

# Seed storage proteins of faba bean (Vicia faba L): current status and prospects for genetic improvement

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# 1 Seed storage proteins of faba bean (*Vicia faba*): current status

# 2 and prospects for genetic improvement

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### 26 ABSTRACT

27 Faba bean (Vicia faba, L.) is one of the foremost candidate crops for simultaneously 28 increasing both sustainability and global supply of plant protein. Its seeds contain about 27% 29 proteins of which more than 80% -consist of globulin storage proteins (vicilin and legumin). 30 For optimum utilization for human and animal nutrition, both protein content and quality 31 have to be improved. Though initial investigations on the hereditability of these traits 32 indicated possibility for genetic improvement, little has been achieved so far partly due to 33 lack of genetic information coupled with the complex relationship between protein content 34 and grain yield. This review reports on the current knowledge on faba bean seed storage 35 proteins; their structure, composition and genetic control and highlights key areas for further 36 improvement of the content and composition of faba bean seed storage proteins on the basis 37 of recent advances in faba bean genome knowledge and genetic tools.

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Key words: *Vicia faba*; sustainability; storage proteins; legumin and vicilin; genetic
 improvement

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### 51 **INTRODUCTION**

### 52 Faba bean production and utilization

Nearly 60% of the global protein supply for human nutrition is sourced from plants <sup>1-2</sup> and 53 about one third of this originates from grain legumes of the Fabaceae family<sup>3</sup>. Besides their 54 55 nutritional significance, legume crops ability to fix atmospheric nitrogen via rhizobial symbiosis makes them invaluable components of sustainable crop production systems <sup>4</sup>. Faba 56 bean (Vicia faba, hereafter Vf), also known as fava bean, broad bean, horse bean or field bean 57 <sup>5</sup> is one of the world's oldest legume crops, its cultivation dating back to the 10<sup>th</sup> millennium 58 BC  $^{6-7}$ . From its origin in the Near East. Vf spread to the rest of the globe <sup>7</sup> and is currently 59 cultivated in nearly 70 countries over the world (Figure 1A), occupying about 2.2 million ha 60 that produce nearly 4 million tons annually <sup>8</sup>. China is the leading Vf producer with 36% of 61 the global output, followed by Ethiopia (20%), Australia (10%) and United Kingdom (6%) 62 63 (Figure 1B). The wide geographical distribution of Vf implies not only a great adaptation to diverse environmental conditions, but also suitability for diverse end uses and trade across 64 65 continents.

Seeds of Vf contain on average about 27% protein <sup>9-11</sup> which provides affordable nutrition for 66 67 millions of people around the world, hence its denomination as "the poor man's meat". While Vf has been traditionally utilized as dry grain for human consumption in developing 68 69 countries, there is growing interest from food industries in developed countries to exploit its protein for the production of protein-rich vegan/vegetarian snacks, <sup>12</sup>, the fortification of 70 71 cereal-based food products such as bread and pasta without significantly affecting their structural and sensory quality <sup>13-14</sup>, or even the production of wholly *Vf*-based bread and 72 pasta products <sup>15</sup>. Vf also represents as significant resource for agro-ecosystem sustainability 73 74 and provision of feed for the growing global livestock inventory. Overall, the global 75 production area for Vf has been increasing in the last two decades (Figure S1A) and a recent Page **4** of **35** 

76 meta-analysis of yield data from 39 legume species indicated that, in the right environment, Vf can be the highest yielding grain legume <sup>16</sup>. Vf also has a high capacity for biological 77 78 nitrogen fixation, to the extent that the amount of N fixed by Vf alone was estimated to be comparable to that of soybean and pea combined  $^{17}$ . For further details on the role of Vf on 79 sustainable cropping systems, readers are referred to Jensen, et al. <sup>18</sup>, Köpke and Nemecek 80 <sup>19</sup>.On the other hand, Vf is yet to be fully exploited as a feedstock for animal production due 81 to presence of some anti-nutrients which limit its optimal inclusion ratio<sup>20-22</sup>. Removal of 82 these anti-nutrients through the development of new low anti-nutrient cultivars or using 83 simple processing techniques like fermentation <sup>13-14</sup> would make this crop a valuable protein 84 85 resource for the animal production industry.

### 86 Faba bean as a sustainable global protein resource

One of the greatest challenges in the 21<sup>st</sup> century is feeding the growing world population 87 which it has been estimated may necessitate a 70% increase in food production by 2050<sup>4</sup>. 88 More than 30% of this increase has to be made via the production of protein-rich foods <sup>1</sup> to 89 90 meet the expected rise in demands due to population growth, increased urbanization and improved incomes in many parts of the world <sup>1, 23-25</sup>. Protein is a critical nutrient required in 91 large quantity by humans (~ 50 g protein per adult per day) to maintain normal body function 92 <sup>26</sup> However,t about one-third of the world population, mainly in Asia, Africa and Latin 93 America, suffers from inadequate intake of proteins, vitamins and minerals<sup>27</sup>. On the other 94 hand, in higher income countries, where daily animal-based protein intake is already high<sup>1</sup>, 95 <sup>25</sup>, continued provision of nutritious feeds for the intensive animal production industry will 96 pose a major challenge in the future. In particular, the livestock production sector in soybean 97 98 non-producing countries will be burdened by the high price of imported soybean and soybean 99 meal. For instance, EU countries have huge deficit in protein-rich feeds with nearly 70% being imported  $^{28}$ . Vf is well-adapted to European climates, as testified by the high yields 100

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101 recorded in this continent for this legume (Figure S1B), and it therefore has the potential to 102 contribute to bridging the gap in animal feed self-sufficiency as part of the EU's policies to increase protein production from locally grown crops  $^{28}$ . Vf is also a candidate crop to meet 103 the protein demands of an emerging consumer category, particularly in developed economies, 104 who are opting for animal meat free life style. For example, Statista<sup>29</sup> reported that 13% of 105 106 European citizens would consider avoiding red meat while nearly 50% of the respondents in another study were willing to replace meat with other sources of proteins  $^{30}$ . 107

108 Considering the projected impact of climate change on global crop production, meeting the 109 nutritional requirements of the current and future generations would necessitate increased 110 exploitation of the global genetic and natural resources for protein production systems based 111 increasingly on biological nitrogen fixation. In this context, the fact that Vf is a high-yielding 112 protein-rich crop with superior N fixation capability makes it a candidate crop for supporting 113 increased protein production while maintaining sustainability of crop production systems.

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### Nutritional constraints to Vf utilization

115 The main determinants of Vf utilization for human food and animal feed include: (i) protein 116 concentration, (ii) protein quality, defined mainly by the content of sulfur-containing amino acids (S-AA) cysteine and methionine, and (iii) concentration of antinutrients in the seeds <sup>5</sup>. 117 118 Protein concentration of Vf, although it can vary greatly between different genotypes (19-39) %) <sup>31-33</sup>, is one of the highest among legumes. However, commercial varieties on the UK 119 120 market contain about 27% protein on average, which is still far less than the protein density 121 of soya meal, and so, further improvements in protein content is required in order for faba 122 bean to displace imported soya in animal feed. The proportion of S-AA in the protein is 123 another crucial quality criterion, particularly in animal feeding. However, like most plant proteins, Vf is poor in certain essential amino acids, namely methionine, cysteine and 124 tryptophan <sup>5</sup>. Though relatively narrow, the genetic variation for the S-AA reported in Vf125

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126 indicates possibility of improving its nutritional quality. So far, the major breeding objectives 127 for Vf, have been the reduction or removal of vicine and convicine (V-C) and tannins: V-C causes favism in humans and have deleterious effects on animals <sup>34-35</sup> while tannins lower 128 protein digestibility <sup>10</sup>. Although these compounds can be removed by processing techniques 129 <sup>36-37</sup>, the most effective approach is probably removing them by breeding. This is now 130 feasible with the availability of molecular markers closely linked to the V-C locus <sup>38</sup> and zero 131 132 tannin gene (zt-1)<sup>39</sup>. Furthermore, the reduction of less significant antinutrients such as trypsin inhibitors, lectins and phytates would improve the nutritional value of Vf based feed 133 134 products.

135 Understanding the genetic basis of the above limiting factors is a prerequisite for the 136 development of new cultivars with desirable agronomic and nutritional attributes. 137 Unfortunately, while scientific interest in Vf was high during 1970's and 1980's, when it 138 became the model species for studying plant cytogenetics and stomatal regulation, Vf can now be considered an orphan crop  $^{40}$ . For instance, less than 5% of the publications on 139 legumes in the years 2004–2013 referred to  $Vf^{9}$ . This is further reflected by the scarcity of 140 141 information on the genetics of many important traits including protein content and quality, for 142 which not a single QTL (Quantitative Trait Loci) has been reported, compared to 160 QTLs 143 from 35 independent studies on soybean protein content <sup>41</sup>. In this context, in order for future 144 work to proceed on a sound basis, we felt there was a need to marry the earlier biochemical 145 literature, where the main species of storage protein were separated and classified, with the 146 later genomic literature, which is replete with unannotated storage protein sequences and 147 implicit map locations. The remainder of this review is devoted to a synthesis of the literature 148 on Vf seed storage proteins, covering sequence, structure, composition and genetic basis for 149 their synthesis and accumulation as well as taking a forward look at how this synthesis might 150 be exploited in future research aiming to increase protein content and/or quality.

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### 151 SEED STORAGE PROTEINS OF FABA BEAN

152 The major storage proteins of legumes are mainly enzymatically inactive proteins deposited 153 in seed cotyledons which provide nutrients needed for seed germination and seedling growth and development <sup>42-43</sup>. Certain seed proteins in legumes including albumins and trypsin 154 155 inhibitors, however, have been identified as antinutritional or allergenic agents and therefore are targeted for removal in breeding programs <sup>44</sup>. Seed storage proteins are classified 156 157 according to the system developed by TB Osborne which is based on their solubility in different solvents <sup>43</sup>. Albumins and globulins are the major storage proteins of legumes and 158 159 are soluble in water and saline solutions, respectively. Globulins alone constitute more than 80% of total seed protein in  $Vf^{45}$  and they are further classified based on their sedimentation 160 coefficients into vicilin-type (7S) and legumin-type (11S)<sup>43</sup>. Both globulin proteins are found 161 162 in nearly all legumes, but their denotations vary across species. For instance, globulins of Vf 163 and pea are often referred as vicilin/convicilin and legumin while they are denoted as 164 conglycinin and glycinin in soybean,  $\beta$  and  $\alpha$  congluting in lupin, while phaseolin (a vicilin-165 like protein) is the only major globulin in common beans. Furthermore, decades of research 166 on legume storage proteins have produced a sufficient database of annotated SDS-PAGE 167 images of various species which facilitates faster identification of major globulin bands 168 without the need for conducting tedious immunoblotting or HPLC procedures. When 169 extracted under reducing conditions, the salt soluble fraction of legume seed proteins can be separated on SDS-PAGE into distinct bands which, based on their molecular weights, are 170 identified as: convicilin (Mr >~60 kDa), vicilin (Mr=~46-55 kDa) and two major legumin 171 subunits (Mr= $\sim$ 38-40 and 23 kDa) (Table 1) 172

173 Legumin and vicilin share notable sequence and structural homology and are believed to 174 originate from a common ancestral gene <sup>46</sup>. Mature legumin is hexameric with a mass of 175 about 330 kDa <sup>45</sup> and is composed of two trimeric subunits (legumin A and B) while vicilin is

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176 a trimeric protein formed by the assembly of three monomers (Figure 2). In contrast to legumin, vicilin lacks cysteine and is usually glycosylated in its C-terminus <sup>46</sup>. These 177 178 structural variations may result in differences in the physiochemical properties of seed 179 storage proteins which in turn determine their nutritional value and utilization. For instance, legumin and vicilin differ in their thermal properties <sup>47-48</sup>, affinity to bind flavor compounds 180 under varying pH conditions <sup>49</sup> and emulsifying ability <sup>48</sup>. Therefore, from a breeding point of 181 182 view, legumin/vicilin ratio could be manipulated to meet certain end-user requirements for 183 protein functionality.

### 184 Structure and composition of *Vf* globulins

Legumin constitutes more than 50% of Vf globulins <sup>45</sup>. It is a hexameric protein with two 185 186 major subunits - the  $\alpha$  and  $\beta$  chains - which are connected by disulphide bonds. Under 187 reducing conditions, these subunits form two bands of molecular weights of about 40 and 24 188 kDa, respectively (Figure 3). These subunits are also referred to as acidic and basic subunits 189 or simply legumin A and B. Polypeptides of both legumins are highly homologous but 190 notably distinguishable by the presence of more methionine residues in the peptide sequences of legumin A subunits <sup>50</sup>. Vf legumin A subunits appear to be more variable and show 191 polymorphic bands between genotypes<sup>51</sup> as is also the case with Medicago legumin A <sup>52</sup>. On 192 the other hand, vicilin-type proteins of Vf are trimeric <sup>45</sup> consisting predominantly of subunits 193 of ~50 kDa while bands of ~66 kDa are referred as convicilin <sup>42, 51</sup>. The classification of 7S 194 195 proteins into vicilin and convicilin was first coined in pea and has been accepted in many 196 legumes including Vf (Table 1). Nonetheless, further investigation into their possible 197 structural and functional differences have concluded that convicilin may be regarded as subunit of vicilin <sup>53</sup>. Such a denotation exists in soybean whereby subunits of 7S protein are 198 categorized into  $\alpha'$  (~76 kDa),  $\alpha$  (~72 kDa), and  $\beta$  (~53) kDa <sup>54-55</sup>. 199

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200 Regarding amino acid composition, nearly 50% of Vf seed protein is accounted for by just a 201 few non-essential amino acids such as glutamic acid, aspartic acid, arginine and leucine while 202 it is low in essential amino acids particularly S-AA (Figure 4). The concentration of S-AA is 203 a critical determinant of the nutritional value of plant proteins destined for human 204 consumption and animal feeding. In humans, dependence on poor quality proteins can result 205 in reduced immunity and underdeveloped mental and physical capacity among young 206 children <sup>56</sup>. Also, animal feeds deficient in critical amino acids can cost farmers in form of animal feed supplements of industrially synthesized S-AA<sup>55</sup>. 207

208 The concentration of S-AA is strongly related to the relative proportions of S-AA rich 209 proteins in the seeds. In Vf and other legumes, it is well accepted that legumins contain relatively higher S-AA compared to vicilin <sup>42, 44, 57-58</sup>. This is further confirmed by 210 211 comparative analysis of coding sequences of vicilin and legumin subunits across legume 212 species which clearly show that legumin subunits contain more residues of cysteine and 213 methionine (Figure 4). This observation leads to the hypothesis that increasing the proportion 214 of legumin subunits relative to vicilin would improve nutritional content of plant proteins. However, considering that vicilin is accumulated in legume seeds earlier than legumin <sup>59-61</sup>, 215 216 their ratios could be easily offset by the prevailing environmental conditions, e.g. soil 217 nutritional status and onset of biotic and abiotic stresses during the plant growth, and in 218 particular, during grain filling. In contrast to globulins, minor legume seed proteins such as 219 elongation factor Tu, citrate synthase, albumin 2 (PA2), defensins 1 and 2 and Bowman–Birk inhibitors (BBI) contain higher S-AA<sup>42, 62</sup>. According to Krishnan, et al.<sup>63</sup>, under higher N 220 221 availability through fertilizer application or symbiotic fixation, S-AA containing proteins like 222 Bowman-Birk protease inhibitor (BBI) were decreased in favour of  $\beta$ -subunits of  $\beta$ -223 conglycinins of soybeans. Similarly, ectopic overexpression of VfAAP1 gene on P. sativum and V. narbonensis resulted in 30% increase in the globulin fraction but no significant effect 224

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on albumin, a S-AA rich protein  $^{64}$ . Hence, it would appear that the negative correlation between high protein and S-AA content in *Vf* <sup>11, 32, 65</sup> may be the result of preferential accumulation of low nutritional quality protein fractions in higher protein lines.

### 228 Genetic control of globulins

229 Globulins are by far the most abundant seed proteins in legumes and, subsequently, their 230 genetic control has been well investigated. In Vf, legumin subunit is encoded by relatively 231 few genes which are classified as legumin A and B genes. A single legumin A gene has been located on the telomeric region of chromosome V of Vf<sup>66</sup>. It is not clear, however, whether 232 the legumin A2 gene (*LegA2*) reported in pea  $^{67}$  also exists in *Vf*, as no up to date information 233 234 is available. Conversely, there are at least five transcribed genes (LeB2, LeB3, LeB4, LeB6, LeB7) for legumin B subunits<sup>66, 68</sup>, of which LeB3 and LeB4 have been mapped to 235 chromosome II and III, respectively <sup>66</sup>. The vicilin coding gene <sup>69</sup> was also located on 236 chromosome II, near the centromere  $^{70-71}$ . While the documented number of genes for Vf 237 238 globulins is relatively small, numerous legumin and vicilin minor subunits with various molecular masses and isoelectric points can be observed in 2D gel electrophoresis analysis<sup>51</sup>, 239 240 suggesting that Vf globulins undergo extensive post-translational processing. A similar occurrence has been found in other legumes including Medicago truncatula <sup>72</sup> and Pisum 241 sativum<sup>73</sup>. 242

There is considerable homology between Vf globulin subunits and those of other legumes (Table S1), and where genome sequences are available, it is now possible to classify and associate seed storage subunits to specific genome locations (Table S2). Considering the lack of genome sequence for Vf, this information is critical for synteny-based mapping of globulin genes and QTLs. For instance, in *M. truncatula*, several genomic regions coding for globulins have been mapped on chromosome I and VII <sup>72</sup> which are notably syntenic with Vf

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chromosome III and V 39-40 where legumin A and B genes were previously located, 249 respectively 66. 250

### 251 Expression of globulin genes

252 Seed protein content can be thought of as the final output of a number of biochemical and physiological processes occurring throughout the crop life cycle, each of which are under the 253 254 control of a regulatory network. Abundance of globulin proteins is regulated by a network of genes involving transcriptional regulation transport and post-translation modifications of 255 storage proteins <sup>72</sup>. Among these are numerous seed specific genes which play profound 256 regulatory roles in the synthesis and accumulation of seed storage proteins <sup>72, 74</sup>. Notably, 257 258 seed specific transcription factors (TFs) such as AB15, LEC1, LEC2, AB13, MYB#2, bHLH#1 and FUS3 are key storage protein regulators <sup>72, 75</sup>. ABA insensitive 5 (ABI5) is expressed 259 during seed filling stages in plants<sup>75</sup> and has been found at the center of the regulatory gene 260 network for storage protein synthesis in *M. truncatula*  $^{72}$ . Specifically, it is a major regulator 261 for vicilin polypeptide abundance with P. sativum abi5 mutants showing nearly 30% decrease 262 in the abundance of vicilin-type globulin <sup>72</sup>. Similarly, *ABI3b* and LEAFY COTYLEDON-1 263 264 (LEC-1) homologs in soybean has been located at the hub of 118 genes related to seed protein content <sup>74</sup>. Given the microsynteny between Vf and the model crop M. truncatula <sup>39</sup>, these 265 266 findings will provide a reference for further discoveries in the genetics of Vf globulins.

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### Synthesis and accumulation of seed storage proteins

268 Globulins are synthesized in the endoplasmic reticulum (ER) sorted in the Golgi body and transported to the protein storage vacuole (PSV) by vesicles <sup>72, 76</sup>. During Vf seed 269 270 development, a diphasic pattern of protein accumulation exists in which proteins synthesized 271 during early developmental stages are only transitorily accumulated and subsequently degraded to sustain the growing embryo while proteins accumulated after heart stage (~12 272

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273 DAP) are mainly destined for storage into cotyledons' protein bodies <sup>77</sup>. During the latter 274 stage, globulin proteins show distinct expression patterns in which vicilin synthesis and 275 accumulation precedes that of legumin and  $\alpha$  chain polypeptides of legumin appear earlier 276 than  $\beta$  chains <sup>59</sup>. Similar pattern of vicilin and legumin gene expression has also been 277 reported in Medicago <sup>78</sup> and soybean <sup>76</sup>.

278 The amount of protein accumulated during seed development can be attributed to various 279 genetic and environmental factors acting on various plant processes ranging from nutrient 280 uptake and transport, photosynthate production and remobilization to protein accumulation 281 rate in the storage organs. However, there are strong indications that mechanisms underlying 282 nitrogen (N) uptake, transport and assimilation could explain the variation in protein content 283 more than any other factor. For instance, in pea, overexpression of the amino acid transporter 284 gene amino acid permease (AAP), has been confirmed to play a critical role in increasing 285 synthesis of seed storage proteins owing to increased leaf and pod phloem loading with free 286 amino acids<sup>79</sup>. A similar mechanism could be attributed to the observed 2-3 times higher free 287 amino acids in the cotyledons of high-protein (HP) Vf genotypes as compared to low-protein genotypes<sup>80</sup>. In rice, a major seed protein content QTL harboring the OsAAP gene was 288 associated with higher uptake of amino acids and their distribution across plant tissues<sup>81</sup>. In 289 290 addition, QTL for N-fixation have been linked to QTL for total N accumulation in common bean <sup>82</sup> and pea <sup>83</sup>. Also, improved capacity for N uptake can be a candidate trait to relax the 291 292 yield-protein negative correlation. In fact, increased genetic capacity for N supply was associated with increased seed size in  $Vf^{64}$  or seed number in pea <sup>79</sup>. These results should be 293 294 taken into consideration when screening for high protein content in Vf.

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### **GENETIC IMPROVEMENT OF PROTEIN CONTENT AND QUALITY**

### 296 Summary of the past work

297 Several studies have focused on the genetic variation for protein content (Table 2) and to 298 what extent protein content was correlated with yield of Vf. One study indicated that protein 299 content was variable between and within varieties (n=33) with broad sense heritability of 0.70 and no significant correlation with seed weight <sup>31</sup>. However, when larger set of 300 germplasm (n=600) was screened, a clear negative relationship between seed weight and 301 302 protein was detected although some large-seeded genotypes with above average protein content were also found <sup>65</sup>. Similarly, after four cycles of selection for protein content. Siödin 303 304 <sup>32</sup> concluded that protein content in Vf could be improved by selection but tended to 305 negatively correlate with number of seeds per plant regardless of thousand seed weight. 306 These early efforts also established the variability for S-AA content (Table 3) and nearly all investigations found a negative correlation between protein and S-AA content <sup>32, 65, 84</sup>. Under 307 308 circumstances where desirable traits of interest are negatively correlated, deeper 309 understanding of the genetic basis of the trade-offs between the traits and availability of 310 appropriate tools to dissect and recombine them is crucial.

### 311 Areas for future focus

### 312 Uncoupling the negative yield-protein correlation

Correlation between traits can arise due to gene linkage or pleiotropy <sup>85</sup>, with the latter being most common in plants, and its resolution requires deeper understanding of both traits. Therefore, several possible mechanisms have been investigated in various crops in order to unlock protein-yield association. It is hypothesized that the negative correlation between the two traits result when the high demand for N during seed filling stage coincides with decline in soil nutrients in the rhizosphere and nitrogen fixation, resulting in re-mobilization of nitrogen from leaves, which in turn shortens grain filling and reduces seed weights <sup>86</sup>. This is

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in line with findings by Egle, et al.<sup>87</sup> who showed that majority of N accumulated during 320 321 seed filling in barley was remobilized from leaves and stems, but that ongoing N uptake 322 could also contribute. Furthermore, wheat genotypes with higher capability for post-anthesis N uptake deviate from grain-protein negative relationship <sup>88-89</sup> and selection for this trait has 323 324 been therefore proposed as a possible criterion for simultaneous improvement of protein 325 content and grain yield. The genetic basis of post-flowering N uptake is not yet fully 326 understood either in cereals or in legumes but could be related to root structure and/or N 327 transport capacity. For instance, pea genotypes with higher mineral nitrogen absorption and symbiotic nitrogen fixation have shown enhanced seed N content and yield <sup>83</sup>. Moreover, 328 329 faster rate and relatively longer duration of N accumulation during seed development has 330 been reported as a possible mechanism for combining high protein and large seed size in soybean <sup>90</sup>. The importance of N uptake capacity for protein content and yield was further 331 demonstrated by Peng, et al.<sup>81</sup> who found major protein content QTL *qPC1* harboring a 332 333 putative amino acid transporter gene (OsAAP6), which they proposed as candidate QTL for 334 simultaneous selection for yield and protein content in rice. These areas of enquiry are 335 amenable for further investigation and can potentially point to QTLs that can be used to 336 improve protein content in Vf without significant yield reduction.

### 337 *Improving S-AA content by modifying legumin: vicilin ratio*

Considering difficulties in genetic improvement of limiting amino acids through conventional breeding approaches, several genetic engineering approaches have been attempted in various crops over recent decades. Detailed information on these strategies and results obtained can be found in Galili and Amir <sup>56</sup>. These included (i) overexpression of genes encoding proteins rich in the limiting amino acid, (ii) *in vitro* modification of genes encoding proteins of interest by adding more residues of the desired amino acid, (iii) introduction of genes coding for protein rich in the limiting amino acid from one species to another target food crop, or by

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345 (iv) modification of biosynthetic and catabolic pathways to directly increase accumulation of 346 target amino acid or indirectly by increasing accumulation of proteins containing the limiting 347 amino acid. Yet, most of these attempts have not succeeded in producing new crop cultivars 348 combining increased protein quality with desired agronomic traits. In rare cases where 349 reasonable success was achieved, commercialization of the improved cultivars was hindered by legal restrictions on GMO release <sup>56</sup> and consumer resistance. Besides these challenges of 350 351 consumer acceptability, the potential of transgenic approaches in Vf is limited by the inherently poor regenerating ability of Vf transgenics  $^{91}$ . 352

353 Alternative strategies include direct selection on QTL for S-AA content or indirectly by 354 selecting for greater relative expression of protein subunits rich in S-AA rich subunits. To our knowledge, soybean is the only legume crop in which QTLs for individual S-AA has been 355 mapped <sup>92-93</sup>. Though total seed content of the S-AA *per se* would be a good indicator, it may 356 357 not be sufficient when considering as selection criteria, due to uncertainty about what 358 percentage of the total S-AA detected is indeed imbedded in the main storage proteins. In Vf 359 and other legumes, since it is observed that the legumin protein subunit have relatively higher S-AA content compared to vicilin<sup>42, 44, 57-58</sup>, increasing legumin subunit in favor of vicilin 360 361 would be expected to enhance the protein quality. In fact, the concept of manipulating 362 legumin: vicilin (L/V) ratio to improve nutritional quality is not new in Vf. It was previously reported that variation in L/V ratio among varieties was consistent across years <sup>94</sup> and 363 environments <sup>95</sup> and concluded that L/V ratio has genetic basis and could be used as a 364 selection criteria to improve nutritional quality in  $Vf^{94-95}$ . To our knowledge, since L/V ratio 365 366 based approach was suggested as a practical breeding strategy for improving nutritional quality in soybean <sup>57</sup>, only study has tried to map QTLs for L/V ratio and showed colocation 367 between some QTLs for structural legumin and vicilin loci and L/V ratio <sup>96</sup>. The recent 368 369 advances in Vf genetics tools such as development of 50 K SNP array and high-density

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linkage map may offer an unprecedented opportunity to discover novel QTLs that couldrepresent targets for improving nutritional quality.

### 372 *Exploiting mutagenesis approaches*

373 Large-scale mutagenesis using physical or chemical mutagenic agents is a well-established 374 method of inducing novel variation to meet human requirements, but which is unlikely to be 375 present in nature. This approach is all the more justified in the case of Vf where the primary 376 gene pool lacks any known wild relatives. Indeed, several mutagenesis efforts have produced new sets of morphological phenotypes in Vf<sup>40, 97-98</sup>. However, no data is available on 377 potential beneficial mutations in the seed composition of Vf. Although Sjödin <sup>97</sup> has reported 378 379 to have identified some high protein content genotypes from a lot of seeds which had been 380 mutagenized he could not ascertain whether the selected plants were genuine mutants or 381 randomly isolated extremes in the original seed lot. There are several potential ways of 382 exploiting induced mutations for improving protein content and/or quality. First, desirable mutations involving photosynthetic and N provision mechanisms can improve protein 383 content. From ethyl methane sulfonate (EMS) mutagenized seeds, Duc <sup>98</sup> discovered a 384 385 supernodulating line with 3-4 times higher number of nodules compared to the parental line. 386 Considering the close relationship between N fixation and protein content, such a trait could 387 be exploited in breeding programs. Secondly, knockdown/knockout or regulatory mutations 388 leading to absence of major protein subunits such as vicilins can result in improved 389 nutritional quality by increasing the ratio of S-AA rich subunits like legumin and albumins. 390 Such mutations could be *cis*-linked to the structural loci themselves or *trans*-acting factors 391 that would need to be mapped *de novo*. For instance, mutants of *PsABI5*, a major *trans*-acting regulator of vicilin abundance in pea, have shown an increased legumin abundance <sup>72</sup>. 392 393 Thirdly, presence or absence of certain subunits can enable dissection of genetic control of individual protein subunits via a QTL mapping approach <sup>55</sup>. Lastly, it is possible via a reverse 394

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395 genetic screen to select non-synonymous mutations that convert non-S-AA residues to S-AA 396 residues in S-AA poor storage proteins such as vicilins, although, only a proportion of codons 397 are available for single base changes that would result in this outcome. Moreover, the 398 physico-chemical properties of cysteine (disulfide bridge-forming) and methionine (hydrophobic) may cause steric constraints <sup>99</sup>. However even a single well-placed additional 399 400 methionine in each vicilin could give rise to a significant step up in S-AA levels and this 401 approach is therefore worth trying. On a more practical level, full exploitation of mutagenesis 402 for the above purposes requires high-throughput and cheap phenotyping methods to screen 403 tens of thousands of plants for nutritional and agronomic traits.

404 In summary, Vf is one of the most important legumes crops with great potential to fulfil 405 multiple nutritional and ecological services for the current and future generations. However, 406 Vf can only play this role if it meets certain producer and end-user expectations which 407 requires plant breeders and research community to address both agronomic and nutritional 408 constraints simultaneously. In drawing together a synthesis of the literature on Vf seed protein 409 content, contribution of different storage protein classes to overall abundance and to varying 410 relative amounts of essential amino acids, globulin structure and globulin-encoding genes, we 411 aim to provide an updated and comprehensive primer for researchers interested in the 412 nutritional optimization of faba beans. We discuss a range of approaches by which protein 413 content could be increased (without compromising yield) and protein quality ameliorated, 414 some of which have successful precedent in related legume species. These include: high 415 resolution mapping of protein, L:V ration and S-AA QTL using powerful modern 416 quantitative genetics methods and genomics technologies; manipulation of known or still-to-417 be-discovered structural and regulatory genes by transformation and screening of mutant 418 libraries to reveal novel structural and regulatory variants not found in nature. In parallel, as 419 genome sequencing become cheaper and more genomic resources for Vf are accumulated, all

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- 420 the above should become ever more efficient, enhancing the prospects of increasing protein
- 421 content and quality in this strategic crop.

### 422 ABBREVIATIONS USED

- 423 Vf, Vicia faba; S-AA, sulfur containing amino acid; V-C, vicine and convicine; QTL,
- 424 quantitative trait loci; SDS-PAGE, sodium dodecyl sulfate
- 425 polyacrylamide gel electrophoresis; HPLC, High Performance Liquid Chromatography; kDa,
- 426 Kilo Dalton; N, nitrogen; GMO, Genetically Modified Organisms; EMS, Ethyl Methane

427 Sulfonate (EMS)

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### 432 SUPPORTING INFORMATION

- 433 Supplementary data including Figure S1 and Tables S1-S3 are provided in MS Word
- 434 document.

### 435 CONFLICT OF INTEREST

436 The authors declare no competing financial interest.

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### 765 FIGURE CAPTIONS

Figure 1. Global distribution of *Vf* cultivation (A) and the major producing countries (B).
Data was sourced from FAOstats and distribution map was generated using Tableau Public
2018.1.

Figure 2. Predicted ribbon structures of Vf globulins. Vicilin (A) is trimeric consisting of 3 protomers (a=light blue, b= magenta and c= green) while legumin is hexameric consisting of legumin A (B) and legumin B (C). Spherical balls in legumin subunits represent disulfide bonds. The models were generated using SWISS-MODEL and processed with PyMOL

- software. Model description details are in Table S3.
- Figure 3. 1D SDS-PAGE showing the major subunits of Vf globulins and the variation in protein band abundance among 11 inbred lines.
- Figure 4. Amino acid composition (g/16 g N) of Vf seed protein (Makkar et al., 1997; Grela et
- al., 2017). It clearly shows the abundance of several amino acids and deficiency of the S-AA
   in *Vf* proteins.
- Figure 5. Relative abundances of limiting amino acids within legumin and vicilin coding sequences of 7 legume species (Table S1). Annotated protein accessions were obtained from
- 781 Uniprot and the amino acid residues were counted using "seqinr" package in R.

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## **Table 1.** Major globulin polypeptides of Vf and related species as annotated on SDS-PAGE

Species	11S legumin-like (~kDa)		7S Vicilin-like (~kDa)		Ref.
	α chain	β chain	vicilin	Convicilin	
	38	22-24	31-65		De Pace, et al. 59
	38-47			64	Liu, et al. 42
Vicia faba	40	20			Gatehouse, et al. 95
vicia juba	35-39	23-25	42-48	66	Tucci, et al. 51
	36-51	19-23			Utsumi, et al. 100
	40	23-24	54	~73	This study
	36-46	23-24	46-47	60-92	Le Signor, et al. 52
Medicago truncatulla	42-46	23	46-47		Gallardo, et al. <sup>61</sup>
	38-41		47	70	Le Signor, et al. 72
	37	20	52-72		Fontes, et al. <sup>101</sup>
Chaine man*	37	20	52-72		Boehm, et al. 55
Giyeine max.	37	20	52-72		Poysa, et al. <sup>102</sup>
	37-44	17-22	53-76		Krishnan, et al. 54

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Pisum sativum *7S subunits of G.max	40-45 40 40 37 consist of α`, α	18-25 24.8  25 α and β polypeptic	53 47.2  43-53 des.	60-88 67.2 >70 70	Bourgeois, et al. <sup>73</sup> Mertens, et al. <sup>103</sup> Rubio, et al. <sup>62</sup> Ladjal E, et al. <sup>104</sup>
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**Table 2.** Genetic variability in seed protein content in Vf

	No. genotypes	Protein content (%)	Reference
	33	22-38	Griffiths and Lawes <sup>31</sup>
	600	19-34	Lafiandra, et al. 65
	125	22-36	Sjödin <sup>32</sup>
	125	29-38	Frauen, et al. <sup>33</sup>
	30	23-39	Griffiths <sup>84</sup>
	12	26-30	Makkar, et al. <sup>10</sup>
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**Table 3.** Genetic variability in sulfur-containing amino acids in Vf(g/16 g N)

No. genotypes	Methionine	Cysteine	Reference
111	0.6-1.0	1.0-1.5	Lafiandra, et al. 65*
125	0.8-1.4	1.3-1.4	Sjödin <sup>32*</sup>
125	0.1-0.2	0.2-0.6	Frauen, et al. 33
12	0.8-1.1	1.1-1.4	Makkar, et al. <sup>10</sup>
50	0.6 - 0.9	1.0 - 1.4	Schumacher, et al. 105
46	0.6-0.9	0.9-1.2	Schumacher, et al. 11

\* S-AA reported as % protein

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861 Figure 1



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- 884 Figure 2



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