

	,
Title	Enhancing the nutritional profile of regular wheat bread while maintaining technological quality and adequate sensory attributes
Authors	Hoehnel, Andrea;Bez, Jürgen;Petersen, Iben L.;Amarowicz, Ryszard;Juśkiewicz, Jerzy;Arendt, Elke K.;Zannini, Emanuele
Publication date	2020-05-17
Original Citation	Hoehnel, A., Bez, J., Petersen, I. L., Amarowicz, R., Juśkiewicz, J., Arendt, E. K. and Zannini, E. (2020) 'Enhancing the nutritional profile of regular wheat bread while maintaining technological quality and adequate sensory attributes', Food and Function, 11(5), pp. 4732-4751. doi: 10.1039/d0fo00671h
Type of publication	Article (peer-reviewed)
Link to publisher's version	10.1039/d0fo00671h
Rights	© 2020, the Authors. Publication rights licensed to the Royal Society of Chemistry. All rights reserved.
Download date	2025-07-31 06:06:45
Item downloaded from	https://hdl.handle.net/10468/10960



Journal Name



ARTICLE TYPE

Cite this: DOI: 00.0000/xxxxxxxxxx

Enhancing the nutritional profile of regular wheat bread while maintaining technological quality and adequate sensory attributes†

Andrea Hoehnel,^a Jürgen Bez,^b Iben Lykke Petersen,^c Ryszard Amarowicz,^d Jerzy Juśkiewicz,^d Elke K. Arendt, *a,e and Emanuele Zannini ^a

Received Date Accepted Date

DOI: 00.0000/xxxxxxxxxx

Plant proteins, and legume proteins in particular, have become the centre of attention moving towards a more sustainable and, therefore, more plant-based human diet. Especially hybrid products, containing wheat and legume proteins, promise a balanced amino acid composition and an upgraded nutritional value of both protein sources. This study investigates a high-protein hybrid bread (HPHB) formulation, where wheat flour was partially replaced by high-protein ingredients from faba bean, carob and gluten. In addition to a detailed characterisation of technological quality and sensory profile, also the formulation's nutritional value was examined in comparison to regular wheat bread. Therefore, macronutrient composition, antioxidant potential, amino acid profile and contents of antinutritional compounds were analysed. Furthermore, protein digestibility was determined in an in vitro model and in vivo. Dough analysis revealed significant differences of the HPHB formulation compared to regular wheat dough. However, results obtained for bread quality characteristics prove HPHB to be equal to regular wheat bread and sensory results and the determined sensory attributes suggest high consumer acceptance. Nutritional analyses of HPHB showed a more favourable macronutrient composition in comparison to regular wheat bread; as well as low contents of antinutritional compounds and high antioxidant potential linked to high levels of phenolics. Also an improved amino acid profile, increased nitrogen utilisation rate (by 69 %) and higher protein efficiency ratio were determined, which are associated with enhanced protein quality. This suggests HPHB, and similar formulations of its kind, as a valuable and healthy food choice, which can contribute to adequate protein supply in predominantly plant-based diets.

2 1 Introduction

Protein from plant sources, next to other trends like digestive health and good carbs/bad carbs, is currently one of the most pop-

- sular and important trends in the food sector. One of the reasons
- 6 for that is an increasing awareness amongst consumers, author-
- 7 ities and industry of the need to create a more sustainable food
- 8 system considering planetary boundaries. ^{2,3} According to many
- ^a University College Cork, School of Food and Nutritional Sciences, College Road, Ireland. Tel: +353 21 490 2064; E-mail: e.arendt@ucc.ie

9 recent reports, this requires a shift to a predominantly plant-based 10 human diet. 2,4 Since we are also facing a growing world popu-11 lation, with a prospect of about 10 billion by 2050, 2 research 12 plays a key role in finding ways to provide high-quality protein 13 from plant sources to cover future protein needs. Even though 14 it is known that current protein consumption exceeds the aver-15 age daily requirement in many parts of the world, this is usually 16 linked to high intakes of animal protein and necessary changes 17 in the food system and human diet are likely to pose a challenge 18 to sufficient protein supply in the future. ^{2,4} In many cases, the overconsumption of protein is associated with a general overconsumption of food and energy intakes exceeding recommended levels⁴ and does not reflect an overconsumption of protein relative to other macronutrients. Furthermore, recommendations for daily protein intakes are based on high-quality protein. When 24 large amounts of protein of lower quality are consumed, intakes might need to be increased in order to meet the body's amino acid requirements.⁵ Apart from sustainability considerations, dietary

^b Fraunhofer Institute for Process Engineering and Packaging, 85354 Freising, Germany.

^c Department of Food Science, University of Copenhagen, Rolighedsvej 26, 1958 Frederiksberg C., Denmark.

^d Institute of Animal Reproduction and Food Research of the Polish Academy of Sciences, Tuwima, St. 10, 10-748 Olsztyn, Poland.

^e APC Microbiome Ireland, Cork, Ireland.

[†] Electronic Supplementary Information (ESI) available: Microbiological shelf life and water activity of reference wheat bread (RWB) and high-protein hybrid bread (HPHB). See DOI: 00.0000/00000000.

reported for a large number of countries and associated with increased health risks. ⁷ Research concerning new alternative plant 91 protein sources is mostly focused on legumes. Due to their ability to grow in a variety of different climates and to fix nitrogen in the soil, they are particularly promising for a local crop cultivation, a considerably reduced use of fertilisers and a food production with a lower carbon and water footprint. 3,8,9 Legumes have little lysine and higher amounts of SAAs. 12,13 Efforts have 101 tein needs of future predominantly plant-based diets. been made to combine both protein sources in "hybrid products" containing cereals and legumes and especially wheat bread has $_{102}$ proven a suitable cereal matrix for the incorporation of legume protein ingredients. ¹⁴ Ideal bread should have a lower glycaemic ₁₀₃ **2.1** Materials index than regular white bread, be an important source of proteins, and contain tolerated dietary fibre, vitamins, magnesium, 104 Three high-protein ingredients (HPIs) were applied in the high-

recommendations advice a reduction of animal protein intake 83 profile with a higher protein content and higher protein quality in in favour of increased plant protein consumption for a healthy 84 particular. Therefore, plant-based high-protein ingredients (HPIs) diet. Many reported adverse effects of high protein intake are 85 from faba beans, carob and wheat, selected based on findings by largely related to proteins from animal sources and the co-intake 66 Hoehnel et al. 30, were incorporated in a regular wheat bread. of sodium, nitrate, nitrite and saturated fatty acids when red meat 87 This high-protein hybrid bread (HPHB) formulation was evaluor dairy products are consumed. ^{2,4,6} Also an overconsumption of 88 ated regarding technological, nutritional and sensory characterisfood carbohydrates, especially refined carbohydrates, has been 89 tics using regular wheat bread as a reference (RWB). The HPHB 90 formulation, containing a dry-processed faba bean HPI as its main source of non-wheat protein, also promises improved sustain-⁹² ability; ³¹ especially when compared to other high-protein bread 93 formulations that are commercially available. These often con-94 tain dairy ingredients as non-wheat protein source. Vogelsang-95 O'Dwyer et al. 32 reported a life cycle assessment (LCA) of the 96 dry-processed faba bean HPI used in this study, which confirmed are naturally rich in protein, which contains high amounts of 97 reduced use of land and water resources as well as lower impact the essential amino acid (AA) lysine but lacks sulphur-containing 98 on climate change (carbon footprint) and aquatic eutrophication amino acids (SAAs).^{8,10,11} This makes legumes particularly inter- 99 in comparison to cow's milk powder. This makes HPHB and foresting for the complementation of cereal based diets, since cereals 100 mulations of its kind even more promising to partially cover pro-

Materials and Methods

trace elements and antioxidants. 15,16 Jenkins et al. 7 state that, 105 protein hybrid bread (HPHB) formulation. Faba bean flour (proin the context of decreased physical activity in our population, 106 tein content 61.25 %DM, fat 3.81 %DM, ash 5.43 %DM, fifoods should possess nutritional density rather than nutrient den- 107 bre 0.35 %DM, carbohydrates by difference 29.17 %DM, tosity. This means that the intake of essential nutrients (macro and 108 tal starch 7.77 %DM; 30 obtained by dry fractionation), which micro) per calorie will need to increase in order to meet require- 109 was experimentally produced and provided by Fraunhofer Instiments at lower caloric intake levels. Legumes are rich in micronu- 110 tute IVV, Freising, Germany; carob germ flour (protein content trients and compounds with antioxidant activity, which could help 111 55.04 %DM, fat 0.20 %DM, ash 7.04 %DM, fibre 17.67 %DM, carto enhance the nutritional value of wheat bread. ^{14,17,18} Also a 112 bohydrates by difference 20.05 %DM, total starch < 0.2 %DM; ³⁰ lowered glycaemic load, increased protein content and improved 113 GRINDSTED VEG PRO S1) from Danisco, UK and vital gluten protein quality could be achieved by the fortification of wheat 114 (protein content 72.38 %DM, fat 0.72 %DM, ash 0.87 %DM, fibre bread with legume proteins. Numerous research articles have in- 115 < 0.1 %DM, carbohydrates by difference 15.31 %DM, total starch vestigated the effects of legume ingredients, from faba bean (Vi- 116 4.95 %DM; 30 NUTRALYS W) from Roquette, France. Wheat flour cia faba) and carob (Ceratonia siliqua) seeds in particular, on both 117 was supplied by Whitworth Bros Ltd, UK; dry yeast by Puratos, the technological as well as nutritional quality of breads. 14,19-23 118 Belgium; salt by Glacia British Salt Ltd, UK; sugar (granulated However, many of these publications report inferior technologi- 119 Irish sugar) by Nordzucker (Ireland) Ltd, Ireland; psyllium (VITAcal and sensory characteristics in favour of increased nutritional 120 CEL P95) by J. Rettenmaier & Söhne, Germany; vegetable oil by quality. Additionally, there are concerns regarding antinutritional 121 Musgrave, Ireland; and xylanase (Biobake 715) by Kerry Group, compounds (ANCs) originating from legumes such as trypsin in- 122 Ireland. For in vivo digestibility trials, the following ingredients hibitors, tannins, lectins and the pyrimidine glycosides vicine and 123 were used for the preparation of diets: casein (C) from Lacpol convicine. Trypsin inhibitors, which can negatively impact pro- 124 Co., Poland; soya protein isolate (SPI) ISOPRO 900 HI charactein digestibility, are present in many plants but are particularly 125 terised as non-GMO protein isolate from EDMIR-POL Co., Poland; important in legumes. 24,25 Vicine and convicine are mainly found $_{126}$ soya flour (SF) SOPRO TB 200 from EDMIR-POL Co., Poland; α in faba beans and can trigger adverse physical conditions like fav- 127 cellulose (C8002) from Sigma-Aldrich, Missouri, USA; soya oil ism. 26-28 This leads to a low popularity of legume ingredients 128 from ZPT Co., Poland; choline chloride from SIGMA, Poland; and cereal/legume hybrid products. 29 Next to an enhanced nutri- 129 cholesterol from PPH Standard Co., Poland; sucrose from POCH tional value, adequate technological quality and sensory proper- 130 SA Co., Poland; and corn starch from Avebe, The Netherlands. ties are essential for a high consumer acceptance of such products 131 Enzymes for in vitro digestion trials were purchased from Sigmaand for an acceleration of the protein transition in our diet. This 132 Aldrich, Missouri, USA: Pepsin from porcine gastric mucosa; EC is why this study proposes and fully characterises a new bread 133 3.4.23.1; P7000; 727 U/mg and pancreatin from porcine panformulation, which was designed to match the technological qual- 134 creas; 4 x USP; P1750. All other chemicals were also purchased 82 ity of regular wheat bread, but promises an improved nutritional 195 from Sigma-Aldrich, Missouri, USA unless stated otherwise.

Technological Analysis 2.2

2.2.1 Flour Analysis

The properties of wheat flour (used for reference wheat bread) and the high-protein (HP) flour mix (used for HPHB) were analysed. The HP flour mix contained wheat flour, the three HPIs (faba bean flour, carob germ flour, gluten) and psyllium in ra- 196 2.2.3 Dough Analysis tios according to HPHB formulation (Table 1). The moisture content of the HP flour mix was calculated considering the moisture determined for each single ingredient. GlutoPeak test -Gluten-aggregation properties of wheat flour and the HP flour mix were investigated following the method previously described by Hoehnel et al. 30 using the GlutoPeak device (Brabender GmbH and Co KG, Duisburg, Germany). In brief, high shear was applied to a flour/water slurry (50:50 ratio, adjusted when moisture of flour differed from 14 %). The device was operated at a paddle speed of 2750 rpm and temperature of 36 °C; torque was recorded over time. Variables Torque maximum (TM, expressed in Brabender units BU) and Peak Maximum Time (PMT, expressed in s) were obtained from the curves. Rapid visco analysis - Examination of pasting behaviour using Rapid Visco Analysis (RVA Super 3, Newport Scientific, Warriewood, Australia) was performed according to AACC 76-21.02. The following heating profile was applied: equilibration at 50 °C for 1 min, heating to 95 °C at 0.2 °C/s, holding at 95 °C for 162 s, cooling to 50 °C at 0.2 °C/s, maintaining at 50 °C for 120 s. The variables peak viscosity (PV), setback and final viscosity (FV) were obtained from the viscograms.

2.2.2 Recipe Adaptation and Bread Production

Bread samples were prepared according to the formulations in Table 1. The HPHB formulation contains HPIs (faba bean flour, 220 2.2.4 Bread Quality Analysis carob germ flour, gluten) and was designed to match the technological quality of the reference wheat bread (RWB). A series of preliminary trials (data not shown) based on the results presented by Hoehnel et al. 30 led to the establishment of the HPHB formulation. A total of 28 different recipes were screened to select a combination of HPIs and to optimise their relative ratios for favourable technological characteristics. Furthermore, the introduction and optimal addition levels of the functional ingredients psyllium, sugar and xylanase were investigated as part of the screening to achieve adequate dough handling characteristics and quality of the end product. For both formulations, the straight dough method was applied. Yeast was activated by dissolving in 30 °C tap water for 10 min. The obtained yeast suspension was added to the remaining, previously weighed ingredients. A total amount of 3600 g dough was prepared. Mixing conditions were the following: RWB - MACPAN MX 10 spiral mixer (MACPAN SNC, Italy) at speed 1 for 6.5 min and speed 2 for 5 min; HPHB -Hobart A200N mixer (Hobart Manufacturing, UK), equipped with hook attachment, at speed 1 for 2 min and speed 2 for 7.5 min. After covering the dough and leaving it to rest for 5 min, it was divided into 7 pieces of 450 g \pm 1 g. The pieces where moulded, put into baking tins and proofed for 90 min at 75 % humidity and 241 Bread crumb was separated from crust, cut into small cubes,

draft throughout the whole baking process. The baking chamber was steamed with 400 mL prior to loading. After baking, breads were removed from tins and left to cool down for 2 h at ambient temperature. The results were obtained from three independently performed baking trials.

197 Doughs for determination of dough properties were prepared as 198 described in section 2.2.2. Rheofermentometer - Formation and 199 retention of gas in the fermenting doughs was analysed using a 200 Rheofermentometer F3 (Chopin, France). A dough piece (300 g) 201 was placed into the sample container and a weight constraint of 202 1.5 kg was applied. The dough fermentation was monitored for 203 3 h at a temperature of 35 °C (matching the proofing temperature 204 used during bread production). The fermentation performance 205 of the doughs was evaluated by the following variables obtained 206 from the generated curves: Total gas volume produced (V_{total}) , 207 volume of CO₂ lost (V_{lost}) and volume of gas retained (V_{ret}) from 208 gaseous release curves; and maximum height of dough devel-209 opment (H_M) from dough development curves. Large deforma-210 tion properties - Extensibility (expressed in mm) and resistance 211 to extension (expressed in g) of the doughs were measured by a 212 texture analyser (TA-XT plus, Stable Micro Systems, Surrey, UK) 213 equipped with a 5 kg load cell and a Kieffer dough and gluten 214 extensibility rig (test settings: pre-test speed 2 mm/s, test speed 215 3.3 mm/s, post-test speed 10.0 mm/s, trigger force 5 g). The uni-216 axial extension test was performed after a dough resting time of 217 20 min (room temperature) inside the dough strip mould. Ten 218 intact strips of dough were measured from each of three batches 219 per formulation.

221 Specific volume (SV) was measured with a Volscan Profiler (Sta-222 ble Micro Systems, Surrey, UK) of 6 loaves per batch. For analysis 223 of crumb structure and hardness, three slices (20 mm) were cut out of the middle of each of 2 bread loaves. A C-Cell Imaging Sys-225 tem (Calibre Control International Ltd, UK) was used to capture 226 images of slices and to determine the variables: number of cells, 227 area of cells and slice brightness. Crumb hardness was analysed 228 with a texture analyser (TA-XT2i, Stable Micro Systems, Surrey, 229 UK) equipped with a 25 kg load cell. A 35 mm cylindrical probe 230 was used to compress the centre of the slice to 40 % of its height 231 as part of a texture profile analysis (TPA): test speed 5 mm/s, 232 post-test speed 10 mm/s, trigger force 0.05 N, waiting time be-233 tween compressions 5 s. TPA of bread slices was repeated on day 234 2 and day 5 after baking to monitor bread staling (whole loaves 235 were stored in plastic bags at ambient temperature in the bak-236 ery and sliced immediately before the measurement). Lightness 237 of crust (L*crust) and crumb (L*crumb) was measured by a Col-238 orimeter CR-400 (Konica Minolta, Japan) using the CIE L*a*b* 239 colour space.

240 2.2.5 Scanning Electron Microscopy

35 °C (KOMA BV Sunriser, Reormond, the Netherlands). Baking 242 frozen at -80 °C and freeze-dried. The dry crumb was further was performed in deck ovens (MIWE Condo, Arnstein, Germany) 243 crushed into small fragments which were mounted onto plain at 220/230 °C top/bottom temperature for 35 min with open 244 aluminium stubs with double-sided carbon adhesive tape. After

Table 1 Recipe for RWB and HPHB

	Referen	ce wheat bread	High-protei	in hybrid bread
Ingredient	% based on flour	% based on recipe	% based on flour	% based on recipe
Wheat flour	100.0	59.70	82.5×	47.22×
Faba bean flour	-	-	$10.0^{ imes}$	$5.72^{ imes}$
Carob germ flour	-	-	5.0×	2.86^{\times}
Gluten	-	-	$2.5^{ imes}$	1.43×
Psyllium	-	-	$2.0^{ imes}$	1.14^{\times}
Sugar	-	-	1.0	0.57
Baker's yeast	2.0	1.19	2.0	1.14
NaCl	2.0	1.19	2.0	1.14
Oil	1.0	0.60	1.0	0.57
Xylanase	-	-	0.0060	0.0034
Water	62.5	37.31	66.70	38.18
Total	167.5	100.00	174.7	100.00

[×] Ingredients are included in HP flour mix

coating with a 5 nm gold-palladium (80:20) layer using a Gold 283 for 10 min and the supernatant transferred to another test tube ating voltage of 5 kV.

2.3 Nutritional Analysis

Analysis of nutritional characteristics of the bread formulations was performed on freeze-dried (according to the procedure described in section 2.2.5) and subsequently milled (laboratory disc mill; Bühler, Brauchschweig, Germany) samples of bread crumb. Results are expressed as contents in fresh bread considering the moisture of freeze-dried and fresh bread crumb unless stated otherwise.

2.3.1 Compositional Analysis

The analysis of the following compositional data was performed 300 2.3.2 Amino Acid Analysis by Concept Life Science Ltd., UK based on the indicated validated Determination of protein amino acid composition was carried out AOAC 1977.992.15; nitrogen-to-protein conversion factor 6.25), ash (removal of organic matter by oxidation at 550 °C, based on nance (NMR), based on MQC-23-35 Oxford Instruments application note), fatty acid profile (GC-FID of fatty acid methyl esters; triglyceride conversion factor 0.956), total dietary fibre (TDF) 75 °amplitude. Hereupon, the sample was centrifuged at 1800 g 321 protein digestibility.

Sputter Coater (BIO-RAD Polaron Division, SEMcoating system, 284 for further processing. Sonication and centrifugation were re-England), they were examined under high vacuum with a JOEL 285 peated (extraction step 2) after adding another 15 mL 80% EtOH scanning electron microscope (SEM) type 5510 (JOEL Technics 286 (at 55 \pm 5 °C) to the pellet. The supernatants of both extraction Ltd., Tokyo, Japan). Images were acquired at a constant acceler- 287 steps were pooled and concentrated using a vacuum centrifuge system (Scanvac Scan Speed 32 with Scanvac VacSafe 15, Labogene ApS, Lynge, Denmark) with the following settings: 2 h at 1500 rpm and 45 °C, followed by 1 h at 2000 rpm and 50 °C; average pressure 15 mbar. The concentrated extract was trans-292 ferred into a 10 mL volumetric flask, which was filled up with 293 ultrapure water containing 50 mg/L NaN3, and filtered through 294 syringe driven polyamide filters (Chro-mafil AO-20/25, pore size: 295 0.20 μm, Machery-Nagel, Düren, Germany). Samples were extracted in duplicates and quantification of the sugars was per-297 formed according to the method described by Ispiryan et al. 33 using a Dionex ICS-5000⁺ system (Sunnyvale, CA) equipped with 299 an electrochemical detector.

methods: energy (calculated considering protein, fat, available 302 by Mérieux NutriSciences CHELAB S.r.l., Italy based on ionic chrocarbohydrates and fibre), protein (Dumas method, modified after 303 matography with postcolumn ninhydrin derivatisation (fluores-304 cence detection; UV detection for tryptophan) after adequate ex-305 traction and protein hydrolysis (separate hydrolysis procedures ISO 936:1998), fat (low resolution proton nuclear magnetic reso- 306 for the determination of tryptophan, sulphur-containing AA and 307 remaining AA).

308 2.3.3 In vitro Protein Digestion

(gravimetric method, based on AOAC 991.43), sodium (flame 309 A previously described static multi-step method for in vitro photometry after removal of organic matter). Moisture was de- 310 protein digestibility (IVPD) 34,35 was used to simulate gastrotermined by air-oven method at 130 °C until constant mass was 311 pancreatic protein digestion. In short, sample amounts containing reached. Total starch content was analysed using the enzyme kit $_{312}$ 50 ± 1 mg protein were weighed in and enzymatic hydrolysis was K-TSTA supplied by Megazyme, Ireland. Mono-, di- and oligosac- 313 started: pepsin digestion at 37 °C and pH 1-2 (1 h) followed by charides were extracted from the freeze-dried product powders 314 sequential pancreatin digestion at 37 °C and pH 7-8 (short-term: as follows: 15 mL of 80/20 (v/v) ethanol/ultrapure water (80% 315 +1 h; medium-term: +3 h; long-term: +24 h). Ratios between EtOH), which was heated to 55 \pm 5 °C, were added to 2 g of sam- 316 enzyme and substrate (w/w) were kept constant at 1:50 (pepsin ple. The mixture was vortexed until the powder was suspended 317 stage) and 1:10 (pancreatin stages). IVPD in % was determined and then subjected to sonication (extraction step 1) utilising a 318 using a trinitrobenzenesulfonic acid (TNBS) assay. Results are ex-BANDELIN Sonoplus HD 3100 homogenizer (Berlin, Germany) 319 pressed as the concentration of free α -amino groups in samples equipped with an MS73 microtip, operated twice for 15 s at 320 in relation to an alanine standard solution representing 100 %

2.3.4 In vivo Nitrogen Balance

The animal protocol used in this study was approved by the local institutional Animal Care and Use Committee (Olsztyn, Poland) and the study was performed in accordance with EU Directive 2010/63/EU for animal experiments. The assessment was conducted on growing male Wistar rats weighing 173.2 g. The rats were randomly divided into groups of seven animals. All animals were housed individually over 14 days in metabolic cages with free access to water and the experimental diets (Table 2). The selection of the animals and their maintenance over the 14day experiment followed common regulations. The environment was controlled with a 12 h light-dark cycle, a temperature of 22±1 °C, relative humidity of 45-65% and 20 air changes per hour. For experimental feeding the following diets were used: a standard control diet based on casein (C) as the main protein source (supplemented with 0.2% DL-methionine), a second control diet based on soya protein isolate (SPI, without any supplementation), a third control diet based on soya flour (SF, without any supplementation) and the experimental diets containing RWB and HPHB. All experimental diets were a modification of the AIN-93G diet for laboratory rodents recommended by the American Institute of Nutrition; 36 the dietary protein level was lowered to approx. 11% to measure the protein digestibility and utilisation rate. During the study, nitrogen (N) digestibility and utilisation tests (balance tests) were carried out. After a 9-day preliminary period, faeces and urine were thoroughly collected for 5 d from all rats (kept in balance cages; Tecniplast Spa, Buguggiate, Italy). 385 Extraction - Phenolic compounds were extracted from the prodeach animal separately (n = 7 per diet group).

2.3.5 Antinutritional Compounds

Trypsin inhibitors were extracted from the lyophilised product powders by adding 2.5 mL sodium acetate buffer (0.1 M, pH4.9) to 350 mg sample and homogenising the mixture for 2 min using an Ultra Turrax. After centrifugation for 5 min at 3000 g (EBA 12 Centrifuge; Hettich Zentrifugen, Tuttlingen, DE), the supernatant was transferred to a new test tube and the extraction procedure was repeated with the same conditions with the pellet. Both supernatants were pooled, stored in the fridge overnight and centrifuged again (5 min, 3000 g) immediately before trypsin inhibitor activity (TIA) analysis. TIA was determined following the method described by Joehnke et al. 34 with some modifications. In brief, TIA levels were measured against a trypsin solution (stock concentration 0.1 mg/mL). A solution of N- α -benzoyl--arginine-4-nitroanilide (L-BAPA) with 0.22 mg/mL was used as substrate. Spectrometric quantification was performed at 410 nm and based on a molar extinction coefficient of the reaction product (4-nitroaniline) of 8800 M^{-1} cm⁻¹. One trypsin inhibitor unit (TIU) is defined as the amount of inhibitor required to reduce the trypsin activity by one unit. One trypsin activity unit (TAU)

Table 2 Composition of diets for in vivo nitrogen balance trials, values given in % of diet

Component of diet	C	SPI	SF	RWB	HPHB
Casein	11.15				
DL-Methionine	0.20				
Soya protein isolate		10.80			
Soya flour			19.69		
Reference wheat bread				67.79	
High-protein hybrid bread					43.85
Cellulose	8.00	8.00	8.00	8.00	8.00
Soya oil	8.00	8.00	8.00	8.00	8.00
Mineral mix ¹	3.50	3.50	3.50	3.50	3.50
Vitamin mix ²	1.00	1.00	1.00	1.00	1.00
Choline chloride	0.20	0.20	0.20	0.20	0.20
Cholesterol	0.30	0.30	0.30	0.30	0.30
Sucrose	5.00	5.00	5.00	5.00	5.00
Corn starch	62.65	62.29	54.31	6.21	30.15

¹ AIN-93G-MX: mineral mixture as specified by Reeves ³⁶ (1997)

377 is defined as the amount of enzyme that catalyses the hydroly- $_{378}$ sis of 1 μ mol L-BAPA into 4-nitroaniline within 1 min at pH 8.2 and 37 °C. Contents of vicine and convicine were determined af-380 ter an extraction of 500 mg of sample with boiling methanol as described by Petersen et al. ³⁷. Quantification was achieved using 382 micellar electrokinetic capillary chromatography as reported by 383 Bjergegaard et al. 38 and with vicine as external standard.

384 2.3.6 Antioxidant Potential

The total N content of each diet as well as each faecal and urinal 386 uct powders using 80/20 (v/v) methanol/water (80% MeOH), sample (collected in the balance period) was analysed in dupli- 387 at a solid to solvent ratio of 1:10 (w/v), for 15 min at 50 °C cate (AOAC 979.09). The rats from each diet group were addi- 388 as described by Amarowicz et al. 39. The extraction was retionally monitored for body-weight (BW) gains (recording BWs 389 peated twice, the supernatants were filtered and pooled, and the at the beginning and end of the study) and diet intake (daily 390 methanol was evaporated under vacuum with a rotary evaporarecord), which enabled calculation of the protein efficiency ra- 391 tor (Büchi Labortechnik AG, Flawil, Switzerland). The remaining tio (PER). All physiological measurements were carried out for 392 aqueous extract was lyophilised. Total phenolic content (TPC) -TPC of phenolic extracts was determined using Folin-Ciocalteu's phenol reagent following a method described by Amarowicz and Raab 40. The results were expressed as mg catechin equivalent. Trolox equivalent antioxidant capacity (TEAC) - TEAC was determined according to the method reported byRe et al. 41. In brief, 398 a ABTS*+ (2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) 399 solution was prepared by mixing an aqueous ABTS stock solution 400 with 2.45 mM (final concentration) sodium persulfate. This mixture was shaken for 12-16 h at room temperature in the dark until 402 a stable oxidative state was reached. The ABTS*+ stock solution was diluted with methanol to an absorbance of 0.720 at 734 nm 404 for subsequent analysis. For the spectrophotometric assay, 2 mL 405 of the diluted ABTS $^{\bullet+}$ solution were mixed with 20 μ l of recon-406 stituted phenolic extract (10 mg/mL in methanol); absorbance 407 was determined at 734 nm at 37 °C for 10 min. A calibration 408 curve was generated using a Trolox standard and the results were 409 expressed as μmol Trolox equivalent. Ferric-reducing antioxidant 410 power (FRAP) - FRAP assay was performed as described by Benzie and Strain 42. The FRAP value was calculated and expressed as μ mol Fe²⁺ using a Fe²⁺ calibration curve. DPPH (2,2-diphenyl-1-413 picrylhydrazyl) assay - The radical scavenging effect of the pheno-414 lic extracts was measured as described in Amarowicz et al. 43. A

² AIN-93G-VX: vitamin mixture as specified Reeves ³⁶ (1997)

Table 3 Sensory attributes and extremes of intensity scales used for QDA $_{433}$ of breads

Attribute	Definition	Extremes
Odour		
Sweet	Odour characteristic of sweet buns produced from wheat flour	None - very intense
Acidulous	Odour characteristic of fermented products (e.g. vinegar, yoghurt)	None - very intense
Appearance		
Beige colour	Crumb colour intensity	Light - dark
Pore size	Visual impression of bread crumb porosity	Small - big
Pore distribution	Regularity of pore distribution in the crumb	Irregular - regular
Texture (manual)		
Elasticity	The extent to which bread crumb returns to its original shape when stretched	Low - high
Texture (oral)		
Chewiness	Extent of chewing necessary to prepare food for swallowing	Low - high
Adhesiveness	Degree of adhesiveness when chewing the food 10 times	Low - high
Moisture	Moisture released by the food after 10 chews	Low - high
Taste		
Rye-wheat bread	Aroma characteristics of commercial rye-wheat bread (retronasal)	None - very intense
Salty	Taste characteristic of NaCl (1 % in water)	None - very intense
Acidulous	Taste characteristics of citric acid (1 % in water)	None - very intense
Aftertaste	Lingering sensation after swallowing the sample	None - very intense
Overall quality	Conclusive evaluation of all attributes and their harmonic balance	Bad - very good

methanolic solution (0.1 mL), containing 0.02-0.10 mg of extract, was mixed with 2 mL of deionised water, and was then added to a methanolic solution of DPPH · (1 mM, 0.25 mL). The mixture was vortexed for 1 min and left to stand at room temperature for 20 min. the absorbance of the solution was measured at 517 nm. The results were expressed as half maximal effective concentration (EC₅₀) of the phenolic extract that scavenged 50% of DPPH radicals.

2.4 Sensory Analysis

ing analysis of variance. Before the sensory analysis, a 28-hour panel training was conducted on various bread samples, including bread from the local supermarkets, with the aim to familiarise the sensory panel with innovative bread samples and their features. A list of sensory attributes was created. Initially, panellists chose characteristics describing the samples individually, followed by a joint agreement on distinguishing attributes and their descriptions (see Table 3). A continuous scale (10 cm long) with the extremes specified in Table 3 was used. Sensory evaluation was carried out in three independent sessions.

2.5 Statistical Analysis

All measurements were performed in triplicate unless stated otherwise. Data analysis was carried out using RStudio, version 1.2.1335 with R version 3.6.1 (RStudio Inc, USA; R Core Team, rproject). One-way analysis of variance (ANOVA) with post-hoc pairwise Tukey's test was used to show significant differences (p < 0.05). When available, values are given as the mean \pm standard deviation or uncertainty (amino acid profile).

3 Results and Discussion

3.1 Technological Characteristics

3.1.1 Flour and Dough Properties

The properties of flours and doughs used for breadmaking have a high impact on the quality of bread products. In addition to the ability to form a stable gluten-network, rheological characteristics such as pasting behaviour, dough extensibility and the dough's proofing performance determine flour and dough quality. Gluten-aggregation and pasting behaviour were evaluated for RWB based on wheat flour and for HPHB based on HP flour mix (Table 1). The aim was to compare measurements, which are commonly performed to determine baking quality of flours, for the two formulations in this study. It was decided to include not only the HPIs in the HP flour mix for flour analyses, but also psyllium, which was expected to have a high impact on rheological properties. Sugar and xylanase were shown to have no significant effect on the performance of the HP flour mix in these tests (preliminary trials, data not shown) and were left out. The GlutoPeak test revealed striking differences in gluten-aggregation properties of the two flours. The variables obtained from the curves are presented in Table 4. Wheat flour exhibits with 68 BU a significantly 472 higher TM than HP flour mix (64 BU), but PMT was detected 14 s earlier for HP flour mix (46 s) than for wheat flour (60 s). When 474 pure wheat flours are measured, a general trend towards earlier and higher gluten peaks for stronger flours with higher gluten 476 contents and/or higher gluten quality has been reported in liter-Descriptive sensory profiling (quantitative descriptive analysis - 477 ature. 44-46 The gluten content in HP flour mix (calculated based QDA) was carried out in order to characterise the bread sam- 478 on composition of ingredients and an average gluten content in ples using an expert panel (n=8). The QDA procedure used in 479 wheat flour protein of 80 %) is about 0.5 % lower than in wheat the study was in accordance with the standard ISO 13299:2016. 480 flour, which could explain the slightly lower TM detected for HP Panellists with appropriate methodological preparation and ex- 481 flour mix. However, Hoehnel et al. 30 showed that the partial reperience in sensory profiling were selected, trained and moni- 482 placement of wheat flour by HPIs leads to complex changes in tored following ISO 13299:2016. Before the sensory analysis, 483 the gluten-aggregation profiles, which do not follow this general the panellists' performance was evaluated using three parameters 484 trend. Therefore, a comparison of gluten-aggregation profiles in - repeatability, discrimination ability and homogeneity by apply- 485 addition to TM and PMT (or other variables obtained from the

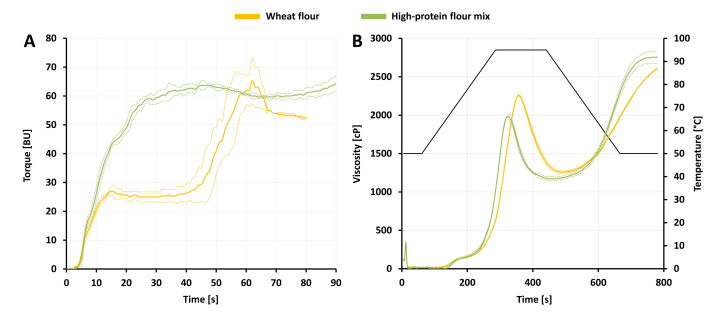


Fig. 1 Flour properties of wheat flour and HP flour mix: (A) Gluten-aggregation profiles obtained by GlutoPeak test; (B) Viscograms obtained from rapid visco analysis describing pasting behaviour of RWB and HPHB with black line representing the applied temperature profile. Dashed curves represent standard deviation.

Table 4 Flour properties of wheat flour (used for reference wheat bread) and HP flour mix (used for high-protein hybrid bread)

Variable	Wheat flour	HP flour mix
GlutoPeak		
Peak maximum time (PMT) [s]	60 ± 4^{a}	46 ± 2^b
Torque maximum (TM) [BU]	68 ± 1^a	64 ± 1^{b}
Rapid Visco Analyser		
Peak viscosity (PV) [cP]	2261 ± 9^a	1989 ± 17^b
Setback [cP]	1350 ± 19^b	1587 ± 59^a
Final viscosity (FV) [cP]	2607 ± 10^b	2756 ± 84^a

Means \pm standard deviation with different letters in the same row were significantly different at p < 0.05

curves) is required (see Figure 1). The profile of wheat flour fol- 505 The corresponding viscograms are displayed in Figure 1. The lows the typical sequence of initial torque increase, equilibrium 506 viscograms suggest a generally similar pasting behaviour of wheat plateau, rapid torque increase, peak maximum and torque de- 507 flour and HP flour mix with only small discrepancies. However, crease due to breakdown of gluten-network. The HP flour mix 508 significant differences have been detected for PV, setback and FV. shows no pronounced equilibrium plateau and the torque in- 509 The PV of HP flour mix is with 1989 cP lower than for wheat flour creases rapidly towards its maximum right in the beginning of 510 with 2261 cP. This can be attributed to the lower starch content the measurement. Instead of a sharp peak with a rapid gluten 511 in HP flour mix and, thus, less gelatinising starch, which has been breakdown, the peak is broad and torque remains high after its 512 previously observed in systems based on wheat flour 50 as well maximum. According to Goldstein et al. 47, a fast build-up of 513 as systems based on rice flour. 51 The presence of psyllium in the gluten-network followed by a sharp peak and rapid breakdown is 514 HP flour mix is expected to increase viscosity of the sample due associated with weak flours. The profile of HP flour mix indicates 515 to its well-known high water absorption and gelling properties a strong and stable gluten-network due to the broad gluten-peak 516 (at low temperatures as well as upon heating). 52,53 This might and delayed gluten breakdown. This could be caused by a co- 517 have partly compensated for the reduced viscosity owing to less networking of gluten with non-wheat proteins from faba bean 518 starch. Hence, only a small difference in PV has been found. In and carob as suggested by Hoehnel et al. 30. The lack of equi- 519 contrast to a lower PV, HP flour mix exhibits higher FV and setlibrium plateau and rapid torque increase at the start can be ex- 520 back compared to wheat flour. Especially the setback expressed plained by the high water absorption of psyllium and gluten 48,49 521 in relation to PV is remarkably high for HP flour mix (wheat flour: resulting in a higher initial viscosity of the sample slurry. Table 4 522 59.7 %, HP flour mix: 79.8 %). A similar pattern was observed shows variables describing the pasting behaviour of the flours. 523 by Hoehnel et al. 30 in a flour blend containing 15 % faba bean flour. Since this ingredient contains a considerable amount of non-wheat starch, high setback and FV could be related to the retrogradation properties of faba bean starch. Dough analyses provide information on rheological and expansion properties of the formulations during proofing. Large deformation properties (Table 5) reveal a reduced extensibility and resistance to extension for the HPHB dough (13.04 mm and 0.475 N, respectively) compared to RWB (16.76 mm and 0.647 N, respectively). According to literature, 54 reduced resistance to extension as well as area under the curve are indicative of weaker doughs. However, also the shape of the curve (see Figure 2), and the ratio of resistance to extension and extensibility (R/E) in particular, seems im-

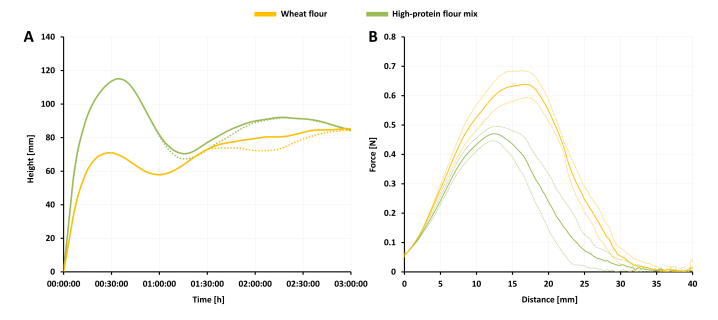


Fig. 2 Dough properties of RWB and HPHB: (A) Gas release curves obtained by Rheofermentometer measurements (dotted line represents gas retained in the dough); (B) Extensibility plots obtained Kieffer rig microextension tests (dashed curves represent standard deviation).

Table 5 Dough properties of reference wheat bread formulation and highprotein hybrid bread formulation

Variable	RWB	НРНВ
Kieffer rig extensibility		
Resistance to extension [N]	0.647 ± 0.059^a	0.475 ± 0.045^b
Extensibility [mm]	16.76 ± 1.25^a	13.04 ± 1.44^b
Rheofermentometer		
Dough development (H_M) [mm]	67.3 ± 5^{a}	61.6 ± 1^a
Total gas volume (V _{total}) [mL]	1982.7 ± 171.1^b	2449.7 ± 102.3^a
Volume of CO_2 lost (V_{lost}) [mL]	58.0 ± 30.5^a	33.0 ± 19.2^{a}
Volume of gas retained (V_{ret}) [mL]	1924.3 ± 192.1^b	2416.3 ± 103.0^{a}

Means + standard deviation with different letters in the same row were significantly different at p < 0.05

portant and provides information on the doughs' viscoelastic bal- 555 in both formulations. This represents the point where easily acance. 54 This ratio is with 0.039 N/mm for RWB and 0.036 N/mm 556 cessible sugars have been consumed by the yeast and further sugfor HPHB very similar for both formulations and suggests similar 557 ars are made available by enzymatic breakdown of starch and expansion properties. Variables describing the proofing perfor- 558 other polysaccharides present in the samples. Gas production at mance of the doughs were obtained by Rheofermentometer mea- 559 the start is the only remarkable difference in an otherwise very surements and are shown in Table 5. No significant difference was 560 similar gas release profile throughout the measurement. Hence, detected for dough development expressed as maximum height 561 the added sugar represents the main factor for the increased V_{total} (H_M) with 67.3 mm for RWB and 61.6 mm for HPHB, which is in 562 of HPHB. Even though the difference observed in V_{lost} is not sigline with the similar expansion properties suggested by microex- 563 nificant, also the curves suggest a tendency towards better gas tension tests. HPHB shows significantly higher total gas volume 564 retention of HPHB dough. This is in accordance with the find-(V_{total}; 2449.7 mL) and retained gas volume (V _{ret}; 2416.3 mL) 565 ings of Courtin and Delcour. 55 They explained a positive effect of than RWB (1982.7 mL and 1924.3 mL, respectively). Also a ten- 566 water-extractable arabinoxylans (AX) on gas retention of doughs dency towards a lower lost gas volume (V_{lost}) for HP flour mix 567 related to a strengthening of liquid films surrounding CO₂ bubwas observed. The gas release curves from Rheofermentometer 568 bles, thereby limiting gas diffusion. The psyllium in HPHB conmeasurements are displayed in Figure 2. The initial gas release 569 tains a considerable amount of AX, of which a small percentage is much more pronounced for HPHB than for RWB. This can be 570 is water-extractable. 56,57 Additionally, xylanase, which degrades explained by the small amount (0.57 %) of added sugar in HPHB, 571 water-unextractable AX (in HPHB from wheat flour 58 and psylwhich leads to higher initial yeast activity and gas production. 572 lium⁵⁷), increases the amount of water-extractable or solubilised The initial peak is followed by a temporary decline in gas release 573 AX present in the dough and their effect on gas retention proper-574 ties. 55 Xylanase degradation of water-unextractable AX has also 575 been reported to lead to a lowered water-binding capacity of AX and redistribution of water in favour of gluten, therefore facilitating gluten-network formation. 49,59 Wang et al. 60 discussed the formation of a secondary network based on AX with the ability to strengthen the gluten-network by entanglement and possibly the creation of diferulic bridges. This is in line with the stability of the gluten-network in HPHB and delayed breakdown indicated by GlutoPeak test results and represents an additional stabilising effect besides potential co-networking of gluten with non-wheat

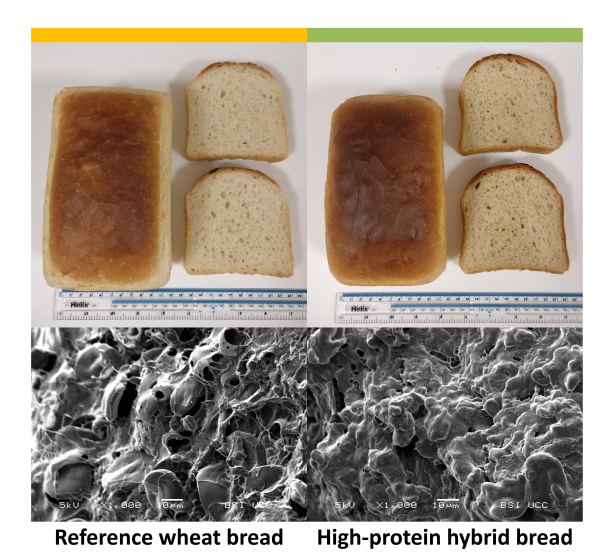


Fig. 3 Photographs and micrographs (obtained by SEM) of RWB and HPHB.

3.1.2 Bread Quality Characteristics

The breads produced from both formulations examined in this study are presented in Figure 3. A visual evaluation reveals little differences in loaf size and crumb structure between RWB and HPHB, but a considerably darker crust and crumb colour for HPHB. The results obtained for bread quality characteristics confirm this general observation and are reported in Table 6. No significant differences between RWB and HPHB have been detected regarding bake loss and SV. The initial crumb hardness on day 0 is with 6.98 N for HPHB slightly higher than for RWB (5.13 N). However, this can only be considered a small difference, especially when compared to previously reported increases in crumb hardness caused by the incorporation of legume ingredients in wheat bread. 22,30,61,62 Additionally, the crumb hardness measured on day 2 and day 5 does not show significant differences between RWB and HPHB. This indicates similar staling properties of both formulations, with a tendency towards less staling for HPHB. Staling rates calculated for day 2 are 1.42 for RWB and 0.78 for HPHB, which represents a by 45 % lower staling rate of HPHB. Staling rates calculated for day 5 are 2.08 and 1.31 for RWB and

605 HPHB, respectively. Also here, HPHB shows a by 37 % lower 606 staling rate. Recrystallysing starch is considered to be the main 607 factor for staling of bread crumb. 63,64 Therefore, the decreased 608 crumb staling in HPHB could be related to its lower starch content and, supposedly, a lower amount of gelatinised starch which

Table 6 Bread quality characteristics of reference wheat bread and high-protein hybrid bread

Variable	RWB	НРНВ
Bake loss [%]	12.3 ± 0.6^{a}	11.9 ± 0.8^a
Specific volume (SV) [ml/g]	3.73 ± 0.07^a	3.75 ± 0.13^a
Hardness day 0 [N]	5.13 ± 0.43^b	6.98 ± 0.60^{a}
Hardness day 2 [N]	12.41 ± 1.43^a	12.41 ± 1.23^a
Hardness day 5 [N]	15.81 ± 0.85^a	16.15 ± 2.06^a
Number of cells	5009 ± 245^b	5563 ± 575^a
Cell area [%]	52.4 ± 0.3^{a}	51.7 ± 0.5^b
Slice brightness	137 ± 4^a	108 ± 3^b
Lightness of crumb (L*crumb)	63.6 ± 2.2^{a}	60.4 ± 3.9^a
Lightness of crust (L*crust)	41.9 ± 5.0^{a}	34.6 ± 2.9^b

Means \pm standard deviation with different letters in the same row were significantly different at p < 0.05.

can recrystallise. Also AX and xylanases have been reported to decrease staling of wheat-based bread formulations. 65,66 The effect has been attributed to a competition for water and therefore a reduced swelling and gelatinisation of starch. 65,66 Maeda and Morita ⁶⁷ observed reduced staling up to 3 days caused by both water-extractable and water-unextractable AX. While their study was focused on wheat AX, Czuchajowska et al. 68 also reported reduced crumb hardness after 72 h when psyllium was incorporated in wheat bread. Specific volume and crumb hardness are generally accepted as the main indicators of bread quality. Therefore, the presented results confirm a technological quality of HPHB similar to RWB. The evaluation of crumb structure reveals small differences between the formulations. A slightly finer crumb structure was observed for HPHB indicated by a higher number of cells (5563) and smaller cell area (51.7 %) compared to RWB with a number of cells of 5009 and a cell area of 52.4 %. This can be related to the higher initial yeast activity and gas production in HPHB. Moulding of dough, in addition to shaping the dough pieces, leads to a division of gas cells produced prior to moulding (during mixing and dough rest). 69 In HPHB, more gas is produced before moulding and a higher number of small gas cells can be generated. Additionally, these gas cells are stabilised by water-extractable and solubilised AX as explained above, which can minimise the coalescence of gas cells as they expand during proofing and lead to a high number of cells in the final product. The higher number of cells and smaller cell area measured for HPHB could also be partially responsible for its slightly higher initial crumb hardness. Values obtained for crumb and crust colour (Table 5) confirm the visually perceivable differences between RWB and HPHB. Slice brightness (obtained by C-Cell imaging) is significantly lower for HPHB with 108 than for RWB with 137. This is in line with the lower lightness of crumb measured for HPHB. A big difference was observed in lightness of crust, which was significantly lower for HPHB (34.6) than for RWB (41.9). The darker crust of HPHB is likely related to its higher protein content and higher presence of reducing sugars (see Table 7), thus, an increased potential for Maillard reaction. 30,70 The micro-structure of the bread crumb of both formulations was captured by scanning electron microscopy (SEM). The resulting micrographs are displayed in Figure 3. While RWB shows a rather porous layer of gluten covering partly intact starch granules, HPHB has a thicker and more continuous layer. This might be due to the presence of non-wheat proteins from faba bean and carob on one hand and psyllium on the other hand. The fact that a very homogenous and continuous layer was formed, further supports the theory of a co-networking of gluten with nonwheat proteins and psyllium AX.

3.2 Nutritional Characteristics

Macronutrient Composition and Sugar Profile

Compositional analysis of both formulations was performed in order to evaluate changes in macronutrient composition caused by the partial replacement of wheat flour by plant-based HPIs in HPHB and addition of psyllium to the formulation (Table 7). The determined bread constituents include all items that are manda-

Table 7 Composition of reference wheat bread and high-protein hybrid bread, contents expressed in % of the fresh bread unless stated other-

Component	RWB	НРНВ
Moisture	45.74 ± 0.06	45.91 ± 0.29
Energy [kcal/100 g]	211.6	209.0
Protein	8.2	13.0
proteinE * [%E]	15.5	24.8
Ash	1.6	2.0
Fat	0.91	1.25
SFA	0.11	0.17
MUFA	0.26	0.36
PUFA	0.50	0.67
Total carbohydrates**	43.5	37.9
Total dietary fibre (TDF)	1.8	2.8
Available carbohydrates**	41.7	35.1
Total starch	36.1 ± 1.2	28.5 ± 0.6
Sodium	0.466	0.440
Sodium expressed as salt (NaCl)	1.16	1.10
Sum of mono- and disaccharides	1.21 ± 0.00	1.13 ± 0.02
Arabinose	< 0.01	< 0.01
Xylose	< 0.01	< 0.01
Galactose	0.01 ± 0.00	0.03 ± 0.00
Glucose	0.02 ± 0.00	0.03 ± 0.00
Fructose	0.02 ± 0.00	0.04 ± 0.00
Sucrose	0.01 ± 0.00	0.01 ± 0.00
Maltose	1.16 ± 0.00	1.02 ± 0.02
Maltotriose	0.03 ± 0.00	0.02 ± 0.00
Raffinose/Stachyose	< 0.01	0.01 ± 0.00
Verbascose	< 0.01	0.02 ± 0.00

Moisture, total starch and sugar profile: means \pm standard deviation

664 tory for nutritional food product labelling according to European 665 food legislation (regulation (EU) No 1169/2011⁷¹). In addition, 666 the sugar profile, total starch content and other important components of the samples were measured or calculated. Protein con-668 tent and content of available carbohydrates represent the main 669 differences in the macronutrient profile of RWB and HPHB. This 670 is essentially caused by the replacement of wheat flour, which is 671 high in starch (72.38 %DM), by HPIs with protein contents of 61.25 %DM (faba bean flour), 55.04 %DM (carob germ flour) 673 and 83.11 %DM (gluten) and starch contents below 10 %DM 674 (protein and starch contents of wheat flour and HPIs previously 675 reported by Hoehnel et al. 30). While the total energy level of the 676 formulations is similar (RWB 211.6; HPHB 209.0), a shift from 677 wheat starch to non-wheat protein characterises the macronutri-678 ent profile of HPHB. This shift is also evident when proteinE values (percentage of calories provided by protein) are compared. 680 In contrast to RWB with 15.5 %E, the HPHB formulation reaches a proteinE of 24.8 %E and therefore qualifies for a "high in pro-682 tein" nutritional claim in accordance with European food legislation (regulation (EC) No 1924/2006⁷²), where a proteinE of 20 % is set as requirement. Bread is a staple food with global importance as source of dietary carbohydrates, protein and fibre. 15 However, within the past 200 years, the consumption of refined-carbohydrate products, including bakery products from 688 refined wheat flour (white bread, white bagels, white buns), has substantially increased. At the same time, significantly less regu-690 lar starchy foods like beans, lentils and wholegrain bakery products are consumed. This is largely associated with generally bet-

^{*} calculated based on energy content, protein content and 4 kcal/g protein

^{**} calculated by difference

ter sensory characteristics of refined-carbohydrate products and potentially higher consumer acceptance due to their sweet taste when starch is rapidly digested by salivary amylase. ^{7,15} The main concern regarding this development is related to high glycaemic indices due to rapidly digestible starch . 2,7,15,73 High-glycaemicload and high-glycaemic-index diets have been associated with elevated risk for diabetes, heart disease and certain types of cancer. 74-78 Due to its reduced content of available carbohydrates, HPHB is expected to have a lowered glycaemic load in comparison to RWB. Even a decreased glycaemic index could be expected, since psyllium has been reported to lower the glycaemic index of foods when added to conventional diets. 79,80 Holt et al. 81 found a significantly lowered blood glucose response of high-protein bread when they compared equal-energy portions of high-protein bread and regular white bread. Furthermore, an isocaloric replacement of refined starch or sugar by protein, like it is the case for HPHB compared to RWB, has been reported to reduce blood pressure and blood lipid concentrations. 2,82 Also the lack of fibre in refined-carbohydrate foods compared to wholegrain alternatives and legumes has been critically discussed. 7,15 Dietary fibre is associated with many health benefits and dietary recommendations advice a daily intake of 25 g or more for adults. ^{2,83} In the present study, HPHB contains with 2.8 % considerably more dietary fibre than RWB with 1.8 %. This is related to the incorporation of faba bean flour, carob germ flour and psyllium in HPHB, which represent ingredients with notable contents of both soluble and insoluble fibre. 30,57 Especially psyllium has been 747 an unbalanced amino acid composition, and to its lack of the reported in literature as dietary fibre with beneficial effects re- 748 indispensable amino acid lysine in particular. 5,12,15 The amino garding the risk of diabetes, obesity, high blood pressure and 749 acid profile of RWB and HPHB was determined and is reported heart disease. 52 Apart from refined carbohydrates, also fat and 750 in Table 8. The results show that the proportions of indispenssalt (sodium cloride) are dietary components which are often 751 able and dispensable amino acids are very similar in both forcritically discussed. 84-86 While HPHB contains with 0.440 % an 752 mulations. Amongst the dispensable amino acids, only the levamount of sodium similar to RWB (0.466 %), it has a slightly 753 els of glutamine/glutamic acid, proline and arginine differ subelevated fat content. However, this increase is mainly caused 754 stantially between RWB and HPHB. While wheat is particularly by higher contents of MUFA and PUFA, which are nutritionally 755 rich in glutamine, glutamic acid and proline but contains little more favourable than saturated fats. ^{2,84} Both formulations con- ₇₅₆ arginine, ¹² faba bean and carob show a complementary pattern tain similar amounts of sugar (mono-and disaccharides) and their 757 for these AA. 32,88 Especially faba bean protein contains relatively sugar profiles reveal little differences. They confirm that sucrose 758 small amounts of glutamine/glutamic acid and is high in argiadded in the recipe of HPHB is fully consumed during yeast fer- 759 nine. This causes a decreased level of glutamine/glutamic acid mentation, which was also evident in the results obtained from 760 and proline but an increased level of arginine in HPHB. Regarding dough analyses. Slightly increased galactose and the presence of 761 the profile of indispensable AA in RWB and HPHB, many minor oligosaccharides like raffinose, stachyose and verbascose can be 762 differences were observed. However, the lysine level is approxiassociated with high contents of galactooligosaccharides (GOS) 763 mately 65 % higher in HPHB than in RWB. Also this change can reported for faba beans. 87 Slightly lower maltose and maltotriose 764 be attributed to faba bean and carob proteins which are naturally

3.2.2 Amino Acid Profile

decrease in high-quality animal protein consumption is taken into 775 not reach the quantity specified as recommended intake (= 1).

Table 8 Amino acid composition of reference wheat bread and highprotein hybrid bread

Content [%Protein]	RWB	НРНВ
Indispensable and conditionally		
indispensable AAs		
Histidine	1.92 ± 0.23	2.23 ± 0.27
Isoleucine	3.94 ± 0.48	3.77 ± 0.46
Leucine	7.33 ± 0.89	7.03 ± 0.85
Lysine	2.36 ± 0.29	3.90 ± 0.48
Cystine	1.99 ± 0.24	1.61 ± 0.19
Methionine	1.08 ± 0.13	0.97 ± 0.12
Cystine + Methionine (SAAs)	3.07 ± 0.37	2.58 ± 0.31
Phenylalanine	4.82 ± 0.59	4.46 ± 0.55
Tyrosine	2.41 ± 0.30	2.23 ± 0.27
Phenylalanine + Tyrosine (AAAs)	7.23 ± 0.88	6.69 ± 0.82
Threonine	2.70 ± 0.33	3.12 ± 0.38
Tryptophan	0.76 ± 0.48	0.78 ± 0.30
Valine	4.03 ± 0.49	4.38 ± 0.53
Total indispensable AAs	43.63 ± 5.71	43.75 ± 5.54
Dispensable AAs		
Asparagine/aspartic acid	4.13 ± 0.50	6.08 ± 0.74
Glutamine/glutamic acid	30.39 ± 3.69	25.71 ± 3.12
Glycine	3.94 ± 0.48	4.10 ± 0.50
Alanine	3.05 ± 0.37	3.40 ± 0.42
Serine	4.97 ± 0.61	4.63 ± 0.56
Proline	10.48 ± 1.27	8.11 ± 0.99
Arginine	3.74 ± 0.46	6.50 ± 0.79
Total dispensable AAs	60.69 ± 7.38	58.53 ± 7.10

Amino acid contents ± uncertainty values

levels in HPHB are potentially related to its lower starch content. 765 richer in lysine than wheat. 12,32,88 Even though the difference 766 of lysine contents expressed in %Protein might seem small, this 767 difference has a big impact on the breads' overall amino acid bal-Many dietary recommendations advice a substantial decrease in 768 ance and, thus, their protein quality. Especially when compared the consumption of animal protein and a shift towards protein 769 to a reference pattern of indispensable amino acids (for adults) from plant sources. 2,4 Even though bread can be considered an 770 recommended by WHO 89 and EFSA 90, the significance becomes important source of plant protein, the poor protein quality of 771 evident. The quantity of indispensable amino acids in RWB and wheat makes regular wheat bread (from both wholegrain or re- 772 HPHB relative to the amino acids in the reference pattern is prefined wheat flour) an inadequate choice to partially compensate 773 sented in Figure 4. The comparison with the reference pattern refuture plant-protein requirements; especially when a substantial 774 veals that in both formulations lysine is the only AA, which does account. The poor protein quality of wheat is mainly linked to 776 Therefore, lysine represents the limiting AA of the protein in RWB

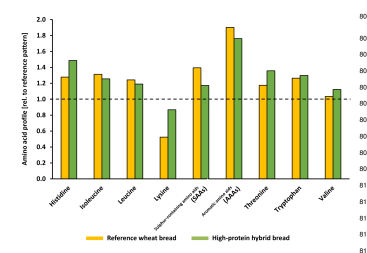


Fig. 4 Profile of indispensable amino acids of reference wheat bread and high-protein wheat bread expressed relative to the requirement pattern (WHO 2007⁸⁹) and based on an average intake of 0.66 g protein/kg

and HPHB. The increased lysine content in HPHB (87 % of lysine in reference pattern) compared to RWB (52 % of reference pattern) leads to a much more balanced AA profile that almost covers the recommended intake of all indispensable AA. The expression of AA levels in a food protein relative to the levels in a reference 824 protein is referred to as amino acid score (AAS). Table 9 shows an overview of AAS and limiting AAs of RWB and HPHB and the in- $\,^{826}$

Table 9 Amino acid scores (AASs) for breads and their raw materials

Protein source	AAS	Limiting AAs	
RWB	0.52	Lysine	
HPHB	0.87	Lysine	
Wheat flour*	0.57	Lysine	
Faba bean flour**	0.66	SAAs	
Carob germ flour*	- (1.02)***	 (Valine)*** 	
Gluten*	0.37	Lysine	

calculated from amino acid composition; determined as for RWB and HPHB (data not

HPHB formulation does not only have an improved AAS compared to RWB, but also in comparison to wheat flour and HPIs. The combination of the ingredients in HPHB leads to an upgrade in nutritional value of most raw materials when AAS is used to evaluate nutritional protein quality. The exception is the protein from carob germ flour, which has a nutritionally favourable AA pattern itself. Since the calculation of AAS is based on a recommended amino acid reference pattern, which considers an average intake of 0.66 g protein/kg bodyweight, this evaluation assumes that RWB or HPHB (or the ingredients) are the sole source of protein in the diet. In a real diet, proteins from other foods can potentially compensate for AA deficiencies. However, the ability of a dietary protein source to fulfil amino acid requirements on its own is regarded as an adequate approach to compare nutritional quality of proteins.

3.2.3 Protein Digestibility and Utilisation

The informative value of AASs is also limited because they do not reflect the protein's digestibility, absorption and utilisation. 91 In the present study, protein digestibility was evaluated in an in vitro model as well as in an in vivo trial with rats (Table 10). In vitro protein digestibility (IVPD) of RWB and HPHB was monitored after 1 h of pepsin digestion and, subsequently, 1 h of pancreatin digestion, which is indicative of the digestibility in the human digestive system. Additionally, IVPDs were measured after a medium term (3 h) and a long term (24 h) pancreatin digestion to evaluate the maximum achievable degradation of the proteins. Both the digestion mimicking gastric conditions (1 h pepsin) as well as the simulated intestinal digestion (1 h pancreatin) yielded higher ratios of degraded protein for HPHB than for RWB, indicated by significantly higher IVPD values. This suggests a slightly improved protein digestibility of HPHB, which is remarkable since legumes, in HPHB specifically faba bean and carob, are often critically discussed regarding their contents of trypsin inhibitors and an associated decrease in protein digestibility. 92 However, due to the incorporation of only 5.72 % of faba bean flour in the whole HPHB formulation (see Table 1), a substantially reduced content of trypsin inhibitors, as compared to the faba bean raw material, is expected. A detailed discussion of the trypsin inhibitor activity (TIA) in HPHB follows in chapter 3.2.4. A higher degree of protein degradation in HPHB could be explained by the higher abundance of lysine and arginine in this formulation. Trypsin, which is a predominant proteolytic enzyme gredients used for their production (wheat flour and HPIs). The 827 in pancreatin, cleaves protein and peptide chains at the carboxyl 828 side of these positively charged AA. Pancreatin also contains chymotrypsin, which cleaves after hydrophobic AA with bulky side chains like phenylalanine, tryptophan and tyrosine. The contents of these AA are very similar in HPHB and RWB. However, abundance of target AA for trypsin and chymotrypsin proteolysis is not the only relevant factor. Also accessibility of such AA in the three-dimensional protein structure is of high importance. This suggests that HPHB contains a higher number of AA accessible for tryptic/chymotryptic digestion. The in vivo protein digestibil-

Table 10 In vitro protein digestibility and in vivo nitrogen balance

Variable	RWB	НРНВ		
In vitro protein digestibility (IVPD) [%]				
Pepsin 1 h	1.1 ± 0.4^b	2.0 ± 0.3^a		
Pancreatin 1 h (short term)	14.2 ± 0.6^b	17.2 ± 0.3^a		
Pancreatin 3 h (medium term)	18.4 ± 1.7^b	22.7 ± 1.2^a		
Pancreatin 24 h (long term)	25.0 ± 0.0^b	31.1 ± 0.1^a		
In vivo nitrogen balance				
N intake [g/5 d]	$1203^b \pm 359$	1556 ± 94^{a}		
N in faeces [mg/5 d]	138 ± 47^{b}	183 ± 12^a		
N faecal [% N intake]	11.4 ± 1.0^{a}	11.8 ± 1.0^a		
N in urine [mg/5 d]	766 ± 206^{a}	733 ± 35^{a}		
N urinary [% N intake]	64.4 ± 3.1^a	47.1 ± 1.8^{b}		
N digestibility [%]	88.6 ± 1.0^{a}	88.2 ± 1.0^a		
N utilisation [%]	24.2 ± 2.7^b	41.0 ± 2.7^{a}		
PER [g/g]	1.13 ± 0.39^b	2.13 ± 0.17^a		

Means \pm standard deviation with different letters in the same row were significantly different at p < 0.05.

^{**} calculated from amino acid composition; determined as for RWB and HPHB and reported by Vogelsang-O'Dwyer et al. 32

 $^{^*}$ not strictly limiting (\geq 1), but represents AA with lowest level relative to reference pattern

Table 11 Contents of antinutritional compounds of reference wheat bread and high-protein hybrid bread, contents refer to fresh bread or dry matter as indicated

Antinutritional compound	RWB HPHB based on fresh bread	RWB HPHB based on dry matter*
Trypsin inhibitor activity (TIA) [TIU/mg] Vicine [%] Convicine [%]	$\begin{array}{ll} \text{n.d.} & 0.21 \pm 0.01 \\ \text{n.d.} & 0.056 \pm 0.005 \\ \text{n.d.} & 0.044 \pm 0.001 \end{array}$	$\begin{array}{cc} \text{n.d.} & 0.39 \pm 0.02 \\ \text{n.d.} & 0.103 \pm 0.009 \\ \text{n.d.} & 0.081 \pm 0.002 \end{array}$

ity trials performed with rats yielded several variables indicative 882 of overall calorie and protein intake on N utilisation and PER has between in vitro and in vivo digestibility data, 93 some legumes 898 compared to RWB. have been found to reach higher digestibility in in vitro experiments than in vivo. 91 This is in agreement with the slightly higher 899 3.2.4 Antinutritional Compounds IVPD observed for HPHB in comparison to RWB in this study. N digestibility is also used to calculate the protein digestibility corrected amino acid score (PDCAAS), which is the most commonly used indicator of nutritional protein quality. Since N digestibility of RWB and HPHB is similar, PDCAAS values follow the same trend as AAS values dicussed in the previous section. Related to its higher lysine content, HPHB reaches a PDCAAS of 0.77, which is 67 % higher than PDCAAS of RWB with 0.46. N utilisation considers N loss in both faeces and urine. Caused by a significantly lower urinary N loss of rats fed with HPHB diet, a by 69 % increased N utilisation was observed for HPHB compared to RWB. This is mainly linked to the improved AA pattern and higher content of lysine in HPHB. It has been shown that the lack of one or more essential AAs (provided by the diet and absorbed after digestion) leads to a plateau in AA retention. Other absorbed essential AA, which are present in excess of the limiting AA according to the required AA pattern, are oxidised in the blood and excreted with the urine. 89,94 In both animal and human studies, correlation was found between level of imbalance of indispensable AA in the diet and inefficient AA utilisation leading to limited protein synthesis. 95,96 Corresponding to the higher N utilisation, also the determined PER was with 2.13 g/g significantly higher for rats with HPHB diet than for rats with RWB diet (1.13 g/g). Protein efficiency ratio is a widely used indicator of protein quality and reflects the protein's ability to fulfil AA requirements for growth (experiment performed with growing rats). An influence

of the breads' nutritional value (Table 10). The most important 883 been discussed. 91,94 Therefore, differences in N utilisation and are N intake, N digestibility, N utilisation and protein efficiency 884 PER between HPHB and RWB in this study might be partially reratio (PER). N intake was monitored as a reference value to cal- 885 lated to the higher N intake (hence, higher calorie intake) that culate relative faecal and urinary N losses. N intake was signif- 886 was observed for HPHB. While both in vitro and in vivo modicantly higher for rats which were fed the diet containing HPHB 887 els have their limitations, especially regarding transferability of (1556 g/5 d) compared to rats with RWB diet (1203 g/5 d). Since ** results to the human digestive and metabolic system, they offer diets were adjusted to contain the same amount of protein, this 889 a valid comparison of proteins and their nutritional quality. 93,97 means that rats consumed significantly more of their whole diet 🕬 Protein digestbility is a matter of the degree of hydrolysis and with HPHB. It is remarkable that N intake with HPHB diet even 891 release of amino acids for absorption. True protein quality is conexceeded that of rats with the control casein diet (1262 g/5 d, 892 sidered a measure of the balance of AA which are absorbed and data not shown). This could be associated with a higher palata- 893 utilised in the human body to achieve defined metabolic actions bility of HPHB diet compared to diets containing RWB or casein. 894 (e.g., growth). 5,94 Even though it is unknown, which AA in par-N digestibility (according to faecal N loss) was similar between 895 ticular are absorbed and utilised in which ratios, the presented the two bread formulations in this study and no significant differ- 896 results (including AA profile, IVPD, N digestibility, N utilisation ences were found. Although literature reports good correlations 897 and PER) conclusively suggest improved protein quality of HPHB

900 Trypsin inhibitors and the pyrimidine glycosides vicine and con-901 vicine are considered antinutritional compounds and their activ-902 ity/contents have been determined for HPHB and RWB in this 903 study (Table 11). It is well known that trypsin inhibitors have 904 the ability to form a complex with the proteolytic enzyme trypsin 905 leading to its inactivation. While this can cause adverse effects 906 like increased pancreatic secretory activity and pancreatic hyper-907 trophy, 24 it is often responsible for substantially reduced pro-908 tein digestibility. ²⁵ No trypsin inhibitor activity (TIA) was de-909 tected for RWB. The TIA of 0.21 TIU/mg measured for HPHB 910 can be considered very low compared to the approximately 10 911 fold higher TIA in the faba bean raw material used for HPHB 912 reported by Vogelsang-O'Dwyer et al. 32. However, this reduc-913 tion of TIA is mainly related to the dilution effect in the bread 914 matrix. While heat treatment is an efficient way to inactivate 915 trypsin inhibitors (changes in active site conformation), baking 916 seems to be considerably less efficient than other thermal pro-917 cessing techniques. 92 In addition to faba bean, also carob germ flour could be a source of trypsin inhibitors in HPHB. 98,99 Ac-919 cording to the determined IVPD of HPHB and RWB, the remain-920 ing TIA in HPHB from faba beans or carob seeds did not lead to 921 a decreased protein digestibility of HPHB compared to RWB. The results do not allow for an interpretation whether this is due to a negligible TIA in the bread matrix or due to the overall improved protein quality compensating for TIA. The ANCs vicine and con-925 vicine are particularly relevant in foods containing faba beans. 100

calculated based on moisture of fresh bread given in Table 7; for comparison purposes

in RWB, HPHB contains 0.056 % vicine and 0.044 % convicine 977 main contributors to the enhanced antioxidant potential of HPHB. et al. ¹⁰¹, G6PD deficient men consumed large quantities (500 g) 979 uated antioxidant potential of breads enriched with carob flours. of faba beans from a low vicine/convicine variety (0.016 % based 980 Also wheat is naturally rich in phenolics. But since these comered safe for individuals with G6PD deficiency. The incorporation 986 cereals (wheat, rye, oats, barley). The incorporation of all whole-HPHB, and formulations of its kind, with regard to nutritional as- 989 was observed when wholegrain rye flour was added, which is simfor the lack of lysine in cereals, when consumed separately.

3.2.5 Antioxidant Potential

Phenolic compounds, and specifically phenolic acids and flavonoids, exhibit many biological activities. They are well known for their antioxidant activity through which they prevent oxidative damage of biomolecules like lipids, proteins and have been associated with the occurrence of both degenerative and neurodegenerative diseases such as cancer, inflammatory and cardiovascular conditions and Alzheimer's disease. 104 It has been velopment of these diseases. 105-108 The total content of phenolics of RWB and HPHB was determined. Additionally, the antioxidant $_{\scriptscriptstyle{1008}}$

Table 12 Antioxidant potential of reference wheat bread and high-protein hybrid bread, contents refer to fresh bread unless stated otherwise

Antioxidant potential	RWB	НРНВ
Total phenolics [mg/100 g]	15.8 ± 0.3^b	66.1 ± 0.3^a
ABTS [mmol Trolox/100 g]	0.08 ± 0.01^b	1.02 ± 0.03^a
FRAP [mmol Fe ²⁺ /100 g]	0.23 ± 0.01^b	0.77 ± 0.01^a
Antiradical activity (DPPH)		
EC ₅₀ [mg extract/mL] [×]	6.22 ± 0.18^a	1.15 ± 0.03^b

Means \pm standard deviation with different letters in the same row were significantly different

When ingested by individuals with glucose-6-phosphate dehydro- 968 potential of the phenolic extracts of the breads was evaluated usgenase (G6PD) deficiency, these compounds can trigger favism, 969 ing ABTS, FRAP and DPPH assays. The results are presented in which leads to acute haemolytic anaemia.²⁸ On average, the 970 Table 12. The total content of phenolics is with 66.1 mg/100 g sum of vicine and convicine accounts for about 1 %DM in faba 971 substantially higher in HPHB than in RWB with only 15 mg/100 g. beans. 27,100 However, efforts in plant breeding have led to cul- 972 Also the assays performed to determine antioxidant activity of tivars with contents of the pyrimidine glycosides as low as 0.01 973 the phenolic extracts conclusively suggest an increased antiox-- 0.02 %DM.²⁷ Vogelsang-O'Dwyer et al.³² reported a content 974 idant potential of HPHB than RWB. High levels of antioxidant (vicine + convicine) of 1.25~%DM in the faba bean flour used for $_{975}$ compounds have been reported for legumes 18 and faba bean and HPHB. While vicine and convicine were, expectedly, not detected 976 carob in particular. 17,43,109 Therefore, they are expected to be the (contents referring to fresh bread). In a recent study by Gallo 978 The same trend was observed by Turfani et al. ²³ when they evalon wet weight as ingested). It was confirmed that this level of 981 pounds are mainly found in the bran fraction, the antioxidant intake was safe and favism was not triggered. Based on the out- 982 potential of breads produced from refined wheat flour is usually comes from Gallo et al. 101 and the results of the present study, 983 low. 110 Ragaee et al. 111 investigated the content of phenolics and the consumption of at least 80 g of HPHB (equivalent to 2 slices 984 antioxidant potential of refined wheat bread when wheat flour of bread with a typical weight of 38 g per slice 102) can be consid- $_{985}$ was partially replaced (30 %) by wholegrain flours from different of faba bean flour in HPHB leads to a substantial dilution of ANCs 987 grain cereals flours increased the breads' antioxidant potential. as compared to the raw material. This underlines the value of 988 The highest content of phenolics of approximately 70 mg/100 g pects. In theory, the separate consumption of legumes and cereals 990 ilar to the content of phenolics reached by HPHB in the present as part of a balanced diet can guarantee a balanced AA intake sim- 991 study. Since the phenolics in a food matrix are present either free ilar to the pattern of HPHB. But the presence of higher amounts 992 or bound to polysaccharides, a prediction whether they can exert of ANCs, which affect protein digestibility and AA bioavailability, 993 antioxidant activity in vivo is difficult. Digestibility of the food, might substantially reduce the capacity of legumes to compensate 994 which determines bioavailability of the phenolics, is an important 995 factor and in vivo antioxidant activity does not always correlate 996 with in vitro data. 112 However, the results in this study clearly 997 show higher antioxidant potential for HPHB than RWB.

998 3.3 Sensory Characteristics

999 Consumer acceptance of food products is highly depending on DNA. 103 Amongst many other factors, such oxidative damages 1000 sensory characteristics, which are in turn related to the products' technological quality. Due to its enhanced nutritional profile and qualification for the nutritional claim "high in protein", 72 HPHB can be considered a functional food. According to consumer surdemonstrated in epidemiological studies that high intake of foods 1004 veys reported in literature, consumers evaluate functional foods containing high levels of compounds with antioxidant activity 1005 the same way they evaluate conventional foods. Functional ben-(e.g., whole-grain foods and legumes) can help to prevent the de- 1006 efits are perceived merely as added value and cannot outweigh 1007 inferior sensory properties. 113 Sensory analysis for the two formulations in this study was performed with a trained panel using selected descriptors for bread quality (Figure 5). Reference wheat bread and HPHB reached similar scores for attributes describing taste and porosity of the crumb. Interestingly, the differences in crumb structure, which were observed in technological analyses of the breads, were not perceived by the panellists. The results for HPHB further indicate an improved crumb texture, which is often perceived as an indicator of freshness amongst consumers. 114,115 Compared to RWB, it scored significantly higher in elasticity and lower in adhesiveness. While elasticity of bread crumb is recognised as a favourable attribute, adhesiveness is often associated with stickiness and an unpleasant mouthfeel. 115 Both formulations reached similar scores in chewiness. This shows that the slightly increased initial crumb hardness for HPHB, which was

Concentration of phenolic extract of breads able to scavenge 50 % of DPPH radicals

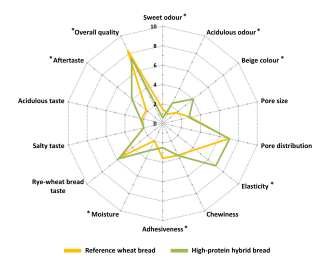


Fig. 5 Sensory characteristics of reference wheat bread and high-protein 1073 hybrid bread; asterisks * indicate attributes which showed significant differences between RWB and HPHB (p < 0.05)

detected in texture profile analysis (TPA), had no perceptible negative impact on the mouthfeel of the bread crumb. Highprotein hybrid bread scored higher than RWB in moisture of 1079 crumb, which is considered another indicator for bread fresh-1080 sumed diets. ness and quality. 115 Significant differences have been found regarding the odour profile of the formulations. While for RWB a slight sweet and almost no acidulous odour was perceived, HPHB 1082 There are no conflicts to declare. had no perceivable sweet odour and slightly stronger acidulous odour than RWB. In accordance with the results of instrumen- 1083 Acknowledgements tal crumb colour measurements, a darker/more beige colour was 1084 The authors want to thank Tom Hannon for technical support, observed for HPHB. Also a moderate increase in aftertaste was 1085 Concept Life Science Ltd. and Mérieux NutriSciences (CHELAB identified in HPHB. However, the overall sensory quality was rated only slightly lower for HPHB than for RWB. This identifies HPHB as a bread formulation with adequate sensory quality when 1088 croscopy imaging. The work for this study has been undertaken compared to RWB, suggesting high consumer acceptance. The 1089 as part of the project PROTEIN2FOOD. This project has received scores of HPHB for sensory attributes like acidulous odour and 1090 funding from the European Union's Horizon 2020 research and beige colour suggest similarities to the typical sensory profile of 1091 innovation programme (grant agreement No 635727). sourdough bread. 116,117 Because of the popularity of sourdough bread amongst consumers, this could further contribute to a high 1092 Abbreviations consumer acceptance of HPHB.

Conclusion

A mixture of HPIs was used to partially replace wheat flour in regular wheat bread to produce a high-protein bread. The HPIs and their ratios were selected based on previous results by Hoehnel et al. 30 to represent both beneficial expected nutritional properties as well as adequate baking properties. In order to match the technological quality of a regular wheat bread, which was used as a reference, also three functional ingredients (psyllium, sugar, xylanase) were added. Dough and bread quality comparable to the reference wheat bread were observed for high-protein hybrid bread (HPHB); mainly mediated by the functional properties of carob and gluten protein as well as psyllium and xylanase. Additionally, a substantially enhanced nutritional profile of the proposed HPHB compared to regular wheat bread was achieved. The macronutrient composition was improved by an isocaloric replacement of refined wheat-starch by plant protein. The protein quality was improved, judging by a better AA profile, increased N utilisation and higher protein efficiency ratio. Mainly due to the dilution effect in the bread matrix, only low levels of ANCs originating from faba bean and carob were measured. Furthermore, determination of phenolics and antioxidant activity indicate high antioxidant potential for HPHB. Apart from favourable technological and nutritional characteristics, the proposed formulation also has high sensory quality which suggests high consumer acceptance. In a time in which we are looking for ways to adequately and sustainably provide enough high-quality plant protein for a future human diet, we cannot afford to focus only on meat and dairy replacement products; especially considering that these applications often require highly purified or additionally functionalised plant proteins obtained by wet-processing. In the proposed high-protein hybrid bread formulation, dry-processed protein ingredients from faba bean and carob were applied and provide a substantial amount of non-wheat protein. The increased content of plant protein with higher protein quality in HPHB and formulations of its kind, could improve the capacity of the staple food bread to cover protein needs in future plant-based diets. The results also suggest that a replacement of regular wheat bread by high-protein hybrid breads could be beneficial in currently con-

1081 Conflicts of interest

1086 S.r.l.) for performing compositional and amino acid analysis, respectively; and Jonas Atzler for his help with single electron mi-

1093 The following abbreviations are used in this manuscript:

AA Amino acid SAA Sulphur-containing amino acids ANC Antinutritional compound HPI High-protein ingredient

High-protein hybrid bread **HPHB RWB** Reference wheat bread LCA Life cycle assessment HP High-protein TM Torque maximum

PMT Peak maximum time PV Peak viscosity FV Final viscosity

 V_{total} Total gas volume produced V_{lost} Volume of CO2 lost V_{ret} Volume of gas retained

Maximum height of dough development H_M

SV Specific volume L*crust Lightness of crust L*crumb Lightness of crumb In vitro protein digestibility **TNBS** Trinitrobenzenesulfonic acid CCasein SF Sova flour SPI Soya protein isolate BW Body weight PER Protein efficiency ratio L-BAPA N- α -benzoyl-L-arginine-4-nitroanilide Trypsin inhibitor unit TIU TAU Trypsin activity unit **ABTS** 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid Total phenolic content TPC **TEAC** Trolox equivalent antioxidant capacity **FRAP** Ferric-reducing antioxidant power DPPH 2,2-diphenyl-1-picrylhydrazyl ODA Quantitative descriptive analysis ANOVA Analysis of variance ΑX Arabinoxylans %DM Percentage based on dry matter proteinE Percentage of calories provided by protein %E Percentage based on energy **SFA** Saturated fatty acids **MUFA** Mono unsaturated fatty acids **PUFA** Poly unsaturated fatty acids %Protein Percentage based on protein AAA Aromatic amino acids AAS Amino acid score Protein digestibility corrected amino acid score **PDCAAS** Trypsin inhibitor activity TIA Half maximal effective concentration EC_{50}

References

1095

1096

1097

1098

1099

1100

1101

1102

1103

1104

1105

1106

1107

1108

1109

1111

1112

1113

1114

1116

1117

1118

1119

1120

1121

1122

1123

- 1 J. Mellentin, 10 Key Trends in Food, Nutrition and Health 1150 2018, 2017.
- 2 W. Willett, J. Rockström, B. Loken, M. Springmann, T. Lang, 1152 S. Vermeulen, T. Garnett, D. Tilman, F. DeClerck, A. Wood, 1153 M. Jonell, M. Clark, L. Gordon, J. Fanzo, C. Hawkes, R. Zu-1154 rayk, J. A. Rivera, W. D. Vries, L. Sibanda, A. Afshin, 1155 A. Chaudhary, M. Herrero, R. Agustina, F. Branca, A. Lartey, 1156 S. Fan, B. Crona, E. Fox, V. Bignet, M. Troell, T. Lindahl, 1157 S. Singh, S. Cornell, S. Reddy, S. Narain, S. Nishtar and 1158 C. Murray, Food in the Anthropocene: the EAT-Lancet Com-1159 mission on healthy diets from sustainable food systems, *The 1160 Lancet Commissions*, 2019, **18**, 1–47.
- 3 E. A. Frison, From Uniformity To Diversity: a paradigm shift 1162 from industrial agriculture to diversified agroecological sys- 1163 tems., International Panel of Experts on Sustainable Food 1164 systems Technical Report 3, 2016.
- 5 M. Friedman, Nutritional Value of Proteins from Different 1170 Food Sources. A Review, *J. Agric. Food Chem.*, 1996, **44**, 6–1171 29.
- 6 A. Etemadi, R. Sinha, M. H. Ward, B. I. Graubard, M. Inoue- 1173 Choi, S. M. Dawsey and C. C. Abnet, Mortality from different 1174 causes associated with meat, heme iron, nitrates, and nitrites 1175 in the NIH-AARP Diet and Health Study: Population based 1176 cohort study, *BMJ*, 2017, **357**, 1–11.
- 7 D. J. Jenkins, C. W. Kendall, A. Marchie and L. S. Augustin,

Too much sugar, too much carbohydrate, or just too much?, *Am. J. Clin. Nutr.*, 2004, **79**, 711–712.

1124

1125

1126

1127

1128

1129

1130

1131

1132

1133

1134

1135

1136

1137

1138

1139

1140

1141

1142

1143

1144

1145

1146

1147

1148

- 8 J. I. Boye, S. Aksay, S. Roufik, S. Ribéreau, M. Mondor, E. Farnworth and S. H. Rajamohamed, Comparison of the functional properties of pea, chickpea and lentil protein concentrates processed using ultrafiltration and isoelectric precipitation techniques, *Food Res. Int.*, 2010, **43**, 537–546.
- 9 E. S. Jensen, M. B. Peoples and H. Hauggaard-Nielsen, Faba bean in cropping systems, *F. Crop. Res.*, 2010, **115**, 203–216.
- 10 M. C. Vaz Patto, R. Amarowicz, A. N. Aryee, J. I. Boye, H. J. Chung, M. A. Martín-Cabrejas and C. Domoney, Achievements and Challenges in Improving the Nutritional Quality of Food Legumes, Crit. Rev. Plant Sci., 2014, 34, 105–143.
- 11 A. Iqbal, I. A. Khalil, N. Ateeq and M. Sayyar Khan, Nutritional quality of important food legumes, *Food Chem.*, 2006, **97**, 331–335.
- 12 F. W. Sosulski and G. I. Imafidon, Amino Acid Composition and Nitrogen-to-Protein Conversion Factors for Animal and Plant Foods, *J. Agric. Food Chem.*, 1990, **38**, 1351–1356.
- 13 M. Henchion, M. Hayes, A. Mullen, M. Fenelon and B. Tiwari, Future Protein Supply and Demand: Strategies and Factors Influencing a Sustainable Equilibrium, *Foods*, 2017, **6**, 53.
- 14 F. Boukid, E. Zannini, E. Carini and E. Vittadini, Pulses for bread fortification: A necessity or a choice?, *Trends Food Sci. Technol.*, 2019, 88, 416–428.
- 15 K. Dewettinck, F. Van Bockstaele, B. Kühne, D. Van de Walle, T. M. Courtens and X. Gellynck, Nutritional value of bread: Influence of processing, food interaction and consumer perception, *J. Cereal Sci.*, 2008, 48, 243–257.
- 16 H. Lopez, A. Adam, F. Leenhardt, A. Scalbert and C. Remesy, Control of the nutritional value of bread, *Ind. des Cereal.*, 2001, **124**, 15–20.
- 17 S. C. Magalhães, M. Taveira, A. R. Cabrita, A. J. Fonseca, P. Valentão and P. B. Andrade, European marketable grain legume seeds: Further insight into phenolic compounds profiles, *Food Chem.*, 2017, 215, 177–184.
- 18 R. Campos-Vega, G. Loarca-Piña and B. D. Oomah, Minor components of pulses and their potential impact on human health, *Food Res. Int.*, 2010, **43**, 461–482.
- 19 C. B. J. Villarino, V. Jayasena, R. Coorey, R. Foley, K. Fanning and S. K. Johnson, The effects of lupin (Lupinus angustifolius) addition to wheat bread on its nutritional, phytochemical and bioactive composition and protein quality, *Food Res. Int.*, 2015, 76, 58–65.
- 20 L.-P. D. Marchais, M. Foisy, S. Mercier, S. Villeneuve and M. Mondor, Bread-making potential of pea protein isolate produced by a novel ultrafiltration/diafiltration process, *Procedia Food Sci.*, 2011, 1, 1425–1430.
- 21 A. Angioloni and C. Collar, High legume-wheat matrices: An alternative to promote bread nutritional value meeting dough viscoelastic restrictions, *Eur. Food Res. Technol.*, 2012, **234**, 273–284.
- 22 Y. Wang, P. Sorvali, A. Laitila, N. H. Maina, R. Coda and

K. Katina, Dextran produced in situ as a tool to improve the 1233 quality of wheat-faba bean composite bread, Food Hydrocoll., 1234 2018, 84, 396-405.

1178

1179

1180

1181

1182

1183

1184

1185

1186

1187

1188

1189

1190

1191

1192

1193

1194

1195

1196

1197

1198

1199

1201

1202

1203

1204

1205

1206

1207

1208

1209

1210

1211

1214

- 23 V. Turfani, V. Narducci, A. Durazzo, V. Galli and M. Carcea, 1236 Technological, nutritional and functional properties of 1237 wheat bread enriched with lentil or carob flours, LWT - Food 1238 Sci. Technol., 2017, 78, 361-366.
- 24 C. Vidal-Valverde, J. Frias, C. Diaz-Pollan, M. Fernandez, 1240 M. Lopez-Jurado and G. Urbano, Influence of Processing on 1241 Trypsin Inhibitor Activity of Faba Beans and Its Physiological 1242 Effect, J. Agric. Food Chem., 1997, 45, 3559-3564.
- 25 G. S. Gilani, C. W. Xiao and K. A. Cockell, Impact of antinutri- 1244 tional factors in food proteins on the digestibility of protein 1245 and the bioavailability of amino acids and on protein quality, 1246 Br. J. Nutr., 2012, 108, S315-S332.
- 26 K. Crépon, P. Marget, C. Peyronnet, B. Carrouée, P. Arese 1248 and G. Duc, Nutritional value of faba bean (Vicia faba L.) 1249 seeds for feed and food, F. Crop. Res., 2010, 115, 329-339. 1250
- 27 K. Khamassi, F. Ben Jeddi, D. Hobbs, J. Irigoven, F. Stoddard, 1251 D. M. O'sullivan and H. Jones, A baseline study of vicine- 1252 convicine levels in faba bean (Vicia faba L.) germplasm, 1253 Plant Genet. Resour. Characterisation Util., 2013, 11, 250-1254 257.
- 28 L. Luzzatto and P. Arese, Favism and glucose-6-phosphate 1256 dehydrogenase deficiency, N. Engl. J. Med., 2018, 378, 60-1257
- 29 R. Coda, J. Varis, M. Verni, C. G. Rizzello and K. Katina, 1259 Improvement of the protein quality of wheat bread through 1260 faba bean sourdough addition, LWT - Food Sci. Technol., 1261 2017, 82, 296-302.
- 30 A. Hoehnel, C. Axel, J. Bez, E. K. Arendt and E. Zannini, 1263 Comparative analysis of plant-based high-protein ingredi- 1264 ents and their impact on quality of high-protein bread, J. 1265 Cereal Sci., 2019, 89, 1-8.
- 31 M. A. Schutyser, P. J. Pelgrom, A. J. van der Goot and R. M. 1267 1212 Boom, Dry fractionation for sustainable production of func- 1268 tional legume protein concentrates, Trends Food Sci. Technol., 1269 2015, 45, 327–335. 1215
- 32 M. Vogelsang-O'Dwyer, J. Bez, I. L. Petersen, M. S. Joehnke, 1271 1216 J. C. Sørensen, A. Detzel, M. Busch, M. Krueger, J. A. 1272 1217 O'Mahony, E. K. Arendt and E. Zannini, Comparison of Faba 1273 1218 Bean Protein Ingredients Produced Using Dry Fractionation 1274 1219 and Isoelectric Precipitation: Techno-Functional, Nutritional 1275 1220 and Environmental Performance, Foods, 2020, 9, 1-25. 1221
- 33 L. Ispirvan, M. Heitmann, A. Hoehnel, E. Zannini and 1277 1222 E. Arendt, Optimization and Validation of an HPAEC-PAD 1278 1223 Method for the Quantification of FODMAPs in Cereals and 1279 Cereal-Based Products, J. Agric. Food Chem., 2019, 67, 1280 1225 4384-4392. 1226
- 34 M. S. Joehnke, A. Rehder, S. Sørensen, C. Bjergegaard, 1282 1227 J. C. Sørensen and K. E. Markedal, In Vitro Digestibility of $_{\rm 1283}$ 1228 Rapeseed and Bovine Whey Protein Mixtures, J. Agric. Food 1284 1229 Chem., 2018, 66, 711–719. 1230
- M. S. Joehnke, R. Lametsch and J. C. Sørensen, Improved 1286 in vitro digestibility of rapeseed napin proteins in mixtures 1232

- with bovine beta-lactoglobulin, Food Res. Int., 2019, 123, 346-354.
- 36 P. G. Reeves, Components of the AIN-93 Diets as Improvements in the AIN-76A Diet, Symposium: Animal Diets for Nutritional and Toxicological Research, 1997, 127, 838S-841S.
- 37 I. L. Petersen, H. C. B. Hansen, H. W. Ravn, J. C. Sørensen and H. Sørensen, Metabolic effects in rapeseed (Brassica napus L.) seedlings after root exposure to glyphosate, *Pestic*. Biochem. Physiol., 2007, 89, 220-229.
- 38 C. Bjergegaard, H. Simonsen and H. Sørensen, Determination of heterocyclic compounds by micellar electrokinetic capillary chromatography, J. Chromatogr. A, 1994, 680, 561-569.
- 39 R. Amarowicz, U. N. Wanasundara, M. Karamać and F. Shahidi, Antioxidant activity of ethanolic extract of mustard seed, Nahrung - Food, 1996, 40, 261-263.
- 40 R. Amarowicz and B. Raab, Antioxidative activity of leguminous seed extracts evaluated by chemiluminescence methods, Zeitschrift fur Naturforsch. Sect. C - J. Biosci., 1997, 52, 709-712.
- 41 R. Re, N. Pellegrini, A. Proteggente, A. Pannala, M. Yang, C. Rice-Evans, R. Verdejo and M. Shaffer, Antioxidant activity applying an improved ABTS radical cation decolorization assay, Free Radic. Biol. Med., 1999, 26, 1231-1237.
- I. F. Benzie and J. J. Strain, Ferric reducing/antioxidant power assay: Direct measure of total antioxidant activity of biological fluids and modified version for simultaneous measurement of total antioxidant power and ascorbic acid concentration, Methods Enzymol., 1999, 299, 15-27.
- Amarowicz, M. Karamać, H. Kmita-Głazewska, 43 R. Troszyńska and H. Kozłowska, Antioxidant activity of phenolic fractions of everlasting pea, faba bean and broad bean, J. Food Lipids, 1996, 3, 199-211.
- 44 T. Amoriello, V. Turfani, V. Galli, F. Mellara and M. Carcea, Evaluation of a new viscometer performance in predicting the technological quality of soft wheat flour, Cereal Chem., 2016, 93, 364-368.
- 45 G. K. Chandi and K. Seetharaman, Optimization of gluten peak tester: A statistical approach, J. Food Qual., 2012, 35, 69-75.
- 46 A. Marti, A. Ulrici, G. Foca, L. Quaglia and M. A. Pagani, Characterization of common wheat flours (Triticum aestivum L.) through multivariate analysis of conventional rheological parameters and gluten peak test indices, LWT - Food Sci. Technol., 2015, 64, 95-103.
- 47 A. Goldstein, L. Ashrafi and K. Seetharaman, Effects of cellulosic fibre on physical and rheological properties of starch, gluten and wheat flour, Int. J. Food Sci. Technol., 2010, 45, 1641-1646.
- 48 L. Day, Wheat gluten: Production, properties and application in: Handbook of Food Proteins, Woodhead Publishing Limited, Cambridge, 2011, ch. 10, pp. 267-288.
- 49 L. Yu, J. Perret, T. Parker and K. G. Allen, Enzymatic modification to improve the water-absorbing and gelling properties

of psyllium, Food Chem., 2003, 82, 243-248.

1287

1288

1289

1290

1291

1292

1293

1294

1295

1296

1297

1298

1302

1303

1304

1324

1325

1326

1327

1328

1329

1330

1335

1337

1338

1339

- 50 E. López, Influence of the addition of lupine protein isolate ¹³⁴² on the protein and technological characteristics of dough ¹³⁴³ and fresh bread with added Brea Gum, *Food Sci. Technol.*, ¹³⁴⁴ 2014, **34**, 195–203.
- 51 C. M. Rosell and A. Foegeding, Interaction of hydroxypropy- 1346 lmethylcellulose with gluten proteins: Small deformation 1347 properties during thermal treatment, *Food Hydrocoll.*, 2007, 1348 **21**, 1092–1100.
- 52 A. Verma and R. Mogra, Psyllium (Plantago ovata) Husk: 1350 A Wonder Food for Good Health, *Int. J. Sci. Res.*, 2015, **4**, 1351 1581–1585.
- 1299 53 A. Farahnaky, H. Askari, M. Majzoobi and G. Mesbahi, The 1353 impact of concentration, temperature and pH on dynamic 1354 rheology of psyllium gels, *J. Food Eng.*, 2010, **100**, 294–301. 1355
 - 54 R. Hoseney, *Rheology of Doughs and Batters in: Cereal Science* 1356 and *Technology*, American Association of Cereal Chemists 1357 Inc., 2nd edn, 1994, pp. 213–228.
- 1305 55 C. M. Courtin and J. A. Delcour, Arabinoxylans and endoxy- 1359 1306 lanases in wheat flour bread-making, *J. Cereal Sci.*, 2002, 1360 1307 **35**, 225–243.
- M. H. Fischer, N. Yu, G. R. Gray, J. Ralph, L. Anderson and 1362
 J. A. Marlett, The gel-forming polysaccharide of psyllium 1363
 husk (Plantago ovata Forsk), Carbohydr. Res., 2004, 339, 1364
 2009–2017.
- 57 V. Van Craeyveld, J. A. Delcour and C. M. Courtin, Ex- 1366
 tractability and chemical and enzymic degradation of psyl- 1367
 lium (Plantago ovata Forsk) seed husk arabinoxylans, Food 1368
 Chem., 2009, 112, 812–819.
- 58 C. M. Courtin, G. G. Gelders and J. A. Delcour, Use of two 1370 endoxylanases with different substrate selectivity for under- 1371 standing arabinoxylan functionality in wheat flour bread- 1372 making, *Cereal Chem.*, 2001, **78**, 564–571.
- 59 M. Wang, G. Oudgenoeg, T. Van Vliet and R. J. Hamer, In- 1374
 teraction of water unextractable solids with gluten protein: 1375
 Effect on dough properties and gluten quality, *J. Cereal Sci.*, 1376
 2003, 38, 95–104.
 - 60 M. Wang, T. Van Vliet and R. J. Hamer, How gluten proper- 1378 ties are affected by pentosans, *J. Cereal Sci.*, 2004, **39**, 395–1379 402.
 - 61 A. Paraskevopoulou, E. Provatidou, D. Tsotsiou and 1381 V. Kiosseoglou, Dough rheology and baking performance of 1382 wheat flour-lupin protein isolate blends, *Food Res. Int.*, 2010, 1383 **43**, 1009–1016.
- 62 C. B. Villarino, V. Jayasena, R. Coorey, S. Chakrabarti-Bell 1385 and S. K. Johnson, The effects of Australian sweet lupin 1386 (ASL) variety on physical properties of flours and breads, 1387 LWT - Food Sci. Technol., 2015, **60**, 435–443.
 - 63 J. A. Gray and J. N. Bemiller, Bread staling: Molecular basis 1389 and control, *Compr. Rev. Food Sci. Food Saf.*, 2003, **2**, 1–21. 1390
 - 64 C. Fadda, A. M. Sanguinetti, A. Del Caro, C. Collar and 1391 A. Piga, Bread staling: Updating the view, *Compr. Rev. Food* 1392 *Sci. Food Saf.*, 2014, **13**, 473–492.
 - 65 M. S. Izydorczyk and J. E. Dexter, Barley β -glucans and 1394

- arabinoxylans: Molecular structure, physicochemical properties, and uses in food products-a Review, *Food Res. Int.*, 2008, **41**, 850–868.
- 66 M. S. Butt, M. Tahir-Nadeem, Z. Ahmad and M. T. Sultan, Xylanases and their applications in baking industry, Food Technol. Biotechnol., 2008, 46, 22–31.
- 67 T. Maeda and N. Morita, Flour quality and pentosan prepared by polishing wheat grain on breadmaking, *Food Res. Int.*, 2003, **36**, 603–610.
- 68 Z. Czuchajowska, B. Paszczynska and Y. Pomeranz, Functional Properties of Psyllium in Wheat-Based Products, *Cereal Chem.*, 1992, **69**, 516–520.
- 69 G. Della Valle, H. Chiron, L. Cicerelli, K. Kansou, K. Katina, A. Ndiaye, M. Whitworth and K. Poutanen, Basic knowledge models for the design of bread texture, *Trends Food Sci. Technol.*, 2014, 36, 5–14.
- 70 E. Purlis, Browning development in bakery products A review, *J. Food Eng.*, 2010, **99**, 239–249.
- 71 European Parliament & Council, Regulation (EU) No 1169/2011 of the European Parliament and of the Council on the provision of food information to consumers, 2011.
- 72 European Parliament & Council, Regulation (EC) No 1924/2006 of the European Parliament and of the Council on Nutrition and health claims made on foods, 2006.
- 73 W. C. Willett and M. J. Stampfer, Rebuilding the Food Pyramid, *Sci. Am.*, 2003, **288**, 64–71.
- 74 J. Salméron, J. E. Manson, M. J. Stampfer, G. A. Colditz, A. L. Wing and W. C. Willett, Dietary Fiber, Glycemic Load, and Risk of Non-insulin-dependent Diabetes Mellitus in Women, J. Am. Med. Assoc., 1997, 277, 472–477.
- 75 D. S. Ludwig, M. A. Pereira, C. H. Kroenke, J. E. Hilner, L. Van Horn, M. L. Slattery and D. R. Jacobs, Dietary fiber, weight gain, and cardiovascular disease risk factors in young adults, J. Am. Med. Assoc., 1999, 282, 1539–1546.
- 76 S. Liu, W. C. Willett, M. J. Stampfer, F. B. Hu, M. Franz, L. Sampson, C. H. Hennekens and J. A. E. Manson, A prospective study of dietary glycemic load, carbohydrate intake, and risk of coronary heart disease in US women, *Am. J. Clin. Nutr.*, 2000, 71, 1455–1461.
- 77 S. Franceschi, L. Dal Maso, L. Augustin, E. Negri, M. Parpinel, P. Boyle, D. Jenkins and C. La Vecchia, Dietary glycemic load and colorectal cancer risk, *Ann. Oncol.*, 2001, 12, 173–178.
- 78 L. Augustin, L. Dal Maso, C. La Vecchia, M. Parpinel, E. Negri, S. Vaccarella, C. Kendall, D. Jenkins and S. Franceschi, Dietary glycemic index and glycemix load, and breast cancer risk: A case-control study, *Ann. Oncol.*, 2001, 12, 1553–1538.
- 79 G. Zhang, Z. Ao and B. R. Hamaker, Controlling the delivery of glucose in foods in: Designing Functional Foods: Measuring and Controlling Food Structure Breakdown and Nutrient Absorption, 2009, ch. 21, pp. 547–571.
- 80 L. (Lucy) Yu, H. Lutterodt and Z. Cheng, Chapter 4 Beneficial Health Properties of Psyllium and Approaches to Improve Its

Functionalities, Adv. Food Nutr. Res., 2008, **55**, 193–220.

1396

1397

1398

1399

1401

1402

1404

1405

1407

1408

1409

1416

1418

1425

1426

1427

1428

1436

- 81 S. H. Holt, J. C. Brand-Miller and P. A. Stitt, The effects of 1451 equal-energy portions of different breads on blood glucose 1452 levels, feelings of fullness and subsequent food intake, J. Am. 1453 Diet. Assoc., 2001, 101, 767-773.
- 82 L. J. Appel, F. M. Sacks, V. J. Carey, E. Obarzanek, J. F. 1455 Swain, E. R. Miller, P. R. Conlin, T. P. Erlinger, B. A. Rosner, 1456 N. M. Laranjo, J. Charleston, P. McCarron and L. M. Bishop, 1457 Effects of protein, monounsaturated fat, and carbohydrate 1458 intake on blood pressure and serum lipids: Results of the 1459 OmniHeart randomized trial, J. Am. Med. Assoc., 2005, 294, 1460 2455-2464.
- 83 EFSA, EFSA Panel on Dietetic Products, Nutrition, and Aller- 1462 gies (NDA); Scientific Opinion on Dietary Reference Values 1463 for carbohydrates and dietary fibre, The EFSA Journal, 2010, 1464 8, 1-77. 1410
- 84 EFSA, EFSA Panel on Dietetic Products, Nutrition, and Al-1466 1411 lergies (NDA); Scientific Opinion on Dietary Reference Val- 1467 1412 ues for fats, including saturated fatty acids, polyunsaturated 1468 1413 fatty acids, monounsaturated fatty acids, trans fatty acids, 1469 and cholesterol, The EFSA Journal, 2010, 8, 1–107. 1415
 - 85 EFSA, EFSA Panel on Dietetic Products, Nutrition, and Aller- 1471 gies (NDA); Dietary reference values for sodium, The EFSA 1472 Journal, 2019, 17, 1-191.
- C. Silow, C. Axel, E. Zannini and E. K. Arendt, Current status 1474 86 1419 of salt reduction in bread and bakery products - A review, J. 1475 1420 Cereal Sci., 2016, 72, 135–145. 1421
- E. J. Landry, S. J. Fuchs and J. Hu, Carbohydrate composi- 1477 1422 tion of mature and immature faba bean seeds, J. Food Com- 1478 1423 pos. Anal., 2016, 50, 55-60. 1424
 - 88 C. Bengoechea, A. Romero, A. Villanueva, G. Moreno, 1480 M. Alaiz, F. Millán, A. Guerrero and M. C. Puppo, Composi-1481 tion and structure of carob (Ceratonia siliqua L.) germ pro- 1482 teins, Food Chem., 2008, 107, 675-683.
- Joint FAO/WHO/UNU Expert Consultation, Protein and 1484 1429 Amino Acid Requirements in Human Nutrition, WHO Tech-1485 nical Report Series, 2007, 935, 1-265. 1431
- 90 EFSA, (European Food Safety Authority) Dietary Reference 1487 1432 Values for nutrients: Summary report, EFSA Support. Publ., 1488 1433 2017, e15121, 1-92.
 - 91 J. Boye, R. Wijesinha-Bettoni and B. Burlingame, Protein 1490 quality evaluation twenty years after the introduction of the 1491 protein digestibility corrected amino acid score method, Br. 1492 J. Nutr., 2012, 108, S183-S211.
- 92 S. Avilés-Gaxiola, C. Chuck-Hernández and S. O. Serna 1494 Saldívar, Inactivation Methods of Trypsin Inhibitor in 1495 1440 Legumes: A Review, J. Food Sci., 2018, 83, 17–29. 1441
- 93 T. Bohn, F. Carriere, L. Day, A. Deglaire, L. Egger, D. Freitas, 1497 1442 M. Golding, S. Le Feunteun, A. Macierzanka, O. Menard, 1498 B. Miralles, A. Moscovici, R. Portmann, I. Recio, D. Rémond, 1499 1444 V. Santé-Lhoutelier, T. J. Wooster, U. Lesmes, A. R. Mackie 1500 1445 and D. Dupont, Correlation between in vitro and in vivo data 1501 on food digestion. What can we predict with static in vitro 1502 108 B. J. Venn and J. I. Mann, Cereal grains, legumes and dia-1447 digestion models?, Crit. Rev. Food Sci. Nutr., 2018, 58, 2239-1503 1448 2261. 1449

- 94 D. J. Millward, D. K. Layman, D. Tomé and G. Schaafsma, Protein quality assessment: Impact of expanding understanding of protein and amino acid needs for optimal health, Am. J. Clin. Nutr., 2008, 87(suppl), 1576S-1581S.
- 95 E. Ha and M. B. Zemel, Functional properties of whey, whey components, and essential amino acids: Mechanisms underlying health benefits for active people (Review), J. Nutr. Biochem., 2003, 14, 251-258.
- 96 B. Wróblewska, J. Juśkiewicz, B. Kroplewski, A. Jurgoński, E. Wasilewska, D. Złotkowska and L. Markiewicz, The effects of whey and soy proteins on growth performance, gastrointestinal digestion, and selected physiological responses in rats, Food Funct., 2018, 9, 1500-1509.
- 97 C. A. Butts, J. A. Monro and P. J. Moughan, In vitro determination of dietary protein and amino acid digestibility for humans, Br. J. Nutr., 2012, 108, 282-287.
- 98 P. Feillet and T. M. Roulland, Caroubin: A gluten-like protein isolated from carob bean germ, Cereal Chem., 1998, 75, 488-492.
- 99 H. D. Belitz, F. Lynen and J. K. Weder, Comparative studies on the inhibitory action of some legume seeds, potato tubers, and bran against human and bovine proteinases, Z. Lebensm. Unters. Forsch., 1982, 174, 442-446.
- 100 H. Khazaei, R. W. Purves, J. Hughes, W. Link, D. M. O'Sullivan, A. H. Schulman, E. Björnsdotter, F. Geu-Flores, M. Nadzieja, S. U. Andersen, J. Stougaard, A. Vandenberg and F. L. Stoddard, Eliminating vicine and convicine, the main anti-nutritional factors restricting faba bean usage, Trends Food Sci. Technol., 2019, 91, 549-556.
- V. Gallo, O. A. Skorokhod, L. F. Simula, T. Marrocco, E. Tambini, E. Schwarzer, P. Marget, G. Duc and P. Arese, G6PDdeficient subjects after ingestion of low No red blood cell damage and no hemolysis in vicine/convicine Vicia faba seeds, Blood, 2018, 131, 1617–1621.
- 102 Fob, The Federation of Bakers: Calories in Bread, Factsheets, 2015, No. 20, 1-3.
- M. Carocho and I. C. Ferreira, A review on antioxidants, 103 prooxidants and related controversy: Natural and synthetic compounds, screening and analysis methodologies and future perspectives, Food Chem. Toxicol., 2013, 51, 15-25.
- 104 A. Scalbert, C. Manach, C. Morand, C. Rémésy and L. Jiménez, Dietary polyphenols and the prevention of diseases, Crit. Rev. Food Sci. Nutr., 2005, 45, 287-306.
- 105 J. W. Anderson, Whole grains protect against atherosclerotic cardiovascular disease, Proc. Nutr. Soc., 2003, 62, 135-142.
- 106 L. Chatenoud, A. Tavani, C. La Vecchia, D. R. Jacobs, E. Negri, F. Levi and S. Franceschi, Whole grain food intake and cancer risk, Int. J. Cancer, 1998, 77, 24-28.
- 107 L. Chatenoud, C. La Vecchia, S. Franceschi, A. Tavani, D. R. Jacobs, M. T. Parpinel, M. Soler and E. Negri, Refined-cereal intake and risk of selected cancers in Italy, Am. J. Clin. Nutr., 1999, **70**, 1107–1110.
- betes, Eur. J. Clin. Nutr., 2004, 58, 1443-1461.

- 109 A. Durazzo, V. Turfani, V. Narducci, E. Azzini, G. Maiani and M. Carcea, Nutritional characterisation and bioactive com-1505 ponents of commercial carobs flours, Food Chem., 2014, 153, 1506 109-113. 1507
- 110 D. W. Hatcher and J. E. Kruger, Simple phenolic acids in 1508 flours prepared from Canadian wheat: Relationship to ash 1509 content, color, and polyphenol oxidase activity, Cereal Chem., 1510 1997, 74, 337-343. 1511
- 111 S. Ragaee, I. Guzar, N. Dhull and K. Seetharaman, Effects of 1512 fiber addition on antioxidant capacity and nutritional quality 1513 of wheat bread, LWT - Food Sci. Technol., 2011, 44, 2147-1514 2153. 1515
- 112 A. Fardet, E. Rock and C. Rémésy, Is the in vitro antioxidant 1516 potential of whole-grain cereals and cereal products well re-1517 flected in vivo?, J. Cereal Sci., 2008, 48, 258-276. 1518
- 113 I. Siró, E. Kápolna, B. Kápolna and A. Lugasi, Functional food. Product development, marketing and consumer 1520 acceptance-A review, Appetite, 2008, 51, 456-467. 1521
- 114 A. Gámbaro, P. Varela, A. Giménez, A. Aldrovandi, S. M. 1522 1523 Fiszman and G. Hough, Textural quality of white pan bread by sensory and instrumental measurements, J. Texture Stud., 1524 2002, 33, 401-413. 1525
- 115 M. J. Callejo, Present situation on the descriptive sensory 1526 analysis of bread, J. Sens. Stud., 2011, 26, 255-268. 1527
- V. Lotong, E. Chambers IV and D. H. Chambers, Determina-1528 tion of the sensory attributes of wheat sourdough bread, J. 1529 Sens. Stud., 2000, 15, 309-326. 1530
- 117 R. L. Heiniö, Sensory Attributes of Bakery Products in: Bakery 1531 Products Science and Technology: Second Edition, John Wiley & Sons, 2008, 2014, ch. 22, pp. 391-407. 1533



ELECTRONIC SUPPLEMENTARY INFORMATION (ESI)

for Food & Function article "Enhancing the nutritional profile of regular wheat bread while maintaining technological quality and adequate sensory attributes"

Andrea Hoehnel, Jürgen Bez, Iben Lykke Petersen, Ryszard Amarowicz, Jerzy Juśkiewicz, Elke K. Arendt, *a,e and Emanuele Zannini a

Microbiological Shelf Life and Water Activity of Reference Wheat Bread (RWB) and High-Protein Hybrid Bread (HPHB)

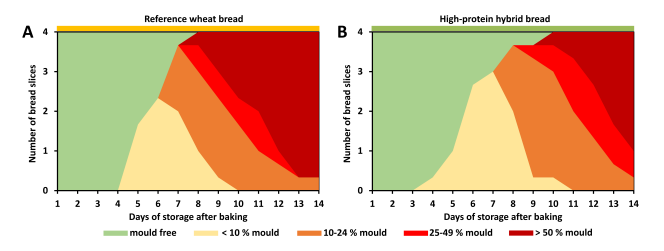


Fig. 1 Microbiological shelf life of (A) RWB and (B) HPHB as indicated by ambient air challenge test. Results represent the mean of three independently performed challenge tests.

Results and Discussion

In addition to crumb staling, the shelf life of bread is affected by microbial deterioration. While also bacteria and yeast can cause bread spoilage, a contamination with fungal spores from the bakery environment after baking is considered the most common reason. Mold growth typically shows a positive correlation with water availability in the food product; the critical water activity, however, varies with fungal species, temperature and substrate. Apart from an unpleasant visual experience for consumers, mould

spoilage can cause the formation of off-flavours, allergenic com-

pounds and mycotoxins, potentially even before visibility of fungal growth.³ It also leads to a substantial amount of food waste - in UK households an estimated 20 % of bread goes to waste due to mould growth. 4,5 Therefore, susceptibility to mould deterioration represents a food safety hazard and indicator for economic loss and should be considered when bread quality is evaluated. The microbial shelf life of both bread formulations was monitored in an ambient air challenge test. The results are presented in Figure 1. A slight tendency towards earlier onset of mould growth for HPHB was observed. The results also suggest a deceleration of mould growth in HPHB represented by later onset of stages 3 - 5 (10 to > 50 % of slices covered in mould). However, these tendencies cannot be considered significant differences and the experiment generally indicated a similar microbial shelf life of HPHB and RWB. This observation can be supported by very similar water activities measured for both formulations (RWB 0.945 ± 0.003 , HPHB 0.943 ± 0.003).

 $[^]a$ University College Cork, School of Food and Nutritional Sciences, College Road, Ireland. Tel: $+353\ 21\ 490\ 2064;$ E-mail: e.arendt@ucc.ie

^b Fraunhofer Institute for Process Engineering and Packaging, 85354 Freising, Germany

^c Department of Food Science, University of Copenhagen, Rolighedsvej 26, 1958 Frederiksberg C., Denmark.

d Institute of Animal Reproduction and Food Research of the Polish Academy of Sciences, Tuwima. St. 10. 10-748 Olsztvn. Poland.

^e APC Microbiome Ireland, Cork, Ireland.

Materials and Methods used for the Determination of Microbiological Shelf Life and Water Activity

Microbiological shelf life of the breads was evaluated using an ambient mould challenge test as described by Dal Bello et al. 6 with some modifications. Bread loaves where sliced in a sterile manner to obtain four slices of 20 mm thickness per loaf. Instead of a treatment with conidial solutions of fungi, each slice was microbiologically challenged by exposure to the bakery ambient air for 5 min on each side. The slices were separately packed in sterile plastic bags which were heat sealed. To guarantee comparable aerobic conditions in all bags, a filter pipette tip was inserted. During a storage period of 14 days (at room temperature), mould growth was visually evaluated. Based on the percentage of slice area covered with fungal growth, slices were sorted into five categories as follows: 0 % - mould free, <10 % mould, 10-24 % mould, 25-49 % mould, >50 % mould. Four slices were monitored from each of three batches per formulation. Water activity of the fresh bread crumb was measured using a water activity meter (HygroLab, Rotronic, Basserdorf, Switzerland).

Abbreviations

The following abbreviations were used:

HPHB High-protein hybrid bread RWB Reference wheat bread

References

- 1 R. A. Knight and E. M. Menlove, Effect of the bread-baking process on destruction of certain mould spores, *J. Sci. Food Agric.*, 1961, **12**, 653–656.
- 2 N. Markova and L. Wadsö, A microcalorimetric method of studying mould activity as a function of water activity, *Int. Biodeterior. Biodegrad.*, 1998, **42**, 25–28.
- 3 N. Magan and G. Keshri, Advances in stored product protection. Proceedings of the 8th International Working Conference on Stored-Product Protection, York, UK, 22-26 July 2002: Use of electronic nose technology for the early detection of spoilage moulds in cereal products, CAB International, Wallingford, United Kingdom, 22nd edn, 2003, pp. 139–143.
- 4 L. Ventour, *Food waste report v2: The food we waste*, Waste & Resources Action Programme, 2008, vol. 2, pp. 1–237.
- 5 C. Axel, E. Zannini and E. K. Arendt, Mold spoilage of bread and its biopreservation: A review of current strategies for bread shelf life extension, *Crit. Rev. Food Sci. Nutr.*, 2017, **57**, 3528–3542.
- 6 F. Dal Bello, C. I. Clarke, L. A. Ryan, H. Ulmer, T. J. Schober, K. Ström, J. Sjögren, D. van Sinderen, J. Schnürer and E. K. Arendt, Improvement of the quality and shelf life of wheat bread by fermentation with the antifungal strain Lactobacillus plantarum FST 1.7, J. Cereal Sci., 2007, 45, 309–318.