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Static *in vitro* digestion model adapted to the general older adult population: an INFOGEST international consensus

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Understanding the mechanisms of food digestion is of paramount importance to determine the effect foods have on human health. Significant knowledge on the fate of food during digestion has been generated in healthy adults due to the development of physiologically-relevant in vitro digestion models. However, it appears that the performance of the oro-gastrointestinal tract is affected by ageing and that a model simulating the digestive conditions found in a younger adult (<65 years) is not relevant for an older adult (>65 years). The objectives of the present paper were: (1) to conduct an exhaustive literature search to find data on the physiological parameters of the older adult oro-gastrointestinal tract, (2) to define the parameters of an in vitro digestion model adapted to the older adult. International experts have discussed all the parameters during a dedicated workshop organized within the INFOGEST network. Data on food bolus properties collected in the older adult were gathered, including food particle size found in older adult boluses. In the stomach and small intestine, data suggest that significant physiological changes are observed between younger and older adults. In the latter, the rate of gastric emptying is slowed down, the pH of the stomach content is higher, the amount of secretions and thus the hydrolytic activities of gastric and intestinal digestive enzymes are reduced and the concentration of bile salts lower. The consensus in vitro digestion model of the older adult proposed here will allow significant progress to be made in understanding the fate of food in this specific population, facilitating the development of foods adapted to their nutritional needs. Nevertheless, better foundational data when available and further refinement of the parameters will be needed to implement the proposed model in the future.

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1 Introduction

Understanding the fate of food in the oro-gastrointestinal tract has been a topic of growing interest over the last years for the scientific community, and particularly for scientists from the INFOGEST international network on food digestion (https://www.cost-infogest.eu). A quick search on the Web of Science shows that the number of peer-reviewed publications having in any field the words "food" and "digest*" has grown from 2439 in 2009 to 8516 in 2021 (no statistics available for a longer

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period). Unravelling the mechanisms of food disintegration during digestion is needed to determine how food structure and composition affect the kinetics of nutrient release in the gut lumen (bioaccessibility) and the proportion of nutrients that are absorbed (bioavailability). These questions are also shared by the scientific community working on drugs and the COST Action UNGAP (European Network on Understanding Gastrointestinal Absorption-related Processes, https://gbiomed.kuleuven.be/english/research/50000715/50000716/ungap) has been very active in investigating the release of drugs in the orogastrointestinal tract and their subsequent absorption.

In both the food and pharma sectors, the digestive process has been investigated using *in vivo* models involving either human volunteers or animals. However, there is currently a general trend tending to limit as much as possible studies involving complex living organisms. Furthermore, *in vivo* studies are cumbersome, time and resource intensive, ethically questionable and exhibit high inter-individual variability. For all these reasons, *in vitro* digestion models, either static or dynamic, have been the centre of interest of many recent studies.

Static in vitro digestion models consist of a series of bioreactors simulating the physicochemical and enzymatic conditions a food or a drug will meet when entering in the different compartments of the oro-gastrointestinal tract. A first bioreactor mimics the food fragmentation exerted by teeth and mandible, moistening of the food by saliva, initiation of starch hydrolysis and the formation of a food bolus, which is subsequently transferred to a second bioreactor mimicking the stomach, where protein and lipid hydrolysis are initiated, and finally to a third reactor simulating the small intestine. Static models are sequential which means that a phase will only start when the previous one has been fully completed; this is different from the physiology since a proportion of a food can still be in the stomach whereas the other part is already in the small intestine. Furthermore, physicochemical conditions and enzyme activities are kept constant throughout digestion in these models whereas parameters such as the pH or the enzyme activities change over time under the physiological conditions. Static digestion models can be used as a prescreening method, when a large number of tests need to be performed, or before moving to more complex systems. Since they simplify the digestive process, they can also allow unravelling mechanisms that occur at a molecular scale. For instance, phospholipids such as phosphatidylcholine released by the gastric mucosa have been shown to interact with globular proteins like β-lactoglobulin to harden its structure and make it more resistant to the action of pepsin.² Finally, static in vitro digestion models can also be relevant to estimate end-point values such as the glycaemic index, protein and lipid digestibility, estimation of micronutrient and secondary plant metabolite release/bioacessibility, among others. Limits and advantages of static in vitro digestion models have been reviewed by INFOGEST scientists.3

A wide variety of static digestion models have been published in the literature, with different parameters (pH and/or

ionic strength, duration of each phase, enzymatic activities...) making the results difficult to compare between studies. To overcome the problem, namely the impossibility of comparing the results between different studies and the need to harmonize a digestion protocol that the entire scientific community can use, the international network of researchers INFOGEST, whose objective is to bring together a community of scientists in the field of digestion, has established a consensus around a static digestion protocol for a healthy adult.^{4,5} Since then, the model has been extensively used to assess food digestibility, nutrient bioaccessibility, food matrix effects, allergen persistence in the GI tract, etc.6 The model is now used all around the world and is about to be recognized as a reference method by International Organization for Standardization (ISO) and International Dairy Federation (IDF) to determine protein digestibility.

Most of the *in vitro* digestion models developed so far simulate the physiological conditions observed in healthy adults. However, significant changes occur over the life course so *in vitro* digestion models must be adapted to the different physiological stages. Static models mimicking the infant gastrointestinal tract have been proposed; among those a model has been proposed as an international consensus by INFOGEST participants and has been, since then, used in more than a hundred studies published by the scientific community. 13–16

Ageing is accompanied by several physiological changes that affect most of the organs of the human body. For example, due to decline in muscular function, impairment in dental status and reduction in salivary flow and modification in composition, there is impairment in oral processing capability which can alter particle size reduction, adversely affecting digestion rate and extent. 17,18 However, other studies suggested an adaptation of the oral processing in older adults leading to the formation of similar boli than those made by younger adults. 19 Several studies have demonstrated that digestive conditions evolve with age. For example, gastrointestinal motor function, food transit, chemical food digestion, and functionality of the intestinal wall have been previously shown to be affected by ageing.20 This evolution has been considered by different groups who proposed static in vitro digestion models mimicking the oro-gastrointestinal tract of older adults.21-25

The use of different older adult *in vitro* digestion models varying in parameters such as pH, enzyme activities, duration *etc.* ends up with data that are not comparable between different studies. Therefore, the objectives of the present work were (1) to conduct an exhaustive literature review in order to find physiological values obtained on older adults for each parameter of the digestion model, (2) to reach an international consensus on the model and propose it to the scientific community. The literature search has been done within the EAT4AGE European project (https://nofima.com/projects/eat4age/) that gathers 6 academic (INRAE, The Norwegian School of Sport Sciences, Nofima, Technion, University of Leeds, and Teagasc Food Research Centre) and 2 industry

(Nortura and GatFoods) partners on the development of "Palatable, nutritious and digestible foods for prevention of undernutrition in active ageing". EAT4AGE aims to prevent undernutrition and avoid impaired muscle function by investigating how age-related changes, such as decline in digestive functions, oral processing, sensory perception, and appetite, can be overcome. Then, based on the proposition made by the EAT4AGE consortium, an international workshop was organised in Cork on the 2-3 of May 2022 gathering 20 experts from 10 countries and 12 institutes. All the parameters of the model have been discussed one by one and only the values for which solid evidence is available has been considered in order to find a consensus based on the existing literature. In the near future, this novel digestion model adapted for the older adult, will help the scientific community and generate comparable data.

In vitro digestion parameters - recommendation and justification

One of the first points was to specify a minimum age to define the starting point of a healthy older adult population. This is a key issue since, for some of the digestion parameters, values at different ages were available in the literature. In 2014, the World Health Organization (WHO) considered that "old people" were over 60-65 years in the developed world.²⁶ Experts in gerontology categorized "old people" into "young old" (60-69 years), the "middle old" (70-79 years) and the "very old" (80 + years)27 whereas others divided the older adults in 3 categories i.e. "young olds" (65-74 years), "middle olds" (75-84 years) and "oldest olds" (85 + years). 28 It is common sense that rather than the chronological age, it is the "physiological" age that matters in terms of digestion, and that physiological ageing can proceed at different rates depending on nutrition, environmental factors, physical activity, access to healthcare, etc. Therefore, in the literature search that guided the discussion of the consensus group, articles were utilized when: 1 - age was mentioned (words such as ageing, old, older, elderly...) in the article title or description of participants; and 2 - the lower value of the age range in the group of older adults was at least 65. One can still wonder whether it could be relevant to develop different in vitro digestion models for different age or health categories of older adults but not enough data are currently available to achieve this goal within the scope of the current paper.

Based on the available data in the literature, the parameters of the healthy older adult in vitro digestion model will be discussed including:

- (1) Oral phase: simulated salivary fluid (SSF) composition, saliva/food dilution, pH, duration, salivary amylase activity, food bolus particle size,
- (2) Gastric phase: simulated gastric fluid (SGF) composition, food bolus/gastric secretions ratio, pH, duration, pepsin and gastric lipase activities,
- (3) Small intestinal phase: simulated intestinal fluid (SIF) composition, chyme/intestinal secretions ratio, pH, duration,

pancreatic lipase and amylase activities, trypsin and chymotrypsin activities, bile salts content.

Oral phase 2

As an introductory note, for the salivary characteristics, we chose whenever possible to select results obtained on stimulated saliva (as opposed to at-rest saliva), which better simulates the situation where food is manipulated in the mouth. The articles quoted below are mostly on saliva obtained after stimulation by chewing parafilm if not stated otherwise.

SSF composition

The ionic composition of older adult's saliva is very poorly documented and the only data available are for at-rest saliva. In an article reporting two distinct studies, a significant increase in K⁺ (by a factor of 1.45) and Cl⁻ concentration (by a factor of 1.50) was observed in old (70–86 years, n = 22) individuals compared to young (20-29 years, n = 23) while the Na⁺ and Ca2+ concentrations increased, but non-significantly.29 However, in the second study, the concentration of K⁺ and Ca²⁺ significantly increased during ageing by a factor of 1.35 and 1.26 respectively, while the Cl⁻ concentration increased in older adults but non-significantly and the Na+ was similar between young (18–24 years, n = 11) and old (60–90 years, n = 11) 18) individuals.²⁹ In contrast, a 27% decrease of calcium concentration was reported between young (20–30 years, n = 20) and old (60–80 years, n = 20) subjects.³⁰

Recommendation: all the values found (though limited in literature) are within close limits of adult SSF composition, so it is recommended to use the SSF composition proposed for the young adult INFOGEST model.5

pН

A cross-sectional study was carried out in 139 adults with a mean age of 79.1 \pm 9.8 years.³¹ A slight increase in pH of saliva was observed when the age increased (p = 0.087) with values of 7.76 ± 0.91 for 60-74 years, 7.86 ± 0.67 for 75-84 years and 8.04 ± 0.89 for people over 85 years. In another study, ³² forty older adult individuals aged 60-86 were divided into two gender-matched groups of 20, according to the use or non-use (control) of medication and the presence or absence (control) of senile dementia. pH values found in both groups were 6.71 ± 0.55 for the medicated group suffering from dementia (mean age 69.6 years) and 6.95 \pm 0.42 for the control group (mean age 68.3 years). In a Swedish study involving 70 years-old 58 men and 53 women, the pH of parafilm-stimulated saliva was found to be 7.2 and 7.1 in males and females respectively.³³ Finally, comparing healthy young (20-35 years) and older (>65 years) adults, no significant difference was reported with values in atrest saliva of 6.58 \pm 0.47 vs. 6.74 \pm 0.40, respectively.³⁴ Based on these four studies, the pH of saliva of older adult is close to neutral.

Recommendation: for all these reasons, it was decided to use a pH of 7.0 for the oral phase in the consensus model of

the old adult identically to the one proposed for the consensus in vitro digestion model of the young adult.

Food/saliva ratio

To our knowledge, only two articles report values of the proportion of saliva incorporated into food during bolus formation in older adults. This concerns two versions (control or enriched in proteins) of two cereal products, brioche and sponge cake^{35,36} tested by 20 subjects with a mean age of 72 years, and four versions of whey-based cheese³⁷ tested by 72 subjects with a mean age of 73.1 years. These results were acquired within the French project ALIMASSENS, where additional insalivation rates i.e. the quantity of saliva incorporated in the food bolus were obtained for cheese, meat products and custard. In addition, in the project REMUS, aiming at designing a dairy product adapted to older adults, 38 insalivation rates were recorded for two versions of custard-type dairy desserts. Table 1 provides a summary of values recorded in vivo on older adults for these different products, where it appears that less saliva is incorporated into products with lower dry matter (e.g. custard and meat). Percentage of saliva was calculated as follows:

(bolus weight in g – food weight in g)/bolus weight in g \times 100.

For cheese, the percentage of saliva incorporated (from 45 to 90%) was higher than in other studies on younger adults, with values of 6 to 19%, 39 23 to 52%, 40 23 to 46% (ref. 41) or 38 to 50%. 42 However, the products used in these different studies were model cheeses with various properties, which makes the comparison of results difficult. For instance, an increase in cheese fat content (from 25% to 50%) induced a decrease (from 41% to 22%) in percentage of saliva incorporated in a middle aged population.40

In the static consensus model of adult digestion,⁵ a ratio 1:1 (weight of SSF or saliva: weight of food) is used whereas in the semi-dynamic consensus model of digestion, 43 a ratio of 1:1 (weight of SSF or saliva added: dry weight of food) was proposed. The results in Table 1 support that this proxy seems equally relevant to the older adult population.

Recommendation: for studies using the static in vitro model of the older adult and when the objective is to obtain data comparable with those obtained with the consensus adult static model, it is recommended to keep the ratio 1:1 (weight of SSF or saliva: weight of food). Nevertheless, the literature review performed for preparing the present article suggests that the ratio of 1:1 (weight of SSF or saliva added: dry weight of food) is more physiologically relevant, and could be considered in future updated versions of the model of adult digestion.

Salivary amylase

Salivary amylase plays a key role in disintegration of starchcontaining foods. It was evidenced in a recent work on bread bolus obtained after deficient mastication and especially in the absence of saliva. Bigger/compacted particles with reduced total and slowly digestible starch were evidenced as demonstrated with FTIR spectroscopy analysis. 18 The enzyme starts hydrolysing starch in the mouth but also in the stomach as long as the pH remains higher than the inactivation pH between 3 and 3.5.44 After food ingestion, the gastric pH is close to that of the ingested food and will decrease slowly due to the gastric emptying and the acidic secretion. The decrease in pH will depend on the amount of food, the food buffering capacity, and the subject's physiology, but the pH conditions can remain favourable to the action of amylase for a long time. Indeed, in a recent work where industrial vs. traditional baguette were submitted to dynamic in vitro digestion, the proportion of partially hydrolysed starch after the oral phase (at t = 2 min) was similar for all foods (about 20%) but continued to increase very rapidly during gastric digestion, reaching a plateau after about 20 min of digestion for all breads. The plateau values were very high for all breads, between 63% and

Table 1 Insalivation rate determined in vivo on elderly people for different food matrices

Mean age (year)	n	Product	Dry matter (%)	Percentage of saliva incorporated $(mean \pm SD)$	Source of data
72	20	Sponge cake	72	79 ± 25	Alimassens (Assad-Bustillos
72	20	Brioche	70	45 ± 11	et al. 2019) ³⁵
73.1	72	Hard cheese	50	56 ± 29	Alimassens (Lorieau et al.,
73.1	72	Soft cheese	50	45 ± 25	$(2021)^{37}$
73.1	72	Whipped cheese	50	90 ± 35	ŕ
73.1	72	Processed cheese	50	45 ± 30	
73.2	73	Hard cheese	50	69 ± 6	Alimassens (unpublished)
73.2	73	Soft cheese	50	65 ± 5	` -
73.2	73	Whipped cheese	50	71 ± 6	
73.2	73	Processed cheese	50	65 ± 6	
74.0	73	Shredded beef	25	36 ± 10	
74.0	73	Laminated beef	30	30 ± 12	
74.0	73	Minced chicken	30	32 ± 10	
74.0	73	Shredded chicken	25	39 ± 10	
73	76	Custard-type dairy dessert	28	32 ± 23	
70.5	31	Commercial custard enriched in proteins	33	26 ± 13	REMUS (unpublished)
70.5	31	Custard enriched in proteins (reformulated)	28	28 ± 17	,

74%, hence confirming the key role of salivary α-amylase's action in the digestion of starch during the gastric phase. 45

Comparing studies on salivary amylase activity is not straightforward since salivary amylase activity can be determined with different assays. For instance, amylase activity can be assessed by quantifying the reduction of 3,5-dinitrosalicylic acid (DNS). In this method, starch is converted into maltose by α-amylase. Maltose released from starch is measured by the reduction of 3,5-dinitrosalicylic acid. Maltose reduces the pale vellow coloured alkaline DNS to the orange-red colour. The intensity of the colour is proportional to the concentration of maltose present in the sample. Alternatively, amylase activity can be monitored by the CNPG3 Kit. In this test, α -amylase hydrolyses the 2-chloro-4-nitrophenyl maltose trioside leading to the formation of chloro-nitro-phenol that can be measured at 405 nm. Finally, the Phadebas® test is also frequently used by the scientific community. The principle behind the test is that Phadebas®, consisting of starch microspheres with a blue dve cross-linked to the starch, are immobilised on filter paper sheets. In the presence of amylase, the starch is digested, releasing the water-soluble dye, which diffuses through the pores of the filter paper. The resulting blue colour is visually observed on the non-reagent side of the Phadebas® paper.

The INFOGEST young adult model recommends the DNSbased method as a reference to measure amylase activity. In a study involving 169 older adults with a mean age of 81.2 years, salivary amylase activity was measured with the Phadebas test at 212.7 ± 168.1 U ml⁻¹.46 A few years later, the same group analysed the saliva of 175 hospitalized patients (age 82 \pm 5.7 years) and 252 outpatients (age 77 ± 5.7 years). 47 Mean values were in the range of 202-216 U ml⁻¹ for hospitalized patients and 111-130 U ml⁻¹ for outpatients. However, using the CNPG3 kit, amylase activity in stimulated saliva of the ALIMASSENS participants (66-89 years) was 15.3 ± 11.5 U ml⁻¹. This illustrates the difficulty of comparing results when they are acquired using different methods. One article measured amylase activity in acid-stimulated saliva of younger (21-49 years, n = 13) and older (64-99 years, n = 10) adultsusing the DNS test, but the assay temperature and the units for expression of results were different from what is advised in the INFOGEST young adult model (ESI of ref. 4). Nevertheless, no significant difference was observed between the younger $(583 \pm 306 \text{ IU} \times 10^{-3} \text{ ml}^{-1} \text{ saliva})$ and the older group $(629 \pm 10^{-3} \text{ ml}^{-1} \text{ saliva})$ $314 \text{ IU} \times 10^{-3} \text{ ml}^{-1}$ of saliva). In at-rest saliva, there was no difference between young (27.8 \pm 2.6 years, n = 20) and older (68.6 \pm 7.4 years, n = 20) adults³⁰ while an approximate twofold increase in older (n = 40 in total) compared to younger (n= 34 in total) adults was measured in the two studies reported in ref. 29.

Recommendation: data are scarce, there is limited scientific evidence suggesting that a modification of the INFOGEST young adult model is needed to mimic the healthy older adult conditions. Therefore, it is recommended to use a value of 75 U mL⁻¹ (using the DNS assay) for salivary amylase *i.e.* the one that is also proposed for the adult model.

Particle size

Ageing is often accompanied by oral deficiencies such as loss of teeth, 49 decrease in salivary secretion, 50 or decreased masticatory muscle' strength.⁵¹ Oral decline and mastication deficiencies cause alteration of food bolus properties and therefore impact on swallowing.

It has been found that individuals differing in age, gender, and ethnicity vary in oral processing time to produce bolus with textural properties optimized to their needs.⁵² For example, older adults (70 \pm 4.3 years, n = 22) produced sausage boli that were softer, more adhesive, less cohesive, and contained more particles than in young adults (22 \pm 2.8 years, n =21). However, ageing did not affect bolus particle size at the swallowing point for this product. In a study comparing 14 young (35.6 \pm 10.6 years) to 14 aged dentate individuals (68.1 \pm 7.0 year) masticating peanuts and raw carrots, the aged individuals produced boli with similar particle size distribution for peanuts, but the distribution was skewed towards bigger particles for carrots.53

The dental status of the subjects is a critical factor in studying the ability of older adults to fragment the food before swallowing. Indeed, the replacement of the natural teeth by removable full dentures impaired mastication of peanuts and carrots, and food boli containing much coarser particles were observed in the aged complete denture wearers (68.1 ± 7.2 years, n = 14) despite an increased number of chewing cycles and greater electromyographic activity.⁵³ However, in a study investigating the comminution progress in 22 subjects wearing removable denture prosthesis (75.1 ± 5.3 years) and 20 young fully dentate subjects (27.6 ± 1.9 years) consuming a combined test meal (cooked rice, sausage, omelet, raw cabbage, and cucumber), a significant difference in particle size between the two groups was observed at the half-mastication point (mean \pm SD = 1.656 \pm 0.098 mm for old adults vs. 1.493 \pm 0.099 mm for young adults, p < 0.05), but not immediately before swallowing.⁵⁴

Overall, these studies suggest that older adults tend to adapt their oral processing, in particular by increasing the number of chewing cycles, to form a food bolus containing particles similar in size to that of young adults. Median particle sizes (D50) and particle size range reported in the literature for different foods after mastication by older adults are summarized in Table 2.

For the oral phase of this static in vitro digestion model, what is needed is a robust protocol to grind the food into particles of similar size to those reported in the literature that have been previously recorded in in vivo bolus collection studies. The protocol must be simple and reproducible. After testing several devices, it was decided to use a manual meat mincer to produce food particles; this kind of device was selected since it is easy-to-use, cheap and identical systems are available everywhere in the world. The consortium tested a simple protocol on four food systems: raw carrots, cooked meatballs, roasted peanuts, and sponge cake. Carrots (90% moisture) and meatballs (>50%) were selected as high moist-

Table 2 Particle size reported in the literature for different food boli after oral processing by older adults

Mean age (year)	n	Product	d 50 (mm)	Particle size range (<i>d</i> 50 mm)	Source of data
72	20	Sponge cake	0.3 ± 0.1		Alimassens (Assad-Bustillos <i>et al.</i> 2019) ³⁵
72	20	Brioche	2.9 ± 4.0		,
75	20	Fortified sponge cake	0.3 ± 0.1	0.14-0.92	Alimassens (Assad-Bustillos et al., 2020) ³⁷
75	20	Fortified brioche	0.8 ± 0.6	0.17-30.8	, ,
74.0	73	Minced beef	1.20 (median)		Alimassens (unpublished)
74.0	73	Laminated beef	3.68 (median)		, ,
74.0	73	Minced chicken	2.36 (median)		
74.0	73	Chopped chicken	3.59 (median)		
72	107	Carrots	1.68 (median)		
68.1	14	Carrots	, ,	1-4	Mishellany-Dutour et al. (2008) ⁵³
68.1	14	Peanuts		0.4-4	. ,
70	22	Hotdog sausages	1.95 ± 0.02		(Aguayo-Mendoza, Martinez-Almaguer, Piqueras-Fiszman, & Stieger, 2020) ⁵²
75.1	22	Mixed foods: cooked rice, sausage, Japanese hard omelet, raw cabbage, raw cucumber	5.5 ± 0.8		(Sugimoto, Tanaka, Kodama, & Minagi, 2020)

d 50 (mm) corresponds to the median particle diameter (portion of particles with diameters smaller and larger than this value are 50%).

ure products, whereas peanuts (~9%) and sponge cake (~30%) were chosen as low moisture products. All these products were studied because bolus granulometry data were available. To simulate the oral processing, food boli were prepared by mixing the various samples with distilled water at a final insalivation ratio of 95%, 70%, 40%, and 10%, for peanuts, sponge cake, meatballs, and carrots, respectively to be consistent with the 1:1 ratio given above. Samples were then minced using a meat mincer (Kitchen Craft No. 5 meat mincer, Leeds) with a 5 cm mincing disc and a 0.5 cm mesh size for one pass. Once the bolus was recovered, 2 g of carrots, sponge cake, meatballs, and peanuts were suspended in 150 mL glycerol and agitated with a magnetic stirrer for 1 h at 200 rpm to allow particle dispersion without damaging the bolus structure. After this time, the bolus particles were imaged using a ChemiDocTM XRS +

System with image LabTM Software (Bio Rad Laboratories, Richmond, CA, USA). Images were acquired in greyscale as it offers more contrast between the particles and the background. For each bolus, a minimum of three images per bolus sample were taken to obtain approximately 100 individual particle images from each sample. ImageJ software (version 1.48r, National Institute of Health, Bethesda, USA) was used to determine the area of the different particles. Particles were considered circular to calculate their corresponding *D*50. Results are presented in Fig. 1. Median particle sizes obtained for the different foods with this simple procedure were fairly similar to the particle sizes reported in the literature (Table 2).

Recommendation: to simulate the oral processing occurring during the first phase of the digestion it is recommended to use of a basic meat mincer (manual, consisting of a mincing

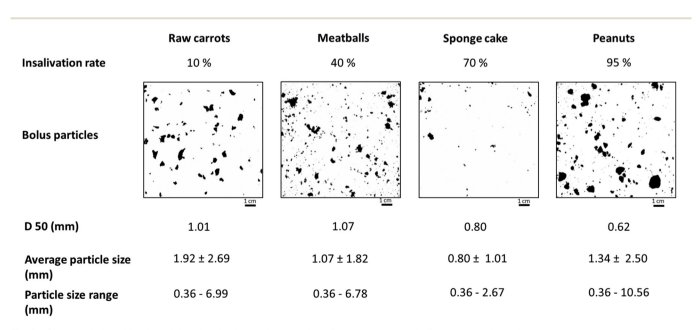


Fig. 1 Characteristics of food particles after in vitro oral processing of raw carrots, meatballs, sponge cake and peanuts.

disk and a blade, only one pass) to produce food particles and then add SSF at pH 7 (with salivary amylase in case of starchcontaining foods) or saliva at a ratio of 1:1 (SSF or saliva added: weight of food or dry weight of food).

Duration

The time of residence of food in the mouth before swallowing highly depends on the rheological properties of the food and in particular the time needed for the teeth to reduce bites into smaller particles. For cheese products, for instance, the average in-mouth resident time before swallowing ranged from 14 s to 28 s depending on the cheese firmness. 19 In case of older adults and for the same type of product, the time of residence observed was slightly longer and ranged from 18 s to 28 s.37 For cereal products, the chewing duration was similar. Older adults are known to adapt their masticatory behaviour by increasing the number of cycles and the chewing duration⁵⁵ and differences in chewing behaviour between healthy fully dentate young (23.7 \pm 1.1 year, n = 10) and older adults (74.1 \pm 1.7 year, n = 10) have been found by electromyographic recordings. 56 In this paper the chewing duration of carrots was evaluated from 10 s to 33 s depending on the cooking procedure (raw or cooked) and it was slightly but significantly higher for older compared to young adults (10-25 s for young vs. 13-35 s for older adults).

In the INFOGEST young adult adult model, the duration of the oral phase has been set to 2 min. This does not reflect the time of residence of the food in the mouth but it is a time long enough to (1) initiate starch hydrolysis as it occurs in vivo and (2) be reproducible when taking samples. Indeed, it has been shown that starch hydrolysis by salivary amylase that starts in the mouth, can continue in the stomach until the gastric pH reaches low values (pH 3-3.5).44 Between 30 to 80% of the starch can be released in the stomach in white bread and pasta, respectively.⁵⁷ Since the INFOGEST in vitro digestion model is static and the pH is set at 3.0, starch will not be hydrolyzed in the stomach. Setting the duration of the oral phase to 2 min at 37 °C allows hydrolysing a significant proportion of starch like it is done in the stomach in vivo.

Recommendation: since the time of residence in the mouth of foods is not very different between young and older adults except for those equipped with a denture, it is recommended to use the same duration of the oral phase for both populations, i.e. 2 min.

Gastric phase

SGF composition

No information regarding possible differences in the gastric fluid composition between old and young adults has been found in the literature despite an exhaustive review. Therefore, the literature was extended to animal models that are traditionally used to mimic what happens in humans i.e. the rat and the pig. Only two references were found on the evolution of gastric secretion output in rats but none of them reported

any information on the possible evolution of the composition of gastric fluid that we could use for the present paper.

Recommendation: it is recommended to use the same SGF described in the young adult INFOGEST model.⁵

pН

In humans, the fasted gastric pH ranges between 1 and 2. After food ingestion, pH increases up to 5-7 depending on the type of food ingested and its buffering capacity. Gastric pH then decreases over time due to emptying of the buffering material and addition of acidic gastric secretions. Gastric pH was recorded in 79 healthy, older men and women (71 \pm 5 years) under both fasted and fed conditions using the Heidelberg capsule technique.⁵⁸ The pH was recorded for 1 h in the fasted state, then a standard liquid and solid meal of 1000 kcal was given to the subjects over 30 min and the pH was finally measured for 4 h postprandially. The measured median fasted gastric pH was 1.3 (1.1-1.6). Following the meal, gastric pH decreased from a peak pH of 6.2 (5.8-6.7) to pH 2.0 within 4 h in most subjects with a gastric emptying half-time ($T_{1/2}$, where 50% of the bolus has been transferred to the small intestine) of 86 min⁵⁸ The observed rate of return was considerably slower than in young healthy subjects. A significant increase in gastric pH with age has also been observed.⁵⁹ They measured the gastric pH in 1615 volunteers classified into four categories of age: 50-59 years (n = 769), 60-69 years (n = 643), 70-79 years (n = 188) and >80 (n = 15). They reported a mean gastric pH of 3.5 \pm 2.3, 3.7 \pm 2.4, 4.4 \pm 2.6 and 4.4 \pm 2.1, respectively.

Recommendation: a strategy to set the gastric pH of the static protocol is to consider the pH at gastric emptying halftime.⁵ Based on these data, the gastric pH should be set at 3.7, which is higher than the values reported in the literature for younger adults (i.e., pH 3.0).

Bolus/secretions ratio

No specific data were found in the literature about the dilution factor of the bolus by gastric secretions in older adults. So, we used the following indirect approach to estimate this parameter. The pH in the stomach highly depends on the amount of acidic secretions (and buffering capacity of the meal). Based on the gastric pH curve reported by Russell et al. (1993),⁵⁸ we estimated the amount of secretions needed to reach a pH of 3.7 at $T_{1/2}$ using the STORM software that monitors the DiDGi® system. 60 At 86 min, the ratio meal/secretions was calculated and a value of 47/53 was obtained.

Recommendation: for the older adult static in vitro digestion model, a 50/50 dilution of the oral bolus in gastric secretions should be used (i.e. the same dilution used in the young adult model).

Duration

Although the impact of ageing on gastric emptying is still controversial, several studies have shown that gastric emptying slows down with age and possibly motor changes in gastric function may include a delay in gastric emptying of liquids

and solids in the older adult. However, these changes are mild.61 Gastric emptying time was assessed on young and older (average 75 years, n = 18) men using ultrasound.⁶² The two groups of volunteers received a 790 kcal test meal consisting of pasta (80 g), beef (100 g), salad (100 g), olive oil (20 g), bread (80 g) and mineral water (200 ml) representing 18% proteins, 52% carbohydrates, 30% fats and 3.42 g of vegetable fibre. Together with the meal, patients swallowed 20 pieces (2 mm × 5 mm in size) of radiopaque markers to determine the transit time. The final gastric emptying time in older subjects was 335 \pm 31 min vs. 245 \pm 25 min in young subjects, corresponding to a 36% increase of the gastric emptying time with age. In another study using ultrasound, the gastric emptying of a whole meal by young (32 \pm 8 years, n = 9) and old adults (77 \pm 3 years, n = 10) was measured. ⁶³ The older participants showed a longer gastric emptying time compared to the younger participants (448.6 \pm 104 vs. 306.6 \pm 57 min, p < 0.002), representing an increase of +46%. Finally, 19 young (23-50 years) and 14 older (70-84 years) volunteers underwent measurements of gastric emptying by scintigraphy after consumption of solid and liquid model meals.⁶⁴ Data showed an increase of +43% of the $T_{1/2}$ for the solid meal and a +34% for the liquid one, when older subjects were compared to the younger group. Based on these data, it appears that the duration of the gastric phase increases by 34-46% with ageing.

Recommendation: to make the protocol simpler, the duration of the gastric phase should be increased by 50% to 3 h in the older adult model.

Gastric enzymes

Studies on enzymatic activity in the ageing stomach are scarce and there is a marked lack of knowledge about pepsin and gastric lipase activities in the postprandial state in older adult populations. Nevertheless, the atrophy of gastric mucosa with age results in a gradual loss of secretory cells (chief cells for pepsin and gastric lipase; parietal cells for gastric acid secretion) that results in the reduced secretion of both enzymes and gastric acid.65,66

Pepsin. In a study involving 206 healthy volunteers (18-98 years), the basal pepsin output and pentagastrin-stimulated pepsin output was reduced by 40% in volunteers over 65 years old (n = 22).65

Recommendation: based on these results, the consortium proposes to set the pepsin activity at 1200 U ml⁻¹ of gastric content in the older adult model (i.e., 60% of the recommended value in the young adult model of Brodkorb et al.⁵).

Gastric lipase. In a study on human gastric mucosal biopsies collected at different locations in the stomach in 28 volunteers, the lipase activity monitored in 22 participants was shown to decline with age, starting from 50 years and reaching down to 80% reduction over 60 years.⁶⁷ However, the number of subjects over 70 years (n = 5) was too low to calculate a precise

Recommendation: for this reason, it is recommended to reduce the gastric lipase activity by 40% in the older adult model compared to the young adult digestion model from

Brodkorb et al., 5 identically to the recommendation for pepsin, i.e. 36 U of lipase per ml of gastric content.

Intestinal phase 4

SIF composition

Electrolyte composition of intestinal fluids in older adults has not been reported precisely so far. However, in a study of the pancreatic exocrine secretions of 180 patients aged from 16 to 83 years, it was shown that calcium concentration was lowest around 41 years and then increased over time, 68 following the equation:

$$[Ca^{2+}] = 0.01 \times age + 0.35$$

Recommendation: if we consider a 65 years old person, the calcium concentration in the SIF should be set at 1 mM rather than 0.6 mM in the young adult INFOGEST model.

Chyme/secretions ratio

This parameter is extremely difficult to assess. Nevertheless, one thing to consider is the decrease of pancreatic secretions with age that, if demonstrated, could eventually limit the dilution of the gastric chyme. Here again, data from the literature are controversial. Fikry (1968) investigated pancreatic exocrine secretions in 23 healthy males aged from 60 to 72 years using the intravenous secretin test considered by investigators in the field of pancreatic diseases as the gold standard.⁶⁹ The data collected showed a two-third reduction in the volume of pancreatic secretions in older compared to young adults. 69 The mean volume of secretions produced over 80 min by the older adults was 55.5 mL (30-81.5 mL) whereas it was around 193 mL (123-310 mL) for the younger group (the mean age of the control population was not provided in this study). This reduced volume of secretions might be attributed to a sole or combined action of three factors: (1) the ageing process itself, (2) the frequency of the chronic fibrosing pancreatitis in the aged population, favoured by increased incidence of gallstone formation, and (3) impairment of the vascular supply to the pancreas. A reduction in pancreatic secretions with age was also observed in another study that reported a linear decrease in secretory volume after 60 years, 70 following this equation:

Pancreatic secretion volume (mL) = $-6.5 \times age + 620.9$

In contrast, other robust studies using similar methodologies reported no differences in the volume of pancreatic secretions with age. Dreiling et al. (1985) found no significant changes in the volume of pancreatic secretions after 50 years on a large group of 1615 subjects. 59 Similarly, in another study involving three groups of volunteers with a mean age of 40 years (n = 30), 64 years (n = 15) and 74 years (n = 10), no significant differences in the volume of pancreatic secretions were observed between the groups.⁷¹

Recommendation: based on these controversial data, it is difficult to determine whether pancreatic secretions tend to decrease during ageing or not. Since the study by Dreiling

et al. (1985) involved the highest number of volunteers (i.e., 1615), including 1034 volunteers over 60 years, the recommendation to keep the gastric chyme/secretions ratio as proposed in the young adult INFOGEST model is based on these results i.e. the 1:1 (v/v) dilution of the gastric content with SIF.

While the duration of the gastric phase can be assessed by recording gastric emptying, the duration of the intestinal phase is not often monitored. An indication of the duration of the intestinal phase of digestion can be obtained by looking at the oro-cecal transit time that corresponds to the sum of the gastric, small and large intestinal phases or at the whole gut transit time. In a recent study, 111 healthy volunteers (21-88 years) had a 602 kcal breakfast consisting in oats/cornflakes, 1 tablespoon raisins/2 teaspoons sugar, skimmed milk, 1 slice wholegrain bread with plant-based margarine and 1 portion jam or ham.⁷² Immediately after having the breakfast, volunteers ingested a 3D-Transit system (Motilis Medica SA, Lausanne, Switzerland) consisting of ingestible electromagnetic capsules which when activated and swallowed emitted an electromagnetic tracking signal that was detected by an external detector plate positioned over the abdomen. The progress of the capsules in the gastrointestinal tract allowed measurement of gastric emptying and small intestinal, colonic and whole gut transit times. Results showed an increase of the gastric emptying time (as already discussed in the Gastric Phase section) and colonic transit time leading to an overall increase of the whole gut transit time (p < 0.01) with age. However, no significant change in small intestinal transit time was observed with age.

Clarkston et al. (1997) measured (1) gastric emptying (by scintigraphy), (2) orocecal transit (through breath hydrogen) and (3) total gut transit (with radiopaque markers) in 19 younger (23-50 years) and 14 older (70-84 years) volunteers.⁶⁴ Gastric emptying $(T_{1/2})$ for solid (182 ± 26 ν s. 127 ± 13 min, p <0.05) and liquid (47 \pm 4 vs. 35 \pm 3 min, p < 0.05) meal components was slower in the older adults. However, there were no significant differences in the orocecal and total gut transit times between the two groups. Ageing seems to be associated with slowing of solid and liquid gastric content emptying (see paragraph on gastric phase duration) but no change in orocecal and total gut transit was observed.⁶⁴

Finally, another study investigating lactose malabsorption did not find any statistical difference in orocecal transit time between three groups of subjects, aged <65 years (45 ± 15 years, n = 33), 65–74 years (69 ± 3 years, n = 17) and >74 years $(81 \pm 4 \text{ years}, n = 34).^{73}$

Recommendation: based on the data, the duration of the intestinal phase of the older adult in vitro digestion model should be set at 2 h, which is the same as in the INFOGEST younger adult model.

Pancreatic enzymes

Pancreatic lipase. Results reported in the literature about pancreatic lipase activity are highly controversial. Two studies

did not find significant differences in pancreatic lipase activity between young and old adults. Fikry (1968) found similar intestinal lipase activities in both age groups (0.8 to 1.4 U ml⁻¹ for young adults vs. 0.1 to 2.4 U ml⁻¹ for older adults).⁶⁹ Similarly, no significant differences were found for intestinal lipase activity between 3 groups of volunteers with a mean age of 40 years (n = 30), 64 years (n = 15) and 74 years (n = 10) with values of 248 \pm 65, 228 \pm 61, and 233 \pm 51 U \times 10³ per 30 min, respectively.⁷¹ Units for expressing pancreatic lipase activity were different between the 2 studies making a comparison impossible.

On the contrary, two other studies found a significant decrease in pancreatic lipase activity with age. In a study involving 180 volunteers (102 males, 78 females) aged 16-83 years, Laugier et al. found that the decrease in lipase activity as a function of age was following the equation:⁶⁸

$$lipase = -8.4 \times age + 1603$$

By dividing the volunteers into two groups (younger and older than 65 years), they reported values of 1256 IU mL⁻¹ and 994 IU mL⁻¹ of pancreatic juice respectively, indicating a 21% decrease in intestinal lipase activity with age. These activities were measured with olive oil as substrate according to the US and European Pharmacopeia assay for pancreatic lipase. INFOGEST recommends another assay with tributyrin as substrate for pancreatic lipase. 5,74 Nevertheless, a conversion factor of 2.8 allows converting USP lipase units into tributyrin units was recently proposed.⁷⁵ The values reported by Laugier et al. for the two groups correspond to 3517 and 2783 U mL⁻¹ of pancreatic juice, and these activity values are in the same range as the activity (4000 U mL-1) recently reported for human pancreatic juice.75 They can be compared with the 2000 U mL⁻¹ currently recommended by INFOGEST for pancreatic lipase in the intestinal phase for healthy adults, i.e. a dilution of pancreatic juice by around a factor 2. Vellas et al. (1988) also observed a decrease in intestinal lipase activity with age. 76 Twenty-seven subjects (36 ± 7.8 years) and 28 subjects (72 \pm 3.2 years), with no clinical or radiological evidence of digestive disease, were selected. Duodenal aspirates (over a 60 min period) were obtained during continuous infusion of secretin (0.5 U kg⁻¹ h⁻¹) and caerulein (75 ng kg⁻¹ h⁻¹). Both lipase output and concentration, measured with olive oil as substrate (Pharmacopeia assay), were significantly reduced in the older adult group by 15.6% and 43.3%, respectively.

Trypsin. Two studies showed opposite results for the effect of ageing on trypsin activity. Fikry (1968) found a 32% decrease of trypsin activity with age with values (calculated as dilution) of 102 (25-200) for older adults against 150 (100-200) for a control group called "normal adults". 69 In contrast, Gullo et al. (1983), found no significant differences in trypsin output between the three groups studied (mean age of 40, 64 and 74 years).71

Chymotrypsin. Three studies assessing chymotrypsin activity in intestinal effluents were found in the literature. Gullo et al. (1983) (see description above) found no statistical difference

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between the chymotrypsin output of three groups of people with increasing mean age (40, 64 and 74 years).⁷¹ In contrast, an 8% decrease in chymotrypsin output was observed by Laugier et al. (1991) in the older group (65-80 years) compared to the control group (20-65 years), 68 whereas Vellas et al. (1988) (see description above) found a 23% decrease in chymotrypsin output with ageing.76

Pancreatic amylase. More data are available in the literature about the effect of ageing on pancreatic amylase secretion and activity than about the other pancreatic enzymes. In these studies, a significant trend showing a decrease in pancreatic amylase output and activity with age is reported. Fikry (1968) observed a 30% decrease in pancreatic amylase activity between old and young adults (554 U vs. 823 U), 69 while Vellas et al. (1988) observed a 13.4% decrease of amylase concentration and a 48% decrease in pancreatic amylase output between both groups.⁷⁶ Similarly, Ishibashi et al. (1991) also found a decrease in amylase output;⁷⁰ according to their results, pancreatic amylase of a 75 years adult would be 30% lower than that of a 40 years adult.

Furthermore, in a rather early study, two groups of healthy men were examined (mean age of the first group 24.7 \pm 3.6 years, n = 10, mean age of the second group 67.2 \pm 6.3 years, n = 10= 10). In all subjects the exocrine pancreatic secretion was

examined after repeated stimulation of the pancreas.⁷⁷ No significant difference in pancreatic amylase output was observed between the two groups after one stimulation of the pancreas. However, repeated stimulations resulted in a significant decrease of about 35% in amylase output in the older group. These findings suggest some exhaustion of the pancreatic secretion function in old age. Finally, Dreiling et al. (1985) did not find any statistically significant difference in duodenal amylase activity with age, although a 25% difference in amylase activity was observed between volunteers in the 50-59 years group compared to volunteers in the 70-79 years group.59

Conclusion for pancreatic enzymes. Although some studies showed no difference between young and older adults in terms of pancreatic enzymes, the general trends indicated a decrease in the activity or output of most of the enzymes. The observed reduction was around 13 to 35%, depending on the enzyme studied.

Recommendation: since pancreatic enzymes are provided by the addition of pancreatin, it is recommended to decrease the amount of pancreatin (expressed by trypsin activity) in the older adult model by 20% compared to the young adult in vitro digestion model, i.e. 80 U trypsin per mL of intestinal content (or 1600 U pancreatic lipase per mL).

Table 3 Parameters for the elderly model are summarized and compared to the adult model

Phase	Parameter	Adult	Elderly
Oral	SSF	See Brodkorb <i>et al.</i> (2019) ⁵	Same
	composition Food : SSF dilution	1:1	1:1 or 1:1 according to DM
	pН	7.0	7.0
	Duration	2 min	2 min
	Chewing protocol	Dilute food with SSF at a ratio of 1:1 (wt/wt) to achieve a swallowable bolus with a paste-like consistency. If necessary, simulate mastication by mincing the food in an electric or manual mincer	Use of a basic meat mincer (manual, consisting of a 5 cm mincing disk, a 0.5 cm mesh size and a blade, only one pass) to produce food particles, then add SSF at pH 7 (with salivary amylase in case of starch-containing foods) or saliva at a ratio of 1:1
	Amylase	75 U ml ⁻¹ (using DNS as substrate, see Brodkorb <i>et al.</i> 2019) ⁵	75 U ml $^{-1}$ (using DNS as substrate, see Brodkorb <i>et al.</i> 2019) 5
Gastric	SGF	See Brodkorb et al. (2019) ⁵	Same
	composition	,	
	Bolus : SGF dilution	1:1	1:1
	рH	3.0	3.7
	Duration	2 h	3 h
	Pepsin	2000 U ml ⁻¹ of gastric content (using haemoglobin as substrate, see Brodkorb <i>et al.</i> 2019) ⁵	1200 U ml ⁻¹ of gastric content (using haemoglobin as substrate, see Brodkorb <i>et al.</i> 2019) ⁵
	Gastric lipase	60 U ml ⁻¹ of gastric content (using tributyrin as substrate, see Brodkorb <i>et al.</i> 2019) ⁵	36 U ml ⁻¹ of gastric content (using tributyrin as substrate, see Brodkorb <i>et al.</i> 2019) ⁵
Intestinal	SIF composition	See Brodkorb <i>et al.</i> (2019) ⁵	Same but with $[Ca^{2+}] = 1 \text{ mM}$
	Chyme : SIF dilution	1:1	1:1
	pН	7.0	7.0
	Duration	2 h	2 h
	Pancreatin	100 U ml ⁻¹ trypsin (using TAME as substrate, see Brodkorb <i>et al.</i> 2019) ⁵	80 U ml $^{-1}$ trypsin (using TAME as substrate, see Brodkorb <i>et al.</i> 2019) ⁵
	Bile salts	10 mM bile salts	6.7 mM bile salts

SSF: simulated salivary fluid, SGF: simulated gastric fluid, SIF: simulated intestinal fluid. DNS: 3,5-dinitrosalicyclic acid. TAME: p-toluenesulfonyl-L-arginine methyl ester.

Bile

Only two studies investigating the effect of ageing on bile concentration and providing interpretable values were found in the literature. In the first one, a 38% decrease in bile acids synthesis with age was reported (1.32 mM per day under 40 years and 0.81 mM per day over 65 years, n = 60).⁷⁸

In another study involving only 24 subjects, a 33% decrease in postprandial conjugated and unconjugated serum bile acids levels was observed with ageing.⁷⁹

Recommendation: based on these limited data, it is recommended to reduce the amount of bile salts in the intestinal phase by 33% in the older adult model compared to the young adult INFOGEST model, i.e. the reduction to 6.7 mM bile salts.

Conclusion

In conclusion, the exhaustive literature search that was conducted within the EAT4AGE consortium, as well as the exchanges that were held in the framework of the INFOGEST international network on food digestion have allowed for design of a static in vitro digestion model representative of bolus properties after oral processing and of the physiology of the gastrointestinal tract of an older, healthy adult. The most important differences relative to the INFOGEST young adult model correspond to different pH and duration of the gastric phase, different activities of the digestive enzymes in the stomach and small intestine and the concentration of bile salts (Table 3). The oral phase might also be different especially for denture wearers or people suffering from xerostomia or dysphagia. Nevertheless, it has to be noted that for some parameters, the values considered in the proposed model were based on a limited number of rather old publications and new data would be of paramount importance to refine the model in future studies. In the coming months, EAT4AGE partners will apply the proposed in vitro digestion model of the older adult to three types of food matrices (cereal-based, dairy and meat products) and compare the data with those obtained with the young adult in vitro model as well as with already published in vivo data, when available.

Conflicts of interest

There are no conflicts to declare.

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