

Modeling the Thermal Denaturation of the Protein–Water System in Pulses (Lentils, Beans, and Chickpeas)

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18 Abstract

19 Thermal treatment applied during the cooking of pulses leads to denaturation and even 20 aggregation of the proteins, which may impact protein digestibility. Thermal transitions of lentil, chickpea and bean proteins were studied using DSC. Protein-enriched samples were 21 22 obtained by dry air classification of dehulled seeds and were heated up to 160 °C, with water contents ranging from 0.2 to 4 kg/kg dry basis. The DSC peaks of the resulting endotherms 23 24 were successfully modeled as overlapping Gaussian functions. The denaturation temperatures 25 were modeled as a function of temperature according to the Flory-Huggins theory. The 26 modeling allows to calculate the degree of protein transition for any temperature and moisture 27 condition. The denaturation diagrams reflect the different protein compositions of lentil, 28 chickpea and bean (particularly the 11S/7S globulin ratio). Chickpea proteins were more thermally stable than those from lentil and bean. Proteins underwent an irreversible transition, 29 30 suggesting that unfolding and aggregation were coupled.

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33 Keywords

34 Protein denaturation, DSC, Mathematical modeling, Pulses, Globulin, Peak desummation

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37 1. Introduction

38 According to the FAO¹, pulses are legume crops that are harvested for their edible dry seeds 39 but are not used for oil extraction. The International Year of Pulses (2016) highlighted their 40 important nutritional benefits as part of sustainable food production targeting food security and 41 nutrition. The seeds are rich in carbohydrates including starch, as well as in proteins and dietary fibers, and are a source of minerals and vitamins.² In particular, their high protein content (15-42 43 30 % dry weight)³ makes pulses an excellent source of plant-based proteins for the formulation 44 of food and ingredients. However, the impact of processing on protein bioavailability and 45 digestibility is important to take into account. In particular, thermal treatment during cooking leads to protein denaturation and aggregation⁴, the latter being one of the main adverse effects 46 on protein digestibility.^{5,6} 47

48 Inter and intra molecular bonds that stabilize the native structure of the protein are allowed to break down during heating, depending on the temperature applied.⁴ This transition from a native 49 50 state to a more disordered arrangement is called denaturation. Pulse proteins, which are 51 oligomeric proteins with quaternary structure, undergo reversible dissociation into monomers.⁷ 52 Both the tertiary and secondary structure are affected with irreversible unfolding that generally remains partial.⁴ Denaturation of pulse proteins occurs between 80 °C and 100 °C in excess 53 water.^{8–10} This spatial reorganization leads to exposure of the reactive groups, especially the 54 hydrophobic amino acid residues embedded in the core of the native protein.⁴ As a consequence, 55 56 the formation of new bonds and interactions between partially unfolded peptide chains is 57 favored. The protein aggregation that follows the denaturation process results in the formation 58 of soluble or insoluble high molecular weight complexes. Protein aggregation reduces access 59 to certain essential amino acids that are specific for protease action, and that are trapped inside the aggregate structures.¹¹ Heat-induced aggregation can thus impair protein digestibility of 60 cooked pulses⁶, to an even greater extent if aggregation is high.⁵ Therefore, a better 61

62 understanding of the denaturation-aggregation status of proteins is needed, as a function of the 63 cooking conditions to enable more accurate monitoring of the nutritional aspect. It should be 64 noted that the presence of other factors that affect digestibility such as heat-sensitive anti-65 nutritional factors (enzymes inhibitors) also requires monitoring.

66 Previous studies on the denaturation of legume proteins mainly focused on thermal stability at 67 constant water content. However, cooking in water implies that pulses undergo hydration. The water content in the seed increases from the initial value to reach equilibrium moisture.¹² Phase 68 69 transitions of other components of the seed, *i.e.* gelatinization of starch, also result in varying 70 hydration rates, meaning the water content in the protein pool changes during cooking. 71 Denaturation-aggregation thus needs to be described as a function of the water content. A 72 similar approach to the one developed for the modeling of gelatinization-melting of starch at increasing temperature and water content¹³ should be suitable. In the present study, the phase 73 transitions of proteins were monitored using differential scanning calorimetry (DSC), a reliable 74 75 and convenient thermo-analytical method to study the endothermic and exothermic changes associated with the denaturation and aggregation of proteins.^{14,15} To avoid the structural 76 77 modifications usually caused by chemical extraction reported in most studies, we chose to work 78 on protein-enriched samples obtained using a physical separation method only, *i.e.* air 79 classification. The study was focused on beans, lentils and chickpeas, which are widely produced and the most consumed pulses in France.¹⁶ 80

81

82 2. Materials and methods

83 2.1 Raw material

Green lentils (*L. culinaris*, var. Anicia), chickpeas (*C. arietinum*, var. Elvar) and navy beans
(*P. vulgaris*, var. Linex) were provided by Cibèle (Saint-Georges-Sur-Arnon, France), Moulin

- 4 -

Marion (Saint-Jean-sur-Veyle, France) and Cavac (La Roche-sur-Yon, France), respectively.
The lentils were harvested in 2017 and the chickpeas and beans in 2018. All the seeds were
stored in a vacuum pack at 7 °C until use.

89 2.2 Sample preparation

90 Protein-enriched samples were prepared by dry fractionation and air classification as described in Lefèvre et al.¹³ Briefly, lentil seeds were dehulled by dry abrasion and residues of the hull 91 92 were removed by sieving through a 2-mm mesh screen. Chickpea and bean seeds were crushed 93 using a cutting mill equipped with a 8-mm sieve and residues of the hull were removed with a vibrating sieve. For all pulses, the removed hulls represented 10-12% of the raw weight, and 94 95 the process induced 6-8% of losses. The dehulled and crushed pulses were ground into flour in 96 a high speed impact mill equipped with a pin mill. The resulting flours were separated into 97 coarse and fine fractions using an ATP air classifier (Hosokawa Alpine, Augsburg, Germany) 98 with the following optimized speeds: 6 500 rpm for lentils, 10 000 rpm for chickpeas and 99 8 000 rpm for beans. The resulting fine fractions were stored at 4 °C until use and are hereafter 100 referred to as protein-enriched samples.

101 2.3 Sample characterization

102 Water content of the protein-enriched samples was calculated on a wet basis by drying 5 g of each sample for 2 h at 132 °C (± 2 °C) using the NF EN ISO 712 standard method (2010). 103 104 Starch content was measured using the enzymatic procedure of Holm *et al.*¹⁷ Protein content 105 was measured using the Kjeldhal method according to NF EN ISO 20483, with 5.4 as nitrogen conversion factor.¹⁸ Lipid content was measured using the gravimetric ether extraction method 106 according to AOAC 2003.05.19 Soluble and insoluble fibers were measured using a TDF 107 Analyzer (Ankom, Macedon, USA) according to AOAC 2011.25 and AACC 32-50.01.²⁰ 108 Phytates were extracted according to a published method²¹ with some modifications. A 100 mg-109

sample was added to a Pyrex vial with 5 mL of 0.5 M HCl. The capped vial was heated under stirring for 6 min in boiling water and then centrifuged for 20 min at 4 500 g. The supernatant was recovered and filtered through a 0.2 µm syringe filter (Acrodisc) and 0.5 M NaOH was added for neutralization and the solution was then diluted (1:10). Phytic acid content was measured by high performance ion chromatography (HPIC) using an ICS-2500 Dionex chromatograph (Sunnyvale, CA, USA). Fiber and phytate measurements were made in triplicate and all other measurements were made in duplicate.

117 2.4 Differential scanning calorimetry

Thermal denaturation of proteins was studied by DSC as described in Lefèvre et al.¹³ Briefly, 118 119 a Perkin DSC 8500 (Perkin Elmer, Norwalk, USA) was calibrated using indium as standard. 120 Protein-enriched samples and deionized water were weighed in stainless steel pans to obtain a water content X ranging from 0.2 to 4 kg kg⁻¹ dry basis (db). This range covers the progressive 121 hydration of proteins while the pulses are cooking. The pans were heated from 20 °C to 160 °C 122 123 at a rate of 10 °C/min, using an empty sealed pan as reference. All measurements were 124 duplicated. A blank thermogram (empty pans in both reference and sample ovens) was recorded 125 daily. The heat flow (mW) of the sample pans minus the variation in the heat flow of the blank 126 during heating was recorded using Pyris Thermal Analysis Software (Perkin Elmer, Norwalk, 127 USA). Table 1 lists the experimental conditions applied for the different water contents. In the 128 study conditions (*i.e.* at $T \le 160$ °C), the lowest water content to be able to detect a complete DSC signal was 0.2, 0.3 and 0.4 kg kg⁻¹ db, for beans, lentils and chickpeas, respectively. When 129 130 the heat flows were similar for close water contents, fewer intermediate water contents were 131 used. In addition, double scans were performed with an intermediate water content of 0.8 kg kg⁻¹ db. Samples were heated from 20 °C to 80 °C, 110 °C or 160 °C at a rate of 132 133 10 °C/min, then cooled to 20 °C and heated a second time from 20 °C to 160 °C at a rate of 134 10 °C/min.

135 [Insert Table 1 here]

136 2.5 Modeling the protein denaturation diagram

137 DSC thermograms record heat flow (φ) as a function of temperature (*T*) at each water content. 138 Phase transitions of the sample components result in peaks in the endotherm. The DSC 139 thermograms obtained from the protein-enriched samples are discussed as if the peaks were 140 only due to the phase transition of proteins. This hypothesis will be confirmed in the results 141 part.

142 2.5.1 Modeling the DSC peaks

143 Thermal denaturation of the protein-enriched samples resulted in single or double peaks in the 144 DSC signal depending on the variety of the pulse concerned. The shape of all the endotherms 145 depended on water content. Therefore, heat flows were analyzed according to the desummation procedure described in Lefèvre et al.¹³ on pulse starches. First, baseline subtraction and non-146 147 dimensionalization were performed on heat flow values in all pulses. In the lentil and chickpea 148 samples, the heat flow was composed of two roughly overlapping endotherms depending on 149 the water content, termed P1 and P2. The dimensionless heat flows ($\overline{\phi}$) were fitted to a sum of 150 two Gaussian functions (i = P1 or P2) as follows:

151
$$\overline{\varphi} = \sum_{i=\text{Pl},\text{P2}} \frac{\beta_i}{\Delta T_i \sqrt{2\pi}} \exp\left(-\frac{1}{2} \left(\frac{T-T_i}{\Delta T_i}\right)^2\right)$$
(1)

where T_i (°C) is the temperature at maximum peak *i*; ΔT_i (°C) controls the width of the peak *i* and is related to the full width at half maximum (FWHM) of the peak *i* according to $2\sqrt{2\ln(2)} \times \Delta T_i = FWHM$; β_i is the dimensionless area of peak *i*. Therefore:

155
$$\int_{-\infty}^{\infty} \overline{\varphi} = \beta_{P1} + \beta_{P2} = 1$$
(2)

156 In bean, the heat flow had a single endotherm (i = P1) irrespective of water content, so $\beta_{P1} = 1$. 157 Eq. (1) was similar to the equation used previously for modeling starch conversion heat flow¹³ 158 which was composed of overlapping gelatinization and melting peaks. The parameters from 159 Eq. (1) depend on the water content X in the samples.

160 2.5.2 Peak temperature

161 The temperature at maximum peak T_i (°C) is the denaturation temperature of the proteins 162 responsible for transition *i* (*i* = P1 or P2). The Flory–Huggins equation²² was used to describe 163 the relation between T_i and the volume fraction of the water (ϕ) in the protein-water mixture:

164
$$\frac{1}{T_i} - \frac{1}{T_{i,0}} = \frac{R}{\Delta h_{i,0}} \frac{v_a}{v_w} \left(\phi - \chi_i \phi^2 \right)$$
(3)

where $\Delta h_{i,0}$ (J mol⁻¹) is the change in the molar enthalpy of denaturation per repeating unit 165 (amino acid); v_a / v_w is the ratio of the molar volume of the repeating unit 166 $(v_a = 74.5 \times 10^{-6} \text{ m}^3 \text{ mol}^{-1})^{23}$ to the molar volume of the water $(v_w = 18.1 \times 10^{-6} \text{ m}^3 \text{ mol}^{-1})$ and, 167 therefore, $v_a / v_w = 4.1$; R is the gas constant (8.31 J mol⁻¹ K⁻¹); $T_{i,0}$ (K) is the denaturation 168 temperature of the pure polymer and χ_i is the Flory interaction parameter. To calculate ϕ , the 169 density of water was taken to be 1 000 kg m⁻³ and the density of protein was attributed an 170 average value of 1 350 kg m⁻³.²⁴ Therefore, ϕ (m³ m⁻³) was expressed as a function of water 171 172 content X:

173
$$\frac{1}{\phi} = 1 + \frac{x_p^{ab}}{1.35X}$$
 (4)

174 where x_p^{db} (kg kg⁻¹ db) is the mass fraction of protein in the protein-enriched samples.

175 2.5.3 Peak width

176 The width-related parameter ΔT_i represents the cooperativity of protein unfolding process and 177 the presence of multiple protein domains with heterogeneous thermal stabilities.^{4,25} ΔT_i was 178 likely to decrease as a function of water content. The following empirical relation was used to 179 describe ΔT_i as a function of *X*:

180
$$\Delta T_{i} = \Delta T_{i,\infty} + \left(\Delta T_{i,0} - \Delta T_{i,\infty}\right) \exp\left(-\frac{X}{\gamma_{i}}\right)$$
(5)

181 where $\Delta T_{i,0}$ (°C) and $\Delta T_{i,\infty}$ (°C) are the values for ΔT_i , with zero and excess water contents, 182 respectively; γ_i is the rate parameter of decrease for the two correlations.

183 2.5.4 Amplitude of the peak area

184 The effect of water content on the relative peak areas β_i was also investigated. With increasing 185 water content, β_{P1} was likely to decrease. The following empirical equation was used to 186 describe this relation for lentil and chickpea:

187
$$\beta_{P1} = \beta_{P1,\infty} + (\beta_{P1,0} - \beta_{P1,\infty}) \exp(-X)$$
 (6a)

188 where $\beta_{P1,0}$ and $\beta_{P1,\infty}$ are the dimensionless area of the first peak, with zero and excess water 189 contents, respectively. β_{P2} was calculated by combining Eq. (2) and Eq. (6a) as follows:

190
$$\beta_{P2} = 1 - \left[\beta_{P1,\infty} + (\beta_{P1,0} - \beta_{P1,\infty}) \exp(-X)\right]$$
 (6b)

191 2.5.5 Degree of protein denaturation

192 The degree of protein denaturation τ was defined for any temperature *T* and water content *X* 193 as the ratio between the change in enthalpy calculated from the beginning of the first peak to *T*, 194 and the change in the whole enthalpy from the beginning of the first peak to the end of the 195 second peak (Eq. (7a)). Denaturation refers to the global transition of proteins, which includes 196 unfolding and potential aggregation.

197
$$\tau = \int_{0}^{T} \overline{\varphi}(T, X) dT / \int_{0}^{\infty} \overline{\varphi}(T, X) dT$$
(7a)

Using the error function to calculate integral Gaussian functions, Eq. (7a) becomes Eq. (7b) for lentil and chickpea (with i = P1 and P2) and for bean (with only i = P1).

200
$$\tau = \sum_{i=\text{P1,P2}} \beta_i \times \frac{1}{2} \left(1 + \operatorname{erf}\left(\frac{T - T_i}{\Delta T_i \sqrt{2}}\right) \right)$$
(7b)

201 Finally, isovalue lines of the degree of protein denaturation τ were represented as a 202 temperature *T* versus water content *X* diagram.

203 2.5.6 Identification of parameters

The modeling of protein denaturation diagram involves 9 parameters for lentil ($T_{i,0}$, $\Delta h_{i,0}$, χ_i , ΔT_i , β_{P1}), 13 parameters for chickpea ($T_{i,0}$, $\Delta h_{i,0}$, χ_i , $\Delta T_{i,0}$, $\Delta T_{i,\infty}$, γ_{P2} , $\beta_{P1,0}$ and $\beta_{P1,\infty}$) and 4 parameters for bean ($T_{P1,0}$, $\Delta h_{P1,0}$, χ_{P1} , ΔT_{P1}), with i = P1 or P2. The overall procedure¹³ was used to identify all the parameters. At the same fitting session, all dimensionless heat flow thermograms were fitted to the model combining Eq. (3), (5) and (6a) in Eq. (1). The curve fitting toolbox (Matlab software, version R2019b, The MathWorks Inc., Natick, USA) with the Levenberg-Marquardt algorithm was used as a solver.

211 2.5.7 Statistical methods

All parameter values presented in this study are given with a 95% confidence interval. The root mean square error (RMSE) was calculated between experimental and predicted dimensionless heat flows for all the experimental conditions.

215

216 3. Results and Discussion

217 *3.1 Characterization of the samples*

218 [Insert Table 2 here]

219 The chemical composition of the protein-enriched fractions obtained after dry fractionation and 220 air classification are listed in Table 2. Proteins were the main component in the fine fractions, with x_p^{db} ranging from 38.2% of db weight for chickpea to 55.8% db for lentil. These values 221 are in good agreement with protein concentration range of 40-60% db reported in literature for 222 air-classification on pulse flours.²⁶⁻²⁸ Protein contents of the dehulled seeds were 24.1% db, 223 19.3% db and 23.2% db in lentil, chickpea and bean, respectively. The dry air-classification 224 225 process can thus be considered as an efficient physical method to produce protein-enriched 226 samples while avoiding the structural modifications that may occur with chemical extraction. Tyler et al.²⁷ already reported good protein separation efficiency using air-classification, 227 particularly for green lentil and navy bean. Insoluble dietary fibers (IDF) were the second 228 229 component in the protein-enriched samples, ranging from 11.1% db for lentil to 18.5% db for 230 chickpea. Other components listed in Table 2 represented all together 25-37% db of the samples 231 and between 2 and 11% db each. The interactions between proteins and those non-proteins 232 components may affect the thermal stability of proteins²⁹ and the water repartition due to the competition for available water³⁰. These interactions may impact the denaturation temperature 233 234 of proteins and were intrinsic to the protein-enriched samples studied here. There have not been 235 investigated further.

236

6 *3.2 DSC characteristics of the protein-enriched fractions*

- 237 [Insert Fig. 1 here]
- 238 [Insert Fig. 2 here]
- 239 [Insert Table 3 here]

For each pulse, Fig. 1 presents a selection of three DSC thermograms at X = 0.4, 1.0 and 2.0 kg kg⁻¹ db. All transition peaks were observed above 60 °C, so the temperature range of interest was between 60 and 160 °C. According to the protein enrichment of the fine fractions

after air-classification in comparison with other components (Table 2), we confirmed the 243 hypothesis that the DSC thermograms were mainly due to protein denaturation. Indeed, the 244 245 second main component of the samples, IDF, underwent no thermal transition below 200 °C.³¹ Denaturation of phytates and alpha-galactosides also occurred at temperatures exceeding the 246 247 conditions of the study, whereas the melting point of lipids was lower than the temperature 248 range of interest. Thus, only the gelatinization of residual starch could lead to additional DSC 249 peaks in the temperature range of interest. However, the starch content was less than 6% db in 250 all samples, resulting in a low contribution to the total enthalpy of the observed transition. The 251 DSC peak were thus mainly attributed to the phase transitions of proteins.

252 The experiments revealed that the thermal behavior of proteins depended on both the water 253 content and the type of pulse (Fig. 1). The DSC thermograms for lentils showed two roughly 254 overlapping peaks P1 and P2, depending on the moisture content of the sample. With lower water content (i.e. $X \le 2 \text{ kg kg}^{-1} \text{ db}$), a biphasic endotherm was observed. With excess water, a 255 256 single broad peak was observed, suggesting that both peaks are merged as modeled by the 257 Gaussian functions (Fig. 1c). The endotherm for chickpea had two distinct peaks, P1 and P2, 258 regardless of the water content. The shape and size of the peaks changed with the moisture 259 content. The endotherm for bean had a single peak, P1, of constant size and shape, regardless 260 of the water content. All DSC thermograms were successfully modeled as shown by the low 261 RMSE between experimental and predicted heat flows (Fig. 1). The parameters presented in 262 Table 3 had low confidence intervals, which allows accurate modeling of experimental data. 263 Modeling is particularly useful to understand the superposition of P1 and P2 endotherms.

The multiple or overlapping peaks observed by DSC showed that the thermal stability of the proteins varied. This is linked to the different protein fractions in the seed. The main proteins in pulses are globulins and albumins in varying proportions depending on the species and variety of the pulse concerned.^{3,32} Almost all pulses contain two main types of globulins with

sedimentation coefficients around 7S and 11S called vicilin and legumin, respectively.^{33,34} The 268 269 temperature, shape and size of the denaturation endotherms presented in Fig. 1 were thus 270 determined mainly by the thermal denaturation of 7S and 11S globulins. Vicilin has a trimeric 271 structure stabilized by hydrophobic interactions, electrostatic and hydrogen bonds. Legumin 272 has a quaternary structure made of three dimers held together by non-covalent bonds. In each dimer, acidic and basic subunits are associated with disulfide bridges.⁷ The covalent bonds give 273 274 legumin higher thermal stability than vicilin, because more energy is needed to break the bonds.³⁵ In excess water, 7S and 11S globulins unfold at 80-85 °C and 90-100 °C, 275 respectively.^{10,36} In lentil, endotherms P1 and P2 were attributed to the denaturation of 7S and 276 11S globulin, respectively, with peak temperatures at 83 °C and 91 °C (with $X = 3 \text{ kg kg}^{-1} \text{ db}$). 277 $\beta_{P_{1,0}}$ and $\beta_{P_{1,\infty}}$ did not differ significantly in lentils, β_{P_1} thus remained constant and equal to 278 279 0.65 irrespective of the water content (Table 3). This value suggests that the P1 transition was 280 slightly predominant than to P2 transition. In chickpea, endotherms P1 and P2 were again 281 attributed to the successive denaturation of 7S and 11S globulin, respectively, with peak temperatures at 83 °C and 103 °C (with $X = 3 \text{ kg kg}^{-1} \text{ db}$). β_{P1} was lower than 0.5, except at the 282 lowest water contents ($X \le 0.5 \text{ kg kg}^{-1} \text{ db}$). This trend suggests that the legumin transition was 283 284 predominant in chickpea. This result is in good agreement with the high 11S/7S ratio commonly reported for several chickpea species.^{37–39} The decreasing function for β_{P1} (Fig. 2e) suggests 285 286 that part of the 7S globulin was solubilized without denaturation with increasing water content. 287 This hypothesis is consistent with the decrease in total enthalpy of phase transition observed before non-dimensionalization (from 21 to 10 J g^{-1} protein when X increased from 0.4 to 288 289 1 kg kg⁻¹ db). In some cases, denaturation peaks of vicilin and legumin in chickpea have been reported to overlap.^{10,37} In bean, the single peak P1 at 92 °C (with $X = 3 \text{ kg kg}^{-1} \text{ db}$) was 290 291 attributed to the predominant vicilin fraction, as already suggested in several bean varieties.^{9,14,40} The diversity of DSC signals between lentil, chickpea, and bean clearly reflects 292

the variations in the 11S/7S ratio between pulses, as already reported.^{26,41} In this sense, DSC analysis can be used to rapidly determine globulin ratios when determination using chromatography and electrophoresis on protein isolates is not possible.

296

3.3 Modeling of peak width and temperature

No significant difference was observed between $\Delta T_{i,0}$ (°C) and $\Delta T_{i,\infty}$ in lentil and bean, the 297 peak widths ΔT_i were thus constant and equal to $\Delta T_{i,\infty}$ (Table 3). This low value (3-4 °C) 298 299 reflected highly cooperative unfolding for vicilin and legumin, irrespective of the water content.²⁵ In chickpea, ΔT_i decreased with a reduction in water content, as shown by the 300 301 decreasing exponential law (Fig. 2 and Table 3). The progressive sharpening of both P1 and P2 302 peaks showed that increasing moisture led to a more cooperative denaturation of chickpea 303 vicilin and legumin.⁴ The heterogeneous composition of the protein-enriched samples may also 304 play a role in the DSC characteristics. The lower protein content in chickpea than in lentil and 305 bean (Table 2) may lead to a broader and more water-sensitive endotherm in the DSC 306 thermogram due to the lower protein purity and hence, possible interaction with residual components, in particular with the higher IDF content.²⁹ 307

The denaturation temperatures T_i of lentil, bean and chickpea proteins decreased with an 308 309 increase in water volume fraction (Fig. 2). Temperatures were modeled using the Flory-310 Huggins theory for a polymer in solution (Eq. 3). The coefficient of determination (R²) was 0.98-0.99 for $T_{\rm Pl}$ and 0.96-0.99 for $T_{\rm P2}$. The fitting strength shows that this model fits both 7S 311 312 and 11S protein denaturation in all pulses studied. Concerning the P1 transition, the denaturation temperatures of pure polymer were similar for lentil and bean, with a $T_{P1,0}$ value 313 of 165.2 °C and 166.2 °C, respectively. $T_{\rm P1.0}$ was 30 °C higher for chickpea. Again for the P2 314 transition, $T_{P2,0}$ was 20 °C higher for chickpea legumin compared to lentil legumin (Table 3). 315

316 The denaturation temperatures of pure protein varied more between pulses than the denaturation temperatures of gelatinization and melting of pure starch, obtained with the same Flory-317 Huggins model.¹³ The change in molar enthalpy of denaturation $\Delta h_{i,0}$ ranged from 41.2 kJ mol⁻ 318 ¹ to 65.8 kJ mol⁻¹ for vicilin and from 41.9 to 45.1 kJ mol⁻¹ for legumin (Table 3). These 319 320 enthalpies were higher than the change in molar enthalpy of gelatinization and melting of starch.¹³ The Flory interaction parameter χ_i was low (0-0.14) compared to the range 0.48-0.51 321 previously reported for cereal, potato or pulse starches.^{13,42–44} These values suggests that the 322 interactions between water and proteins were low.⁴⁴ To conclude, all the values determined for 323 Flory-Huggins parameters $(T_{i,0}, \Delta h_{i,0}, \chi_i)$ differ from those previously obtained for starch 324 325 gelatinization and melting modeling. The Flory-Huggins theory is thus an accurate and useful 326 tool for predicting the impact of moisture on the denaturation temperatures of pulse proteins.

327

3.4 Relevance of the predicted denaturation temperatures

Denaturation temperatures T_i modeled using the Flory-Huggins parameters described above 328 329 were consistent with denaturation temperatures already reported for lentil, chickpea and bean proteins. For lentil in excess water ($X = 10 \text{ kg kg}^{-1} \text{ db}$), two overlapping peaks were found at 330 77.9 °C and 83.7 °C, respectively. At the same water content and also in green lentil proteins, 331 Barbana and Boye³⁶ observed a similar result with a broad peak at 79.6 °C. The denaturation 332 333 temperatures of green lentil proteins were lower than those of red lentil proteins, which have been reported to range from 87 °C in excess water to 123 °C with limited water.^{36,45} For 334 chickpea, T_{P_1} and T_{P_2} were predicted to be 83.3-84.9 °C and 101.9-103.6 °C, respectively, when 335 X ranged between 3 and 4 kg kg⁻¹ db. These values are in good agreement with the denaturation 336 337 temperatures of chickpea proteins reported in previous studies for the same range of water content.^{46–48} These authors reported broad peaks ranging from 85 °C to 100 °C, depending on 338 339 the varieties and on the extraction methods used. Our results differ slightly from the two peaks

observed by Chang et al.³⁷ at 79.3 °C and 96.3 °C. At higher water content ($X = 9 \text{ kg kg}^{-1} \text{ db}$), 340 Withana-Gamage et al.¹⁰ reported a vicilin transition at 80-85 °C depending on the variety of 341 chickpea, which is similar to our T_{P1} at 80.5 °C. The same authors reported that the legumin 342 transition occurred at 89.2-93.2 °C, which is lower than our $T_{\rm P2}$ predicted at 99.0 °C. For bean 343 in excess water ($X = 10 \text{ kg kg}^{-1} \text{ db}$), we found a single peak at 87.1 °C. This temperature was 344 slightly lower than the peak temperature of 90.6 °C reported by Rui et al.9 in white bean, the 345 same variety as the navy bean we used in the present study. Yin et al.⁴⁰ and Law et al.⁴⁹ also 346 reported higher peak temperatures for red bean and rice bean, respectively (92.5 °C and 91.9 °C, 347 respectively). Meng and Ma¹⁴ observed a major peak at 86.4 °C close to our result, and a 348 349 shoulder at 92.2 °C for red bean globulin, at the same water content. Their measurements on 350 fractionated fractions showed that the DSC signal could be split into a biphasic endotherm for 351 7S vicilin (peaks at 88 °C and 94 °C) due to non-glycosylated and glycosylated subunits, and a minor endotherm for 11S legumin (peak at 89.5 °C), overlapping the first one. In the present 352 study, it is possible to hypothesize that a peak at a higher temperature than T_{p_1} but with a lower 353 354 amplitude may be hidden under P1 endotherm, corresponding with the denaturation of residual 355 11S legumin.

356 The denaturation temperature of proteins can differ considerably among pulse species, and also 357 between varieties of the same species. In this study, the denaturation temperature of green lentil 358 and navy bean proteins was lower than the denaturation temperatures reported for red lentil and 359 for red and rice beans, respectively. A difference of up to 10 °C has been reported among the denaturation temperatures of proteins in different bean varieties.^{9,49} This variability is linked to 360 the different proportions of storage proteins.³² It is important to underline that the DSC 361 characteristics are also affected by environmental and processing conditions, and protein 362 extraction methods.^{15,36,45} Studying pure vicilin and legumin fractions or a globulin mixture 363 may also lead to slight variations in the denaturation temperatures.^{14,15,37} The purity degree of 364

the protein sample is important because the thermal stability of proteins may be affected by interactions between protein and non-protein components.²⁹ All these factors can explain the variations in denaturation temperatures, number and shape of DSC peak between studies on a similar pulse cultivar.

369 *3.5 Thermal aggregation of proteins*

370 [Insert Fig. 3 here]

371 Double scans were performed in order to determine if the observed transitions were reversible, 372 *i.e.* still observable on the DSC signal after a preheating treatment. The first heating treatments 373 are hereafter referred to as preheating. The DSC signals of the second heating treatments are 374 presented in Fig. 3 and showed that a peak disappears when the temperature of the preheating 375 exceeds the temperature of the transition considered. For all pulses, a preheat treatment to 80 °C 376 did not damage the protein, since the DSC thermograms were similar to the control. This is 377 consistent with the higher denaturation temperatures described above. When lentil and chickpea samples were preheated to 110 °C (Fig. 3a and b), only one transition was observed at a 378 379 temperature corresponding to the P2 transition of the untreated sample (115-120 °C). This 380 means that the vicilin transition P1 was no longer detected, while the legumin peak P2 was still 381 observed. In addition, the legumin peak was no longer detected when the sample was preheated 382 to 160 °C. These results confirm that 11S globulin is more thermally stable than 7S globulin. 383 Also in bean, vicilin was irreversibly damaged at 110 °C (Fig. 3c). In conclusion, both types of 384 protein were irreversibly damaged when the thermal treatment exceeded their denaturation temperature. Ricci et al.²⁹ also observed irreversible denaturation process using DSC. The 385 386 authors reported additional reversible glass transition for lentil, bean and chickpea (X = 0.15kg kg⁻¹) ranging from 150 to 170 °C. The differences in water content, heating temperatures, 387 388 protein purity and pulse variety may explain why a similar glass transition was not observed on 389 the DSC thermograms in Fig. 3.

390 The irreversibility of the transition suggests that thermal denaturation involves both unfolding of the structure and aggregation between protein chains.^{10,11} Partially denatured proteins may 391 form a soluble and insoluble complex with new interactions. German et al.⁸ showed that the 392 393 dissociation of 11S purified soy proteins during heating was followed by aggregation of the 394 released basic subunits into insoluble complexes. In the presence of 7S proteins, basic 11S 395 subunits and 7S subunits can form soluble complexes. Denaturation and aggregation seemed to be coupled in the present study, or at least not separable using DSC analysis. Sirtori et al.³⁵ also 396 397 reported irreversible denaturation of pea globulins. These authors reported that the vicilin peak 398 disappeared from the DSC thermogram when samples were pre-heated to 200 °C, or heated at 399 100 °C for a long time (120-300 min). The 11S peak was visible in all conditions, even after 400 harsh treatment at 200 °C, but its detergent solubility decreased.

401 *3.6 Modeling the denaturation diagram of proteins*

402 [Insert here] Fig. 4.

403 The results obtained with desummation modeling allowed us to calculate the degree of protein 404 denaturation (including unfolding and aggregation) for any temperature T and water content X. 405 Some isovalue lines of degree of protein denaturation are presented as a T versus X diagram in 406 Fig. 4. The differences we observed in the denaturation temperatures between the three pulses, 407 related to their respective protein composition, are still visible in the diagram. The higher 408 11S/7S ratio for chickpea led to isovalue lines of denaturation degree at higher temperatures 409 than for bean and lentil proteins. For example, to achieve 99% protein denaturation at $X = 4 \text{ kg kg}^{-1}$ db, lentil and bean should be heated to 96-98 °C while chickpea should be heated 410 411 to 111 °C. The fact that the thermal behavior depends on the species and variety of pulse means 412 that extrapolating this diagram to other legumes will be tricky, in contrast to extrapolating the starch gelatinization diagram.¹³ The specific denaturation characteristics of each pulse need to 413 414 be known to accurately predict the denaturation state of the proteins.

In addition, some studies^{35,50} demonstrated that the length of the thermal treatment and the moisture conditions have an effect on the degree of denaturation of pulse proteins. This result suggests that protein denaturation-aggregation should be considered as a kinetic process. From this point of view, our diagram represents a short heating time (8 minutes heating at between 80 °C and 160 °C). With longer heating times, the final diagram would probably be different.

420

421 To conclude, the DSC experiments on protein-enriched fractions of pulses and the desummation 422 method of the thermograms were suitable for modeling the thermal denaturation of pulse 423 proteins. The temperature, shape and size of the DSC peaks were consistent with the protein 424 composition of each pulse, in particular the 11S/7S globulin ratio. The single transition 425 observed for bean was characteristic of 7S vicilin denaturation, which is the main globulin in 426 this pulse. However, an endotherm with two transitions was observed for chickpea and lentil. 427 Its thermal characteristics are in good agreement with the denaturation of 7S vicilin and then 428 11S legumin at higher temperatures. Irreversible denaturation of both protein types was 429 observed, suggesting that unfolding and aggregation were coupled under the heating conditions 430 used in the study. The three obtained diagrams of protein denaturation enable monitoring of the 431 phase status of lentil, chickpea and bean proteins, in different T and X conditions. However, the 432 diagrams cannot be used to generalize denaturation temperature ranges to all cultivars and has 433 to be studied with a case-by-case approach as composition may vary.

Since the unfolding and aggregation caused by processing, such as heating, modifies the functionality and nutritional quality of pulse proteins, denaturation diagrams are key parameters to improve the cooking process of pulses. The next step in this work will be to combine the phase transition models for starch and for proteins to provide a model of physicochemical changes at the seed scale. However, the present results should be interpreted with caution, as the proteins may behave differently when purified or mixed with other components of the seeds.

440 Abbreviations used

db	Dry basis
DSC	Differential scanning calorimetry
FWHM	Full width at half maximum (°C)
P1	First peak of the DSC thermogram
P2	Second peak of the DSC thermogram
R	Gas constant (J mol ⁻¹ K ⁻¹)
RMSE	Root mean square error
Т	Temperature (°C)
T_i	Temperature at maximum peak <i>i</i> (°C)
Х	Water content (kg kg ^{-1} dry basis)
x_p^{db}	Mass fraction of protein (kg kg $^{-1}$ dry basis)
Greek sym	bols
eta_i	Dimensionless area of peak <i>i</i>
$eta_{ ext{P1,0}}$	Dimensionless area of first peak (P1) when water content is zero
$eta_{ ext{P1},\infty}$	Dimensionless area of first peak (P1) in excess water
$\Delta h_{i,0}$	Change in molar enthalpy of transition i per repeating unit (J mol ⁻¹)
ΔT_i	Parameter related to the width of peak i (°C)
$\Delta T_{i,0}$	Parameter related to the width of peak i when water content is zero (°C)
$\Delta T_{i,\infty}$	Parameter related to the width of peak i in excess water (°C)
v_a	Molar volume of repeating unit amino acid (m ³ mol ⁻¹)
$\mathcal{V}_{_W}$	Molar volume of water (m ³ mol ⁻¹)
τ	Degree of protein denaturation
φ	Heat flow (W)
\overline{arphi}	Normalized dimensionless heat flow
ϕ	Volume fraction of water in protein-water mixture (m ³ m ⁻³)
χ_i	Flory interaction parameter for <i>i</i> transition

441

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- 610

611

612 Figure captions

Fig. 1. DSC thermograms of lentil, chickpea and bean protein-enriched fractions for water contents $X = 0.4 \text{ kg kg}^{-1}$ db (a), $X = 1.0 \text{ kg kg}^{-1}$ db (b) and $X = 2.0 \text{ kg kg}^{-1}$ db (c). The experimental dimensionless heat flows (dots) were modeled using the overall procedure described in Lefèvre *et al.*¹³ The predicted dimensionless heat flows and P1 and P2 endotherms are represented by the solid, dashed, and dotted lines, respectively. RMSE were calculated between experimental and predicted heat flows for the different water contents.

Fig. 2. Parameters of the Gaussian functions T_i (a,b), ΔT_i (c,d) and β_{P1} (e) represented as a function of water volume fraction (ϕ) or water content (X) in the water-protein mixture for the lentil, chickpea and bean protein-enriched fractions. Dots represent the primary parameters of the Gaussian function obtained using the desummation method by fitting the experimental dimensionless heat flows to Eq. (1) for each water content. The solid lines represent the modeling of the Gaussian parameters using the overall identification procedure for every water contents in the same fitting session.

626 Fig. 3. DSC thermograms of protein-enriched fractions of lentils (a), chickpeas (b) and beans 627 (c) using the double scan procedure, for $X = 0.8 \text{ kg kg}^{-1}$ db. Samples were pre-heated from 628 20 °C to 80 °C, 110 °C or 160 °C followed by standard heating from 20 °C to 160 °C. A control 629 sample (only standard heating from 20 °C to 160 °C) is presented for the sake of comparison.

Fig. 4. Modeled denaturation diagram of lentil, chickpea and bean proteins obtained with the desummation procedure. Isovalue lines showing the degree of protein denaturation (unfolding and aggregation coupled) are presented in a temperature T versus water content X diagram.

633

Tables

$X(\text{kg kg}^{-1} \text{db})$	Lentil	Chickpea	Bean
0.2	nd	nd	×
0.3	×	nd	_
0.4	×	×	×
0.6	×	×	×
0.8	×	×	×
1.0	×	×	×
1.2	_	×	×
1.4	_	×	×
1.6	_	×	×
1.8	_	×	×
2.0	×	×	×
2.2	_	_	_
2.4	_	_	_
2.5	_	×	×
2.6	_	_	_
2.8	_	_	_
3.0	×	×	×
3.5	×	_	×
4.0	×	_	_
Total measurements	18	22	26

Table 1. Experimental conditions of the database according to the type of pulse: water content

(X)	of protein	samples	analyzed	with	DSC	from	20	°C to	160	°C.
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×: measured in duplicate.

nd: incomplete signal detected by DSC at $T \le 160$ °C;

-: not tested.

Table 2. Chemical composition of protein-enriched fractions as a fine fraction obtained after

Pulse	Moisture	Starch	Proteins	IDF	SDF	Lipids	Phytates	Alpha- galactosides
Lentil	7.9 ± 0.1	3.0 ± 0.6	55.8 ± 0.1	11.1 ± 1.4	3.1 ± 0.6	2.2 ± 0.1	3.5 ± 0.1	6.3 ± 0.5
Chickpea	5.3 ± 0.1	5.6 ± 0.7	38.2 ± 1.5	18.5 ± 1.1	2.7 ± 0.8	11.5 ± 0.1	2.2 ± 0.1	5.2 ± 0.8
Bean	5.8 ± 0.1	4.2 ± 0.3	49.4 ± 0.1	13.9 ± 3.1	3.7 ± 1.4	2.7 ± 0.2	2.8 ± 0.1	6.3 ± 0.1

dry fractionation and air classification (g/100g dry weight basis)

IDF: insoluble dietary fibers; SDF: soluble dietary fibers

Table 3. Parameters of Eqs. (3), (5) and (6) used to describe the heat flows (Eq. (1)) for lentil, chickpea and bean protein-enriched samples (mean

values ± 95 % confidence interval). The values were used to calculate the degree of protein denaturation $\tau = \sum_{i=P1,P2} \beta_i \times \frac{1}{2} \left(1 + \operatorname{erf}\left(\frac{T - T_i}{\Delta T_i \sqrt{2}}\right) \right)$ with

Protein- enriched sample	$\frac{1}{T_{i}} - \frac{1}{T_{i,0}} = \frac{R}{\Delta h_{i,0}} \frac{v_{g}}{v_{w}} (\phi - \chi_{i} \phi^{2})$					$\Delta T_{i} = \Delta T_{i,\infty} + \left(\Delta T_{i,0} - \Delta T_{i,\infty}\right) \exp\left(-\frac{X}{\gamma_{i}}\right)$				$\beta_{\mathrm{Pl}} = \beta_{\mathrm{Pl},\infty} + \left(\beta_{\mathrm{Pl},0} - \beta_{\mathrm{Pl},\infty}\right) \exp\left(-X\right)$			
	<i>Т</i> _{Р1,0} (°С)	T _{P2,0} (°C)	$\Delta h_{\mathrm{P1,0}}$ (kJ mol ⁻¹)	$\Delta h_{ m P2,0}$ (kJ mol ⁻¹)	$\chi_{ m P1}$	$\chi_{ extsf{P2}}$	$\Delta T_{\rm P1,0}$ (°C)	$\Delta T_{\mathrm{P2,0}}$ (°C)	$\Delta T_{\mathrm{Pl},\infty}$ (°C)	$\Delta T_{\mathrm{P2},\infty}$ (°C)	γ_{P2}	$eta_{ ext{P1},0}$	$eta_{ ext{Pl},\infty}$
Lentil	165.2 ± 0.4	$\begin{array}{c} 201.1 \\ \pm 3.3 \end{array}$	57.6 ± 0.4	45.1 ± 2.9	$\begin{array}{c} 0.00 \\ \pm \ 0.05 \end{array}$	$\begin{array}{c} 0.05 \\ \pm \ 0.05 \end{array}$	4.2 ± 0.1	4.0 ± 0.2	4.2 ± 0.1	$\begin{array}{c} 4.0 \\ \pm \ 0.2 \end{array}$	1	0.65 ± 0.01	$\begin{array}{c} 0.65 \\ \pm \ 0.01 \end{array}$
Chickpea	$\begin{array}{c} 198.1 \\ \pm 8.5 \end{array}$	$\begin{array}{c} 224.7 \\ \pm \ 4.6 \end{array}$	41.2 ± 5.2	41.9 ± 2.5	0.12 ± 0.05	$\begin{array}{c} 0.14 \\ \pm \ 0.03 \end{array}$	$\begin{array}{c} 6.5 \\ \pm \ 0.3 \end{array}$	7.0 ± 1.6	$\begin{array}{c} 2.2 \\ \pm \ 0.2 \end{array}$	3.9 ± 0.1	0.3 ± 0.1	$\begin{array}{c} 0.59 \\ \pm \ 0.02 \end{array}$	$\begin{array}{c} 0.12 \\ \pm \ 0.01 \end{array}$
Bean	166.2 ± 0.3		$\begin{array}{c} 65.8 \\ \pm \ 0.3 \end{array}$		$\begin{array}{c} 0.00 \\ \pm \ 0.05 \end{array}$		3.5 ± 0.1		3.5 ± 0.1			1	1

two peaks (P1 and P2 for lentil and chickpea) or one peak (P1, bean).

 ϕ : volume fraction of water (m³ m⁻³)

X : water content (kg kg⁻¹ db)

 $\gamma_{\rm P1}$ was fixed at 1.

 $\beta_{\rm P2} = 1 - \beta_{\rm P1}$

Figures

Fig. 1.



Fig. 2.



Fig. 3.



Fig. 4.



Graphic for Table of Contents



Label : For Table of Contents Only