliva in the mouth; (g) the air phase in the nose; (h) the mucosa layer in the nose; and (i) the variation over time of saliva volume in the mouth. For each figure (except i), the insets focus on the first 3 seconds.
Figure 6. Comparison between individual experimental and simulated release kinetics of: (a) ethyl propanoate ($m/z$ 75); (b) (Z)-3-hexenol ($m/z$ 83); and (c) 2-nonanone ($m/z$ 143) in the nasal cavity. Experimental release kinetics were obtained by PTR-MS measurements during the inhalation of gaseous sample by one panellist using the NMS protocol. d) Values of the parameters that were changed for model fitting (with respect to reference values).