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Pea protein extraction method impacts the protein (micro)structural organisation and *in vitro* digestion kinetics

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There is increasing interest in including pulse proteins into food products due to their nutrient-rich and sustainable character. However, little is known regarding the consequences of different extraction approaches on the pulse protein structure and the subsequent protein (micro)structural organization and protein digestion kinetics. Therefore, three green pea protein extracts were created: (i) cooking followed by cotyledon cell isolation, (ii) alkaline extraction followed by isoelectric precipitation, or (iii) salt extraction, and compared to the original pea flour as well as to sodium caseinate. The results showed that encapsulated, denatured protein inside pea cotyledon cells presented the (s)lowest digestion, while accessible and more native protein (e.g., pea flour, pea protein salt extract) presented much faster and higher digestion. Moreover, the alkali extracted pea protein was denatured to some extent, significantly lowering *in vitro* digestion kinetics. In the second part, three different *in vitro* approaches were applied to digest the salt extracted pea protein. Semi-dynamic gastric digestion approaches simulate *in vivo* conditions more closely which especially impacted the rate of digestion.

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1 Introduction

Pulse seeds are sustainable crops rich in nutrients such as protein. Given the topical protein transition, pulse seeds can be seen as valuable sources of alternative protein.¹ Additionally, the increasing demand for innovative and functional food products resulted in the exploration of plant-based protein as food ingredients (e.g., flours, concentrates and isolates).²

Industrially, proteins are majorly extracted under alkaline conditions followed by isoelectric precipitation and spray drying to obtain a protein-rich powder. Both the alkaline conditions and spray drying process can enhance hydrophobic interactions between proteins which in turn can negatively affect functional properties such as protein solubility.⁵

Therefore, research has been focusing on other extraction techniques and how they impact techno-functional properties. In that context, several studies showed that the protein extraction method used, affects the protein types and/or structures present in the extract which in turn has consequences for the techno-functional properties such as foaming, gelling, or emulsifying potential.^{3,5–10}

Next to the optimization of techno-functional aspects of protein extracts, the nutritional functionality should not be forgotten. In this context, our research group recently showed that a beverage containing a commercial pea protein isolate was only *in vitro* digested to a limited extent. However, when increasing protein dispersibility by high pressure homogenisation, *in vitro* protein digestion kinetics were improved.¹¹ Until now, only a limited number of studies focused on the *in vitro* evaluation of proteolysis rate and extent of non-commercial protein extracts. These nutritional consequences are of utmost importance to investigate and to understand so the industry can further optimize its production processes to generate pulse protein ingredients with desired techno-functional as well as nutritional properties.

Various methods have been employed to assess *in vitro* digestion.¹² The most commonly used in this context are static *in vitro* models based on the human gastrointestinal physiology. These models are simpler, more economical, and offer higher throughput compared to *in vivo* models.^{12,13} Their

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main limitation is the absence of any time dependency of the digestion processes within a compartment.¹³ Important physiological factors that directly affect digestion kinetics include the dynamic nature of secretions (*e.g.* digestive enzymes, pH) and gastric emptying. These factors also alter, *e.g.* nutrient structural organization, enzymatic activity, and substrate-enzyme contact time. In response to this, semi-dynamic *in vitro* models were developed to study digestion phenomena under more realistic conditions. In 2020, a consensus semi-dynamic *in vitro* gastric digestion model was published by the INFOGEST consortium.¹⁴ This method gives specific attention to mimicking the transient nature of gastric secretions and emptying by use of a single reactor.¹⁴ Furthermore, a semi-dynamic method to introduce dynamic secretions of digestive fluids and enzymes both during the gastric and small intestinal phase was implemented by our research group.¹⁵ However, based on these existing models, strategic selections on which relevant dynamic factors to include into a digestion simulation cannot be made. Currently, limited knowledge exists on how these different semi-dynamic approaches impact *in vitro* nutrient digestion kinetics.

Peas are considered interesting plant-based protein alternatives due to their limited number of allergens compared to other protein alternatives such as soy.³ Besides, peas contain a relatively high amount of protein (20–25% on dry a basis), but are also rich in starch, micronutrients, and bioactives such as phenolic acids.⁴ The main pea proteins are the salt-soluble globulins (55–80%) and water-soluble albumins (18–25%).³ Therefore, the aim of this study was twofold. Firstly, we aimed to investigate the impact of different protein extraction techniques on the extract composition, protein structure, protein solubility and *in vitro* proteolysis kinetics of green peas. To reach this goal, three extraction techniques were compared: (i) hydrothermal treatment combined with mechanical disintegration and wet sieving to recover an extract containing intact cotyledon cells encapsulating protein, (ii) alkaline extraction in combination with isoelectric precipitation to generate an industrial-like protein-rich extract yet freeze dried and not heat treated, and (iii) salt extraction to generate a protein-rich extract in which the native protein structure was retained as much as possible. The original non-cooked pea (flour) sample was used as a reference sample as well as an animal-based protein extract, sodium caseinate. Secondly, three *in vitro* digestion approaches were compared for one specific protein extract (*i.e.*, pea protein salt extract) to gain more detailed insight into how the *in vitro* digestion approach can impact digestion kinetics.

2 Materials and methods

2.1 Materials

Mature dry green peas (*Pisum sativum*) were kindly donated by Casibbeans (Melsele, Belgium), sorted, cleaned from foreign material, and stored at -40°C . Sodium caseinate (CAS) from bovine milk was acquired from Sigma Aldrich (Diegem,

Belgium). Porcine pepsin (2681 U mg^{-1}), trypsin (247.8 U mg^{-1}), and chymotrypsin (55.8 U mg^{-1}) were purchased from Sigma Aldrich (Diegem, Belgium). All chemicals and reagents used in this work were of HPLC or analytical grade.

2.2 Preparation of samples

2.2.1 Pea flour. Dry green peas were thawed and ground using a laboratory mill (Cyclotec 1093 Sample Mill, FOSS, Högenäs, Sweden) with a sieve pore mesh size of 0.5 mm. The green pea flour (PF) obtained was stored in a desiccator at room temperature until further use.

2.2.2 Pea cotyledon cells. Dry green peas were soaked (16 h, room temperature) and cooked (45 min, 95°C) exactly as described before by Guevara-Zambrano *et al.*¹⁶ Subsequently, after mechanical disintegration of the cooked peas (2 min, 8000 rpm, IKA® T25 ultra-turrax, Janke and Kunkel, Germany), pea cotyledon cells were isolated using a vibratory wet sieve shaker (AS200, Microtract Retsch, Haan, Germany) at an amplitude of 2.5 mm for 3 min. The pea cotyledon cells (PC) were collected from the sieves with a 40, 80, and $125\text{ }\mu\text{m}$ pore size,^{16,17} lyophilized (Alpha 2–4 LSC plus, Christ, Osterode, Germany) and stored in a desiccator at room temperature.

2.2.3 Pea protein alkali extract. Pea protein was isolated using two distinct extraction procedures. The procedure described by Yang *et al.*¹⁸ was used with few adaptations for the alkaline extraction, and followed by isoelectric precipitation.

Green pea flour (section 2.2.1) was dispersed in Milli-Q water (1 : 10 w/v) for 1 h at room temperature. Hereafter, the pH of the mixture was adjusted to 11 using 1 M NaOH. The mixture was stirred for 2 h at room temperature and centrifuged at $8000g$ for 15 min at 4°C (Avanti JXN-26, Beckman Coulter Inc., Indianapolis, IN, USA). The supernatant was filtered and adjusted to a pH of 4.5 using 1 M HCl to precipitate protein. After centrifugation with the same parameters as described above, the pellet was collected, resuspended in Milli-Q water, and neutralized (pH 7) using 1 M NaOH. The protein fraction obtained was dialyzed for 48 h against demineralized water (Spectra/Por®, Molecular weight cut-off = 6–8 kDa), lyophilized (Alpha 2–4 LSC plus, Christ, Osterode, Germany), and stored in a desiccator at room temperature. The pea protein fraction obtained after alkali extraction will be referred to as PPAE.

2.2.4 Pea protein salt extract. A second protein isolation procedure using salt extraction conditions was based on the methods described by Sun and Arntfield¹⁹ and Tanger, Engel and Kulozik.²⁰

Green pea flour (section 2.2.1) was dispersed in a 0.1 M phosphate buffer containing 0.3 M NaCl (1 : 5 w/v) for 2 h at room temperature. Hereafter, the mixture was centrifuged at $4260g$ for 15 min at 4°C (Avanti JXN-26, Beckman Coulter Inc., Indianapolis, IN, USA). The supernatant was filtered, dialyzed for 48 h against demineralized water (Spectra/Por®, Molecular weight cut-off = 6–8 kDa), lyophilized (Alpha 2–4 LSC plus, Christ, Osterode, Germany), and stored in a desiccator at room temperature. The pea protein fraction obtained after salt extraction will be referred to as PPSE.



2.3 Sample characterisation

To determine the composition of all samples (PF, PC, PPAE, PPSE, CAS), the powdered samples were first ball milled (Ball mill MM400, Retsch GmbH, Haan, Germany) to disrupt any physical barriers. All analysis were performed at least in duplicate.

Starch content of all samples was determined using a Total Starch Assay Kit K-TSTA (AA/AMG, Megazyme Inc., Bray, Ireland). An automated Dumas analyser (CHNS-O Elemental Analyzer, CE instrument, Thermo Fischer Scientific, Waltham, MA, USA) was used to evaluate the nitrogen content of all samples. For all pea-derived samples, a conversion factor of 5.4 was applied to determine the protein content,²¹ while for CAS a conversion factor of 6.38 was used.²² The total lipid content was determined gravimetrically, according to the procedure of Janssen *et al.*²³ The ash content of all samples was determined as the mass difference upon complete incineration at 550 °C in a muffle furnace (Controller P330, Nabertherm, Lillenthal, Germany). The moisture content of all samples was determined as the mass difference upon vacuum oven drying at 70 °C (UniEquip 1445-2, Planegg, Germany).

A Q2000 heat flux differential scanning calorimeter with Advanced Tzero™ technology (TA Instruments, New Castle, DE, USA) was used to determine enthalpy changes which can be related to protein denaturation according to the procedure described by Chigwedere *et al.*²⁴ Lastly, sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) under reducing conditions was used to study the protein profile of the different samples under investigation.²⁵

2.4 Preparation of protein dispersions

For the suspensions made from PF or PC, a 5% protein concentration (dry basis) was considered due to the relatively low protein, yet high dry matter content in those samples. For the suspensions made from the protein extracts, PPAE, PPSE, and CAS, a 10% protein concentration (dry basis) was considered.

Briefly, 5% (PF, PC) or 10% (PPAE, PPSE, CAS) protein (w/v) was weighed from the respective dry powder, suspended in 0.1 M sodium phosphate buffer (pH 7), and stirred overnight at room temperature.

2.5 Protein solubility

To determine protein solubility, protein dispersions (section 2.4) as well as diluted dispersions according to the simulation conditions of the static gastric phase without enzyme addition (section 2.7.1) were considered. All samples were centrifuged at 20 000g for 30 min at 20 °C (Optima XPN-80 Ultracentrifuge, Beckman Coulter Inc., Indianapolis, IN, USA). The supernatants were collected with a pipette and lyophilized (Alpha 2–4 LSC plus, Christ, Osterode, Germany). An aliquot of the non-centrifuged samples was lyophilized as well to determine the exact amount of nitrogen in the original dispersion. The protein content of all samples was quantified using an automated Dumas analyser and applying a correction factor as described in section 2.3. Lastly, protein solubility was calcu-

lated according to eqn (1). The analysis was performed in duplicate per sample type.

$$\text{Protein solubility (\%)} = \frac{\text{protein}_{\text{supernatant}}}{\text{protein}_{\text{initial dispersion}}} \times 100 \text{ (dry basis)}. \quad (1)$$

2.6 Protein secondary structure

To gain some insight in the protein secondary structure of PPAE and PPSE, Fourier transform infrared spectroscopy was used. Transmission of each sample was measured from 4000 cm⁻¹ to 400 cm⁻¹ at a resolution of 4 cm⁻¹. A total of 100 scans were taken per sample and peaks were deconvoluted before spectra were analysed.²⁶ Existing literature was used to identify the Gaussian peaks obtained. In this sense, peaks around 1638 cm⁻¹ were related with β -sheet structures, 1654 cm⁻¹ with α -helices, 1663 cm⁻¹ with β -turns, and 1680 cm⁻¹ with antiparallel β -sheets.²⁷ Additionally, peaks between 1625 and 1610 cm⁻¹ were related with intermolecular β -sheets,²⁸ and peaks between 1695 and 1690 cm⁻¹ to amino acid side chains.²⁹ To calculate the contribution of each protein secondary structure, its corresponding peak area was divided over the total peak area.

2.7 In vitro digestion simulation

Three different *in vitro* digestion approaches were applied in this study. Firstly, all samples were digested using the commonly used static *in vitro* digestion procedure of INFOGEST.³⁰ Secondly, one selected sample, PPSE, was also digested using more advanced semi-dynamic *in vitro* digestion conditions: (i) an approach without gastric emptying (MuReDi approach¹⁵) and (ii) an approach with gastric emptying (INFOGEST approach¹⁴).

2.7.1 Static *in vitro* digestion. The static *in vitro* digestion protocol published by the international INFOGEST network was followed.³⁰ In the oral phase, however, the recommendations described in the semi-dynamic INFOGEST protocol were applied since this creates a more relevant bolus.¹⁴ Briefly, a 1 : 1 dilution of the dry matter content of the food to the salivary fluids (including simulated salivary fluid, 0.3 M CaCl₂, and Milli-Q water) was applied. Hereafter, simulated gastric fluid, 0.3 M CaCl₂, pepsin (2000 U mL⁻¹ chyme), 2 M HCl (to reach pH 3), and Milli-Q water were added according to the ratios described by Brodkorb *et al.*³⁰ To simulate the small intestinal phase, simulated intestinal fluid, 0.3 M CaCl₂, trypsin (100 U mL⁻¹ digest), chymotrypsin (25 U mL⁻¹ digest), 1 M NaOH (to reach pH 7), and Milli-Q water were added as described by Brodkorb *et al.*³⁰ Digestion was simulated in glass tubes which rotated head-over-heels (40 rpm) in an incubator set at 37 °C. Sixteen individual tubes were used to simulate the gastric and small intestinal phase. The enzymatic activity in each tube was thermally stopped (5 min, 95 °C) at different pre-determined moments (after 5; 10; 20; 30; 45; 60; 90; 120; 125; 130; 140; 150; 165; 180; 210; and 240 min) to be



able to study the time dependent digestive behaviour (*i.e.*, rate and extent) of all samples.³¹

2.7.2 Semi-dynamic *in vitro* digestion. The MuReDi approach as described in our previous work¹⁵ was employed to study the semi-dynamic digestion behaviour of the PPSE dispersion without including gastric emptying.

The oral phase was mimicked as described in section 2.7.1. The gastric phase was mimicked semi-dynamically exactly as described before.¹⁵ Briefly, 90% of the gastric fluid master mix (pH = 0.45) and 90% of the pepsin solution were added gradually as a function of gastric digestion time to reach a final pH of 2 and an enzyme activity of 2000 U mL⁻¹ chyme. The subsequent small intestinal phase was mimicked statically. In that way, the effect of the semi-dynamic gastric phase on the digestion kinetics could be studied.

Digestion was simulated in glass reactor vessels, each having individual pH, temperature, and stirring control. Every vessel was connected with 4 individual feeding lines which are used to pump the appropriate digestive solutions at a set speed in each vessel. For more details regarding the BioXplorer 100 equipment and the MuReDi approach, we kindly refer to our previous publication.¹⁵ The enzymatic activity in each vessel was thermally stopped (5 min, 95 °C) at different pre-determined moments (after 5; 10; 20; 30; 45; 60; 90; 120; 125; 130; 140; 150; 165; 180; 210; and 240) to be able to study the time dependent digestive behaviour.

2.7.3 Semi-dynamic *in vitro* digestion with gastric emptying. The standardised semi-dynamic *in vitro* digestion method with gastric emptying¹⁴ was followed to study the impact of gastric emptying on the proteolysis kinetics both in the simulated gastric and small intestinal phase.

The gastric phase with gastric emptying was executed exactly as described by Mulet-Cabero *et al.*¹⁴ A sample volume of 30 mL was started from, and eight emptying moments were considered, meaning that each 6.63 min an aliquot was emptied from the vessel taking into account a gastric emptying rate of 2 kcal per min. The total gastric phase lasted 53 min. The vessel with a thermostat jacket (Metrohm) was shaken using an orbital shaker (Stuart mini gyro-rocker, SSM3). The gastric fluid master mix (pH = 0.45) and pepsin solution (2000 U mL⁻¹ chyme) were added gradually as a function of simulated gastric digestion time using the pumps of a BioXplorer 100 equipment (H.E.L Group, London, U.K.) and winISO software. The pH was monitored by the BioXplorer 100 and the corresponding winISO software.

To analyse the samples taken during the gastric phase, enzymatic activity of each emptied aliquot was stopped by thermal inactivation (5 min, 95 °C).

To perform a static *in vitro* small intestinal phase after gastric emptying, all emptied aliquots were combined in a vessel containing simulated intestinal fluid and bile salts. This vessel was connected to the BioXplorer 100 equipment, which allowed to bring and keep the pH around 7 in a controlled manner, inactivating pepsin. Thermal inactivation was not considered since it could alter protein structure before small intestinal digestion. At the end of the gastric phase, the

remaining amount of Milli-Q water was added to reach the same dilution as described in the static INFOGEST protocol. Aliquots of this mixture (8.75 mL) were brought into test tubes to which 1.25 mL of enzyme mixture was added, containing trypsin (100 U mL⁻¹ digest) and chymotrypsin (25 U mL⁻¹ digest). Hereafter, all test tubes were incubated at 37 °C and rotated head-over-heels (40 rpm). Again, the enzymatic activity was stopped (5 min, 95 °C) at different pre-determined moments (after 5; 10; 20; 30; 45; 60; 90; and 120 min of enzyme addition in the small intestinal phase) to study the time dependent digestive behaviour. The simulation of semi-dynamic *in vitro* digestion with gastric emptying was performed in duplicate since sampling in the gastric phase was done from one vessel yielding dependent data.

2.8 Protein digestion quantification

The readily bioaccessible protein are described as the fraction of protein that can be absorbed as such at the brush-border of the small intestine.³² This protein fraction predominantly contains amino acids, dipeptides, and tripeptides.

All digested samples were centrifuged (10 min, 2000g, 4 °C, Sigma 4–16 KS, Sigma, Osterode am Harz, Germany) to harvest the soluble protein fraction under these centrifugation conditions. Next, 5% trichloroacetic acid (TCA) was added to each sample to precipitate larger peptides.³² Lastly, the *o*-phthalaldehyde spectrophotometric assay was used to quantify the readily bioaccessible fraction (NH₂(TCA)) in each sample.^{32–34} The amount of readily bioaccessible protein is expressed according to eqn (2). The analysis was performed in duplicate per sample point.

$$\begin{aligned} \text{Readily bioaccessible protein (\%)} \\ = \frac{\text{NH}_2(\text{TCA}) - \text{NH}_2(\text{initial})}{\text{NH}_2(\text{total}) - \text{NH}_2(\text{initial})} \times 100 \end{aligned} \quad (2)$$

2.9 Particle size analysis

The mean particle size and particle size distributions were assessed in duplicate for all initial samples as well as during digestion using a laser diffraction equipment (Beckman Coulter Inc., LS 13 320, FL, USA). Additionally, the microstructure was visualized by means of an optical microscope (Olympus BX-51, Olympus, Optical Co. Ltd, Tokyo, Japan) equipped with an Olympus XC-50 digital camera and an image-analysis software (cellSens, Olympus, Tokyo, Japan). Fluorescent microscopy (X-Cite Fluorescence Illumination, Series 120Q EXFO Europe, Hants, United Kingdom) was used to visualize protein (micro)structures in the dispersions based on the intrinsic protein fluorescence.

2.10 Data analysis

Significant differences among samples were determined by one way ANOVA and a Tukey's Studentised Range *Post-hoc* test with a 95% level of significance ($P < 0.05$).

The protein digestion kinetics of each sample were determined using eight individual recipients in each digestive



Table 1 Chemical composition (%) of all protein samples. Different letters within the same column indicate significant differences ($p < 0.05$)

	Starch (%)	Protein (%)	Lipids (%)	Ash (%)	Moisture (%)
Pea flour	45.37 ± 1.36 b	18.00 ± 0.37 d	4.49 ± 0.01 b	1.61 ± 0.17 cd	7.73 ± 0.09 a
Pea cells	57.69 ± 1.39 a	18.92 ± 0.11 d	1.28 ± 0.08 c	0.81 ± 0.13 d	7.88 ± 0.02 a
Pea protein alkali extract	1.50 ± 0.13 c	77.03 ± 0.03 b	8.23 ± 0.01 a	4.34 ± 0.16 a	3.97 ± 0.13 d
Pea protein salt extract	1.48 ± 0.23 c	74.63 ± 0.96 c	7.54 ± 0.38 a	1.98 ± 0.44 bc	5.95 ± 0.04 b
Casein	0.23 ± 0.08 c	87.79 ± 0.62 a	0.92 ± 0.06 c	2.85 ± 0.17 b	5.07 ± 0.05 c

phase. Consequently, each recipient can be considered to be an independent, consecutive evaluation of the same digestion curve. The experimental data obtained were therefore integrated using nonlinear regression (JMP Pro16, SAS Institute Inc., Cary, NC, USA). For this, an empirical, first order fractional conversion model was applied as shown in eqn (3). In eqn (3), C (%) represents the predicted readily bioaccessible protein at time t (min), C_i (%) represents the initial amount of readily bioaccessible protein, C_f (%) represents the final, plateau value reached under the given conditions, k (min^{-1}) represents the reaction rate constant, and t (min) represents the time in the simulated gastric or small intestinal phase.^{31,35} Confidence intervals (95%) were used to determine significant differences among parameter estimates.

$$C = C_f + (C_i - C_f) \times e^{-kt} \quad (3)$$

3 Results and discussion

3.1 Characterization of all protein samples

The composition of all samples was determined and displayed in Table 1. The original PF contained around 45% starch, 18% protein, and 4.5% lipids. These values are in line with results previously reported.⁴ The macronutrient composition of PC (yield of 44.7 ± 1.2%) is the most similar to the one of the original PF. The starch and protein content slightly increased which can be explained by seed coat removal, compound leaching, and/or concentration of the cells during soaking, cooking, and wet sieving to isolate cells that predominantly contained starch and protein.³⁶ Oppositely, the protein extracts PPAE (yield of 12.2 ± 0.5%) and PPSE (yield of 9.5 ± 1.2%), only contained very small amounts of starch (around 1.5%), and were rich in protein (74–77%). The lipid content of these protein extracts was around 7.5 to 8.2% which can be explained by not defatting the PF before protein extraction.⁶ CAS was very rich in protein (around 88%) and contained only minor amounts of other nutrients such as starch and lipids (Table 1).

In addition to the nutrient composition, all samples were also analysed using a differential scanning calorimeter to detect potential enthalpy changes which could be related to protein denaturation (Table 2). Only for PF, PPAE, and PPSE a peak could be detected around 89 °C. When comparing the energy input needed for protein denaturation between PPAE and PPSE, PPSE needed significantly more energy to denature protein. This most probably means that the protein of PPSE is

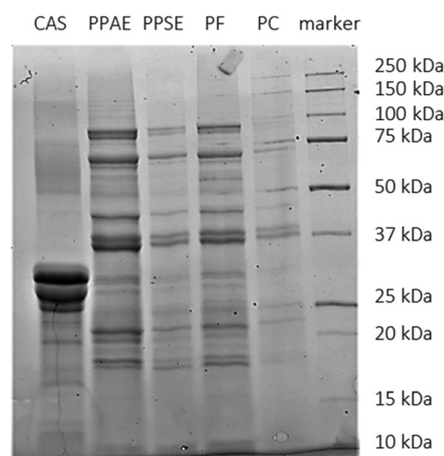
Table 2 Enthalpy measurement of all protein samples (n.d.: not detected). Different letters within the same column indicate significant differences ($p < 0.05$)

	Enthalpy (J kg^{-1})
Pea flour	0.96 ± 0.01 b
Pea cells	n.d.
Pea protein alkali extract	0.79 ± 0.01 b
Pea protein salt extract	1.84 ± 0.10 a
Casein	n.d.

more native or underwent less denaturation than in PPAE as a result of the different extraction conditions. It was shown before that extracting protein under alkaline conditions could cause (partial) protein denaturation.⁹ For PC, the cooking process most probably caused complete denaturation of the protein present.¹⁶

3.2 Evaluation of protein profile and secondary structure

SDS-PAGE analysis was performed (Fig. 1) to study the polypeptide mass profile of the different protein samples. Generally, the pea protein samples presented several bands between 10 and 100 kDa. The main ones were located at a molecular weight of around 75–80 kDa, 70 Da, 40 kDa, 35–37 kDa, 20–22 kDa, 18 kDa and 10 kDa. This profile is in line with previous SDS-PAGE analysis under reducing conditions of green

**Fig. 1** SDS-PAGE profile of the different protein extracts (CAS: sodium caseinate, PPAE: pea protein alkali extract, PPSE: pea protein salt extract, PF: pea flour, PC: pea cells).

pea protein.^{20,37} Bands around 70–80 kDa are associated with convicilin, while bands around 18, 35, and 50 kDa are related with dissociated vicilin trimers.²⁰ The bands at 20 and 40 kDa are linked to β -legumin and α -legumin, respectively.^{6,37} These two subunits are normally connected with disulphide bonds which were broken due to the reducing conditions used. In all pea protein samples, bands related to albumins could also be observed (<15 kDa).²⁰ However due to the dialysis step (molecular weight cut-off of 6–8 kDa), some smaller albumins could have been lost. Overall, little differences were observed in terms of the protein profile among the pea protein samples since in all samples globular protein were the main protein structures present.

Since some structural differences were expected among the two pea protein rich extracts due to the distinct extraction conditions applied, the secondary structure was studied as well (Table 3). For both samples, (intermolecular) β -sheets were the major structures present. Additionally, substantial amounts of α -helices and β -turns were detected while anti-parallel β -sheets and amino acid side chains were present in minor amounts. Significant differences among PPAE and PPSE were detected in terms of the relative amount of β -sheets and α -helices. This profile was in line with previous research on the structural and functional properties of the different storage protein in pulses.⁶ Our results indicated that PPAE had slightly, but significantly more β -sheets while PPSE presented significantly more α -helices. A negative relation between the β -sheet content and protein digestibility was shown before for legumes like chickpea, lentil, and soybean. This was explained by the hydrophobic character of these β -sheet structures which can also negatively affect protein solubility.³⁸

3.3 Protein solubility of the initial protein dispersions and under simulated gastric conditions

Protein solubility was evaluated for all samples (Table 4), except PC. It can be assumed for PC, that the protein solubility will be very low since all protein are encapsulated within the cell structures. The original PF had a native protein solubility of 66% in the 0.1 M sodium phosphate buffer (pH 7) used for sample preparation. The two protein-rich extracts, PPAE and PPSE, presented a completely different behaviour. PPAE presented a very low solubility (around 27%) in the phosphate buffer, while 77% of the PPSE protein was considered soluble. This observation is

Table 3 Secondary protein structures present in the two pea protein rich extracts. Samples indicated with an asterisk are significantly different ($p < 0.05$)

	Pea protein alkali extract	Pea protein salt extract
Intermolecular β -Sheets (%)	30.8 \pm 1.3	30.3 \pm 2.3
β -Sheets (%)	37.8 \pm 1.9	34.8 \pm 1.0*
α -Helix (%)	8.7 \pm 1.0	11.9 \pm 2.0*
β -Turn (%)	14.1 \pm 0.5	14.4 \pm 0.6
Anti-parallel β -sheets (%)	5.3 \pm 0.3	5.4 \pm 0.4
Amino acid side chains (%)	3.3 \pm 0.5	3.3 \pm 0.3

Table 4 Protein solubility of the original sample dispersed in a 0.1 M sodium phosphate buffer (pH 7) versus under conditions of the simulated static gastric phase. Different letters within the same column indicate significant differences among samples, while asterisks indicate significant different among the conditions evaluated ($p < 0.05$)

	Protein solubility (%)	
	Original sample	Gastric phase conditions
Pea flour	66.08 \pm 3.83 c	43.68 \pm 2.30 c*
Pea protein alkali extract	27.13 \pm 0.03 d	35.90 \pm 1.98 d*
Pea protein salt extract	76.91 \pm 0.82 b	76.87 \pm 0.78 a
Casein	87.59 \pm 2.98 a	49.10 \pm 0.80 b*

most probably related to differences in protein structures (Tables 2 and 3) present in the extract due to the different extraction conditions used. Previous research on pea protein already showed that using salt extraction led to a higher protein solubility in water while protein solubility in water was lower when using alkaline conditions.²⁰ It is hypothesized that PPSE possesses more native protein in comparison to PPAE, solubilizing more easily due to its less hydrophobic character. The lab-scale produced PPAE had a higher solubility than determined by other researchers for commercial pea protein isolates which also underwent spray drying, leading to higher protein denaturation than freeze drying.³ Protein solubility for the pea protein rich extracts PPSE and PPAE, could have been improved by defatting the flour before protein extraction, reducing hydrophobic protein–lipid interactions.³ Also, co-extracted phytic acids have shown to negatively affect protein solubility of soybean.³⁹ Additionally, protein solubility of pulse protein dispersion can be improved by processing, such as high pressure homogenisation.^{11,40} The milk protein, sodium caseinate, presented a relatively high protein solubility of around 88%.

To gain more insight in the phenomena occurring during the simulated gastric phase, protein solubility was also evaluated under these increased salt and acidic conditions, yet without enzyme addition. Generally, some changes were observed in comparison to the original samples. For example, a significant decrease from 66% to 44% of protein solubility for PF was noticed. Protein solubility is highly depending on pH and salt conditions.^{6,20} The pH of the (static) gastric phase, pH 3, is rather close to the pI of protein very often reducing protein solubility in comparison to neutral pH conditions.⁶ However, for PPAE, protein solubility significantly increased from 27% to 36%. Nonetheless, protein solubility was still considered low for PPAE under gastric phase conditions. Protein solubility of PPSE remained the same as in the phosphate buffer (77%). Lastly, solubility of sodium caseinate significantly decreased from 88% to 49% under the gastric phase conditions. This milk protein is known for its low solubility at pH conditions between 3 and 5, especially at room temperature.⁴¹

3.4 Microstructural evaluation of all initial protein dispersions and during *in vitro* digestion

Particle size (distributions) of all initial, undigested samples as well as during *in vitro* digestion were evaluated to gain



insight in potential changes which could affect proteolysis kinetics (Table 5 and Fig. 2, 3). Additionally, the observations of the particle size analysis were complemented with micrographs to visually display the microstructure (Tables 6 and 7).

Different particle size distributions were observed for the initial protein extracts. PF presented the highest average particle size, and a very heterogenous particle size distribution. The different peaks observed can be related to different microstructures present within the sample. In this sense, the region between 10 and 50 μm was linked to individual starch granules, protein bodies, and cell wall fragments released during cell breakage. The broad region between 50 and 1000 μm was related to seed coat fragments and other large tissues.³⁶ The presence of this mix of microstructures is also visually observed in the micrograph (Table 6). The PC sample presented a uniform peak around 100–110 μm . This corresponds with the size of individual cotyledon cells which were extracted after wet sieving of cooked peas as reported before.¹⁶ Also, the micrograph mainly displayed individual cotyledon cells, next to some individual starch granules (Table 6). Previous research on individual cotyledon cells of lentils showed that the presence of the cell structure retards proteolysis in comparison to freely accessible protein.³⁶ This more complex structural organisation of pea protein might eventually impact the *in vitro* proteolysis kinetics. Despite the high protein solubility presented in the original dispersion (section 3.3), CAS presented a relatively large average particle size. However, the micrograph presented a nicely dispersed microstructure for CAS. The two other protein-rich extracts, PPAE and PPSE, presented an average particle size around 19 and 12 μm , respectively. This is much smaller than the sizes detected for the other samples. However, this difference in particle size between PPAE and PPSE could still impact the digestion kinetics. The (fluorescent) micrographs do not completely align with the observations made during particle size analysis. PPSE presents some (weak) aggregation which might have broken up

during particle size analysis, while PPAE presents a mix of densely packed protein (brighter green oval shapes) and dissolved protein (darker green background) (Table 7).

Particle size and microstructure were analysed at three different moments during the simulated gastric or small intestinal phase. For all pea protein-based samples, little differences were observed during the simulated static gastric phase in comparison to their respective initial sample (Table 5). For CAS, a drastic increase was observed when the sample was brought under the digestive conditions of the gastric phase. Previous studies already showed that milk protein mainly composed of casein, is pH sensitive and coagulates at pH values close to its pI.⁴² The aggregates formed decreased significantly in size as a function of gastric digestion time, yet the gastric samples presented much larger sizes than the original CAS dispersion. Overall, it can be stated that all pea protein-related samples showed much smaller particle sizes than CAS in the low pH environment of the gastric phase which might offer possibilities for usage in acidic food products. When the gastric phase was simulated in a semi-dynamic way, slightly larger particles were observed during the gastric phase which tended to decrease again in size towards the end of the simulated gastric digestion. For most samples, the particle sizes remained in the same order of magnitude during the simulated static small intestinal phase (Table 5). One exception was CAS, which presented a drastic decrease in particle size when shifting from the low pH conditions of the gastric phase towards neutral pH conditions in the small intestinal phase, related to a change in protein solubility.⁴¹

3.5 *In vitro* protein digestion

3.5.1 Static *in vitro* protein bioaccessibility of various pea protein extracts as affected by the extraction method. For the different protein samples, clear differences in digestion behaviour could be observed both in the simulated gastric and small intestinal phase (Fig. 4). For instance, in the *in vitro* gastric

Table 5 Evolution of the volume-based mean particle sizes of all protein dispersions as well as the pea protein salt extract digested using different *in vitro* digestion approaches (GP: gastric phase; SIP: small intestinal phase; GE: gastric emptying)

	Undigested	GP 20 min	GP 60 min	GP 120 min
Casein	62.92 \pm 4.70	647.92 \pm 16.29	345.30 \pm 17.41	221.03 \pm 2.52
Pea flour	233.89 \pm 13.01	268.76 \pm 20.13	302.22 \pm 59.54	309.24 \pm 43.36
Pea cells	132.27 \pm 1.38	133.34 \pm 0.32	134.71 \pm 0.09	134.88 \pm 0.22
Pea protein alkali extract	19.13 \pm 0.22	15.70 \pm 1.22	15.78 \pm 0.03	15.97 \pm 0.33
Pea protein salt extract static	12.27 \pm 4.05	6.65 \pm 0.44	7.23 \pm 0.06	6.08 \pm 1.51
Pea protein salt extract without gastric emptying		29.38 \pm 0.46	21.57 \pm 0.43	18.69 \pm 0.30
		GE 1	GE 2	GE 3
Pea protein salt extract with gastric emptying		13.56 \pm 0.76	23.04 \pm 0.67	11.93 \pm 1.46
	Undigested	SIP 20 min	SIP 60 min	SIP 120 min
Casein	62.92 \pm 4.70	6.86 \pm 1.77	6.44 \pm 0.76	7.08 \pm 0.63
Pea flour	233.89 \pm 13.01	231.59 \pm 40.44	202.42 \pm 45.85	201.33 \pm 38.63
Pea cells	132.27 \pm 1.38	140.45 \pm 0.32	134.90 \pm 0.45	134.42 \pm 0.07
Pea protein alkali extract	19.13 \pm 0.22	21.66 \pm 2.31	20.41 \pm 0.74	17.11 \pm 1.04
Pea protein salt extract static		20.85 \pm 1.19	18.77 \pm 1.69	19.00 \pm 0.60
Pea protein salt extract without gastric emptying	12.27 \pm 4.05	28.90 \pm 0.37	35.55 \pm 2.22	91.34 \pm 11.24
Pea protein salt extract with gastric emptying		14.55 \pm 0.55	15.25 \pm 0.96	15.04 \pm 2.81



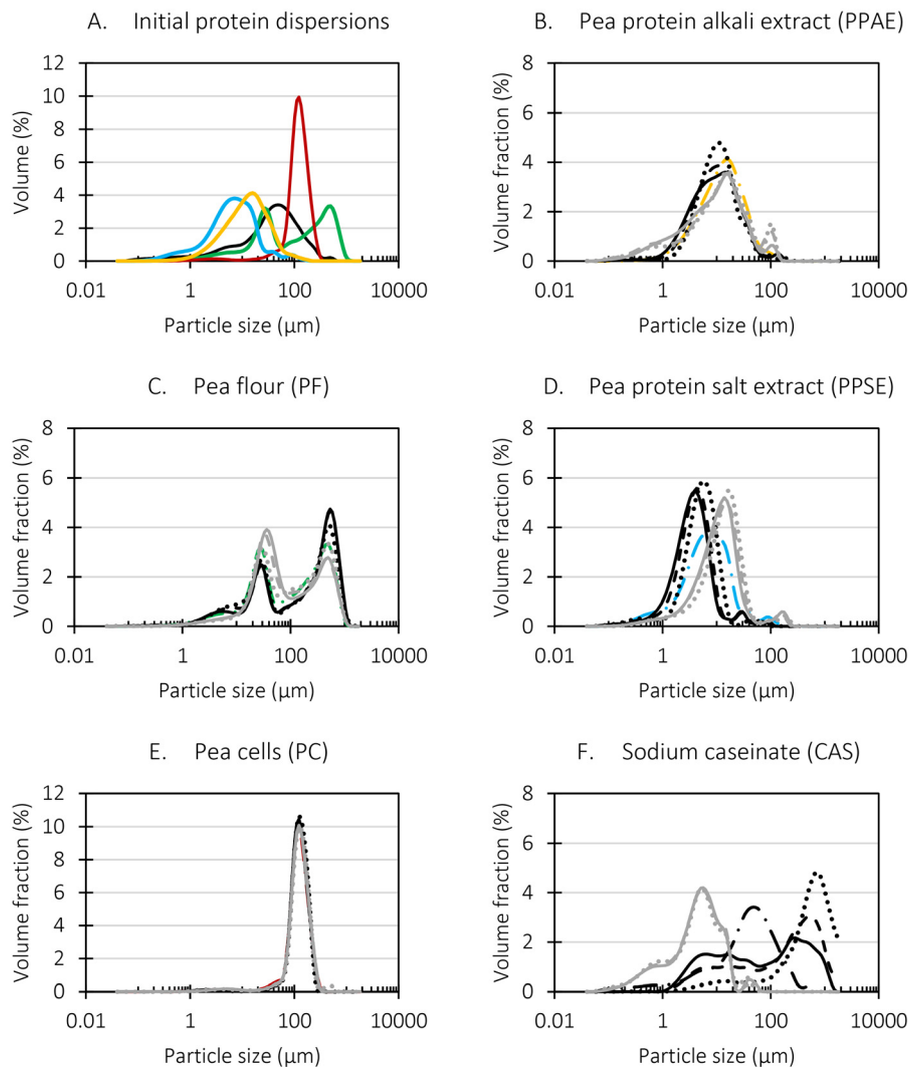


Fig. 2 Particle size distribution of the (A) initial protein dispersions (— pea flour (PF), — pea cells (PC), — pea protein alkali extract (PPAE), — pea protein salt extract (PPSE), — sodium caseinate (CAS)) and (B–F) initial protein dispersion distributions versus protein dispersion distributions during the static simulated digestive tract (coloured, dashed-dotted lines: initial samples, black lines: gastric phase, grey lines: small intestinal phase, dotted lines: after 20 min, dashed lines: after 60 min, full lines: after 120 min of simulated digestion).

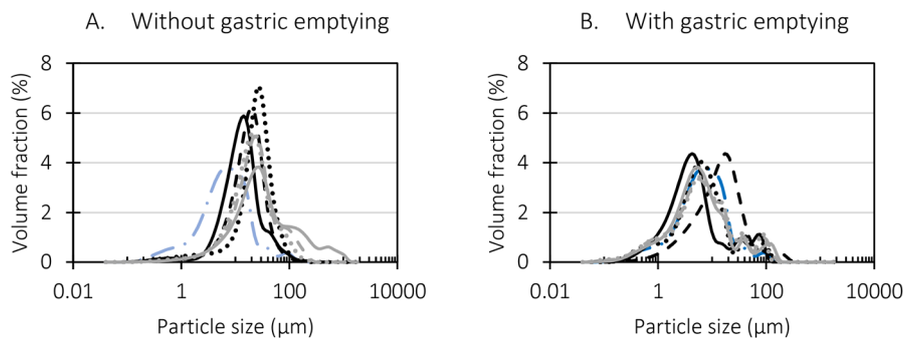
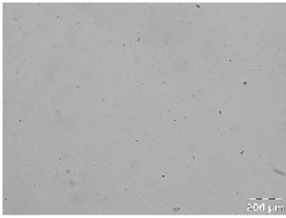
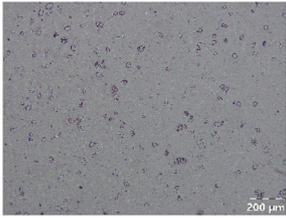
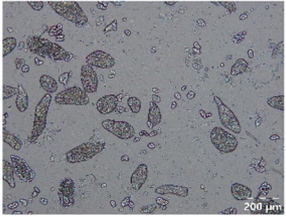
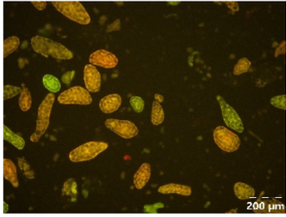
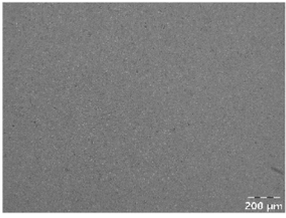
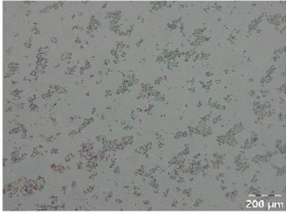
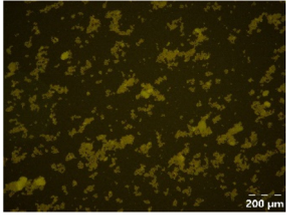


Fig. 3 Particle size distribution of the pea protein salt extract dispersion digested using an *in vitro* semi-dynamic digestion approach (A) without gastric emptying and (B) with gastric emptying (black lines: gastric phase, grey lines: small intestinal phase, dotted lines: after 20 min, dashed lines: after 60 min, full lines: after 120 min of simulated digestion). It must be noted that for the simulated gastric phase with gastric emptying (black lines in (B)), the samples were analysed after 6.6 min (dotted line), 26.5 min (dashed line), and 53 min (full line) since the total length of the gastric phase was only 53 min in this case.



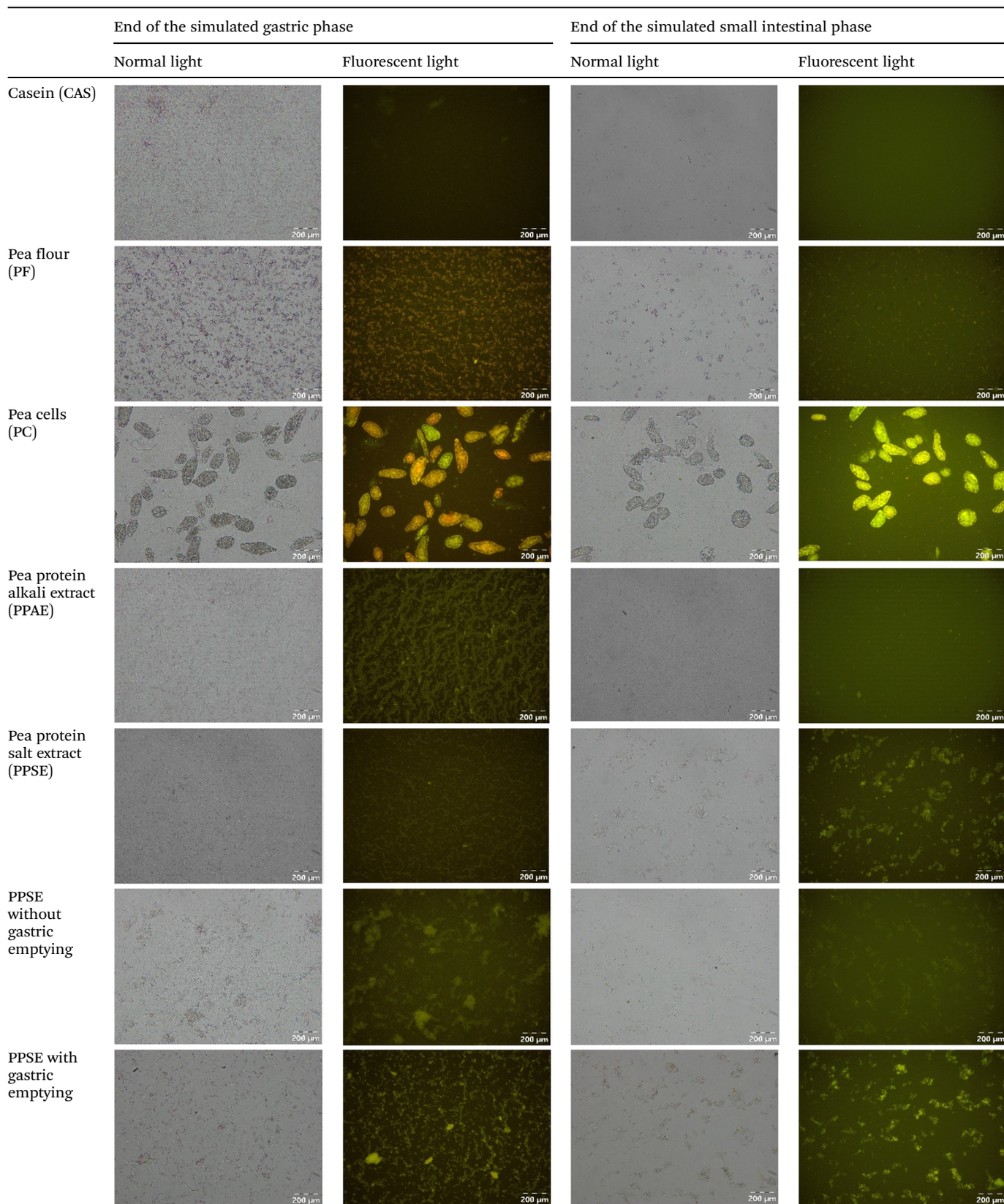
Table 6 Microscopic evaluation of the (micro)structure of all initial samples under investigation. The scale bar represents a length of 200 μm

	Initial samples Normal light	Fluorescent light
Casein (CAS)		
Pea flour (PF)		
Pea cells (PC)		
Pea protein alkali extract (PPAE)		
Pea protein salt extract (PPSE)		

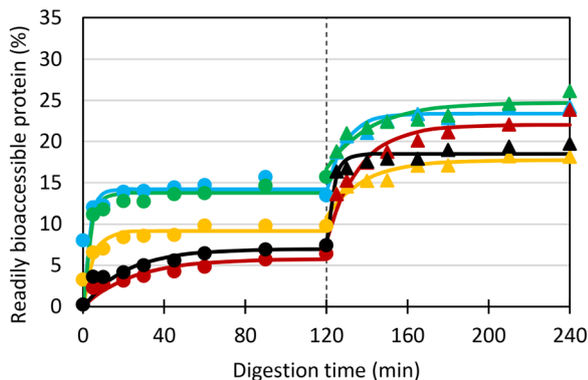
phase, PF presented the highest and fastest protein digestion (Table 8). PF was characterized by an open microstructure with relatively easily accessible protein, despite the moderate protein solubility. The presence of other nutrients such as (non-gelatinised) starch and lipids, seemed to not form a barrier for protein enzymatic hydrolysis. Additionally, PF was not thermally treated in this work, so the protein structure could be considered native which was rapidly digested and to a relatively high extent. These insights are in line with a recent study of Rivera del Rio *et al.*⁴³ on the effect of processing on pea protein digestibility. PPSE presented highly accessible, soluble, and native proteins as well. However, some aggregation was observed during particle size analyses. Hence, this might explain why PPSE showed very similar *in vitro* protein digestibility kinetics as PF, yet slightly (s)lower. PPAE pre-

sented significantly (s)lower *in vitro* proteolysis kinetics in the gastric phase compared to PPSE and PF. It can be hypothesized that the structural differences among the PPAE and PPSE sample induced by the different extraction conditions, significantly impacted the *in vitro* digestion kinetics. The different extraction conditions used in this study impacted other functional properties as well which should also be taken into account for the application potential of pea protein (extracts). PC presented the slowest and lowest proteolysis kinetics in the simulated gastric phase of all pea protein-based samples. It is known that when the protein accessibility for enzyme interaction is restricted, for example by an intact cell wall barrier, digestion rate and extent is reduced in comparison to samples with a higher accessibility for enzymes.^{16,36} A higher level of structural organization was also observed for



Table 7 Microscopic evaluation of the (micro)structure of all samples under investigation at the end of the simulated gastric and small intestinal phase. The scale bar represents a length of 200 μm 

- A. All protein dispersions digested using static *in vitro* conditions



- B. Dispersion of pea protein salt extract digested using three different *in vitro* approaches

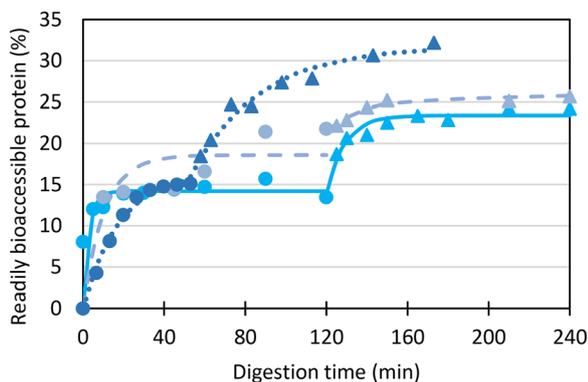


Fig. 4 Time dependent release of readily bioaccessible protein (%) quantified spectrophotometrically as a function of *in vitro* digestion time. (A) Different protein dispersions digested under static *in vitro* conditions (— pea flour (PF), — pea cells (PC), — pea protein alkali extract (PPAE), — pea protein salt extract (PPSE), — sodium caseinate (CAS), ● gastric phase, ▲ small intestinal phase). (B) The dispersion of pea protein salt extract (PPSE) digested using three different *in vitro* approaches (full lines: static approach, dashed lines: semi-dynamic approach without gastric emptying (MuReDi), dotted lines: semi-dynamic approach with gastric emptying).

the CAS sample in the gastric phase. Also for CAS a relatively (s)low protein digestion was observed. This was in line with previous research on milk protein.⁴² CAS is called a “slow protein” due to coagulation occurring under low pH conditions of the gastric phase.¹⁴ It has been shown that the rate of proteolysis can impact amino acid uptake in the blood,⁴⁴ and eventually prolong the feeling of fullness after food/meal ingestion.⁴⁵ In the simulated small intestinal phase, very similar trends were observed among the pea protein samples with PF and PPSE presenting a significantly higher percentage of bioaccessible protein than PPAE and PC. Oppositely, the CAS sample reached the plateau value rather rapidly. The changing conditions in the small intestinal phase reversed the coagulation of CAS (Table 7), reducing the level of protein structural organisation and thus increasing the accessibility of CAS protein for the digestive enzymes added during small intestinal digestion. Overall, these results showed that modulating the protein structural organisation is an useful tool to steer protein digestion kinetics, not only prior to digestion but also during digestion. Combining different structural organisations and/or protein sources might potentially be an interesting strategy to tune satiety and satiation responses.

3.5.2 *In vitro* protein bioaccessibility of a pea protein salt extract as affected by the digestion simulation approach. Three different *in vitro* digestion approaches were used to digest PPSE to gain insight in potential differences in digestion kinetics related to the *in vitro* digestion approach applied. The results related to these proteolysis kinetics are given in Fig. 4 and Table 8.

Firstly, when looking at the experimental data of the simulated gastric phase, the semi-dynamic approaches presented the highest final extent of bioaccessible protein. Oppositely, the static approach presented a significantly lower amount of bioaccessible protein at the end of the gastric phase. Additionally, differences were observed related to the reaction rate constant (k). It could be clearly observed that the static approach reached the plateau the fastest, followed by the semi-dynamic approaches. The semi-dynamic approach without gastric emptying did not reach the plateau within the time frame of the simulated gastric phase. This can be related to

Table 8 Estimated kinetic parameters for *in vitro* protein digestion in the gastric and small intestinal phase of the different protein dispersions evaluated using the spectrophotometric approach. C_f is the final extent of bioaccessible protein estimated by the model in the gastric or small intestinal phase (%), C_i (%) is the initial extent of bioaccessible protein at the start of the small intestinal phase and k is the reaction rate constant of the release of bioaccessible protein in the gastric or small intestinal phase (min^{-1}). Different small or capital letters in the same column indicate significant differences among parameter estimates according to their confidence intervals (95%) (PPSE: pea protein salt extract)

	Gastric phase			Small intestinal phase			
	C_f	k	R^2	C_0	C_f	k	R^2
Sodium caseinate	7.00 ± 0.35 c	0.047 ± 0.008 c	0.96	7.52 ± 0.95 d	18.49 ± 0.39 d	0.279 ± 0.071 a	0.95
Pea flour	13.75 ± 0.44 a	0.289 ± 0.068 a	0.95	16.70 ± 0.93 a	24.71 ± 0.78 a	0.046 ± 0.016 c	0.90
Pea cells	5.79 ± 0.50 d	0.041 ± 0.010 c	0.92	7.90 ± 1.30 d	22.03 ± 0.86 c	0.062 ± 0.016 c	0.94
Pea protein alkali extract	9.18 ± 0.30 b	0.198 ± 0.037 b	0.95	10.82 ± 0.79 c	17.75 ± 0.57 e	0.055 ± 0.018 c	0.91
Pea protein salt extract static	14.21 ± 0.35 a/C	0.329 ± 0.066 a/A	0.97	13.85 ± 0.84 b/C	23.39 ± 0.41 b/C	0.111 ± 0.025 b/A	0.95
PPSE without gastric emptying	18.59 ± 1.54 A	0.096 ± 0.038 B	0.86	21.50 ± 0.32 A	25.83 ± 0.27 B	0.047 ± 0.011 B	0.96
PPSE with gastric emptying	16.48 ± 0.57 B	0.056 ± 0.005 C	0.99	15.98 ± 0.82 B	31.76 ± 1.08 A	0.030 ± 0.006 C	0.97



the combined effect of gradual enzyme addition and acidification of the chyme¹⁵ which allows a gradual increase of pepsin activity. In contrast, in the case of static *in vitro* digestion, all enzymes are added from the start of the simulated phase.

Besides, the pH is set at 3 from the start of the gastric phase which is not optimal but allows pepsin activity from the onset of the gastric phase. It can thus be concluded that the *in vitro* digestion approach used will impact the digestion kinetics obtained, especially the path followed to the end of (gastric) digestion. Similar conclusions were made in our previous research on lentils, yet more similar final digestions extents were observed.¹⁵

Slightly different observations were made in the simulated small intestinal phase. The digestion behaviour of the static and MuReDi approach were very similar, reaching similar extents despite to the differences at the end of the simulated gastric phase which seemed to be compensated by the action of trypsin and chymotrypsin. The static small intestinal simulation followed after gastric emptying, went much slower, reaching a final amount of bioaccessible protein slightly higher than the other approaches. The much slower reaction rate constant in the former case might be explained by the mix of digested aliquots present at the start of the small intestinal phase. In more detail, to simulate the static small intestinal phase after gastric emptying, all emptied samples were pooled and pepsin activity was stopped by increasing pH to around 7. This means that each emptied sample had a different extent of proteolysis. In other words, some pea protein were only hydrolysed to a very low extent (*e.g.*, first emptied sample) while the samples emptied towards the end of the gastric phase were most likely hydrolysed to a much higher extent which might impact enzyme action in the subsequent small intestinal phase. In particular, it is possible that trypsin and chymotrypsin potentially prefer to hydrolyse larger protein structures first, slowing down the overall release of bioaccessible protein.

4 Conclusions

In this work, different extraction methods were used to create pea protein extracts with distinct structural properties, impacting their structural organization in dispersion. More specifically, individual pea cells were isolated from cooked peas using wet sieving, leading to denatured, encapsulated protein with a high level of structural organization. Besides, two extraction conditions were used, alkaline conditions followed by isoelectric precipitation (PPAE) *versus* salt extraction (PPSE), to create pea protein rich extracts with distinct structural properties. The pea protein profile between those two extracts was highly similar, yet some significant differences were observed in the secondary structure. In this context, more protein-protein hydrophobic interactions were detected for PPAE than for PPSE. Additionally, enthalpy changes related to protein denaturation were higher for PPSE than PPAE, meaning that the protein of PPSE were more native than the ones of PPAE. Lastly, protein solubility in a phosphate buffer was much lower

for PPAE (27%) than for PPSE (77%). However, limited differences were detected in particle size (distribution) among the two samples when dispersed in a phosphate buffer. In terms of *in vitro* protein digestibility, both the rate constant and extent of proteolysis were much lower for the pea cells and alkali extract than for the salt extract. Besides, the salt extract had a very similar digestion behaviour as the original, uncooked pea flour. This means that the differences in protein structure significantly impacted the structural organization of protein in dispersion and subsequently *in vitro* digestion kinetics.

Lastly, also the *in vitro* digestion approach used significantly impacted the digestion behaviour. Static digestion approaches are the most simple approaches delivering a quick screening tool to compare different food design properties. However, when more realistic insight in the digestion behaviour, especially rate, needs to be obtained, semi-dynamic approaches seem to be more appropriate to use since these approaches significantly affect the digestion kinetics.

Overall, the insights obtained in this work should be further extended with different extraction and production conditions. Additionally, the potential of such new pulse-based ingredients should be investigated in more realistic food products with varying pH and salt concentrations since this will largely affect its application potential. Besides, *in vivo* consequences of incorporating protein with different (combinations) microstructures could be studied as well. In conclusion, this work displayed the need to better understand structure–function–digestion relationships of new pulse-based ingredients or foods to be able to steer the (digestive) functionality and potentially physiological responses.

Abbreviations

CAS	Sodium caseinate
PC	Pea cells
PF	Pea flour
PPAE	Pea protein alkali extract
PPSE	Pea protein salt extract
TCA	Trichloroacetic acid

Author contributions

S.H.E. Verkempinck: conceptualization; data curation; formal analysis; funding acquisition; investigation; methodology; visualization; writing – original draft. D. Duijsens: methodology; writing – review & editing. A. Mukherjee: investigation; methodology. P.J. Wilde: supervision; writing – review & editing.

Conflicts of interest

There are no conflicts to declare.



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