

UCC Library and UCC researchers have made this item openly available. Please let us know how this has helped you. Thanks!

Title Membrane filtration and isoelectric precipitation technological approaches for the preparation of novel functional and sustainable protein isolate from lentils		
Author(s)	Alonso-Miravalles, Loreto; Jeske, Stephanie; Bez, Jurgen; Detzel, Andreas; Busch, Mirjam; Krueger, Martina; Wriessnegger, Clara L.; O'Mahony, James A.; Zannini, Emanuele; Arendt, Elke K.	
Publication date	2019	
Original citation Alonso-Miravalles, L., Jeske, S., Bez, J., Detzel, A., Busch, M., Kr M., Wriessnegger, Clara L., O'Mahony, J. A.; Zannini, E. and Are E. K. (2019) 'Membrane filtration and isoelectric precipitation technological approaches for the preparation of novel, functional and sustainable protein isolate from lentils', European Food Research a Technology, In Press		
Type of publication	Article (preprint)	
Link to publisher's version	https://link.springer.com/journal/217 Access to the full text of the published version may require a subscription.	
Rights	© 2019. This is a pre-print of an article to be published in European Food Research and Technology	
Item downloaded http://hdl.handle.net/10468/7868 from		

Downloaded on 2019-12-02T15:00:37Z



University College Cork, Ireland Coláiste na hOllscoile Corcaigh



Draft Manuscript for Review

Membrane filtration and isoelectric precipitation technological approaches for the preparation of novel functional and sustainable protein isolate from lentils

Journal:	European Food Research and Technology
Manuscript ID	EFRT-19-0094.R1
Manuscript Type:	Original paper
Date Submitted by the Author:	20-Apr-2019
Complete List of Authors:	Alonso-Miravalles, Loreto ; University College Cork, Food and Nutritional Sciences Jeske, Stephanie; University College Cork School of Food and Nutritiona Sciences Bez, Juergen; Fraunhofer-Institut fur Verfahrenstechnik und Verpackung Detzel, Andreas.; Institut fur Energie- und Umweltforschung Heidelberg GmbH Busch, Mirjam; Institut fur Energie- und Umweltforschung Heidelberg GmbH Krueger, Martina; Institut fur Energie- und Umweltforschung Heidelberg GmbH Wriessnegger, Clara Larissa.; Institut fur Energie- und Umweltforschung Heidelberg Heidelberg GmbH O'Mahony, James A. ; University College Cork, Food and Nutritional Sciences Zannini, Emanuele; University College Cork, Food and Nutritional Sciences Arendt, Elke; University College Cork, Food and Nutritional Sciences; University College Cork APC Microbiome Institute,
Keywords:	Lentil protein isolate, Ultrafiltration, Isoelectric precipitation, Physicochemical properties, Protein functionality, Life cycle assessment

SCHOLARONE[™] Manuscripts

Membrane filtration and isoelectric precipitation technological approaches for the preparation of novel, functional and sustainable protein isolate from lentils

- Loreto Alonso-Miravalles¹ and Stephanie Jeske¹, Juergen Bez², Andreas Detzel³, Mirjam Busch³, Martina
 Krueger³, Clara Larissa Wriessnegger³, James A. O'Mahony¹, Emanuele Zannini¹, Elke K. Arendt^{1,4}
- 5 ¹School of Food and Nutritional Sciences, University College Cork, Cork, Ireland
- 6 ²Fraunhofer Institute for Process Engineering and Packaging, Giggenhauser Str. 35, D-85354 Freising, Germany
- 7 ³IFEU Institut für Energie- und Umweltforschung Heidelberg GmbH, Im Weiher 10, 69121 Heidelberg
- 8 ⁴APC Microbiome Institute Ireland, University College Cork, Ireland
- 9 Corresponding author e-mail: e.arendt@ucc.ie; tel: +353 21 490 2064

10 Abstract

Isoelectric precipitation and ultrafiltration were investigated for their potential to produce protein products from lentils. Higher protein concentrations were obtained when ultrafiltration was used (>90%), whereas isoelectric precipitation resulted in higher contents of dietary fibre and some minerals (i.e., sodium and phosphorus). Differences in the functional properties between the two ingredients where found as the isoelectric precipitated ingredient showed lower protein solubilities over the investigated pH range (from 3 to 9) which can be linked to the slightly higher hydrophobicity values (2688.7) and total sulfhydryl groups (23.9 μ M/g) found in this sample. In contrast, the protein ingredient obtained by ultrafiltration was superior with regard to its solubility (48.3%; pH 7), fat-binding capacity (2.24 g/g), water holding capacity (3.96 g/g), gelling properties (11%; w/w), and foam-forming capacity (69.6%). The assessment of the environmental performance showed that both LPIs exhibited promising properties and low carbon footprints in comparison to traditional dairy proteins.

21 Keywords

Lentil protein isolate, Ultrafiltration, Isoelectric precipitation Physicochemical properties, Protein functionality, Life cycle assessment.

24 1 Introduction

The expected continued growth of the global population to 9.6 billion people by 2050 is creating a need to identify and develop solutions for the provision of high-quality food [1, 2]. In addition, the high demand for healthy, sustainable and cost-effective food protein ingredients by consumers is driving the investigation of new and innovative protein sources [3, 4]. Agriculture is one of the main contributors to climate change, and cattle farming faces particular sustainability challenges [5]. The conversion factor of feed protein to milk protein is about 14%, while the remaining 86% is "lost" for human nutrition [6]. Plant-based protein ingredients can serve as an alternative to animal-derived protein, due to their contribution to environmental sustainability, their role in addressing food security challenges and their cost-effectiveness [7]. However, replacing animal-based protein ingredients with those of plant origin is not easy, as significant differences exist between ingredients from both sources in composition, taste, digestibility and techno-functional properties. Nevertheless, research is advancing and several plant ingredients have been applied in a wide range of products [8, 9].-For instance, using extrusion, soya protein was processed into a highly fibrous texture simulating that of meat [10].- In addition to soy, other plant-based proteins have been studied as meat replacers such as canola, rapeseed, wheat gluten, peas and beans [11], bread guality and nutritional profile was improved with the addition of fermented faba bean [10], and sensory evaluation of a strawberry flavoured lupin based yogurt-like product showed good acceptability sensory properties [11].

In that respect, legumes are gaining increased attention, as they contain high amounts of protein, typically ranging between 20 and 40%, and are rich sources of essential amino acids such as lysine [12, 13]. Traditionally, they are consumed as whole, split or milled products [14] and approximately 5,481,120 ha are harvested and 6,315,858 tonnes of lentils are produced globally each year [15]. Lentil seeds are showing promising results for the preparation of protein flours, concentrates and isolates due to the lack of allergens and anti-nutritional compounds (e.g., isoflavones found in soya) and also as they are an affordable, sustainable and abundant raw material [16]. Various techniques and approaches such as wet fractionation (e.g., ultrafiltration and isoelectric precipitation) are used to separate and concentrate high levels of protein from other constituents [17, 18] in cereals and legumes. The physicochemical properties and functionality of these isolated protein ingredients are essential in the processing and formulation of food products, providing texture, taste and nutrition for a desirable and pleasurable product. These properties depend not only on the nature of the protein but also on the processing and isolation techniques used. Most studies focused on this subject have been conducted on dairy and soya [19, 20]

 and increasingly also on legumes in recent years [21–25]. The results of these studies indicate that the methods applied for isolation affect the composition and the physicochemical characteristics of <u>the</u> extracted protein ingredients. Jarpa-Parra et al. [26] highlights the need to establish a deeper connection between the extraction conditions of lentil protein and their influence on lentil protein functionality.

The aim of this work was to produce novel lentil protein isolates using two different technological approaches and to study the techno-functional properties (e.g., solubility, emulsifying, gelling properties) and environmental sustainability (life cycle assessment) of the ingredients based on the same raw material. The results obtained in this study will provide much needed information about the sustainability of the two different approaches and potential applications of the resultant ingredients in the development of novel, healthy and sustainable food product formulations.

63 2 Materials and Methods

2.1 Raw materials and chemicals

For extraction of lentil proteins, brown lentils of commercial quality (*Lens culinaris cv. Itaca*), provided by
 Agroservice Spa, San Severino Marche, Italy, were used as raw material. All chemicals used were purchased from
 Sigma-Aldrich (St Louis, Missouri, USA), unless otherwise stated.

68 2.2 Preparation of protein isolates

Lentil seeds were dehulled in an underrunner disc sheller (Streckel & Schrader GmbH, Germany) and the kernels and hulls were separated in an air classifier (Turboplex, Hosokawa Alpine AG, Augsburg, Germany). Kernels were milled using an impact mill (UPZ, Hosokawa Alpine AG, Germany) to a mean particle size (D50) of 21 µm. For extraction of protein, lentil flour was suspended in water at pH 7.5 to extract the high molecular weight proteins. The insoluble dietary fibre and lentil starch were then separated from the soluble high molecular weight proteins by decanting. Lentil protein isolate (LPI) was recovered from the resulting protein extract either by isoelectric precipitation (IEP) or by ultrafiltration (UF), as shown in Figure 1. LPI-IEP was isolated from the aqueous protein extract by acid precipitation at pH 4.5, which coincides with minimum solubility of lentil proteins [27]. Subsequently, the precipitated proteins were separated in a disc separator and the sediment was neutralized with 3 M NaOH, pasteurised (65°C, 30 min) and spray dried (Tm: 180°C, Tout: 75°C) to obtain the protein isolate powder. LPI-UF was extracted at 50°C using a polysulfone membrane with a molecular weight cut-off of 10 kDa followed by diafiltration with demineralized water (retentate:water; 1:1.7) to enrich the protein content of the retentate. The resulting retentate was pasteurized (65°C, 30 min) and spray dried (T_{in}: 180°C, T_{out}: 75°C). The protein isolates were stored at room temperature until further analysis.

2.3 Compositional analysis

Total nitrogen content of the LPIs was analysed according to the Kjeldahl method (MEBAK 1.5.2.1) [28] using a nitrogen-to-protein conversion factor of 6.25. Fat content was measured following the Soxhlet method (AACC Method 30-25.01) [29]. Ash content was determined by dry ashing in a muffle furnace at 500°C for 5 h (AOAC 923.03) [30]. Moisture was determined by oven drying at 103°C for 5 h (AOAC 925.10) [31]. Total starch (AOAC Methods 996.11 and AACC Method 76-13.01) [32, 33] content was determined using an enzymatic kit (Megazyme, Bray, Co. Wicklow, Ireland). Minerals were analysed using inductively coupled plasma-optical emission spectrophotometry [34]. The soluble and insoluble fibre content of the samples was analysed in accordance with the AOAC method 991.43 [35].

92 2.4 Protein profile analysis

Protein profile was assessed using sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) using precast gels (Mini-PROTEAN TGX, Bio-Rad Laboratories, CA, USA) under non-reducing and reducing conditions as described by Alonso-Miravalles & O'Mahony [36]. The sample loading buffer contained 65.8 mM Tris-HCl (pH 6.8), 26.3% (w/v) glycerol, 2.1% SDS and 0.01% bromophenol blue. The running buffer (10x Tris/Glycine/SDS, Bio-Rad Laboratories, CA, USA) had a composition of 25 mM Tris, 192 mM glycine and 0.1% SDS (w/v), pH 8.3. The staining solution used was Coomassie Brilliant Blue R-250 (Bio-Rad Laboratories, CA, USA). The target final protein concentration was 1 mg/mL and 8 μ L¹ of sample solution was loaded into each well of the gel and the gels were run at a constant voltage of 150 V.

101 2.5 Protein secondary structure

102 Information about secondary structure of the proteins was obtained using circular dichroism (CD) 103 spectrophotometry (Chirascan, Applied Photophysics, Leatherhead, UK). Protein solutions of 1 mg/mL were 104 prepared in 10 mM sodium phosphate buffer (pH 7) and solubilized overnight at 4°C using magnetic stirring at 105 250 rpm. Subsequently, samples were filtered (0.25 μ m) and the CD spectra was measured with a path length of 106 0.1 mm in the range 180-260 nm at a bandwidth of 1 nm and spectral resolution of 1 nm and data acquisition rate 107 of 1 point/s. The average of three spectra was obtained and a 5-point smoothing algorithm was applied.

2.6 Scanning electron microscopy

Protein powders were mounted on aluminium stubs using double-sided adhesive carbon tape, and sputter coated with a 5 nm layer of gold/palladium (Au:Pd = 80:20) using a Quorum Q150R ES Sputter Coating Unit (Quorum Technologies Ltd., Sussex, U-K-). The coated samples were loaded into a sample tube and examined using a JSM-5510 scanning electron microscope (JEOL Ltd, Tokyo, Japan), operated at an accelerating voltage of 5 kV.

114 2.7 Particle size distribution

Particle size distribution of protein dispersions was measured using static laser light diffraction (Mastersizer 3000, Malvern Instruments Ltd, Worcestershire, UK). For the preparation of samples, the protein isolate powders were mixed with ultrapure water at a concentration of 1% protein (w/v), pH adjusted to 7, and stirred overnight at 4°C. The refractive index of protein was set at 1.45 [27] and the absorption and dispersant refractive indices used were 0.1 and 1.33, respectively. LPI dispersions, equilibrated at 22°C, were introduced into the dispersing unit using ultrapure water as dispersant until a laser obscuration of 12% was achieved.

2.8 Hydrophobicity

Surface hydrophobicity (S₀) of protein particles was measured according to Hayakawa and Nakai [37] using 1-anilino-8-naphthalenesulfonate (ANS) with slight modifications as described by Karaca et al. [16]. Protein solutions were serially diluted with 10 mM phosphate buffer (pH 7) ranging from 0.0006–0.015% (w/v). ANS (10 μ LJ; 8.0 mM in 0.1 M phosphate buffer, pH 7) were mixed with 2 mL of diluted sample and left in darkness for 15 min. Fluorescence was measured ($\lambda_{excitation}$ 390 nm, $\lambda_{emission}$ 470 nm) and corrected by a blank measured without ANS. The results are presented as the slopes (R² \geq 0.98) of the absorbance versus protein concentration.

2.9 Sulfhydryl groups

Sulfhydryl groups were determined using Ellman's reagent (5,5'-dithio-bis-(2-nitrobenzoic acid) according to the method of Van der Plancken et al. [38]. The protein samples were diluted to 2 mg/mL with 10 mM phosphate buffer (pH 7) for free sulfhydryl groups, while for total sulfhydryl groups a buffer containing 6 M urea and 0.5 M SDS was used. Ellman's reagent (80 μ L) was added to 2.5 mL of diluted sample and absorbance was measured at 412 nm after 15 min. For the reagent blank, the protein samples were replaced by the sodium phosphate buffer and mixed with 80 μ L of Ellman's reagent. Sulfhydryl groups were quantified as follows:

135 (1) $\mu m SH/g \ protein = (A_{412} - A_{412B}) \cdot \frac{1,000,000}{\varepsilon} \cdot C$

where A_{412} is the absorbance at 412 nm, A_{412B} is the absorbance at 412 nm for the blank, ε is the extinction coefficient, which was taken as 13,600 M⁻¹ cm⁻¹, and C is the protein concentration in mg/mL of the diluted sample.

139 2.10 Protein solubility

The solubility of proteins as influenced by pH, was determined by adjusting the pH of protein dispersions from 3.0 to 8.0 at 0.5 units intervals using 0.1 and 1 M HCl or NaOH. Protein samples (1% w/v) were hydrated at 4°C. The pH was re-adjusted before measurements. Samples were centrifuged at 5,000 *g* for 30 min. The protein contents of the supernatants were analysed using the Kjeldahl method as described in Section 2.3. The results were expressed as % of the total protein content.

145 2.11 Zeta potential

56146The zeta potential of protein solutions at the same pH values as for protein solubility analysis were determined57147using a Zetasizer nano-Z (Malvern Instruments Ltd; UK). Samples were prepared as described for the protein59148solubility, excluding the centrifugation step, and diluted with ultrapure water to a concentration of 0.1% (w/v) and

pH was readjusted. The measurement was performed using an automatic voltage selection and zeta potential was calculated using the Smoluchowski model. Refractive and absorption indices of 1.45 and 0.001 were used, respectively.

152 2.12 Water holding capacity

153 Analysis of water holding capacity (WHC) of proteins was determined according to AACC method 56-30.01 154 [39] with some modifications. Samples (1.000 g \pm 0.005 g) were mixed with 30 mL¹ of distilled water using an 155 Ultra-Turrax equipped with a S10N-5G dispersing element (Ika-abortechnik, Janke and Kunkel GmbH, Staufen, 156 Germany) for 15 s and then shaken for 30 min at 1,000 rpm using a platform shaker (UNI MAX 1010, Heidolph, 157 Schwabach, Germany). Subsequently, the mixture was centrifuged at 2,000 g for 10 min. WHC was expressed as 158 grams of water retained per gram of protein isolate.

159 2.13 Fat absorption capacity

Fat absorption capacity (FAC) was determined following the method described by Boye et al. [13] with slight modifications. Powder (1 g) and <u>sunflower oil</u> (6 g) were weighted into a 15 mL centrifuge tube (Sarstedt, Nümbrecht, Germany), mixed with a vortex for 3 min and centrifuged at 4,000 g for 30 min. The oil was removed from the tube carefully and weighed again. FAC was expressed as grams of <u>fat water</u> retained per gram of protein isolate.

165 2.14 Foaming properties

Protein dispersions (20 mL¹) with a protein concentration ranging from 0.1 to 3.3% (w/v) in ultrapure water were frothed using an Ultra-Turrax equipped with a S10N-10G dispersing element (Ika-Labortechnik, Janke and Kunkel GmbH, Staufen) at high speed for 30 s. The height of the sample (liquid and foam phase) was measured over 60 min. The foaming capacity was taken as sample expansion at 0 min, while foam stability was expressed as sample expansion after 60 min. Foam expansion was calculated according to the following equation:

$$(2) Foam expansion = \frac{Sample height after foaming - Initial sample height}{Initial sample height} \cdot 100$$

2.15 Emulsifying properties

Protein solutions (1%, w/v) were hydrated with ultrapure water using a magnetic stirrer at 250 rpm overnight at 4°C and pH 7. The next day samples were adjusted to room temperature and the pH was re-adjusted if necessary and pre-emulsions were prepared as follows: 20 mL of <u>sunflower</u> oil was added to 180 mL of 1% protein (w/v) solution and homogenized for 3 min at 10,000 rpm using an ultraturrax (T 25 digital Ultra-Turrax, Staufen,

Germany). Emulsifying activity (EAI) and stability (ESI) indices were determined using the method described by Pearce and Kinsella [40], with slight modifications. In brief, 250 μ and μ emulsion were taken from the bottom of the homogenized sample after 0 and 120 min and diluted (1:100, v/v) in 0.1% sodium dodecyl sulphate (SDS) solution. The absorbance at a wavelength of 500 nm was read using a spectrophotometer. EAI and ESI were calculated using the following equations:

$$(3) EAI\left(\frac{m^2}{g}\right) = \frac{2 \cdot 2.303 \cdot A_0 \cdot DF}{C \cdot \theta \cdot 10000}$$

$$(4) ESI (min) = \frac{A_0}{A_0 - A_{120}} \cdot \underline{120} + 10$$

where DF is the dilution factor (100), C is the initial concentration of protein (0.01 g/mL), θ is the fraction of oil used to form the emulsion (0.1), and A₀ and A₁₂₀ are the absorbance of the diluted emulsion at 0 and 120 min, respectively.

187 2.16 Gelation characteristics

188 2.16.1. Least gelling concentration

The least gelling concentration (LGC) is defined as the lowest concentration required to form a selfsupporting gel. The LGC test was performed according to the method of Sathe et al. [41] with some modification. LPI dispersions ranging from 6 to 16% (w/v) were prepared in 0.01 M phosphate buffer at pH 7.0. These suspensions were heated in 15 mL test tubes (Sarstedt, Nümbrecht, Germany) in a water bath at 90°C for 30 min, after which they were cooled rapidly under running water and stored at 4°C overnight. LGC was determined visually as the minimum concentration of protein at which the contents of the tube did not flow.

195 2.16.2. Texture profile analysis

Texture profile analysis (TPA) of the LPI gels was performed using a TA.XT Plus[™] texture analyser (Stable Microsystems Ltd., Crawley, UK) to determine their mechanical properties. Protein gels (25%; w/v) were prepared by heating LPI dispersions as described above. Gels were cut into small cylinders of 8.2 mm in diameter and 8.0 mm in height. The gel pieces were compressed twice to 30% of their original height at a constant speed of 0.3 mm/s using a cylindrical probe with 20 mm diameter. The TPA parameters of hardness, cohesiveness, adhesiveness, gumminess and springiness were calculated according to the definitions of Bourne [42].

202 2.17 Life cycle assessment

 Environmental performance of LPIs was examined by means of life cycle assessment (LCA) using Umberto
5.5 software. LCA is carried out as an attributional cradle-to-gate LCA and includes the individual processes
associated with LPIs shown in Figure 1. Impact assessment methods are based on Umweltbundesamt Berlin [43].

206 2.18 Statistical analysis

All analyses were carried out in triplicate, with exception of analyses of fibre and minerals, which are performed following a validated method and therefore analysed just once and reported without standard deviation. The other data generated was subjected to student's T-test to determine statistically significant differences (p < 0.05) between mean values for the different samples, at a 95% confidence level. The statistical program used was Excel (Microsoft Office 365 ProPlus, version 1809).

212 3 Results and Discussion

213 3.1 Compositional analysis

The macro- and micro-nutrient composition of the LPIs is shown in Table 1. The protein content of LPI-UF (93.7%) was significantly higher (p < 0.05) than that obtained for LPI-IEP (85.1%). The reason for this higher protein content can be explained by the ultrafiltration process, where specific pore sized membranes are used leading to higher protein levels in the final ingredient [25]. Additionally, by diafiltration more soluble substances (e.g. sugars, minerals) permeate the membrane thereby further purifying the protein. (4.49%). Regarding the fat content, no significant differences ($p \le 0.05$) were found between LPI-UF (4.40%) and LPI-IEP (4.49%). The ash content for LPI-IEP (5.46%) was significantly higher than for LPI-UF (3.51%) which was expected since, with the former approach, NaOH and HCl are used to solubilize and precipitate the proteins [17]; this can be seen in the determined sodium content of LPI-IEP. Interestingly, high values of magnesium and calcium were obtained in LPI-UF. An explanation for these high values might be the retention of these minerals in the retentate along with the protein during the UF process. These differences in the mineral profile can play an important role in the functionality of these protein ingredients such as the solubility, emulsifying and gelling properties [44]. For example, in dairy proteins, especially caseins, calcium plays an important role in determining their gelation behaviour, facilitating linkages between proteins [45]. Also, other authors have studied the role of calcium on gelation properties of a soya drink, finding coagulation of soya proteins when the ionic calcium concentration was increased [46, 47]. The fibre content, mostly soluble dietary fibre, was higher in the LPI-IEP (1.8%) than in LPI-UF (<0.1%). A reason for the higher fibre content in LPI-IEP could be that a part of the fibres were precipitated together with the protein and/or were only partially removed by the centrifugation step; the lower protein content of LPI-IEP is an indicator of this.

233 3.2 Structural properties

234 3.2.1 SDS-PAGE

SDS-PAGE analyses under non-reducing and reducing conditions of the two LPIs are shown in Figure 2.
SDS-PAGE analyses under non-reducing and reducing conditions of the two LPIs are shown in Figure 2.
Both samples showed similar protein profiles, with several common bands under non-reducing and reducing
conditions. Proteins with molecular weight (MW) of ~50, ~37 and ~20 kDa under non-reducing conditions were
observed. The bands at MW ~50 kDa may correspond to vicilin subunits, which composes a 7S trimeric protein,
one of the major globulins, together with legumin found in many pulses. Each trimer of vicilin has a MW of 150
kDa without disulphide bridging [48]. The bands at 37 and 25 kDa correspond to the acidic and basic subunits of

legumin, in accordance with previous studies [49, 50]. Legumin, an 11S globulin, is an hexameric protein formed by subunits with MW ~60 kDa, which consist of an acidic (~40 kDa) and a basic (~20 kDa) subunit linked by disulfide bonding [14, 24]. Under reducing conditions, similar profiles were observed, although bands at 37 and 25 kDa were slightly more intense, with the disappearance of some high MW bands at ~50 kDa. This can be correlated with the dissociation of legumin into its acidic (MW ~40 kDa) and basic (~20 kDa) subunits by the dissociation of the disulphide bond when a reducing agent (DTT) is applied.

247 3.2.2 Secondary structure

Furthermore, far-UV CD spectroscopic measurements were performed to gain information about the secondary structure of LPIs. Amide groups are optically active and absorb circular polarized far-UV light. Depending on their conformation, i.e., their secondary structure, characteristic CD spectra are obtained [51]. As shown in Figure 3, both LPIs exhibited a positive peak at 185 nm, and a broad negative peak with a minimum at 208 nm, indicating a defined secondary structure of α -helix [51]. Only slight differences can be observed in the spectra. Similar spectra for lentil flour and isolated proteins using IEP were found by Aryee and Boye [52], indicating that secondary structure conformational changes were limited during the extraction of the proteins.

255 3.2.3 Scanning electron microscopy

Representative micrographs of the LPI powders are given in Figure 4. In general, a heterogeneous mixture of rounded particles with smooth, shrivelled, hollow and wrinkled surfaces were observed in both LPIs. These features are typical for spray dried powders and have been attributed to rapid evaporation of water during the spray-drying process [53–55]. Joshi et al. [24]) also observed similar folded and wrinkled surfaces in LPI powders obtained by spray drying. The sizes of the powder particles, as seen from the scale bars, were generally between 10 and 50 µm. LPI-IEP and LPI-UF showed similar powder characteristics, although the LPI-IEP primary particles are in a closer arrangement than the LPI-UF particles where the powder particles seem more dispersed.

3.2.4 Particle size distribution

The particle size observations obtained by SEM can be correlated with the particle size distribution (PSD) determined using laser diffraction (Figure 5). Both LPIs showed a monomodal size distribution with a size range of 10 to 100 μ m. The volume-weighted mean particle diameter (D_{4,3}) values of LPI-UF and LPI-IEP were 32.8 μ m and 29.4 μ m, respectively. The LPI-IEP also had significantly lower values for surface-weighted mean particle diameter (D_{3,2}), D (50) and D (90) (Table 2). Similar profiles were observed by Crowley et al. [56]) in high-protein 269 (90%) milk protein concentrates after 24 h of rehydration, with particle sizes ranging from 10 to 100 μ m, 270 classifying them as large and poorly-dispersible particles.

271 3.2.5 Surface hydrophobicity

Hydrophobic groups exposed to the surface of the proteins enable hydrophobic interactions, and adsorption to interfaces; hence having an influence on many properties, such as emulsification and foaming [57]. These values are shown in Table 2. The LPI-IEP had a significantly higher surface hydrophobicity with a value of 2688 in comparison to LPI-UF with a value of 2411. However, the differences were not major, but significantly different, indicating that the extraction method had no major impact on the surface hydrophobicity of the proteins. Comparable studies found a value of 2200 for legumin-like proteins isolated from lentils [14], while Joshi et al. [58] found a considerably higher value of 568 determined for mg/mL, which translates to 5680 using the same protein concentration units as in this study.

280 3.2.6 Sulfhydryl groups

Results of sulfhydryl groups measured as free and total are shown in Table 2. The concentration of free and total sulfhydryl groups were found to be higher for LPI-IEP, with 6.04 and 23.9 µmol/g protein, respectively, compared to 5.88 and 22.5 µmol/g protein, respectively for LPI-UF. Literature considering the sulfhydryl groups of lentil proteins is scarce and diverse; Li & Lee [59] reported disulphide contents of 0.31 µmol/g, and free sulfhydryl groups of 0.032 µmol/g, being considerably lower than the values found in this study. On the other hand, Ladjal-Ettoumi et al. [49] found comparable values; they reported 16.1 µmol/g and 31.0 µmol/g for free and total sulfhydryl groups, respectively. In both cases, the relatively low amount of free sulfhydryl groups indicated the formation of aggregates, being characteristic for globular proteins, and can be linked also to the relatively large particle size. In general, both LPIs showed similar values for hydrophobicity and also sulfhydryl groups. However, LPI-IEP showed a trend with significant difference (p < 0.05) to higher values, indicating a slightly more open structure with higher surface active groups.

- 292 3.3 Functional properties
 - 293 3.3.1 Solubility and zeta potential

Both isolates showed similar solubility and zeta potential values across the pH range, as shown in Figure 6,
achieving the highest solubility at acidic and alkaline pH values. Similarly, at the extreme low and high pH ranges,
the lentil protein particles showed the highest positive and negative charge, respectively. LPI-IEP showed lower
solubility values across the pH range compared with LPI-UF. This may be explained by the removal of soluble

proteins in the supernatant during the extraction process and, therefore, more insoluble protein fraction is present in LPI-IEP (Figure 1). High solubility values and positive charge (+ 30 mV) were observed at pH 3, followed by minimum solubility and a net charge of 0 mV for both isolates at pH 4.5, indicating that the isoelectric point was reached. Solubility was higher again at pH 6 and pH 6.5 for LPI-UF and LPI-IEP, reaching a value of 43% for both isolates at pH 7 and the surface charge decreased, reaching values between -20 and -30 mV. Karaca et al. [16], found similar values for surface charge (-22.6 mV) for LPI-IEP at pH 7. The highest solubility was obtained for the LPI-UF at pH 9, at 54.7%, while LPI-IEP reached 50.18%. LPI-UF showed also a higher solubility at acidic conditions with a value of 39.4% at pH 3.5. The LPI-IEP had lower solubility at lower pH's, having a value of 11.9% at pH 3.5. The general profiles of the observed solubility and zeta potential curves are characteristic for lentil proteins, as previously reported by Boye et al. [13], who studied the solubility of different plant-based protein isolates finding high solubilities for lentil and pea proteins in comparison with chickpea. Lee et al. [60] analysed the protein solubility of commercial soya products including flours, concentrates and isolates and found generally lower solubility at low pH's. Solubility is one of the most important properties of proteins, influencing for example the ability to form and stabilise foams, emulsions and gels. Although insoluble proteins can be used in meat preparations, highly soluble proteins provide the most versatility for substitution and extension of animal proteins [61].

314 3.3.2 Foaming properties

Foaming is, in many product applications, a desired property of proteins, providing structure and stability. The foaming properties of LPIs, as a function of protein concentration, are shown in Table 3. The foaming capacity was low for both isolates at 0.1% (w/v) at 9.42 and 6.52% for LPI-UF and LPI-IEP, respectively. With increasing concentration, the foaming capacity increased, reaching 69.5 and 57.2% for 3.3 % (w/v) LPI-UF and LPI-IEP, respectively. Likewise, the foam stability increased from 0% for both LPIs to 44.9% and 39.1% for LPI-UF and LPI-IEP, respectively. The LPI-UF showed significantly better foaming properties at the high protein concentrations compared to the LPI-IEP. In addition, other studies found that protein isolates of various sources prepared by UF were superior to those obtained by precipitation, especially in terms of protein solubility and foaming characteristics [13, 62]. The results obtained show a high ability of lentil proteins to create foam with high stability, indicating its potential for application in food processing. Compared to other commercial proteins from potato (36.9%), pea (10.6%), carob (17.2%), lupin (13.9%) and soya (36.4%) analysed by Horstmann et al. [63] for their application in bread, both LPIs showed better foam capacities; even potato protein, known to have good foaming ability, showed lower values. These results underline the great prospect of LPIs being used in
 bakery products, ice cream or other dairy formulations, where foaming properties are desired.

329 3.3.3 Emulsification properties

Proteins can act as emulsifiers by forming a film/skin around oil droplets dispersed in an aqueous medium, thereby stabilizing emulsions and preventing structural changes such as coalescence, creaming, flocculation or sedimentation [13]. The emulsifying activity (EAI) and emulsifying stability index (ESI) of LPI-UF and LPI-IEP are shown in Table 4. EAI and ESI are two indices often used to evaluate the emulsifying properties of proteins. ESI values were found to be higher for LPI-UF (63.8 min), compared to LPI-IEP (51.0 min). The EAI values were quite similar for both LPIs; however, higher values were found for LPI-IEP (16.5 m^2/g) in comparison with LPI-UF (14.3 m²/g). These higher values for LPI-IEP can be related to the higher surface hydrophobicity compared to LPI-UF. The EAI values were lower in comparison to other studies where different protein-to-fat ratios or high-pressure homogenization were applied [13, 64]. For example, M. Joshi et al. [58] found that the EAI increased 3-fold when the concentration of protein increased from 10 mg/mL¹ to 30 mg/mL¹, whereas in this study the protein concentration was maintained at 10 mg/mL¹. In addition, high pressure homogenization, as applied to cow's milk, can help to unfold globulins (which are known for having high MW and compact structures) and enable them to migrate to the interface in order to form a stable emulsion [16]. These factors can be taken into consideration for further studies in order to enhance the emulsification properties.

3.3.4 Water holding and fat absorption holding capacity

Water holding and fat absorption-holding_capacity (WHC and FHAC) of proteins is an important functionality, since it influences structure, mouth feel and flavour retention of food formulations. The ability of protein to retain oil or water can be important in food applications, such as ground meat formulations, doughnuts and bakery products. Values are shown in Table 2. Significant differences were found for the WHC between the two different LPIs, showing a higher value of 3.96 g/g for the LPI-UF, compared to 2.60 g/g for LPI-IEP. Compared to other studies, both isolates showed a relatively high WHC; Boye et al. [13] reported values of 0.6 and 2.7 g/g for protein concentrates isolated from several legumes. However, the authors found no considerable effect of the preparation method, possibly due to the comparatively low protein contents of their samples. Horstmann et al. [63], found values ranging from 0.0 g/g for a potato and soya, and up to 2.66 g/g for a pea protein ingredient. They associated the protein content to be negatively correlated with the WHC, i.e., other constituents affect the values to a substantial degree. In contrast, in this study it was found that a higher protein content

European Food Research and Technology

356correlateds with a higher WHC. Results for FHAC showed also significantly higher values for LPI-UF (2.24 g357oil/g protein) in comparison to LPI-IEP (2.09 g oil/g protein). The value obtained for LPI-UF is comparable to358that obtained by Boye et al. [13] for red lentil protein with a FAC of 2.26 g oil/g protein. In addition, this author359found the highest FAC value for LPI in comparison with yellow pea and kabuli chickpeas proteins.

3.3.5 Gelation characteristics

Heat induced gelation occurs when proteins aggregate to form a three-dimensional network. The ability to do so depends on the state and surface conformation of the proteins, e.g., free sulfhydryl groups, hydrophobicity, charge and correspondingly the electrostatic interaction between proteins, and their ability to associate to form a continuous network throughout the matrix [65]. LGC was measured as an indicator of the gelation capacity. The LPI-UF formed a gel, resisting flow when inverted, at a concentration of 11% (w/v), whereas for the LPI-IEP 16% was needed (Table 4). The lower concentration needed for a firm gel to be formed by the LPI-UF may be associated with the higher protein solubility, which is known to be an important factor in gel formation [24]. Likewise, Boye et al. [25] found that various legume proteins isolated by UF have lower LGC in comparison to IEP methods. They found comparable values, with 10% for LPI prepared by UF and 12% for isolates prepared by IEP. The LPI-UF also formed a much stronger gel, which is evident in the values obtained from the TPA test: the hardness for LPI-UF was three-fold higher than that of LPI-IEP. Likewise, gumminess and chewiness were also significantly higher for LPI-UF than for LPI-IEP gels. On the other hand, other parameters, such as adhesiveness, springiness, resilience and cohesiveness were not significantly different. The higher gelling properties of LPI-UF may be linked to the higher calcium levels, which has been shown to enhance hydrophobic coagulation of heat treated milk and soya proteins [46]. The ability to form strong gels upon heating is a desirable propertyfunctionality in bakery products: when heat is applied to the dough, its viscosity increases , which gives stability to expanding gas cells, resultsing in a higher gas retention during baking and a higher desirable specific volume of the product is reached [66]. Furthermore, in non-traditional ways,-also meat, yoghurt and cheese alternatives may be produced from heat-set gels, providingfacilitating to the product with a gel-like matrix.

3.4 Life cycle assessment

Environmental performance of LPI obtained by IEP and UF was examined by means of life cycle assessment (Table 5). Indicators such as aquatic eutrophication, photochemical oxidant formation, stratospheric ozone depletion, phosphorus use and land use showed lower potential environmental impacts for LPI-UF in comparison to LPI-IEP. For the remaining indicators studied, the ranking was switched. Especially the contribution of the

lentil cultivation stage affects the outcome of these indicators. An overview on main contributors is exemplarily given in Figure 7 for four indicators - the remaining indicators followed one out of those four illustrated result contribution patterns. The higher the contribution of the lentil cultivation stage, the more important is the protein yield advantage from LPI-UF, as less lentil seeds were required per kg protein isolated. On the other hand, lower process energy was required for processing of LPI-IEP, which leads to lower potential environmental impacts for the remaining indicators including climate change. The net nitrogen benefit due to air nitrogen fixation by lentil plants (as they are legumes) in the growth phase is up to 20% of the total environmental impact depending on the indicator for both LPIs.

Further, LPIs showed promising carbon footprints within the portfolio of soya-based and cow's milk-based protein isolate food ingredients: The production of both LPIs potentially releases a quarter of carbon dioxide equivalents (3.5 to 4.2 kg CO²-e/kg) than caseinate or whey protein production (19 kg CO²-e/kg and 20 kg CO²e/kg, respectively) as examined in an attributional LCA by Thrane, Paulsen, Orcutt, & Krieger [67]. Compared to soya protein isolate, depending on the literature source chosen, LPIs showed similar [67] or up to 4-fold lower values [68] for their potential release of carbon dioxide equivalents.

The environmental impact profiles of LPIs were also compared with traditional cow's milk protein as illustrated in Figure 8. Two different scenarios for the production of cow's milk protein were taken into consideration: the environmental impact, high or low, of the milk protein was set up depending on the theoretical amount of protein that is fed to the cow i.e. cow feed per kg milk and share of concentrate versus silage feed components within the feed mix were the parameters set to low and high for those ranges. Indicator results of LPIs are lower (and thus favourable) or equal for all of the examined indicators except the land use indicator. The latter is related to comparatively high agricultural yields of feed crops in comparison with relatively low yields for lentils. It should be noted that feed crops have undergone long-time optimisation of agricultural practices in order to reach relatively high yields. Lentils on the other hand have not been cultivated in comparable amounts on global scale than animal feed crops. Therefore, related optimization might take place along with increased interest in lentils in the future.

 412 4 Conclusion

gases compared to cow's milk protein.

Overall, the environmental impact of both LPIs was lower, contributing e.g. to a reduction of greenhouse

Various physical and functional properties of two LPI isolated by IEP and UF were investigated, indicating that they could contribute different desirable attributes to a wide range of food products. The results suggest that, in general, UF resulted in a product with better functional properties, such as higher protein solubility, WHC, greater gelling and foaming properties and emulsion stability. Differences in functional properties between the isolates under investigation were attributed to differences in the extraction methods, resulting in different compositions. Both isolates contained high levels of protein; however, LPI prepared by UF contained significantly higher values of protein, calcium and magnesium, whereas LPI prepared by IEP had higher levels of other minor constituents such as fibre, sodium and phosphorous. The life cycle assessment showed that the two main drivers for the environmental impact of LPIs were the cultivation stage and the protein isolation process. Overall, both LPIs exhibited promising environmental performance, especially if compared to traditional cow's milk proteins. These favorable favorable functional, nutritional and environmental properties of LPIs could be exploited in the preparation and development of diverse food products and may also be suitable for the substitution of soya or animal derived proteins. Further studies are required to investigate protein functionality and applicability of these in food systems as well as life cycle assessments of the food products thereof.

427 5 Declarations of interest

The authors declare no conflicts of interest.

429 6 Acknowledgement

The work for this publication has been undertaken as part of the PROTEIN2FOOD project. This project has received funding from the European Union's Horizon 2020 research and innovation programme under grant agreement No 635727. We thank our colleagues Dave Waldron, Tom Hannon and Michael Cronin for technical assistance and expertise that greatly assisted the work.

1 2			
3	434	7 I	References
4 5	435	1.	Henchion M, Hayes M, Mullen AM, Fenelon M, Tiwari B (2017) Future Protein Supply and Demand:
6 7	436		Strategies and Factors Influencing a Sustainable Equilibrium. Foods 6:53.
8 9	437		https://doi.org/10.3390/foods6070053
10 11 12	438	2.	United Nations (2015) Revision of World Population Prospects. In: United Nations.
13 14	439		https://population.un.org/wpp/Graphs/Probabilistic/POP/TOT/. Accessed 1 Dec 2018
15 16	440	3.	Day L (2013) Proteins from land plants - Potential resources for human nutrition and food security. Trends
17 18 19	441		Food Sci Technol 32:25-42. https://doi.org/10.1016/j.tifs.2013.05.005
20 21 22 23	442	4.	Don C (2017) Future Proteins: Functionality, Processing, and Sourcing. AACC Int 62:144-148
24	443	5.	Poore J, Nemecek T (2018) Reducing food's environmental impacts through producers and consumers.
25 26 27	444	Science (80-) 360:987-992. https://doi.org/10.1126/science.aaq0216	
28	115	(Share A Fahrle New F Mile P 2010 France dance in finder for dance in fficiencies in
29 30	445	6.	Shepon A, Eshel G, Noor E, Milo R (2016) Energy and protein feed-to-food conversion efficiencies in
31 32	446 447		the US and potential food security gains from dietary changes. Environ Res Lett 11:105002.
33 34	https://doi.org/10.1088/1748-9326/11/10/105002		
35 36	448	7.	Aiking H (2011) Future protein supply. Trends Food Sci Technol 22:112–120.
37 38 39	449		https://doi.org/10.1016/j.tifs.2010.04.005
40 41	450	8.	Coda R, Varis J, Verni M, Rizzello CG, Katina K (2017) Improvement of the protein quality of wheat
42 43	451		bread through faba bean sourdough addition. LWT - Food Sci Technol 82:296-302.
44 45	452		https://doi.org/10.1016/J.LWT.2017.04.062
46 47 48	453	9.	Jiménez-Martínez C, Hernández-Sánchez H, Dávila-Ortiz G (2003) Production of a yogurt-like product
49 50	454		from Lupinus campestris seeds. J Sci Food Agric 83:515-522. https://doi.org/10.1002/jsfa.1385
51 52	455	10.	Lin S, Huff HE, Hsieh F (2002) Extrusion Process Parameters, Sensory Characteristics, and Structural
53 54	456		Properties of a High Moisture Soy Protein Meat Analog. J Food Sci 67:1066-1072.
55 56 57	457		https://doi.org/10.1111/j.1365-2621.2002.tb09454.x
57 58 59 60	458	11.	Riaz MN (2011) Texturized vegetable proteins. In: Phillips GO, Williams PA (eds) Handbook of food

1 2						
3 4	459		proteins, 1st ed. Woodhead Publishing Limited, pp 395-418			
5 6	460	12.	Duranti M (2006) Grain legume proteins and nutraceutical properties. Fitoterapia 77:67-82.			
8 9	9					
10 11	462	13.	Boye JI, Aksay S, Roufik S, Ribéreau S, Mondor M, Farnworth E, Rajamohamed SH (2010) Comparison			
12 13	463		of the functional properties of pea, chickpea and lentil protein concentrates processed using ultrafiltration			
14 15	464		and isoelectric precipitation techniques. Food Res Int 43:537-546.			
16 17 18	465		https://doi.org/10.1016/j.foodres.2009.07.021			
19 20	466	14.	Jarpa-Parra M, Bamdad F, Tian Z, Zeng H, Temelli F, Chen L (2015) Impact of pH on molecular structure			
21 22	467		and surface properties of lentil legumin-like protein and its application as foam stabilizer. Colloids			
23 24	468		Surfaces B Biointerfaces 132:45-53. https://doi.org/10.1016/J.COLSURFB.2015.04.065			
25 26						
27	469	15.	FAOSTAT Database (2016) Food and Agriculture Organization. http://www.fao.org/faostat/en/#data/QC.			
28 29	470		Accessed 15 Nov 2018			
30 31	471	16.	Karaca AC, Low N, Nickerson M (2011) Emulsifying properties of chickpea, faba bean, lentil and pea			
32 33	472		proteins produced by isoelectric precipitation and salt extraction. Food Res Int 44:2742-2750.			
34 35	473		https://doi.org/10.1016/j.foodres.2011.06.012			
36 37						
38 39	474	17.	Arntfield SD, Maskus HD (2011) Peas and other legume proteins. In: Phillips GO, Williams PA (eds)			
40 41	475		Handbook of Food Proteins, 1st ed. Woodhead Publishing, Cambridge, UK, pp 233-266			
42 43	476	18.	Dijkstra D, Linnemann A, Van Boekel T (2003) Towards Sustainable Production of Protein-Rich Foods:			
44 45	477		Appraisal of Eight Crops for Western Europe. Part II: Analysis of the Technological Aspects of the			
46	478		Production Chain. Crit Rev Food Sci Nutr 43:481–506. https://doi.org/10.1080/10408690390246332			
47 48						
49 50	479	19.	Nishinari K, Fang Y, Guo S, Phillips GO (2014) Soy proteins: A review on composition, aggregation and			
51 52	480		emulsification. Food Hydrocoll 39:301-318. https://doi.org/10.1016/j.foodhyd.2014.01.013			
53 54	481	20.	Fox PF, Uniacke T, McSweeney PLH, O'Mahony JA (2015) Dairy Chemistry and Biochemistry, 2nd ed.			
55 56	482 Springer International Publishing, New York					
57 58 59	483	21	Panalamprou FM Dovastakis GL Bilizdaris CG Vissesaglou V (2000) Influence of properation wether to			
60	400	21.	Papalamprou EM, Doxastakis GI, Biliaderis CG, Kiosseoglou V (2009) Influence of preparation methods			

2			
3 4	484		on physicochemical and gelation properties of chickpea protein isolates. Food Hydrocoll 23:337-343.
5 6	485		https://doi.org/10.1016/j.foodhyd.2008.03.006
7 8	486	22.	Peng W, Kong X, Chen Y, Zhang C, Yang Y, Hua Y (2016) Effects of heat treatment on the emulsifying
9 10 11	487		properties of pea proteins. Food Hydrocoll 52:301-310. https://doi.org/10.1016/j.foodhyd.2015.06.025
12 13	488	23.	Sánchez-Vioque R, Clemente A, Vioque J, Bautista J, Millán F (1999) Protein isolates from chickpea
14 15	489		(Cicer arietinum L.): Chemical composition, functional properties and protein characterization. Food
16 17 18	490		Chem 64:237–243. https://doi.org/10.1016/S0308-8146(98)00133-2
19 20	491	24.	Joshi M, Adhikari B, Aldred P, Panozzo JF, Kasapis S (2011) Physicochemical and functional properties
21 22	492		of lentil protein isolates prepared by different drying methods. Food Chem 129:1513-1522.
23 24 25	493		https://doi.org/10.1016/j.foodchem.2011.05.131
26 27	494	25.	Boye J, Zare F, Pletch A (2010) Pulse proteins: Processing, characterization, functional properties and
28 29 30	495		applications in food and feed. Food Res Int 43:414–431. https://doi.org/10.1016/j.foodres.2009.09.003
31	496	26.	Jarpa-Parra M (2018) Lentil protein : a review of functional properties and food application . An overview
32 33 34	497		of lentil protein functionality. Int J Food Sci Technol 53:892–903. https://doi.org/10.1111/ijfs.13685
35 36	498	27.	Johnston SP, Nickerson MT, Low NH (2015) The physicochemical properties of legume protein isolates
37 38	499		and their ability to stabilize oil-in-water emulsions with and without genipin. J Food Sci Technol 52:4135-
39 40 41	500		4145. https://doi.org/10.1007/s13197-014-1523-3
42 43	501	28.	MEBAK (2011) MEBAK- Würze, Bier, Biermischgetränke, Methodensammlung der Mitteleuropäischen
44 45	502		Brautechnischen Analysenkommission, Selbstverlag der MEBAK. In: Wort, beer and beer-based
46 47 48	503		beverages.
49 50	504	29.	AACC, Approved Methods of the American Association of Cereal Chemists, Method 30-25.01 - Crude
51 52	505		Fat in Wheat, Corn, Soy Flour and Feed Mixes (2001). In: International Approved Methods of Analysis,
53 54 55	506		11th ed. St. Paul, MN
55 56 57	507	30.	AOAC, Official Methods of Analysis of AOAC International, AOAC Method 923.03 - Ash of Flour
57 58 59 60	508		(2005). In: Official Methods of Analysis, 18th ed. AOAC International Publisher, Gaithersburg, MD

http://mc.manuscriptcentral.com/efrt

 2 3 509 31 AOAC Official Methods of Analysis of AOAC International AOAC Method 925 10 Solids 							
4	509	31.	AOAC, Official Methods of Analysis of AOAC International, AOAC Method 925.10, Solids (Total) and				
5 6	510	510 Moisture in Flour (2005). In: Official Methods of Analysis, 18th ed. AOAC Internationa					
7 8	511		Gaithersburg, MD				
9 10 11	512	32.	AOAC, Official Methods of Analysis of AOAC International, AOAC Method 996.11 - Total Starch in				
12	513		Cereal Products (2005). In: Official Methods of Analysis, 18th ed. AOAC International Publisher,				
13 14 15	514		Gaithersburg, MD				
16 17	515	33.	AACC, Approved Method of the American Association of Cereal Chemists, Method 76-13.01 - Total				
18 19 20	516		Starch Assay Procedure (2001). In: International Approved Methods of Analysis, 11th ed. St. Paul, MN				
21 22 23	517	34.	US EPA Method 6020 Method 6020, Inductively Coupled Plasma/Mass Spectrometry				
24 25	518	35.	AOAC, Official Methods of Analysis of AOAC International, AOAC Method 991.43 - Total, Soluble and				
26 27	519		Insoluble Dietary Fibre in Foods (2005). In: Official Methods of Analysis, 18th ed. AOAC International				
28 29 30	520		Publisher, Gaithersburg, MD				
31 32	521	36.	Alonso-Miravalles L, O'Mahony JA (2018) Composition, Protein Profile and Rheological Properties of				
33 34 35	522		Pseudocereal-Based Protein-Rich Ingredients. Foods 7:1-18. https://doi.org/10.3390/foods7050073				
35 36 37	523	37.	Hayakawa SS, Nakai S (1985) Relationships of Hydrophobicity and Net Charge to the Solubility of Milk				
38 39	524		and Soy Proteins. J Food Eng 50:486–491. https://doi.org/10.1111/j.1365-2621.1985.tb13433.x				
40 41	525	38.	Van Der Plancken I, Van Loey A, Hendrickx MEG (2005) Changes in Sulfhydryl Content of Egg White				
42 43 44	526	Proteins Due to Heat and Pressure Treatment. https://doi.org/10.1021/jf050289+					
45 46	527	39.	AACC, Approved Methods of the American Association of Cereal Chemists, Method 56-30.01 - Water				
47 48	528		Hydration Capacity of Protein Materials (2001). In: International Approved Methods of Analysis, 11th				
49 50 51	529		ed. St. Paul, MN				
52 53	530	40.	Pearce KN, Kinsella JE (1978) Emulsifying properties of proteins: evaluation of a turbidimetric technique.				
54 55 56	531		J Agric Food Chem 26:716-723. https://doi.org/10.1021/jf60217a041				
50 57 58	532	41.	Sathe SK, Deshpande SS, Salunkhe DK (1982) Functional Properties of Lupin Seed (Lupinus mutabilis)				
59 60	533		Proteins and Protein Concentrates. J Food Sci 47:491-497. https://doi.org/10.1111/j.1365-				

2 3 4	534		2621.1982.tb10110.x						
5 6	535	42.	Bourne MC (2002) Food Texture and Viscosity - Concept and Measurements. In: Food Texture and						
7 8 9	536	Viscosity, 2nd ed. Food Science and Technology, Elsevier, p 416							
9 10 11	537	43.	Umweltbundesamt Berlin (2016) Prüfung und Aktualisierung der Ökobilanzen für						
12 13	538		Getränkeverpackungen. UBA-Texte 19/2016. Berlin						
14 15 16	539	44.	Foegeding EA, Davis JP (2011) Food protein functionality: A comprehensive approach. Food Hydrocoll						
17 18	540		25:1853–1864. https://doi.org/10.1016/j.foodhyd.2011.05.008						
19 20	541	45.	Farrell HM, Bleck GT, Brown EM, Butler JE, Creamer LK, Hicks CL, Hollar CM, Swaisgood HE (2004)						
21 22 23	542		Nomenclature of the Proteins of Cows ' Milk — Sixth Revision. J Dairy Sci 87:1641–1674						
24 25	543	46.	Pathomrungsiyounggul P, Lewis MJ, Grandison AS (2010) Effects of calcium-chelating agents and						
26 27	544	40.	pasteurisation on certain properties of calcium-fortified soy milk. Food Chem 118:808–814.						
28 29	545								
30 31	https://doi.org/10.1016/j.foodchem.2009.05.067								
32 33	546	47.	Joshi M, Timilsena Y, Adhikari B (2017) Global production, processing and utilization of lentil: A review.						
33 34 35	547		J Integr Agric 16:2898–2913. https://doi.org/10.1016/S2095-3119(17)61793-3						
36	549	40							
37 38	548	48.	Dagorn-Scaviner C, Gueguen J, Lefebvre J (1987) Emulsifying properties of pea globulins as related to						
39 40	549		their adsorptions behaviours. J Food Sci 52:335–341						
41 42	550	49.	Ladjal-Ettoumi Y, Boudries H, Chibane M, Romero A (2016) Pea, Chickpea and Lentil Protein Isolates:						
43 44	551		Physicochemical Characterization and Emulsifying Properties. Food Biophys 11:43-51.						
45 46 47	552		https://doi.org/10.1007/s11483-015-9411-6						
47 48 49	553	50.	Barbana C, Boye JI (2011) Angiotensin I-converting enzyme inhibitory properties of lentil protein						
50 51	554		hydrolysates: Determination of the kinetics of inhibition. Food Chem 127:94-101.						
52 53	555		https://doi.org/10.1016/j.foodchem.2010.12.093						
54 55 56	556	51.	Greenfield N (2007) Using circular dichroism spectra to estimate protein secondary structure. Nat Protoc						
57 58	557		1:2876-2890. https://doi.org/10.1038/nprot.2006.202.Using						
59 60	558	52.	Aryee AN., Boye JI (2015) Comparative study of the effects of processing on the nutritional,						

http://mc.manuscriptcentral.com/efrt

2 3	559		physochochemical and functional properties of lentil. J Food Process Preserv 41:1–13.					
4 5	560		https://doi.org/10.1111/jfpp.12824					
6 7	500		https://doi.org/10.1111/jtpp.12024					
8 9	561 53. Kim EHJ, Chen XD, Pearce D (2009) Surface composition of industrial spray-dried milk powders. Eff							
10	of spray drying conditions on the surface composition. J Food Eng 94:169-							
11 12 12	563		https://doi.org/10.1016/j.jfoodeng.2008.10.020					
13 14 15	564	54.	Kelly GM, O'Mahony JA, Kelly AL, Huppertz T, Kennedy D, O'Callaghan DJ (2015) Influence of					
16 17	565		protein concentration on surface composition and physico-chemical properties of spray-dried milk protein					
18	566		concentrate powders. Int Dairy J 51:34–40. https://doi.org/10.1016/j.idairyj.2015.07.001					
19 20	200							
21 22	567	55.	Amagliani L, O'Regan J, Kelly AL, O'Mahony JA (2016) Physical and flow properties of rice protein					
23 24	568		powders. J Food Eng 190:1-9. https://doi.org/10.1016/j.jfoodeng.2016.05.022					
25 26	560							
27	569	56.	Crowley S V., Desautel B, Gazi I, Kelly AL, Huppertz T, O'Mahony JA (2015) Rehydration					
29								
30 31	571		https://doi.org/10.1016/j.jfoodeng.2014.09.033					
33								
34 35	573		with surface properties of proteins. Biochim Biophys Acta - Protein Struct 624:13-20.					
36 37	574		https://doi.org/10.1016/0005-2795(80)90220-2					
38 39								
40 41	575	58.	Joshi M, Adhikari B, Aldred P, Panozzo JF, Kasapis S, Barrow CJ (2012) Interfacial and emulsifying					
42 43	576		properties of lentil protein isolate. Food Chem 134:1343–1353.					
44 45	577		https://doi.org/10.1016/j.foodchem.2012.03.029					
46	578	59.	Li M, Lee T-C (2000) Effect of Extrusion Temperature on the Solubility and Molecular Weight of Lentil					
47 48	579		Bean Flour Proteins Containing Low Cysteine Residues. J Agric Food Chem 48:880–884.					
49 50	580		https://doi.org/10.1021/jf990328f					
51 52								
53 54	581	60.	Lee K, Ryu H, Rhee K (2003) Protein solubility characteristics of commercial soy protein products. J Am					
55 56	582		Oil Chem Soc 80:85-90. https://doi.org/https://doi.org/10.1007/s11746-003-0656-6					
57 58	583	61.	Boland MJ, Rae AN, Vereijken JM, Meuwissen MPM, Fischer ARH, van Boekel MAJS, Rutherfurd SM,					
59 60	584	01.	Gruppen H, Moughan PJ, Hendriks WH (2013) The future supply of animal-derived protein for human					
	507		Shappen 11, moughan 19, menanks with (2015) The future suppry of annual-derived protein for numan					

European Food Research and Technology

2 3 4 5	585		consumption. Trends Food Sci Technol 29:62-73. https://doi.org/10.1016/j.tifs.2012.07.002			
5 6 7	586	62.	Fuhrmeister H, Meuser F (2003) Impact of processing on functional properties of protein products from			
8 9	587		wrinkled peas. J Food Eng 56:119-129. https://doi.org/10.1016/S0260-8774(02)00241-8			
10 11 12	588	63.	Horstmann SW, Foschia M, Arendt EK (2017) Correlation analysis of protein quality characteristics with			
13 14	589		gluten-free bread properties. Food Funct 8:2465–2474. https://doi.org/10.1039/C7FO00415J			
15 16	590	64.	Avramenko NA, Low NH, Nickerson MT (2013) The effects of limited enzymatic hydrolysis on the			
17 18	591		physicochemical and emulsifying properties of a lentil protein isolate. Food Res Int 51:162-169.			
19 20 21	592		https://doi.org/10.1016/j.foodres.2012.11.020			
22 23	593	65.	Kinsella J (1983) Relationships between structure and functional properties of food proteins. In: Fox P,			
24 25 26	594		Condon S (eds) Food proteins. Applied Science, London, p 51			
27 28	595	66.	Zhou J, Liu J, Tang X, Tang CX (2018) Effects of whey and soy protein addition on bread rheological			
29 30 31	596	6 property of wheat flour. J Texture Stud 49:38–46. https://doi.org/10.1111/jtxs.12275				
32 33	597	67.	Thrane M, Paulsen PV, Orcutt MW, Krieger TM (2017) Soy Protein: Impacts, Production, and			
34 35	598		Applications. In: Nadathur SR, Wanasundra JPD, Scanlin L (eds) Sustainable Protein Sources, 1st ed.			
36 37 38	599		Academic Press, London			
39 40	600	68.	Berardy A, Costello C, Seager T (2015) Life Cycle Assessment of Soy Protein Isolate. Proc Int Symp			
41 42	601		Sustain Syst Technol 3:13. https://doi.org/10.6084/m9.figshare.1517821			
43 44 45 46	602					
47						
48 49						
50 51						
52						
53 54						
55 56						
57						
58 59						
60						

Composition [g/100 g]	LPI-UF	LPI-IEP
Protein	93.7 ± 0.34^{a}	85.13 ± 0.76^{b}
Fat	4.40 ± 0.13^{a}	$4.49\pm0.37^{\text{a}}$
Starch	*N.D.	*N.D.
Moisture	5.63 ± 0.02^{a}	$4.87\pm0.08^{\rm b}$
Ash	3.51 ±0.11 ^a	$5.46\pm0.04^{\rm b}$
Insoluble dietary fibre	< 0.1	<0.1
Soluble dietary fibre	< 0.1	1.8
Minerals [mg/kg]		
Chlorine	2.0	2.4
Sodium	1300	11000
Zinc	57	48
Calcium	2200	710
Magnesium	2300	750
Iron	150	170
Phosphorous	6100	9400

Table 1. Macro- and micro-nutrient composition of lentil protein isolates obtained by ultrafiltration (LPI-UF) or isoelectric precipitation (LPI-IEP).

lare a superv. Values within a column that share a superscript are not significantly different from one another (p<0.05).

*N.D. = Not Detected

http://mc.manuscriptcentral.com/efrt

Table 2. Particle size distribution parameters of 1% (w/v) protein solutions, surface hydrophobicity,
sulfhydryl groups, water and oil holding capacity of lentil protein isolates obtained by ultrafiltration (LPI-
UF) and isoelectric precipitation (LPI-IEP).

	LPI-UF	LPI-IEP
Particle size distribution [µm]		
D _{4,3}	$32.8\pm3.21^{\text{a}}$	$29.4\pm0.64^{\text{a}}$
D _{3,2}	$23.3\pm0.91^{\text{a}}$	18.1 ± 1.37^{b}
Dv (10)	$12.5\pm0.26^{\rm a}$	$9.02\pm0.24^{\rm b}$
Dv (50)	$26.9\pm1.31^{\rm a}$	$19.0 \pm 1.41^{\text{b}}$
Dv (90)	$62.8\pm9.03^{\rm a}$	56.6 ± 5.81^{a}
Surface Hydrophobicity	$2411\pm49.5^{\rm a}$	2688 ± 92.8^{b}
Free Sulfhydryl groups [µM/g protein]	$5.88\pm0.01^{\text{a}}$	$6.04\pm0.58^{\rm a}$
Total Sulfhydryl groups [µM/g protein]	$22.5\pm0.15^{\text{a}}$	$23.9\pm1.42^{\mathrm{a}}$
Water holding capacity [g water/g protein]	3.96 ± 0.2^{a}	$2.6\pm0.11^{\text{b}}$
Fat binding_holding_capacity [g oil/g protein]	2.24 ± 0.16^{a}	$2.09\pm0.23^{\text{a}}$

Values within a column that share a superscript are not significantly different from one another

(p < 0.05).

Table 3. Foaming properties of protein solutions of lentil protein isolates obtained by ultrafiltration (L	PI-
UF) and isoelectric precipitation (LPI-IEP)	

Protein concentration (w/v)	Foam capacity (%)		Foaming stabilit	y after 60 min (%)
	LPI-UF	LPI-IEP	LPI-UF	LPI-IEP
0.1	9.42 ± 1.26^{a}	$6.52\pm0.00^{\text{b}}$	$0.00\pm0.00^{\mathrm{a}}$	0.00 ± 0.00 a
0.5	18.1 ± 1.26^{a}	18.8 ± 1.26^{a}	6.52 ± 5.75 a	5.80 ± 3.32 a
1.0	33.3 ± 2.51^{a}	33.3 ± 11.2^{a}	15.9 ± 4.53 a	12.3 ± 1.26 ^a
3.0	$58.7\pm9.48^{\mathrm{a}}$	$51.4\pm6.28^{\rm a}$	43.5 ± 5.75 a	31.8 ± 5.47 a
3.3	69.6 ± 3.77^{a}	57.2 ± 5.47^{b}	44.9 ± 1.26^{a}	39.1 ± 5.75 a

Values within a column that share a superscript are not significantly different from one another (p < 0.05).

1	
2	
3	
4	
5	
6	
7	
, 8	
-	
9	
10	
11	
12	
13	
14	
15	
16	
17	
18	
19	
20	
21	
22	
23	
24	
24	
26	
27	
28	
29	
30	
31	
32	
33	
34	
35	
36	
37	
38	
39	
40	
40	
42	
43	
44	
45	
46	
47	
48	
40 49	
49	
50	
51	
52	
53	

60

1

Table 4. Gelling and emulsifying properties of protein solutions of lentil protein isolates obtained by ultrafiltration (LPI-UF) and isoelectric precipitation (LPI-IEP).

	LPI-UF	LPI-IEP
Emulsifying properties		
Emulsifying activity [m ² /g]	14.3 ± 1.22^{a}	$16.5\pm0.03^{\rm b}$
Emulsifying stability [min]	$63.8\pm6.70^{\mathrm{a}}$	$51.0\pm0.96^{\text{b}}$
Least gelation concentration % [w/v]	$11.0\pm0.00^{\mathrm{a}}$	16.0 ± 0.00^{b}
Texture profile analysis of LPI gels		
Hardness [mN]	$2055\pm114^{\text{a}}$	669 ± 20.2^{b}
Adhesiveness [mN/s]	$\textbf{-98.7} \pm 9.02^{a}$	-83.7 ± 1.53^{b}
Springiness	$0.47\pm0.08^{\rm a}$	$0.32\pm0.04^{\rm a}$
Cohesiveness	$0.30\pm0.01^{\rm a}$	$0.30\pm0.02^{\rm a}$
Resilience	$0.05\pm0.01^{\rm a}$	$0.03\pm0.00^{\mathrm{a}}$
Gumminess	623 ± 53.7^{a}	210 ± 4.36^{b}
Chewiness	257 ± 83.1^{a}	76.3 ± 7.09^{b}

Values within a column that share a superscript are not significantly different from one another (p < 0.05).

Ce perez

	LPI- IEP	LPI-UF
Environmental impact potentials (LCA):		
Climate Change [kg CO ² -e/kg PI]	3.53	4.17
Aquatic Eutrophication [g PO ₄ -e/kg PI]	111	103
Terrestrial Eutrophication [g PO ₄ -e/kg PI]	1.57	1.77
Acidification [g SO ₂ -e/kg PI]	14.5	18.2
Photochemical Oxidant Formation [g O ₃ -e/kg PI]	2.22	2.17
Fine Particulate Matter [g PM2.5-e/kg PI]	11.9	14.9
Stratospheric Ozone Depletion [mg CFC11-e/ PI]	58.2	55.1
Additional indicators at the inventory level (LCI):		
Phosphorus Use [g/kg PI]	245	229
Cumulative Energy Demand, non-renewable [MJ/kg PI]	45	59
Blue Water (process) [kg/kg PI]	42	49
Land Use [m ² /kg PI]	57	53

Table 5. Environmental impact profile of lentil protein isolates per kg isolate (PI), prepared by ultrafiltration (LPI-UF) and isoelectric precipitation (LPI-IEP)

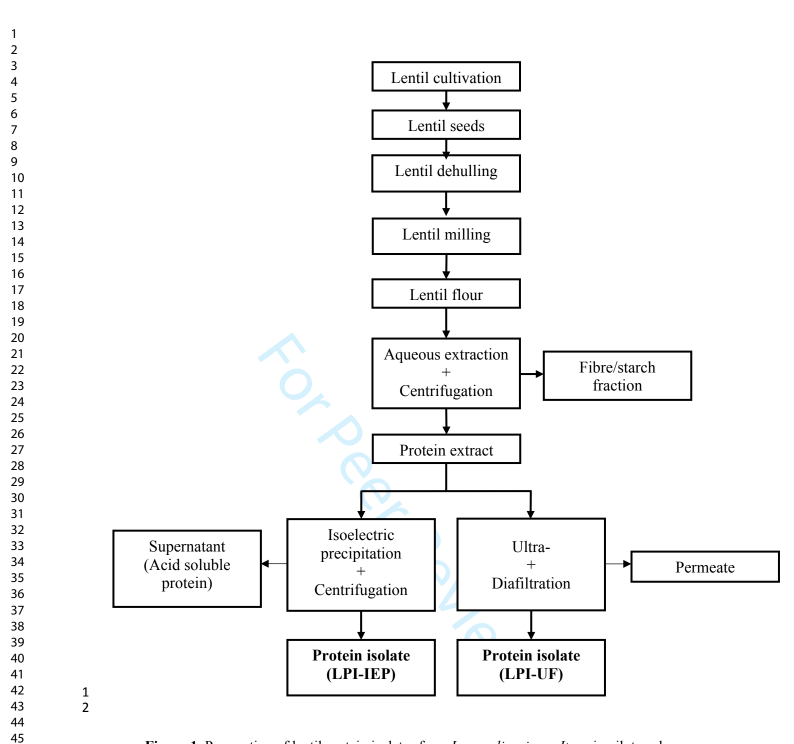


Figure 1. Preparation of lentil protein isolates from Lens culinaris cv. Itaca in pilot-scale.

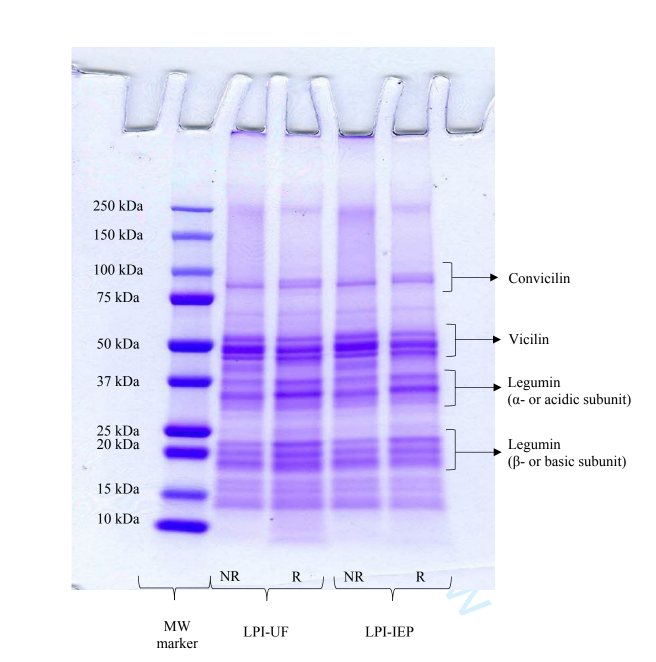


Figure 2. Representative sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) pattern of lentil protein isolates obtained by ultrafiltration (LPI-UF) and isoelectric precipitation (LPI-IEP) under non-reducing (NR) and reducing (R) conditions. The first lane of the gel contains the molecular weight marker.

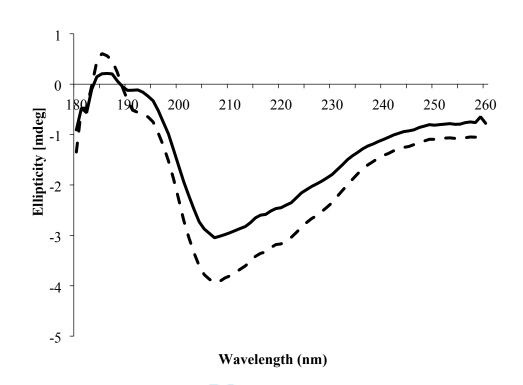


Figure 3. Far-UV circular dichroism (CD) spectra (smoothened curve) of lentil protein isolates obtained by ultrafiltration (--) or isoelectric precipitation (--).

Per.ez

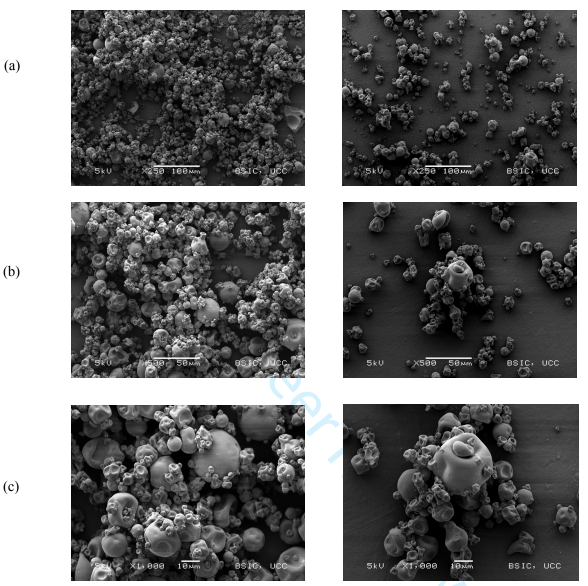
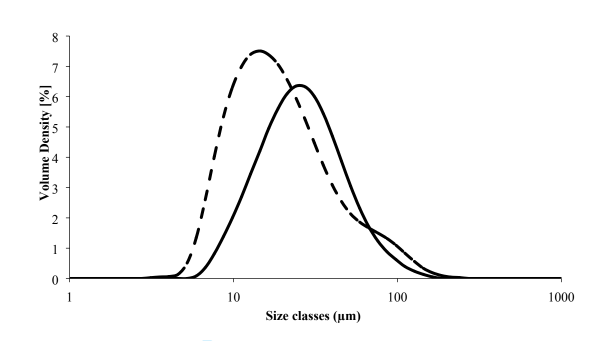


Figure 4. Scanning electron micrographs of isoelectric precipitated (column 1) and ultrafiltrated (column 2) lentil protein isolate powder ingredients. Magnification of row (a) 250, (b) 500 and (c) 1000. Scale bars $10 \mu m$.

http://mc.manuscriptcentral.com/efrt



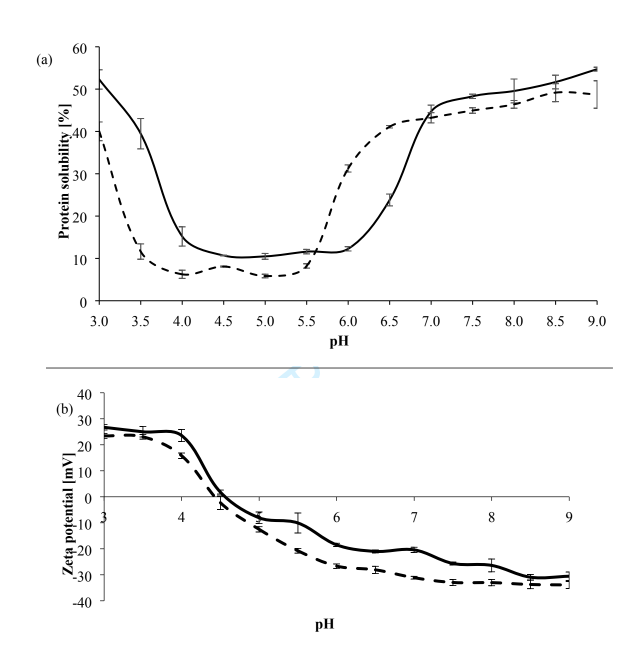


Figure 6. Protein solubility (a) and zeta potential (b) values at different pH ranges of lentil protein isolates obtained by ultrafiltration (---) and isoelectric precipitation (---).

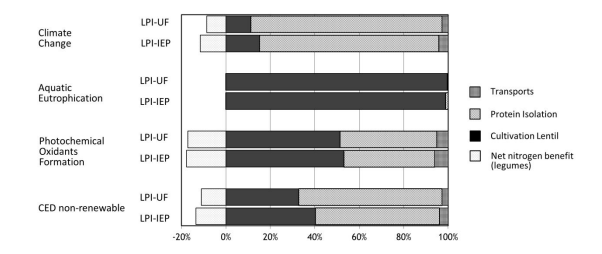


Figure 7. Contributions of main life cycle steps to environmental impact profiles of lentil protein isolates prepared by ultrafiltration (LPI-UF) and isoelectric precipitation (LPI-IEP). CED = Cumulative primary energy demand

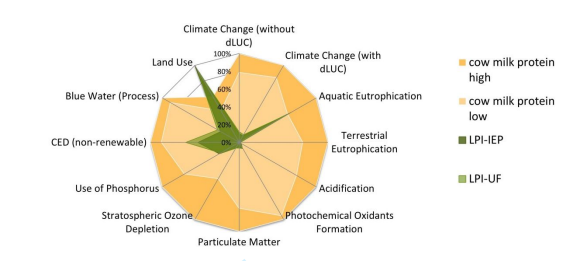


Figure 8. Comparison of environmental impact profiles of lentil protein isolates versus cow milk protein ranges, lentil protein isolates are prepared by ultrafiltration (LPI-UF) and isoelectric precipitation (LPI-IEP). Highest result is set to 100%. dLUC = direct land use change