Nutritional evaluation of mixed wheat–faba bean pasta in growing rats: impact of protein source and drying temperature on protein digestibility and retention

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Abstract
This study aimed to evaluate the nutritional value of pasta enriched with legume or wheat gluten proteins and dried at varying temperature. A total of four isonitrogenous experimental diets were produced using gluten powder/wheat semolina (6/94, g/g) pasta and faba bean flour/wheat semolina (35/65, g/g) pasta dried at either 55°C (GLT and FLT, respectively) or 90°C (FVHT and GVHT, respectively). Experimental diets were fed to ten 1-month-old Wistar rats (body weight = 176 (SEM 15) g) for 21 d. Growth and nutritional, metabolic and inflammatory markers were measured and compared with an isonitrogenous casein diet (CD). The enrichment with faba bean increased the lysine, threonine and branched amino acids by 97, 23 and 10%, respectively. Protein utilisation also increased by 75% (P<0.01) in FLT in comparison to GLT diet, without any effect on the corrected faecal digestibility (P>0.05). Faba bean pasta diets’ corrected protein digestibility and utilisation was only 3-5 and 9%, respectively, lower than the CD. Growth rate, blood composition and muscle weights were not generally different with faba bean pasta diets compared with CD. Corrected protein digestibility was 3% lower in GVHT than GLT, which may be associated with greater carboxymethyllysine. This study in growing rats clearly indicates improvement in growth performance of rats fed legume-enriched pasta diet compared with rats fed gluten–wheat pasta diet, regardless of pasta drying temperature. This means faba bean flour can be used to improve the protein quality and quantity of pasta.

Key words: Faba bean pasta: Drying temperature: Protein digestion: Protein retention: Protein structure

Proteins are fundamental macronutrients in human and animal diet. Their primary function is to provide amino acids required for growth and maintenance of body tissue after digestive hydrolysis. Nutritional value of proteins varies according to their essential/non-essential amino acid composition, total digestion and rate of digestion in the gastrointestinal tract and the body’s utilisation of the absorbed minimally oxidised amino acids1,2. Nutritional value of proteins is highly dependent on food component and processing conditions.

Animal products, chiefly meat, are a natural part of human diet recognised as a source of high-quantity, high-quality protein. Meat provides 16·3–23·3 g/100 g protein3 with a good essential amino acid profile1,4. Meat supplied 40% of human-diet protein in the world in 2013 (FAOSTAT, 2017). Meat consumption reached 34 kg/capita per year in 2013–2015 and is projected to increase by 6·5% in developed countries by 20255. However, it is recommended to decrease meat consumption, due

Abbreviations: ATTD, apparent total tract digestibility; BV, biological value; CD, control diet; CFD, corrected faecal digestibility; CML, carboxymethyllysine; FLT, diet based on faba bean–wheat pasta dried at low temperature; FN, faecal nitrogen; FVHT, diet based on faba bean–wheat pasta dried at very high temperature; GLT, diet based on gluten–wheat pasta dried at low temperature; GVHT, diet based on gluten–wheat pasta dried at very high temperature; LM, lean mass; LT, low temperature; MCP-1, monocyte chemoattractant protein-1; NI, nitrogen intake; NPU, net protein utilisation; PFD, protein-free diet; UN, urinary nitrogen; VHT, very high temperature.

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to adverse effects of excessive intake on the environment\(^\text{60}\) and on health, notably an increased risk of CVD\(^\text{7}\).

This context has spurred growing interest in the development and production of vegetable-based protein foods that are easy to prepare, rich in good quality proteins and appreciated by consumers. Pasta, a highly popular product, partly meets these requirements. It is consumed worldwide and widely appreciated as a relatively inexpensive, easy-to-prepare, long-shelf-life food staple. From a nutritional point of view, dry wheat pasta is a source of protein (12–13\%) and slowly digestible carbohydrates (28\%)\(^\text{50}\) that result in a low glycaemic index\(^\text{9,10}\). Despite this valuable nutritional quality, pasta – like most cereal products – is deficient in lysine and threonine\(^\text{11}\). Furthermore, it generally has lower protein content than animal products (around 4\% protein in cooked pasta). Several studies have been carried out to improve the nutritional quality of pasta by incorporating non-traditional ingredients such as legume flours\(^\text{11–14}\). Legumes are a valuable source of dietary proteins (20–40\%)\(^\text{15}\). They contain 11–20\% (w/w) resistant starch and are rich in slowly digestible carbohydrates\(^\text{16}\). In addition, legumes are rich in lysine and have a complementary amino acid profile to wheat proteins\(^\text{15}\). We have recently shown that the incorporation of 35\% faba bean flour increased pasta protein content and enhanced lysine score by 52\% in comparison to classical wheat pasta\(^\text{11}\). Enrichment with legume (faba bean) was also reported to increase the \textit{in vitro} digestibility of the protein fraction, due to a specific weaker structure of the pasta protein\(^\text{11}\).

Besides pasta composition, variation in the process parameters, notably drying temperature, was also reported to affect pasta protein digestibility due to a modification of its protein network structure\(^\text{17,18}\). Industry pasta makers generally dry the pasta at high temperature (65–94\°C) to save time and energy. High-temperature drying was also shown to improve the technological quality (resilience and overall integrity) of legume-enriched pasta\(^\text{19}\). However, applying temperatures higher than 85\°C strongly aggregates the component proteins, leading to lower \textit{in vitro} protein digestibility. This phenomenon is not observed when pasta is dried at low temperature (LT)\(^\text{17,18}\).

There is little data on the impact of the enrichment of pasta with legumes on \textit{in vitro} protein digestibility\(^\text{13,14,20}\), and no data available on the effect of both improving protein composition and using different drying temperatures on the \textit{in vivo} digestibility and nutritional value of pasta proteins. To address this gap, this study in growing rats aimed to compare the nutritional value and degree of protein digestibility and retention of mixed legume (faba bean)–wheat pasta dried at two drying temperatures (low and very high) with gluten–wheat pasta and casein.

### Methods

#### Pasta manufacture and diet preparation

Wheat (\textit{Triticum durum}) semolina, wheat (\textit{Triticum aestivum})–gluten powder and faba bean (\textit{Vicia faba}) flour were provided by Panzani, Syral and GEMEF Industries, respectively. Composition of raw material is presented in Table 1. The moisture content was determined in duplicate by drying at 130\°C for 2 h in an oven\(^\text{21}\). The ash content was determined in duplicate by calcinations at 900\°C, according to the French method NF 03-720\(^\text{22}\). Crude protein content was determined in duplicate with an enzymatic assay kit (Megazyme Co.; AACC method 76-13.01)\(^\text{23}\). Fibres and lipids were determined by ISHA using the JORF (1986) method\(^\text{23}\) and by Inovalys according to the NF ISO 6492\(^\text{20}\) procedure, respectively.

Pasta was manufactured and cooked as recently described by Laleg et al.\(^\text{11}\). Briefly, wheat semolina was enriched with 6\% gluten powder or 35\% faba bean flour to obtain the same protein content in the blend (approximately 15 g/100 g) and then processed into spaghetti form using a continuous pilot-scale pasta extruder (Bassano). All spaghetti were dried at LT (55\°C, 70–90\% relative humidity, for 15 h) or at very high temperature (VHT, 90\°C, 70–90\% relative humidity, for 3 h) in a pilot-scale drier (AFREM).

To elaborate rat diets, pasta were cooked, dried at LT (40\°C, 70\% relative humidity) for 24 h to reach approximately 12\% moisture and then ground into a fine powder. A total of four experimental diets were then formulated by adding cellulose,

### Table 1. Composition of raw materials

<table>
<thead>
<tr>
<th>Composition of raw material (g/100 g)</th>
<th>Wheat semolina</th>
<th>Faba bean flour</th>
<th>Gluten powder</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>14±6</td>
<td>10±8</td>
<td>5±9</td>
</tr>
<tr>
<td>Protein</td>
<td>11±2</td>
<td>21±4</td>
<td>75±1</td>
</tr>
<tr>
<td>Starch</td>
<td>66±4</td>
<td>51±4</td>
<td>9±9</td>
</tr>
<tr>
<td>Ash</td>
<td>1±1</td>
<td>2±6</td>
<td>0±9</td>
</tr>
<tr>
<td>Fibres</td>
<td>2±0</td>
<td>10±4</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Lipids</td>
<td>1±5</td>
<td>1±7</td>
<td>&lt;1</td>
</tr>
</tbody>
</table>

*The moisture content was determined in duplicate by drying at 130\°C, for 2 h, in an oven\(^\text{21}\). Protein content was determined in duplicate using the Kjeldahl procedure (digestion units: BUCHI K-350, INST-0060-ANALYS; distillation unit: BUCHI K-350, INST-XXXX-ANALYS; NF V 03-050, 1970)\(^\text{24}\) with a conversion factor of 5.7 for wheat and 6.25 for legume proteins. Starch content was determined in duplicate with an enzymatic assay kit (Megazyme Co.; AACC method 76-13.01)\(^\text{23}\). The ash content was determined in duplicate by calcinations at 900\°C according to the French method NF 03-720\(^\text{22}\). Fibres and lipids were determined by ISHA (Lonjumeau) using the JORF (1986) method\(^\text{23}\) and by Inovalys (Nantes) according to the NF ISO 6492\(^\text{20}\) procedure, respectively. Moisture: s<0.3, protein: s<0.06, starch: s<0.5, ash: s<0.01.
soya fat and mineral and vitamin mix to pasta powders (Table 2): 

- FLT diet: based on faba bean–wheat pasta dried at LT
- FVHT diet: based on faba bean–wheat pasta dried at VHT
- GLT diet: based on gluten–wheat pasta dried at LT
- GVHT diet: based on gluten–wheat pasta dried at VHT

These diets were compared with a control diet based on casein and maize starch (CD). In addition, a protein-free diet (PFD) mainly made of maize starch was studied (Table 2). This diet was used to estimate the endogenous and bacterial faecal N (EBFN) and the endogenous urinary N (EUN) of rats (N loss in faeces and urine that are not related to the N content of the food).

Total starch content was determined in each diet in duplicate using an enzymatic assay kit (Megazyme Co.; AACC method 76-13.01)(23). Crude protein content was determined in duplicate using the Kjeldahl procedure (digestion units: BÜCHI K-350, INST-0060-ANALYS; distillation unit: BÜCHI K-350, INST-XXXX-ANALYS; NF V 03-050, 1970) with a conversion factor of 5·7 for wheat and 6·25 for legume proteins 6·25(26). Total starch content was determined in duplicate using an enzymatic assay kit (Megazyme Co.; AACC method 76-13.01)(23). Fat and fibres were determined by Qualitec using the JORF (1977)(27) and the AOAC (1990)(28) methods, respectively. Total energy was calculated using estimates of the energy content of fat, starch and fibres, and protein of 37·7, 16·7, and 16·7 kJ/g, respectively. Protein:SD = 0·2, starch:SD = 1·1, fibres:SD = 2·9, lipids:SD = 0·2.

### Biological assays

#### Ethical considerations.

All animal procedures were approved by the institution’s animal welfare committee (Comité d’Éthique en Matière d’Expérimentation Animale Auvergne; permit number: C2EA-02) and conducted in accordance with European guidelines on the care and use of laboratory animals (2010-63UE). The animals were housed in the research unit’s own animal facilities (accreditation no. C634514). At the end of the experiment, the rats were killed under isoflurane anaesthesia. Every effort was taken to minimise animal suffering.

#### Experimental design

In all, 1-month-old male Wistar rats (n = 60, body weight = 176 (SEM 15) g) were purchased from JANVIER, and housed in individual cages in an environmentally controlled room (temperature 22±0·5°C, humidity 50±5 %, and 12/24 h light cycle). The rats had ad libitum access to chow and water. The rats were fed a CD during the 2-week acclimatisation period (AIN-93M, safe diets) before the study began. A total of sixty rats were randomly allotted to one of the six diets, for a total of ten animals per diet. Each group received ad libitum one experimental diet (GLT, GVHT, FLT or FVHT), CD or PFD. The study lasted 21 d, during which rat food intake and weight gain were recorded every week. During the last 4 d, rats were transferred into metabolic cages where food intake was weighed every day. Faeces and urine were also collected, weighed and then stored at 4°C until N determination at LaSalle (Bauvais), according to the Dumas method. At the end of the experiment, the rats were fasted for 12 h before being killed by decapitation. Blood samples were collected after decapitation into EDTA tubes (Veno-safe®) and centrifuged at 1300 g for 10 min at 4°C to separate the plasma, which was then snap-frozen in liquid N₂ and stored at –80°C until analysis. Blood was also collected into dry tubes

### Table 2. Composition of the experimental and control diets

<table>
<thead>
<tr>
<th>Ingredient (g/100 g)</th>
<th>Experimental diet</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FLT</td>
</tr>
<tr>
<td>Pasta</td>
<td>90·0</td>
</tr>
<tr>
<td>Casein</td>
<td>0·0</td>
</tr>
<tr>
<td>Maize starch</td>
<td>0·0</td>
</tr>
<tr>
<td>Cellulose</td>
<td>0·9</td>
</tr>
<tr>
<td>Soya fat</td>
<td>4·6</td>
</tr>
<tr>
<td>Minerals and vitamins</td>
<td>4·5</td>
</tr>
<tr>
<td>Total</td>
<td>100·0</td>
</tr>
<tr>
<td>Nutritional value of the diet (g/100 g)*</td>
<td></td>
</tr>
<tr>
<td>Protein</td>
<td>14·1</td>
</tr>
<tr>
<td>Starch</td>
<td>57·1</td>
</tr>
<tr>
<td>Fibres</td>
<td>6·6</td>
</tr>
<tr>
<td>Lipid</td>
<td>5·3</td>
</tr>
<tr>
<td>Energy (kJ/100 g)</td>
<td>1503</td>
</tr>
</tbody>
</table>

* Total protein content was determined in duplicate using the Kjeldahl procedure (digestion units: BÜCHI K-350, INST-0060-ANALYS; distillation unit: BÜCHI K-350, INST-XXXX-ANALYS; NF V 03-050, 1970) with a conversion factor of 5·7 for wheat and 6·25 for legume proteins 6·25(26). Total starch content was determined in duplicate using an enzymatic assay kit (Megazyme Co.; AACC method 76-13.01)(23). Fat and fibres were determined by Qualitec using the JORF (1977)(27) and the AOAC (1990)(28) methods, respectively. Total energy was calculated using estimates of the energy content of fat, starch and fibres, and protein of 37·7, 16·7, and 16·7 kJ/g, respectively. Protein:SD = 0·2, starch:SD = 1·1, fibres:SD = 2·9, lipids:SD = 0·2.
(Venosafe®) and, after incubation for 20 min at room temperature, centrifuged at 1300 g for 10 min at 4°C (Jouan GR 412, DBJ Labcare), to separate the serum, which was then snap-frozen in liquid N2 and stored at −80°C until analysis.

Immediately after killing, the plantaris, soleus and tibialis muscles were removed from each side. Liver, peri-renal adipose tissue (PAT) and subcutaneous adipose tissue (SAT) were also collected. All tissues were weighed, frozen in liquid N2 and stored at −80°C until analysis.

Body weight and body composition

Body weight was recorded at the beginning and end of the experiment. Body composition (lean mass (LM) and fat mass (FM)) of each rat was determined at the beginning and end of the experiment by MRI using Echo MRI (Echo Medical Systems). The results were expressed as percentage of body weight.

Protein quality evaluation of treatments diets

Rate of growth (g/d) was calculated as the weight gain between the beginning and end of the study ratioed to number of study days. Food and protein efficiency were calculated according to Proll et al. (31) by the equations:

\[
\text{Food efficiency} = \frac{\text{Rate of growth}}{\text{Food intake}} \quad (\text{g of weight gain/g food intake})
\]

\[
\text{Protein efficiency} = \frac{\text{Rate of growth}}{\text{Protein intake}} \quad (\text{g of weight gain/g protein intake})
\]

Protein quality of each diet was evaluated. Apparent total tract digestibility (ATTD), corrected faecal digestibility (CFD), net protein utilisation (NPU) and biological value (BV) were calculated according to Proll et al. (31) using the following equations:

\[
\text{ADF} \% = \frac{\text{NI} - \text{FN}}{\text{NI}} \times 100
\]

\[
\text{CFD} \% = \frac{\text{NI} - (\text{FN} + \text{EBFN})}{\text{NI}} \times 100
\]

\[
\text{NPU} \% = \frac{\text{NI} - (\text{FN} + \text{UN}) + \text{EBFN} + \text{EUN}}{\text{NI}} \times 100
\]

\[
\text{BV} \% = \frac{\text{NPU}}{\text{CFD}} \times 100
\]

EBFN is determined using the faecal N of PFD-group rats, EUN is determined using the urinary N of PFD-group rats, FN is the faecal N, NI is the N intake and UN is the urinary N.

Biochemical analyses

Tissue protein content was measured using a Micro BCA Protein Assay Kit (Thermo Fischer Scientific), according to the manufacturer’s instructions. Blood glucose, NEFA, total cholesterol, TAG, albumins and total proteins were measured using a Konelab 20 chemical analyser (Thermo Scientific). Plasma inflammatory markers, that is, monocyte chemotactant protein-1 (MCP-1), fibrinogen, and orosomucoid, and plasma metabolic markers, that is, insulin, adiponectin and leptin were measured using an ELISA kit (PromoKine), according to the manufacturer’s instructions.

Statistical analysis

Individual rat was considered as experimental unit. The sample size was calculated for an expected difference of 20–25% for corrected faecal protein digestibility between the control (casein) group and the gluten pasta diet group. A variance of 10% was considered according to the previous studies (15,20) for the selected parameter (corrected faecal protein digestibility). The type I error (α) was set at 5%, the type II error (β) was set at 5%, leading to 95% power. All analyses were performed using Statistica 8.0 software. All parameters were first analysed for normality. All data were tested by ANOVA with one factor: diet (one-way ANOVA), followed by a Fisher’s least significant difference test to compare means at the 5% significance level. Tendency towards statistical significance is considered if 0.05 < P < 0.1. The specific relationships between plasma (inflammatory and hormones) markers and metabolic markers, body weight and body composition were evaluated using Pearson’s correlation coefficient with 95% CI. In addition, all data on pasta diets (FLT, FVHT, GLT and GVHT) were subjected to two-way ANOVA using ‘formulation’ (faba bean–wheat or gluten–wheat) and ‘drying’ (LT or VHT) as factors. These results are given in the online Supplementary materials.

Results

Amino acid composition of diets

Table 3 reports the amino acid composition of diets. Essential amino acid profile differed considerably according to the protein source. All essential amino acid contents were lower in GLT compared with CD, notably lysine, threonine and branched amino acids. Among branched amino acids, leucine was 20% lower in GLT compared with CD. In FLT, the enrichment with faba bean increased the lysine content by 97%, threonine content by 23% and branched amino acid content by 10%, in comparison to GLT. Leucine content was increased by 7%, whereas the sulphur amino acid content decreased by 27% in comparison to GLT. VHT drying gave the same pattern of results. In addition, lysine content was 6 and 4% lower, and tryptophan content was 17 and 7% lower in GVHT and FVHT diets compared with GLT and FLT diets, respectively. CML was 98% higher in FHT compared with GLT diets. The use of VHT increased CMl by 67% in faba bean pasta diets and by 79% in gluten pasta diets in comparison to LT.

Impact of the type of diet on growth parameters and body composition

Throughout the study, rats consumed an equivalent amount of diet (Table 4). Table 4 reports food efficiency, protein
efficiency, rate of growth and final body weight of the rats. Regardless of the drying temperature (LT or VHT), rats receiving a faba bean pasta diet were characterised by a 1.7–2.0-fold higher food efficiency, protein efficiency and rate of growth, and a 1.3-fold higher body weight compared with rats receiving a diet composed of gluten pasta. FLT and FVHT were as food efficient and protein efficient as CD. The rate of growth and body weight was not significantly different between rats fed faba bean pasta diet and CD. Rats fed GLT and GVHT diets were characterised by a 1.6–1.9-fold lower food efficiency, protein efficiency and rate of growth, and 1.3-fold lower body weight compared with rats fed CD.

There was no difference in growth parameters of rats between LT and VHT drying regardless of the protein source (faba bean–wheat or gluten–wheat).

There were no significant differences between rats fed FLT and GLT in FM to body weight and LM to body weight ratios (Table 4), whereas rats fed GVHT had significantly lower FM and higher LM than their FVHT counterparts. Body composition of all rats fed pasta diets except GVHT was characterised by a higher FM to body weight ratio (44–61%) and a lower LM to body weight ratio (5–7%) than rats fed the CD. A significant drying effect was observed only in the gluten pasta diet group, with 21% lower FM and 4% higher LM in GVHT than GLT-fed rats.

Table 5 reports the muscle weight of each group. Regardless of pasta drying temperature, rats fed faba bean pasta diets had 27–34% heavier *plantaris*, 13–21% heavier *tibialis* anterior and 22–28% heavier *soleus* compared with rats fed gluten pasta diets. Rats fed faba bean pasta diets had generally a close muscle weight to rats fed CD, except for *soleus* which was 13% lower in FVHT compared with CD. Conversely, rats fed with gluten pasta diets were characterised by significantly lower muscle weight (24, 23 and 20% for *plantaris*, *tibialis* and *soleus*, respectively) compared with CD.

Rats fed FLT and FVHT had a 12% higher liver weight to body weight ratio compared with GLT and rats fed GVHT. Liver weight was 7 and 17% higher in rats fed CD than in rats fed faba bean and gluten pasta diets, respectively. There was no significant difference in PAT and SAT masses between rats fed FLT and FVHT.

### Table 3. Essential amino acid profile and carboxymethyl-lysine (CML) composition of the diets

<table>
<thead>
<tr>
<th>Amino acids (mg/g of protein)*</th>
<th>FLT</th>
<th>FVHT</th>
<th>GLT</th>
<th>GVHT</th>
<th>CD</th>
</tr>
</thead>
<tbody>
<tr>
<td>His</td>
<td>25.8</td>
<td>25.4</td>
<td>21.9</td>
<td>21.8</td>
<td>26.4</td>
</tr>
<tr>
<td>Ile</td>
<td>39.2</td>
<td>39.9</td>
<td>36.5</td>
<td>37.3</td>
<td>48.9</td>
</tr>
<tr>
<td>Leu</td>
<td>78.4</td>
<td>80.0</td>
<td>73.6</td>
<td>72.6</td>
<td>88.4</td>
</tr>
<tr>
<td>Val</td>
<td>46.9</td>
<td>48.0</td>
<td>39.9</td>
<td>40.7</td>
<td>66.5</td>
</tr>
<tr>
<td>Lys</td>
<td>37.4</td>
<td>35.9</td>
<td>19.0</td>
<td>17.9</td>
<td>75.5</td>
</tr>
<tr>
<td>Sulphur-amino acids</td>
<td>32.1</td>
<td>31.7</td>
<td>43.8</td>
<td>44.0</td>
<td>46.1</td>
</tr>
<tr>
<td>Aromatic amino acids</td>
<td>77.4</td>
<td>77.2</td>
<td>75.4</td>
<td>76.2</td>
<td>88.9</td>
</tr>
<tr>
<td>Thr</td>
<td>33.1</td>
<td>33.7</td>
<td>26.9</td>
<td>26.4</td>
<td>39.8</td>
</tr>
<tr>
<td>Trp</td>
<td>11.4</td>
<td>10.6</td>
<td>10.9</td>
<td>9.1</td>
<td>N/A</td>
</tr>
<tr>
<td>Branched amino acids</td>
<td>164.5</td>
<td>167.9</td>
<td>150.0</td>
<td>150.6</td>
<td>203.8</td>
</tr>
<tr>
<td>CML (µg/g protein)†</td>
<td>212</td>
<td>354</td>
<td>107</td>
<td>192</td>
<td>N/A</td>
</tr>
</tbody>
</table>

FLT, diet based on faba bean–wheat pasta dried at low temperature; FVHT, diet based on faba bean–wheat pasta dried at very high temperature; GLT, diet based on gluten–wheat pasta dried at low temperature; GVHT, diet based on gluten–wheat pasta dried at very high temperature; CD, casein diet based on casein and maize starch; N/A, not analysed.

* Amino acids were determined by Agrobio (Rennes) according to the EC Regulation 152/2009 (2009) method. The relative standard deviation of the method is lower than 3%.

† CML was determined by liquid chromatography–MS/MS according to the method developed by Niquet-Léridon & Tessier. The relative standard deviation of the method is lower than 5%.

### Table 4. Food intake, food and protein efficiency, rate of growth, body weight and body composition of rats fed treatment diets

<table>
<thead>
<tr>
<th>Diet</th>
<th>Food intake (g/d)*</th>
<th>Food efficiency (g gain/ intake)*</th>
<th>Protein efficiency (g gain/ g intake)*</th>
<th>Rate of growth (g/d)*</th>
<th>Body weight (g)*</th>
<th>FM (% body weight)†</th>
<th>LM (% body weight)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLT</td>
<td>28.7±</td>
<td>0.33a</td>
<td>2.3a</td>
<td>9.3±</td>
<td>327±</td>
<td>14.1±</td>
<td>78.7±</td>
</tr>
<tr>
<td>FVHT</td>
<td>29.1±</td>
<td>0.32a</td>
<td>2.3a</td>
<td>9.3±</td>
<td>328±</td>
<td>15.8±</td>
<td>77.2±</td>
</tr>
<tr>
<td>GLT</td>
<td>31.2±</td>
<td>0.18b</td>
<td>1.3b</td>
<td>5.4±</td>
<td>257b</td>
<td>14.2±</td>
<td>78.4±</td>
</tr>
<tr>
<td>GVHT</td>
<td>28.4±</td>
<td>0.17b</td>
<td>1.2b</td>
<td>4.6±</td>
<td>237b</td>
<td>11.2±</td>
<td>81.3±</td>
</tr>
<tr>
<td>CD</td>
<td>27.3±</td>
<td>0.33a</td>
<td>2.3a</td>
<td>8.9±</td>
<td>320a</td>
<td>9.8±</td>
<td>82.9±</td>
</tr>
<tr>
<td>Pooled SEM</td>
<td>3.1</td>
<td>0.02</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

FM, fat mass; LM, lean mass; FLT, diet based on faba bean–wheat pasta dried at low temperature; FVHT, diet based on faba bean–wheat pasta dried at very high temperature; GLT, diet based on gluten–wheat pasta dried at low temperature; GVHT, diet based on gluten–wheat pasta dried at very high temperature; CD, casein diet based on casein and maize starch.

* Mean values within a column with unlike superscript letters were significantly different (P<0.05).

† Food intake (g/d) was measured every week. Body weight was recorded at the beginning and the end of the experiment. Rate of growth (g/d) was calculated as the weight gain between the beginning and the end of the study ratioed to the number of study days. Food efficiency (g of weight gain/g food intake) was calculated using the equation:

\[
\text{Food efficiency} = \frac{\text{Rate of growth}}{\text{Food intake}}
\]

Protein efficiency (g of weight gain/g protein intake) was calculated using the equation:

\[
\text{Protein efficiency} = \frac{\text{Rate of growth}}{\text{Protein intake}}
\]

† LM and FM were measured by MRI using Echo MRI (Echo Medical Systems).

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and GLT. PAT and SAT were significantly higher in both FLT and GLT compared with rats fed CD.

There was no effect of drying temperature on any of the above parameters, other than a significant 21% lower PAT in rats fed GVHT compared with rats fed GLT.

### Tissue protein content

Table 6 reports the rat muscle and liver protein contents. There were no significant differences between rat groups in protein content of soleus muscle, tibialis muscle and liver \( (P = 0.6-0.9) \). A faba bean pasta diet tended to increase the protein content of plantaris muscle compared with the other diets but without significant effect \( (P = 0.07) \). There was no drying temperature effect on any of the muscle or liver protein contents.

### Protein quality evaluation of treatments diets

The higher FN content of rats fed faba bean pasta diet led to a lower ATTD compared with rats fed gluten pasta diet (Table 7). All pasta diets except GLT resulted in a lower (4–8%) ATTD compared with CD. Drying pasta at a VHT led to a significant decrease in the ATTD of rats fed gluten pasta diet (3%) compared with LT drying, but no impact on faba bean pasta diet. CFD (Table 7) did not vary according to protein composition of the pasta diet but was decreased by 5–6% in rats fed pasta diet compared with CD rats. Rats fed GVHT had 3% reduced CFD of protein compared with rats fed GLT, but this was not the case for rats fed FLT and FVHT.

In accordance with the UN results, the NPU and BV (Table 7) were significantly increased (1.5–1.8-fold) in rats fed FLT and FVHT compared with rats fed GLT and GVHT diets, without any effect of drying temperature. Rats fed CD conserved the highest NPU and the best BV. Interestingly, rats fed the FLT diet had a BV that was close to rats fed the CD.

### Metabolic parameters

Table 8 reports the recorded blood parameters. Regardless of the processing temperature, rats fed faba bean diet have 6–4% greater albumin and 11–7% greater total protein in the blood compared with rats fed gluten diet. There were no significant differences between rats fed faba bean pasta diet and rats fed CD in blood albumin and total protein concentrations, whereas rats fed gluten pasta diet had 7% lower blood albumin and 12–13% lower total blood proteins compared with rats fed CD. Blood glucose was greater in rats fed FLT or FVHT compared with rats fed CD. There was no significant difference in glucose between rats fed GLT and GVHT and rats fed CD.

---

**Table 5.** Muscle, liver and adipose tissues (peri-renal and sub-cutaneous) weights of rats fed treatment diets

<table>
<thead>
<tr>
<th>Diet</th>
<th>Plantaris (g)</th>
<th>Soleus (g)</th>
<th>Tibialis (g)</th>
<th>Liver (g/100 g body weight)</th>
<th>PAT (g/100 g body weight)</th>
<th>SAT (g/100 g body weight)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLT</td>
<td>0.59&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.29&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.05&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.00&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.12&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.38&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>FVHT</td>
<td>0.56&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.27&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.99&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.18&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.59&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>GLT</td>
<td>0.44&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.24&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.86&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.67&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1.07&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.49&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>GVHT</td>
<td>0.44&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.24&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.83&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.88&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.85&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.37&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>CD</td>
<td>0.58&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.31&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.07&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.21&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.74&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.98&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Pooled SEM</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.01&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

**Table 6.** Protein composition<sup>*</sup> of muscles (soleus, tibialis and plantaris) and liver of rats fed treatment diets

<table>
<thead>
<tr>
<th>Diet</th>
<th>Soleus (%)</th>
<th>Tibialis (%)</th>
<th>Plantaris (%)</th>
<th>Liver (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLT</td>
<td>14.7&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.8&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.7&lt;sup&gt;a&lt;/sup&gt;</td>
<td>18.8&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>FVHT</td>
<td>15.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>16.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>19.7&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>GLT</td>
<td>14.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>15.5&lt;sup&gt;b&lt;/sup&gt;</td>
<td>19.3&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>GVHT</td>
<td>14.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>15.5&lt;sup&gt;b&lt;/sup&gt;</td>
<td>19.3&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>CD</td>
<td>15.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>18.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>16.5&lt;sup&gt;b&lt;/sup&gt;</td>
<td>19.8&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Pooled SEM</td>
<td>1.0&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1.6&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.5&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.0&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a,b</sup> Mean values within a column with unlike superscript letters were significantly different. (P<0.05).
Table 7. Evaluation of protein quality of treatment diets
(Mean values with their standard errors, n 10 rats for all experiments)

<table>
<thead>
<tr>
<th>Diet</th>
<th>FN/NI (g/g)*</th>
<th>UN/NI (g/g)*</th>
<th>ATTD (%)*</th>
<th>CFD (%)*</th>
<th>NPU (%)*</th>
<th>BV (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLT</td>
<td>0.15a</td>
<td>0.01b</td>
<td>85.3c</td>
<td>98.3b</td>
<td>76.4b</td>
<td>77.9b</td>
</tr>
<tr>
<td>FVHT</td>
<td>0.15a</td>
<td>0.53b</td>
<td>85.1c</td>
<td>97.2b,c</td>
<td>72.3b</td>
<td>74.3b</td>
</tr>
<tr>
<td>GLT</td>
<td>0.09c</td>
<td>0.75a</td>
<td>91.1a</td>
<td>98.5b</td>
<td>43.7c</td>
<td>44.3c</td>
</tr>
<tr>
<td>GVHT</td>
<td>0.11b</td>
<td>0.68a</td>
<td>88.5d</td>
<td>95.6c</td>
<td>46.6d</td>
<td>48.7c</td>
</tr>
<tr>
<td>CD</td>
<td>0.08c</td>
<td>0.39a</td>
<td>92.0a</td>
<td>101.9a</td>
<td>83.8a</td>
<td>82.2a</td>
</tr>
<tr>
<td>Pooled</td>
<td>0.02</td>
<td>0.06</td>
<td>1.6</td>
<td>2.1</td>
<td>5.7</td>
<td>5.5</td>
</tr>
<tr>
<td>SEM</td>
<td>&lt;0.01</td>
<td>&lt;0.05</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

FN, faecal N; NI, N intake; UN, urinary N; ATTD, apparent total tract digestibility; CFD, corrected faecal digestibility; NPU, net protein utilisation; BV, biological value; FLT, diet based on faba bean; FVHT, diet based on faba bean–wheat dried at low temperature; GLT, diet based on gluten–wheat; GVHT, diet based on gluten–wheat dried at high temperature; CD, casein diet based on casein and maize starch; EBFN, endogenous and bacterial faecal N; EUN, endogenous urinary N.

a,b,c Mean values within a column with unlike superscript letters were significantly different (P<0.05).

* FN and NI were measured according to the Dumas method, in triplicate for each rat. ATTD, CFD, NPU and BV were calculated according to Proll et al. using the equations:

\[
\text{ATTD} \% = \frac{\text{NI} - \text{FN}}{\text{NI}} \times 100
\]

\[
\text{CFD} \% = \frac{\text{NI} - (\text{FN} + \text{EBFN})}{\text{NI}} \times 100
\]

\[
\text{NPU} \% = \frac{\text{NI} - (\text{FN} + \text{NI}) + \text{EBFN} + \text{EUN}}{\text{NI}} \times 100
\]

\[
\text{BV} \% = \frac{\text{NPU} \times \text{CD}}{\text{CFD}} \times 100
\]

The EBFN and the EUN of rats were determined using the FN and the UN obtained from rats fed protein-free diet, respectively.

Table 8. Blood concentration of metabolic markers of rats fed treatment diets
(Mean values with their standard errors, n 10 rats for all experiments)

<table>
<thead>
<tr>
<th>Diet</th>
<th>Albumin (g/l)*</th>
<th>Total protein (g/l)*</th>
<th>Glucose (g/l)*</th>
<th>Cholesterol (g/l)*</th>
<th>TAG (g/l)*</th>
<th>NEFA (mg/l)*</th>
<th>Free glycerol (mg/l)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLT</td>
<td>33.4a</td>
<td>56.4a</td>
<td>1.06a</td>
<td>0.74b</td>
<td>0.81a</td>
<td>276b</td>
<td>243a</td>
</tr>
<tr>
<td>FVHT</td>
<td>33.4a</td>
<td>56.1a</td>
<td>1.04a</td>
<td>0.74b</td>
<td>0.85a</td>
<td>299b</td>
<td>255b</td>
</tr>
<tr>
<td>GLT</td>
<td>31.4b</td>
<td>50.5b</td>
<td>0.99b</td>
<td>0.86b,c</td>
<td>0.43b</td>
<td>281b</td>
<td>243b</td>
</tr>
<tr>
<td>GVHT</td>
<td>31.3c</td>
<td>51.2c</td>
<td>0.92b</td>
<td>0.95a</td>
<td>0.47b</td>
<td>331a</td>
<td>252a</td>
</tr>
<tr>
<td>CD</td>
<td>33.7a</td>
<td>58.3a</td>
<td>0.89b</td>
<td>0.54c</td>
<td>0.70b,k</td>
<td>314b</td>
<td>236a</td>
</tr>
<tr>
<td>Pooled</td>
<td>0.8</td>
<td>1.9</td>
<td>0.08</td>
<td>0.12</td>
<td>0.19</td>
<td>38</td>
<td>29</td>
</tr>
<tr>
<td>SEM</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>NS</td>
<td>NS</td>
</tr>
</tbody>
</table>

FLT, diet based on faba bean–wheat pasta dried at low temperature; FVHT, diet based on faba bean–wheat pasta dried at very high temperature; GLT, diet based on gluten–wheat pasta dried at very high temperature; GVHT, diet based on gluten–wheat pasta dried at high temperature; CD, casein diet based on casein and maize starch.

a,b,c Mean values within a column with unlike superscript letters were significantly different (P<0.05).

* Blood glucose, NEFA, total cholesterol, TAG, albumins and total proteins were measured using a Konelab 20 chemistry analyser (Thermo Scientific).

Cholesterol concentration was 37–76% higher in all rats fed pasta diets compared with rats fed CD. Interestingly, cholesterol concentration in rats fed FVHT was 22% lower compared with rats fed GVHT. There was no significant difference in blood TAG between rats fed pasta diet and rats fed CD. Blood TAG content was lowered by 88 and 80% in GLT and GVHT, respectively, compared with FLT and FVHT. There was no significant diet-type effect (P>0.05) on blood NEFA and free glycerol. Drying temperature significantly decreased the blood NEFA in GVHT compared with GLT (online Supplementary Table S5). There was no drying temperature effect on any of the other parameters (albumin, total protein, glucose, cholesterol, TAG and free glycerol).

Plasma and inflammatory markers

The role of chronic inflammation in the degenerative diseases of modern society has stimulated research into the influence of diets on inflammatory markers. Dietary pattern was reported to be one of the main strategies to reduce inflammation. Legume consumption may contribute to reductions in C-reactive protein concentrations, a marker of systemic inflammation. Here, the inflammatory profiles resulting from legume, gluten or casein diets consumption were investigated. Blood fibrinogen (Fig. 1A) were not significantly different in all rats (P=0.3). There was no significant correlation between fibrinogen and metabolic markers, body weight and composition. Blood oosomocoid (Fig. 1B) was 57–75% higher in rats fed faba bean pasta compared with rats fed gluten pasta diet and CD. There was no difference in blood oosomocoid content between rats fed CD, GLT or GVHT diets. Oosomocoid content was significantly positively correlated with total body weight (r 0.55), FM (r 0.59), blood glucose (r 0.45) and blood TAG (r 0.50), and negatively correlated with LM (r -0.59). MCP-1 (Fig. 1C) was at equivalent levels between rats fed faba bean pasta diet and CD. Rats fed GLT and GVHT had approximately 15% lower MCP-1 content compared with rats fed CD and faba bean pasta diet. MCP-1 was significantly correlated with body...
weight (r 0·58), FM (r 0·30) and blood TAG (r 0·43), and negatively correlated with blood cholesterol (r −0·34). There was no drying temperature effect on fibrinogen, orosomucoid and MCP-1 composition.

Blood insulin (Fig. 1(D)) was higher in rats fed FLT compared with rats fed GLT. There was no drying temperature effect on blood insulin. Overall, faba bean–enriched pasta diet led to a blood insulin level equivalent to CD. Insulin was significantly positively correlated with body weight (r 0·63), FM (r 0·44), TAG (r 0·64), orosomucoid (r 0·50) and MCP-1 (r 0·28) and negatively correlated with LM (r −0·44). Adiponectin content was 29 and 41 % lower under FLT and FVHT diets compared with GLT and GVHT diets (fig. 1(E)). VHT drying led to a substantial 36 % increase in adiponectin content in rats fed gluten pasta diet. Adiponectin contents were not significantly different in rats fed FLT and CD. Adiponectin was significantly positively correlated with cholesterol (r 0·49) and negatively correlated with body weight (r −0·54). Blood leptin concentrations (Fig. 1(F)) were not significantly different in rats fed FLT and GLT but significantly higher (55 %) in rats fed FVHT than GLT rats. Blood leptin was higher in rats fed faba bean pasta diet compared with CD (38 % higher for FLT and 51 % higher for FVHT). Leptin concentration was not significantly different in rats fed FLT and GVHT and rats fed CD. Blood leptin was significantly positively correlated with body weight (r 0·54), FM (r 0·81), blood TAG (r 0·67), blood insulin (r 0·63), MCP-1 (r 0·30) and orosomucoid (r 0·61), and significantly negatively correlated with LM (r −0·80). There was no significant effect of drying temperature on leptin.

Discussion

The primary objective of this study in growing rats was to assess the impact of a diet based on mixed wheat–faba bean pasta dried at LT (55°C) or VHT (99°C), on key indicators of dietary protein quality, body composition, tissue protein content and blood metabolic and inflammatory markers. Diets were composed of either plant proteins, that is, the classical gluten–wheat pasta and mixed faba bean–wheat pasta, or animal proteins, that is, casein, all delivering the same food protein content (approximately 14%). Casein diets were considered as controls. The pasta used in this study (gluten–wheat pasta and faba bean–wheat pasta) were subjected to the same multiscale structure determination as in our previous work(11). The relationships between their protein structure and digestibility are discussed below.

Despite an equivalent food intake and food protein content, diets based on faba bean pasta (FLT and FVHT) resulted in a higher rate of growth compared with diets based on gluten pasta (GLT and GVHT). This could be related to the higher essential amino acid content in FLT and FVHT in comparison to GLT and GVHT diets. The food efficiency ratios of faba bean diets observed here (approximately 0·53) were higher than those reported for diets containing 20% of purified chickpea and lentil proteins (0·21)(36). Mixed wheat–legume meals could be an effective way to improve the low nutritional performance of diets with legume protein as the sole protein source and to achieve a similar protein efficiency as a casein control diet (CD).

Results of the present study support those of Torres et al.(13,14,20), where protein efficiency ratio in pasta enriched with 10% legume (germinated or fermented pigeon pea, or α-galactoside-free lupin flours) was 64–73 % higher than that in wheat pasta. In our study, we were able to obtain a 77 % increase in protein efficiency between FLT diet and GLT diet using a higher inclusion of legume flour (35 v. 10 %). The enhancement in protein efficiency ratio likely depends on legume type, enrichment level and preprocessing of the raw material.

ATTD was lower in rats fed faba bean than in rats fed gluten and casein diets. This was related to the higher faecal N loss in rats fed faba bean pasta diet compared with rats fed gluten and
casein diets. The presence of trypsin inhibitors in legume diets leads to the formation of the irreversible trypsin enzyme–trypsin inhibitor complex, causing a drop in trypsin in the intestine\(^ {37}\). Digestive enzymes are considered as an important source of endogenous N, which can increase total faecal N, thus decreasing ATTD. The amount of trypsin inhibitor in faba bean pasta (0–61 mg/g) measured in our previous work\(^ {39} \), even though substantially reduced once the pasta is processed and cooked compared with raw legume flour (7·8 mg/g faba bean flour)\(^ {38} \), likely caused a slight decrease in ATTD in the faba bean diets. The amount and type of fibres can also affect ATTD. Compared with gluten pasta diet, faba bean pasta diet provided an extra approximately 2·2 g/100 g diet of fibres (Table 2). The presence of fibres could play a role in increasing the excretion of endogenous N. Shah et al.\(^ {39} \) reported that the incorporation of fibres (such as cellulose, pectin, lignin and guar gum) in a 10% casein diet reduced protein digestibility in growing rats by increasing endogenous N excretion (digestive gland secretion, desquamated cells from active replacement of gastrointestinal mucosal lining and a small amount of plasma proteins). Therefore, the digestion of faba bean pasta diet components likely required a higher production of endogenous enzymes due to the high level of constitutive fibres compared with gluten pasta diet. The specific type of fibre brought by legume flour could also affect faecal N and ATTD. Faba bean flour and wheat semolina contain 2·36 %\(^ {38} \) and 1·54 % (K Laleg and V Micard, unpublished results) of arabinose, respectively. Faba bean flour is richer in galactose compared with wheat semolina (2·63 v. 0·69 %, dry basis). Galactose is a constituent of the arabinogalactans\(^ {38} \). Whereas wheat semolina is richer in xylose compared with faba bean (2·03 v. 0·74 %, dry basis). Xylose is a major component of arabinoxylans, the major polymers of wheat endosperm cell walls\(^ {40} \). Van Laere et al.\(^ {41} \) reported that arabinogalactans were generally fermented in vitro by twelve of the eighteen bacterial intestinal species studied, whereas arabinofuranoses were only fermented by two intestinal bacteria with the same bacterial profile. Here, a different activity of gut bacteria could partially explain the increase in bacterial N, contributing to a higher faecal N and a lower ATTD in faba bean pasta compared with gluten pasta diets.

Aside from endogenous protein secretion, modifications of the dietary and non-dietary protein occur at the hindgut of the animal. The human large intestine exhibits substantial proteolytic activity, partly attributable to colonic microbes. In vitro and in vivo studies showed that proteolysis occurs rapidly, resulting in the formation of protein fermentation metabolites such as SCFA, ammonia and various phenolic acids\(^ {42,43} \).

Here, total amount of dietary absorbed N (corrected faecal digestibility) was not significantly different in rats fed gluten and faba bean pasta diets. In all, 40% of the protein network structure in faba bean pasta was demonstrated to interact through weak bonds (mainly hydrogen, hydrophobic and electrostatic interactions), contrary to gluten pasta protein which is 90% linked by covalent bonds mainly disulphide. This structural difference resulted in a higher in vitro ileal protein hydrolysis (17·8 %) of faba bean pasta\(^ {11} \) – a difference not observed in vitro here. Among the few studies investigating the effect of enriching pasta with legumes on corrected faecal protein digestibility, only Torres et al.\(^ {13,14} \) reported an increase (6–13%) in faecal digestibility by substituting 10% of wheat semolina with pigeon pea flour. This enhancement in corrected faecal protein digestibility was most likely due to the pretreatment of legume flour (germination or fermentation) that could facilitate protein digestion.

Rats fed faba bean pasta diet showed lower urinary N compared with rats fed gluten pasta diet. At a similar protein intake, urinary N excretion may be affected by several parameters, chiefly essential amino acid composition\(^ {44} \). The essential amino acid composition of FLT and FVHT proteins was closer to the CD and to the rat needs determined by National Research Council for growing rats\(^ {45} \) compared with the GLT and GVHT proteins, notably for lysine, threonine, branched amino acids, histidine and tryptophan. This higher protein quality of faba bean pasta proteins could be responsible for the noticeable improvement in the efficiency of amino acid utilisation and in the BV of the diet compared with gluten pasta proteins that were poorer in some essential amino acids, for example, lysine.

In addition to essential amino acid composition, several studies reviewed by Jha & Berrocoso\(^ {46} \) have shown that the increase in dietary fibre can shift N from urinary to faecal excretion. The higher level of dietary fibre in FLT and FVHT diet could be another reason for the increase in faecal N excretion.

Rats fed faba bean pasta diet still present a higher urinary N and a lower protein utilisation compared with the rats fed CD. This could be related to the relatively lower quantity of essential amino acids in FLT and FVHT compared with the CD and to the increase in protein catabolism induced by certain bioactive compounds, notably trypsin inhibitors as demonstrated by Cenarruzabeitia et al.\(^ {47} \) in rats fed a field bean (V. faba) diet.

Alongside essential amino acids, another factor that affects urinary N excretion and subsequent protein utilisation is the rate of protein digestion/absorption. Amino acids from a faster digested legume (soya) are directed toward deamination pathways more than milk amino acids\(^ {44} \). This higher rate of the soya protein digestion/absorption process has been attributed to their faster intestinal transit compared with casein. In the acidic stomach conditions, legume (pea) protein aggregates are of smaller particle size (80 μm) than casein (2 mm) and showed faster intestinal bioavailability than casein\(^ {48} \). These data suggest that proteins in our faba bean pasta diet could be digested more rapidly than casein. A rapid absorption of amino acids is associated with both enhanced amino acid catabolism and transiently higher protein synthesis\(^ {24,49} \). The fact that faba bean protein acts as a rapidly digested protein could be beneficial in specific subpopulations such as older people. Muscle protein synthesis is altered in older people, mainly due to a decrease in the specific anabolic response to amino acids. It has been suggested that protein synthesis in older people could be resistant to amino acid availability, and that high aminoacidemia is needed to improve postprandial protein gain\(^ {50,51} \). Hence, a fast digestive protein, like whey protein, was more efficient than an isonitrogenous casein protein in increasing leucine balance and protein utilisation in old men\(^ {52} \). This improvement in protein gain was related to potentially higher hyperaminoacidaemia induced by rapid digestion of whey protein in comparison to casein. Thus, with respect to its potentially faster kinetics of digestion, faba bean-enriched pasta
could be a promising way to increase aminoauxaemia and subsequent postprandial protein retention in older subjects.

Rats fed FLT and FVHT showed a higher level of global growth parameters (rate of growth, liver and muscle weights) compared with GLT and GVHT, which was related to the higher protein retention with FLT and FVHT diets. Branched amino acids were also reported to be able to dose-dependently stimulate muscle protein synthesis in growing rats. The higher (9–10%) branched chain amino acid content in faba bean pasta diet compared with gluten pasta diet could be involved in higher muscle protein synthesis in rats fed faba bean. Compared with the CD, the faba bean pasta diet led to a slightly lower LM and liver weight (ratioed to body weight) associated with the relatively lower protein utilisation and branched amino acids. Rate of animal growth between these two groups was nevertheless equivalent because of the higher FM in faba bean group rats compared with the CD group rats. Huang et al. also reported a 10–13% decrease in liver weight in rats fed 18–20% gluten or soya proteins compared with rats fed casein. Alonso et al. reported that in addition to the liver, both gastrocnemius and tibialis anterior muscle weight and protein content were also reduced after consumption of a approximately 13% protein diet based on raw or extruded pea compared with a casein diet. This was linked to the amino acid deficiencies in the diet and to the action of anti-nutritional factors. We did not observe any effect of faba bean pasta diet consumption on muscle weight or protein content, likely due to a better balanced amino acid composition in the mixed protein source food (compared with the legume alone) and to the reduction of certain bioactive compounds (trypsin inhibitors, α-galactosides) by pasta processing and cooking steps.

Fasting blood albumin and total protein were observed to be lower with GLT and VHT than with FLT, FVHT and CD. Albumin is the major secretory protein synthesised by the liver. Hypoalbuminaemia is the result of both inadequate protein and energy intake. In our study, feeding GLT and GVHT did not increase blood inflammatory markers (orosomucoid, fibrinogen and MCP-1). As all diets were isoenergetic, the quality of dietary protein was probably involved in decreasing the plasma albumin. Indeed, the removal of certain essential amino acids (leucine, isoleucine and tryptophan) from a rat liver culture medium induced a decrease in total protein and albumin liver synthesis. Thus, the dietary protein and its amino acid composition were likely at the origin of this low plasma albumin in the GLT and GVHT groups.

Eating pasta diets (except GVHT) increased FM compared with eating the CD. Glucose was also slightly increased under faba bean pasta diet, and tended to be higher under gluten pasta diet than under the CD diet (P = 0.07). This slightly higher glucose content in rats eating pasta diets could stem from the different structures of starch within pasta and CD. The pasta was cooked until total starch gelatinisation, whereas the maize starch used to prepare the CD was in its native ungelatinised form. Highly gelatinised starch is more available to the digestive enzymes, causing an increase in the rate of starch digestion. Consequently higher plasma glucose in rats eating pasta diets probably led to higher adipose tissue deposition through de novo lipogenesis. Blood leptin was positively correlated with body FM, adipose tissue weight and blood TAG content. Leptin contributes to the regulation of body weight by modulating appetite and energy expenditure. It stimulates fatty acid oxidation and prevents lipid accumulation in non-adipose tissues. Obese humans showed higher leptin levels, suggesting a resistance to its action. The increase in leptin content observed here could be related to the adipose tissue, as demonstrated in mice by Frederich et al. This increase in leptin and blood glucose was more pronounced after feeding faba bean pasta than when fed gluten pasta diet, and it was also accompanied by a higher blood insulin and orosomucoid inflammatory marker levels in rats receiving faba bean pasta than those receiving gluten pasta diet. Orosomucoids are acute-phase plasma proteins synthesised by the liver. Inflammatory disorders can increase the concentration of these compounds. The increase in orosomucoids could indicate that FLT and FVHT consumption could have induced an inflammatory response. Although faba bean pasta diets contained higher fibres shown to decrease inflammation, and a more adequate essential amino acids profile for rat growth compared with gluten pasta (based on the National Research Council requirement), it still presents limiting sulphur amino acids that could be responsible for the inflammatory disorders observed in rats fed faba bean.

The second objective of our study was to evaluate whether the use of VHT drying during pasta processing affects protein digestibility and retention and the subsequent growth, nutritional and metabolic parameters. Among all the end points determined, VHT drying only increased faecal N and decreased faecal protein digestibility in rats fed gluten pasta diet. This difference in the digestive behaviour of GVHT could not be attributed to the amount of covalent interactions or to the conformation of the pasta protein, as we previously showed that proteins in gluten pasta dried at LT and VHT had the same amount of covalently linked protein and the same secondary conformation. The low faecal protein digestibility of GVHT could be attributed to some Maillard products. Indeed, Maillard reactions occur during high-temperature treatment of wheat pasta and were shown to decrease the in vitro ileal release of amino acids. In addition, in line with our results, the consumption of Maillard-rich products was shown to increase faecal N excretion in humans and decrease apparent and true protein digestibility without effect on protein utilisation or serum biochemical variables. Here, among Maillard products, CML was higher in all VHT pasta diets, and its increase was greater with gluten pasta (79%) than with faba bean pasta (67%) diets, which could explain the lower effect of VHT drying on faba bean pasta diet digestibility.

In conclusion, this study clearly demonstrated that consumption of mixed wheat–legume pasta diet leads to improved faecal protein digestibility and protein utilisation, muscle weight, blood nutritional markers and rat growth compared with the classical isoprotein gluten–wheat pasta diet consumed alone in growing rats. Combining wheat with legume as a complementary plant source led to a more adequate essential amino acid profile for growth processes that likely explains the improved nutritional and growth parameters. The weakening in the force of protein–protein interactions induced by enriching wheat pasta with faba beans did not affect corrected faecal protein digestibility. We also
showed that the high drying temperature could decrease amino acid bioavailability through structural protein modifications induced by the formation of certain Maillard products.

Mixed wheat–legume pasta diet led to much closer nutritional and growth parameters as a high-nutritional-value reference protein, that is, casein. In addition, the proteins in faba bean-enriched pasta diet likely acted as relatively rapidly digested protein in the digestive tract compared with casein protein. This physicochemical behaviour of legume pasta protein could be involved in modulating the rate of protein digestion, affecting the whole-body protein metabolism. This characteristic would be beneficial to promote protein deposition in settings marked by an age-related loss of muscle.

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S. W., V. M., K. L., J. S., Y. B. conceived and designed the experiments. K. L., A. B., J. S., V. P., P. D., C. G., S. W. and V. M. performed the experiments. F. J. T., A. G. and M. H. were involved in the analysis of CML. K. L., V. M. and S. W. analysed the data and performed statistical analysis. K. L. wrote the paper. S. W. acquired the funding.

The authors declare that there are no conflicts of interest.

Supplementary material

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