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1       **PHYSIOLOGICAL BASES OF CULTIVAR DIFFERENCES IN AVERAGE GRAIN**  
2       **WEIGHT IN WHEAT: SCALING DOWN FROM PLOT TO INDIVIDUAL GRAIN IN**  
3       **ELITE MATERIAL**

4

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13

14 **Abstract**

15

16 In recent decades, increases in wheat yield have been achieved mainly through increases in grain  
17 number per m<sup>2</sup> (GNM2) rather than through increases in average grain weight (AGW). Using AGW  
18 as a lever to increase yield would require avoidance of the negative relationship between GNM2  
19 and AGW. It is usually supposed that this trade-off arises from an increase in the proportion of  
20 small grains as GNM2 rises. The proportional increase in small grains being the result either of (1)  
21 an increase in the proportion of secondary tillers in the spike population or (2) of an increase in the  
22 proportion of grains located in distal positions within each spike. Either or both of these two  
23 populational effects would tend to mask any true genotypic differences in AGW. The existence of  
24 these constitutive differences has already been proposed, but without considering the full extent of  
25 the populational confounding effects. Identification of a component of the constitutive genetic  
26 determinism of AGW - one that is truly independent of GNM2 - could contribute to cultivar  
27 developments that would lead to further increases in grain yield under future target environments.

28 To address this question, we analysed populational effects on AGW in four, modern, well-adapted  
29 bread-wheat cultivars. The four chosen cultivars show similar grain yields but contrasting AGWs.  
30 The analysis of populational effects was carried out at three hierarchical levels (the plot, the spike  
31 and the single grain) and under two contrasting environmental conditions (well-watered *vs* water-  
32 deficit conditions).

33 Regardless of the environment, no (or only slight) differences in individual spike size were  
34 observed between cultivars. Furthermore the weak relationship between spike size and  
35 AGW demonstrates that AGW differences between cultivars cannot be attributed to spike-level  
36 populational effects. Meanwhile, the analysis of individual grain mass distributions, showed that the  
37 differences in AGW between cultivars, originated from shifts in the whole grain-mass distribution,  
38 rather than from shape changes in the grain-mass distribution. This clearly indicates that AGW  
39 differences between cultivars cannot be attributed to populational effects at the individual grain  
40 level.

41 The analysis carried out at both spike and individual grain levels indicates that the AGW  
42 differences between cultivars are largely constitutive, so that increases in grain yield through AGW  
43 may be considered independently of the GNM2 : AGW trade-off.

44 Taken together, these findings offer a new perspective for the genetic improvement of wheat, and  
45 one that should lead to further increases in yield.

46 **Keywords:** Wheat, grain number per m<sup>2</sup>, average grain weight, trade-off, individual grain weight;  
47 genetic determinism.  
48

## 50 **1. Introduction**

51 Yield of wheat can be decomposed into two major components: grain number per m<sup>2</sup> (GNM2)  
52 and average grain weight (AGW), often expressed as the thousand grain weight. The GNM2 is the  
53 main determinant of grain yield variation resulting from variations in environmental and/or genetic  
54 factors. Over the last fifty years grain yield increases have been achieved mainly through increases  
55 in GNM2 - rather than through increases in AGW (e.g., Fischer, 2008; Mondal et al., 2020; Slafer et  
56 al., 2014). Further increases in GNM2 are probably still possible under projected future climate  
57 conditions but the specific targeting of AGW is also a worthwhile option under climate conditions  
58 that may become less favourable and especially if these include greater levels of heat stress  
59 (Calderini and Ortiz-Monasterio, 2003; Quintero et al., 2018).

60 Support for the strategy of obtaining a genetic yield increase through direct manipulation of AGW  
61 is found in the observation that grain weight has a strong and heritable genetic determinism (Sadras,  
62 2007). Indeed, some molecular-physiological studies have identified genes involved in the various  
63 processes that govern grain growth and grain filling and the effects of their allelic variations on final  
64 grain size and weight (Laudencia-Chingcuanco et al., 2007; Nadaud et al., 2010; Simmonds et al.,  
65 2016; Brinton et al., 2018; Golan et al., 2019). Moreover, numerous quantitative trait loci (QTLs)  
66 for grain weight have been identified (e.g., Breseghello and Sorrells, 2007; Gegas et al., 2010; Farré  
67 et al., 2016; Brinton et al., 2016). However, for an increase in AGW to translate into a yield  
68 increase, requires a better understanding of how the mechanisms involved in grain weight  
69 determinism (Brinton and Uauy, 2019) and, in particular, of those allowing avoidance of the critical  
70 trade-off between GNM2 and AGW under variations in both environment and genotype (Slafer et  
71 al., 2015; Slafer and Savin, 1994). This trade-off could arise from limitations imposed by the source  
72 organs and/or by the sink organs.

73 A source limitation (i.e., competition for assimilates between sinks) could occur either before or  
74 after anthesis. Before anthesis, reduced assimilate availability can restrict individual grain ovary  
75 size, generating a physical constraint to growth (Ugarte et al. 2007) and thus reducing final grain  
76 size (Calderini et al., 1999; 2001; 2021). After anthesis, during the grain filling phase, a grain's  
77 final size may also be constrained by its share of a limited amount of assimilate - competition  
78 between grains (Sinclair and Jamieson, 2006). Even though the exact nature on the relationship is  
79 uncertain (Calderini et al., 2021) the idea that source limitation should affect final grain size  
80 through a limitation in ovary size seems unlikely, since the relation between ovary size and final

81 grain size appears mainly to be correlative, not causative (Benincasa et al., 2017). On the other  
82 hand, source limitation during grain filling has usually been dismissed as a main driver of average  
83 final grain size, at least in temperate environments (Borras et al., 2004; Borrill et al., 2015; Serrago  
84 et al., 2013; Elias et al., 2016 and references therein; Reynolds et al., 2022).

85 The idea of a trade-off originating from sink limitation, is based on the observation that an increase  
86 in GNM2 usually leads to a higher proportion of grains of low potential weight (Acreche and Slafer,  
87 2006; Chen et al., 2019; Feng et al., 2019). The higher proportion of small grains could be driven by  
88 processes occurring at two different levels. First, it could result from an increase in the number of  
89 secondary tillers and, consequently, an increase in the proportion of small spikes, that carry smaller  
90 grains (Xu et al., 2015) and, second, it could result from an increase in the number of grains per  
91 spike. These additional grains may be located in the basal or the apical spikelets, or in the more  
92 distal positions within each spikelet. All these locations are associated with lower potential weights  
93 and, thus, will tend to contribute to an increase in the proportion of small grains within the spike  
94 (e.g., Baillot et al., 2018; Calderini and Ortiz-Monasterio, 2003; Ferrante et al., 2017; Xie et al.,  
95 2015). These relationships seem to offer the most plausible explanation for the apparent trade-off  
96 produced by the introgression of Rht dwarfing genes (Miralles et al., 1995a). Both the processes  
97 described above, would lead to a higher proportion of small grains and so will be referred to below  
98 as ‘populational’ effects. The important roles of these populational effects on the trade-off between  
99 GNM2 and AGW is supported by a large number of studies (Acreche and Slafer, 2006; Chen et al.,  
100 2019; Feng et al., 2019; Ferrante et al., 2017; García et al., 2013; Li et al., 2016; Miralles and  
101 Slafer, 1995b; Quintero et al., 2018).

102 Given the importance of the trade-off, it appears necessary to distinguish between the part of the  
103 AGW genetic determinism that occurs *via* the populational effects described above and the part  
104 having a constitutive nature that affects each grain individually. Most quantitative genetic studies  
105 focusing on AGW do not explicitly consider the trade-off between GNM2 and AGW (e.g., Gegas et  
106 al., 2010; Brinton et al., 2018). The analytical framework of only one recent study seeks to take this  
107 trade-off into account (Quintero et al., 2018). This suggests the genetic determinism of individual  
108 grain weight is at least partly independent of the populational effects. However, their demonstration  
109 is incomplete: (i) because the study was restricted to particular spike and grain models (i.e.,  
110 proximal grain (G2) and distal grain (G4) sampled on the middle spikelet of five median spikes per  
111 experimental treatment) and (ii) because neither the number of grains per spike/spikelet nor the  
112 proportions of small vs. large grains were explicitly considered.

113 Thus, the aim of our study was to confirm the existence of a constitutive variability in AGW due  
114 only to genetic determinism, working with different well-characterised spike and grain populations.  
115 This requires we characterise the precise distribution of individual grain weights and their  
116 association with AGW at (1) plot scale, (2) spike scale and (3) single-grain scale. With this aim, we  
117 compared four modern, well-adapted, high-yielding wheat cultivars, that express robust differences  
118 in AGW. To help ensure that any differences we might observe in AGW and its putative  
119 determinism were not specific to any particular cultivar  $\times$  environment interactions, our analyses  
120 were carried out under two contrasting environmental conditions, (a) water-sufficient and (b) water-  
121 deficient.

## 122 **2. Materials and methods**

### 123 **2.1 Field experiment**

#### 124 *2.1.1. Choice of cultivars*

125 Four cultivars expressing either low AGW (hereafter LCvs) or high AGW (hereafter HCvs) were  
126 chosen from among a range of registered, modern, awned and high-yielding winter-wheat cultivars.  
127 This choice was made by analysing a dataset compiled from a large number of post-registration  
128 field trials (J.-C. Deswartes, P. du Cheyron, Arvalis-Institut du végétal, personal communication,  
129 2016) in which the mean yield components were calculated for each cultivar. Additional  
130 precautions were taken to minimise phenological differences between cultivars. However, it was not  
131 possible to identify cultivars that were within the exact same precocity class and still commercially  
132 available. Therefore, we selected Altigo (Nickerson, 2007) and Solehio (Momont, 2009) that exhibit  
133 higher AGWs and lower grain numbers per m<sup>2</sup> than Cellule (Florimond Desprez, 2012) and SY  
134 Moisson (Syngenta, 2012) (Table 1).

135

136 **Table 1.** Average yield components for selected winter bread wheat cultivars in multi-environment field trials (mean  $\pm$  Standard Deviation)

<b>Yield component</b>	<b>Altigo</b>	<b>Solehio</b>	<b>Cellule</b>	<b>SY Moisson</b>
<b>Number of trials</b>	56	135	126	57
<b>Years</b>	2007 - 2014	2009 – 2017	2012 - 2018	2012 – 2018
<b>Spike number per m<sup>2</sup></b>	494 $\pm$ 92	487 $\pm$ 72	555 $\pm$ 83	560 $\pm$ 86
<b>Grain number per spike</b>	37 $\pm$ 9	39 $\pm$ 5	43 $\pm$ 6	43 $\pm$ 6
<b>Grain number per m<sup>2</sup></b>	17607 $\pm$ 2729	18732 $\pm$ 2688	23473 $\pm$ 3269	23740 $\pm$ 2831
<b>Average grain weight (mg grain<sup>-1</sup>)</b>	52 $\pm$ 4	50 $\pm$ 5	41 $\pm$ 4	42 $\pm$ 4
<b>Grain Yield (t ha<sup>-1</sup> at 15% humidity)</b>	9.1 $\pm$ 1.4	9.3 $\pm$ 1.3	9.6 $\pm$ 1.4	9.9 $\pm$ 1.0

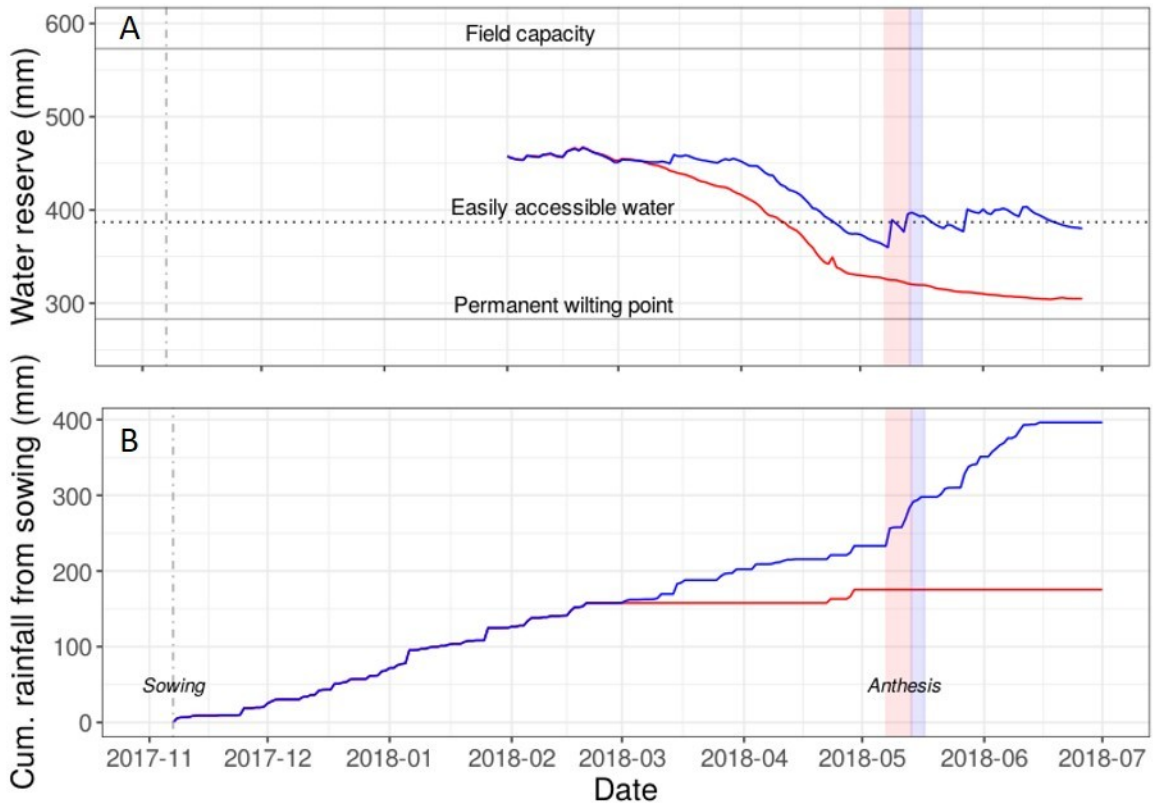


137 2.1.2. *Experimental design*

138 The field experiment was conducted at INRAE, Clermont-Ferrand (45°78' N, 3°08' E, 401 m asl)  
139 using the Pheno3C high-throughput, field-phenotyping platform. Over the last 20 years, mean  
140 annual air temperature at this site has been 11.7°C and mean annual precipitation 570 mm. The  
141 weather conditions during the experiments are representative of the mean climate at the site (Fig.  
142 S1) and no acute environmental stress events were noticed. The tillable horizon is a clay soil with  
143 42, 38 and 20% of clay, silt and sand, respectively and with high organic matter content (3.3%).

144 The Pheno3C facility consists of eight independent blocks. Four blocks are reserved for controls,  
145 maintained under the local climate, and these can be irrigated if required. The remaining four blocks  
146 can be covered by automatic rain-out shelters to generate a controlled water stress. All cultivars  
147 were sown on 10 November 2017 in plots within each of the eight blocks at a density of 250 grains  
148 m<sup>-2</sup>. Each plot (eight rows, 100 cm long, 17 cm inter-row spacing, 2.1 m<sup>2</sup> harvestable area) was  
149 managed following local agronomic practices. Due to a high residual mineral N after winter (150 kg  
150 N ha<sup>-1</sup>) a single N application was made to all plots at Zadock's stage Z32 (50 kg N ha<sup>-1</sup>). Weed and  
151 pest management included one herbicide application (at Z26), two fungicide applications (at Z51  
152 and Z69) and one insecticide application (at Z69).

153 In the four water-deficit blocks, the rain-out shelters of the Pheno3C facility were set in action from  
154 1 March 2018 to divert all incoming rainfall, except for one 17 mm rain event occurring during the  
155 last week of April. From sowing to harvest, the well-watered and water-deficit plots received 396  
156 and 175 mm of rainfall, respectively. Calculated from the start of the drought treatment, cumulative  
157 rainfall was 238 and 17 mm in the well-watered and water-deficit blocks, respectively (Fig. 1A). In  
158 the water-deficit blocks, the water stress was well established from the start of April, and deepened  
159 in intensity thereafter until harvest (Fig. 1B). The well-watered blocks were maintained at the onset  
160 of water stress (or above) by irrigation.



161

162 **Fig.1.** Dynamics of soil water reserve (A) and cumulative rainfall from sowing date (B) for the well-watered (blue line)  
 163 and water-deficit (red line) conditions. Rectangles indicate the maximum range of observed anthesis dates for the four  
 164 cultivars under well-watered (pale blue) and water-deficit (pink) conditions. Horizontal lines indicate the soil water  
 165 reserves corresponding to field capacity, easily accessible water and permanent wilting point.

166

167 We emphasise that the aim of the study was not to analyse responses to water deficit but merely to  
 168 create two contrasting background environments to explore the consistency of any differences that  
 169 might appear in AGW and its determinants, so bringing a greater degree of confidence and  
 170 robustness to our conclusions.

171

## 172 2.2 Plant sampling and measurements

173 Within each plot, the number of spikes per m<sup>2</sup> (SNM2) was determined after anthesis, at about  
 174 anthesis + 500°C days (base 0°C) by counting all spikes within two, 100 cm long, transects in the  
 175 two central rows.

176 At physiological maturity, in July 2018, 100 spikes per plot were sampled from the central 1.5 m of  
 177 the two central rows to avoid possible edge effects. To avoid bias in the selection of spikes, the 100

178 spikes were sampled as four separate samples of 25 adjacent spikes. All spikes were oven-dried to  
179 constant weight (60°C, 48 h) and for each individual spike, average grain weight ( $AGW_{spike}$ ) and the  
180 number of grains per spike ( $NGS_{spike}$ ) were determined with an automatic seed-weighing/-counting  
181 device (Opto Agri2, Optomachine, France). The weight of each individual grain within each spike  
182 was then measured with an automatic seed-weighing device developed in our laboratory. This  
183 device comprises a six-axis robotic arm (Yaskawa MH3, Yaskawa, Japan) integrated with a  
184 precision weight module (WMC15-SH, Mettler-Toledo, Switzerland).

185 Additional variables were calculated at the plot level. Mean number of grains per spike per plot  
186 ( $NGS_{plot}$ ) and AGW per plot ( $AGW_{plot}$ ) were calculated as the means of the 100 spikes of the sample.  
187 GNM2 was estimated as the product of SNM2 and  $NGS_{plot}$  and grain yield (GY) was calculated as  
188 the product of GNM2 and  $AGW_{plot}$ .

189

## 190 **2.3 Statistical analyses**

191 Results were analysed at different scales depending on the variable considered. Cultivar effects on  
192 mean yield components (SNM2,  $NGS_{plot}$ , GNM2,  $AGW_{plot}$  and GY) were analysed using mean  
193 variable values at the plot scale. Average grain weight per spike ( $AGW_{spike}$ ) and the individual grain  
194 weight per spike were analysed using individual spike values considering the nesting effect of plots.  
195 All analyses were carried out separately for the two environmental conditions.

196

### 197 *2.3.1. Cultivar effects on mean yield components at plot scale*

198 Within each environmental condition, comparison of the mean yield components (SNM2,  $NGS_{plot}$ ,  
199 GNM2,  $AGW_{plot}$ , GY) between cultivars was carried out using Tukey's *post-hoc* test following an  
200 ANOVA. For each environmental condition, the ANOVA model corresponded to a randomised  
201 complete block design with cultivar and block as fixed effects (n=4):

$$202 \quad Y = \mu + X_1Cv + X_2B + \epsilon \quad (\text{Eq.1})$$

203 where Y is the vector of each mean yield component value,  $\mu$  is the mean effect, Cv the fixed effect  
204 of each cultivar, B the fixed effect of the block,  $X_i$  the incidence matrix and  $\epsilon$  is the residual  
205 following a normal distribution (0,  $\sigma^2_\epsilon$ ).

### 207 2.3.2. Cultivar effects on average grain weight ( $AGW_{spike}$ ) at spike scale

208 To compare the differences in average grain weight per spike ( $AGW_{spike}$ ) between cultivars at spike  
 209 scale, a linear model was developed including the effect of the covariable  $NGS_{spike}$ . The  
 210 experimental design was also considered. To test the effect of the  $AGW$  genotypic level on  
 211  $AGW_{spike}$ , the four cultivars were pooled into two groups: HCvs (i.e., Altigo and Solehio) and LCvs  
 212 (i.e., Cellule and SY Moisson).

213 The following linear model was then used with the package ASREML (Gilmour et al., 1995):

$$214 Y = \mu + X_1Cov + X_2G + X_3Cv + X_4B + X_5Cov \times G \times B + \epsilon \quad (Eq.2)$$

215 where  $Y$  is the vector of average grain weight per spike ( $AGW_{spike}$ ) values,  $\mu$  is the intercept,  $Cov$   
 216 the fixed effect of the covariable ( $NGS_{spike}$ ),  $G$  the fixed effect of the group of cultivars,  $Cv$  the fixed  
 217 effect of each cultivar nested within each group of cultivars,  $B$  the fixed effect of the block,  $X_i$  the  
 218 incidence matrix and  $\epsilon$  is the residual following a normal distribution ( $0, \sigma^2_\epsilon$ ). A Wald test was  
 219 computed on this linear model to estimate the sizes of the different effects. Based on coefficients  
 220 calculated with the linear model, slopes and intercepts were calculated for each cultivar and under  
 221 each environmental condition.

222

### 223 2.3.3. Cultivar effects on individual grain weight

224 To analyse the putative populational effects associated with  $AGW$  differences requires going  
 225 beyond just the differences in means, to try to understand how the distributions actually differ.  
 226 Comparing quantiles is well suited to this purpose as they provide informative, robust and intuitive  
 227 information (Rousselet et al., 2017). Rousselet et al. (2017) proposed plotting a ‘shift function’ that  
 228 represents the differences between the quantiles of the distributions of two groups as a function of  
 229 the quantiles of one of the groups. Quantiles are estimated with the Harrell-Davis estimator (Harrell  
 230 and Davis, 1982 *in* Wilcox, 1995) and then quantiles differences between groups and their  
 231 associated confidence interval are estimated using a bootstrap method (as described in Rousselet et  
 232 al., 2017). In our study we extended this approach to provide a graphical representation of each  
 233 cultivar distribution to compare with the distribution of a pooled population comprising all of the  
 234 four cultivars. This offered a visual representation of the cultivar differences in spike size ( $NGS_{spike}$ )

235 and individual grain weight. The same approach was used to compare individual grain weight  
236 distribution of the two groups of cultivars (HCv and LCv).

237 To complete the graphical comparison, we carried out an analysis using the deciles as a categorical  
238 variable with the following linear model using the package ASREML (Gilmour et al., 1995):

$$239 \quad Y = \mu + X_1\text{GrD} + X_2\text{G} + X_3\text{Cv} + X_4\text{B} + X_5\text{GrD} \times \text{G} \times \text{Cv} \times \text{B} + \epsilon \quad (\text{Eq. 4})$$

240 where Y is the vector of individual grain weight values,  $\mu$  is the intercept, GrD the fixed effect of  
241 the grain decile, G the fixed effect of the group of cultivars, Cv the fixed effect of each cultivar  
242 nested within each group of cultivars, B the fixed effect of the block,  $X_i$  the incidence matrix and  $\epsilon$   
243 is the residual following a normal distribution  $(0, \sigma^2_\epsilon)$ . A Wald test was computed on this linear  
244 model to estimate the sizes of the different effects.

245 All statistical analyses were carried out with R statistical software (R version 3.5.0, R Development  
246 Core Team (2005)).

247

248 **3. Results**

249 **3.1. The four cultivars achieved similar grain yields through different combinations of AGW**  
250 **per plot and GNM2**

251        Within each environmental condition (well-watered *vs* water-deficit), there were no  
252 differences in grain yield among the four cultivars (Table 2). These results agree with those of the  
253 multi-environment trials used to select the cultivars for the experiment (Table 1). As expected,  
254 water-deficit led to a decrease in grain yield (~29% averaged among the four cultivars; Table 2).  
255 The expected small differences in cultivar precocity (Table 1) were also observed. Under well-  
256 watered conditions, the anthesis date difference between the earliest and the latest cultivars was  
257 only four days (Table 2). Under water-deficit conditions, anthesis dates of the four cultivars  
258 occurred over a seven-day period (Table 2).

260 **Table 2** – Average yield and yield components for the four selected winter bread wheat cultivars under well-watered or water-deficit conditions.

Condition	Group of cultivars <sup>1</sup>	Cultivar	Flowering date (days from sowing)	SNM2 <sup>2</sup> (spike per m <sup>2</sup> )	NGS <sub>plot</sub> <sup>2</sup> (grains spike <sup>-1</sup> )	GNM2 <sup>2</sup> (grains m <sup>-2</sup> )	AGW <sub>plot</sub> <sup>2</sup> (mg grain <sup>-1</sup> )	GY <sup>2</sup> (t ha <sup>-1</sup> )
Well-watered	HCv	Altigo	137	368 ± 24 a <sup>3</sup>	35.9 ± 0.6 b	13182 ± 837a	51.96 ± 0.77 d	7.0 ± 4 a
		Solehio	137	430 ± 84 ab	34.5 ± 1.3 a	14821 ± 3007 ab	49.83 ± 0.79 c	7.5 ± 14a
	LCv	Cellule	133	378 ± 32 ab	40.5 ± 0.8 c	15336 ± 1209 ab	40.43 ± 0.86 a	6.3 ± 4a
		SY Moisson	136	485 ± 78 b	35.5 ± 1.0 ab	17146 ± 2478 b	42.54 ± 0.82 b	7.5 ± 10a
Water-deficit	HCv	Altigo	134	362 ± 18 a	28.7 ± 1.2 a	10397 ± 648 a	48.77 ± 2.44 b	5.1 ± 5 a
		Solehio	134	344 ± 45 a	30.2 ± 1.8 a	10440 ± 1892 a	48.62 ± 1.03 b	5.1 ± 9 a
	LCv	Cellule	127	394 ± 50 a	33.1 ± 1.0 a	13042 ± 1645 b	38.43 ± 1.52 a	5.0 ± 7a
		SY Moisson	130	380 ± 21 a	31.2 ± 4.8 a	11793 ± 1141 ab	40.20 ± 1.34 a	4.8 ± 5a

261 <sup>1</sup>: HCv: high-AGW cultivar; LCv low-AGW cultivar262 <sup>2</sup>: SNM2: Spike number per m<sup>2</sup>; NGS<sub>plot</sub>: number of grains per spike determined on a plot basis; GNM2: grain number per m<sup>2</sup>; AGW<sub>plot</sub>: Average Grain Weight  
263 determined on a plot basis; GY: Grain Yield.264 <sup>3</sup>: Per condition, different letters indicate statistical significance at level P=0.05 within the same column (according to a Tukey post-hoc test following ANOVA).

265 The homogeneity among cultivars for yield, masked strong differences in the individual  
266 yield components between cultivars. Again, as expected from the cultivar choices, strong  
267 differences in AGW between cultivars were observed irrespective of the environmental conditions.  
268 Based on  $AGW_{plot}$ , under well-watered conditions, two classes of cultivars could be distinguished:  
269 on the one hand Altigo and Solehio (HCvs) had an average  $AGW_{plot}$  of  $\sim 51$  mg grain<sup>-1</sup> and on the  
270 other hand Cellule and SY Moisson (LCvs) had an average  $AGW_{plot}$  of  $\sim 41$  mg grain<sup>-1</sup> (Table 2).  
271 This distinction was emphasised under the water-deficit conditions with  $AGW_{plot}$  averages of  $\sim 48.7$   
272 mg grain<sup>-1</sup> for HCvs and  $\sim 39.0$  mg grain<sup>-1</sup> for LCvs (Table 2).

273 Given the trade-off between AGW and GNM2, strong differences in GNM2 were also  
274 expected between genotypes. Even though exhibiting a tendency, these differences in GNM2 were  
275 not always significant within the groups created on a  $AGW_{plot}$  basis. For example, within the well-  
276 watered experimental conditions, Solehio (HCv) and Cellule (LCv) showed similar values of  
277 GNM2 but different ones of  $AGW_{plot}$ . Similarly, under water-deficit condition, SY Moisson (LCv)  
278 showed GNM2 values not significantly different from those of Altigo or Solehio (HCvs) (Table 2).

279 No clear trend was observed for the source of cultivar variation in GNM2. The differences  
280 could be attributed either to genotypic variation of NGS at plot scale ( $NGS_{plot}$ ) that varied between  
281 34.5 and 40.5 grains per spike under well-watered conditions or to SNM2 depending on the  
282 genotype and the environmental conditions (Table 2).

283 Therefore, with similar grain yields but contrasting  $AGW_{plot}$ ,  $NGS_{plot}$  and SNM2 under both  
284 environmental conditions, the four cultivars allow exploration of the main sources of variability in  
285 cultivar  $AGW_{plot}$  and, in particular, the putative populational effects on variability of cultivar AGW.

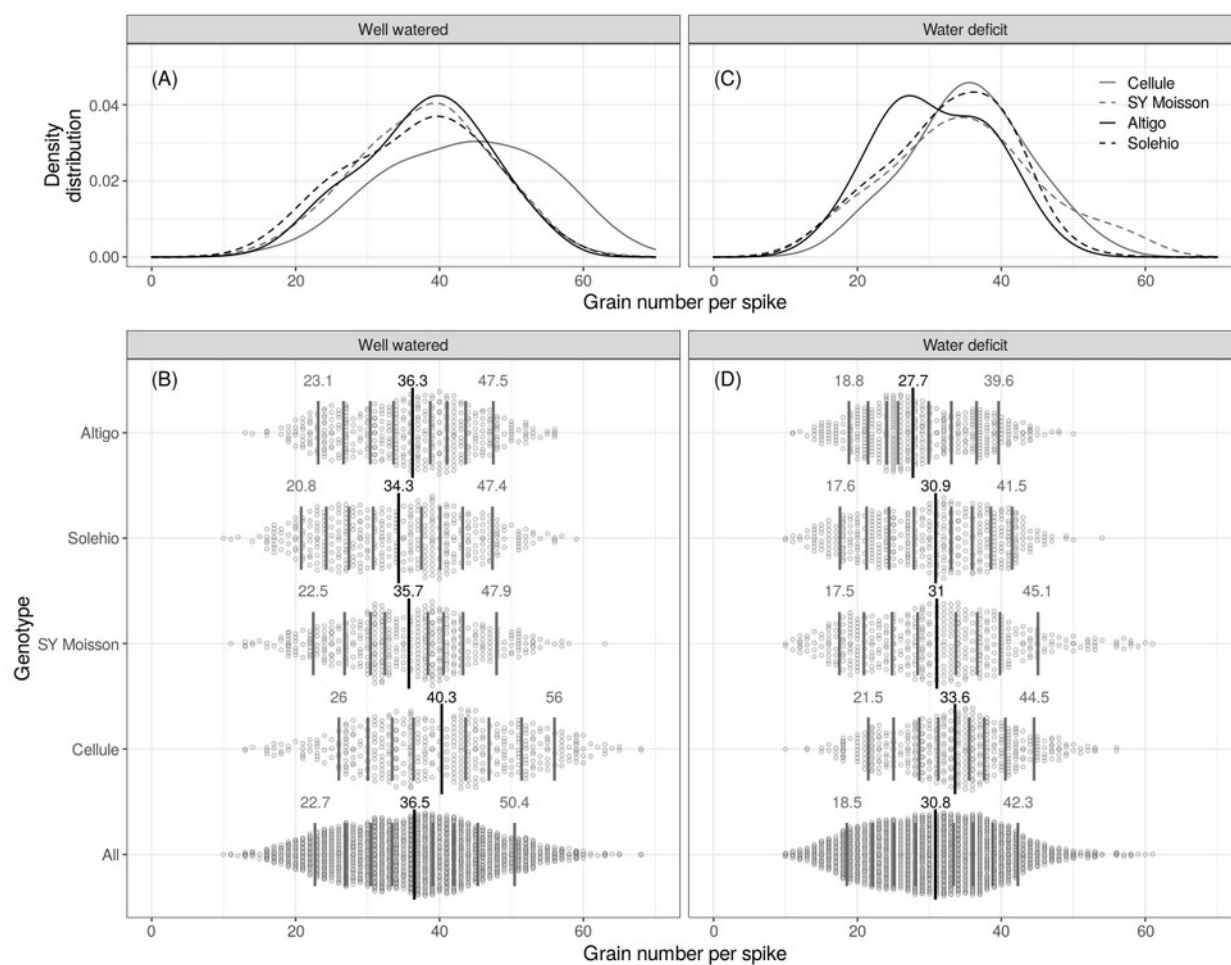
286

### 287 **3.2. $AGW_{spike}$ cultivar differences were not explained by differences in the distributions of** 288 **spike size**

289 To explore spike population effects on AGW our methodology requires information at the  
290 individual spike level. To this end, 100 individual spikes were sampled for each experimental unit,  
291 and  $NGS_{spike}$  and  $AGW_{spike}$  were determined for each spike sampled. This information obtained at the  
292 single spike scale allows us to study the contributions of spike population to cultivar  $AGW_{plot}$   
293 following a two-step evaluation. First, the distributions of spike size between cultivars were



294 compared using  $NGS_{spike}$ . Second, we examined whether between-spike variations in  $NGS_{spike}$   
 295 significantly affected  $AGW_{spike}$ .

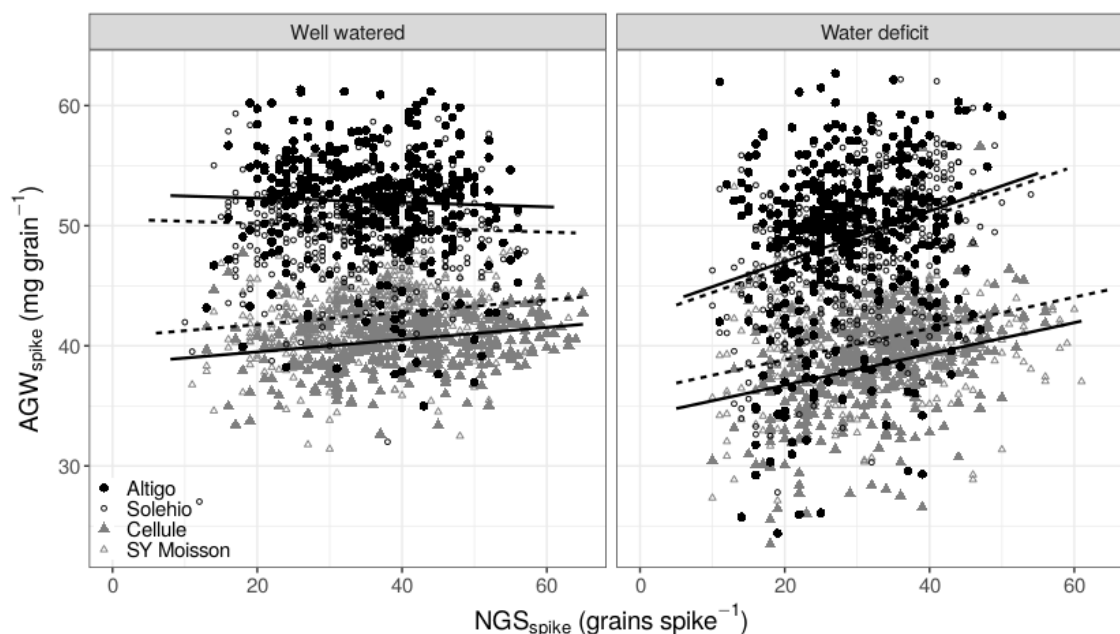


296

297 **Fig. 2:** Distributions of the grain number per spike ( $NGS_{spike}$ ) of the four cultivars under well-watered (A, B) and water-  
 298 deficit (C, D) conditions. Top panels (A, C) represent the density distribution of  $NGS_{spike}$  per cultivar. Distribution  
 299 parameters (mean, variance, skewness) for each condition and each cultivar: (A): Altigo (mean= 35.85, variance=  
 300 85.00, skewness= -0.02); Solehio (34.50, 105.36, 0.01); Cellule (40.53, 134.67, 0.06); SY Moisson (35.47, 92.11, -  
 301 0.04); (D): Altigo (mean= 28.71, variance= 64.95, skewness= 0.24); Solehio (30.17, 81.29, -0.11); Cellule (33.11,  
 302 76.79, 0.02); SY Moisson (31.28, 115.18, 0.33). Bottom panels (B, D) show the decile values of  $NGS_{spike}$  distributions  
 303 for the four cultivars and the whole spike population pooling the four cultivars (All). Figures in the bottom panels stand  
 304 for the values of the first decile, median and last decile.

305 For the well-watered condition, all distributions of  $NGS_{spike}$  were close to symmetrical (i.e.,  
 306 exempt from strong negative skewness) and were relatively similar except Cellule, for which  
 307  $NGS_{spike}$  values were shifted towards higher values (Fig. 2A) for all deciles (Fig. 2B). Regardless of  
 308 Cellule specificity, no link between SNM2 differences and  $NGS_{spike}$  distributions could be inferred.  
 309 For example, SNM2 was 32% higher in Altigo (HCv) than in SY Moisson (LCv) (Table 2) whereas  
 310 the  $NGS_{spike}$  distributions of these two cultivars were nearly identical (Fig. 2A).

311 Similar conclusions can be drawn under the water-deficit condition (Fig. 2C and 2D). The  
 312 distributions of  $NGS_{spike}$  were very similar for the four cultivars with only a small shift of Altigo  
 313 towards smaller spikes. This small difference does not appear to be correlated with NSM2, as  
 314 differences among the genotypes were marginal and not significant (Table 2).



315

316 **Fig.3.** Relationship between  $NGS_{spike}$  (grains spike<sup>-1</sup>) and  $AGW_{spike}$  (mg grain<sup>-1</sup>) for each cultivar under well-watered (A)  
 317 and water-deficit conditions (B). For each condition, slope and intercept are calculated following the model described in  
 318 the Material and Methods section: (A) Altigo (slope= - 0.018; intercept= 52.65); Solehio (-0.018, 50.55); Cellule  
 319 (0.051, 38.48); SY Moisson (0.051, 40.75); (B) Altigo (slope= 0.21, intercept= 42.82); Solehio (0.21, 42.34); Cellule  
 320 (0.13, 34.13); SY Moisson (0.13, 36.25). The  $R^2$  of the models are 0.67 (A) and 0.56 (B). Altigo and Solehio are High-  
 321 AGW cultivars (HCv); Cellule and SY Moisson are Low-AGW cultivars (LCv).

322 There were no relationships (Fig. 3A) between  $AGW_{spike}$  and  $NGS_{spike}$  for the well-watered  
 323 conditions and only weak positive (Fig. 3B) ones for the water-deficit conditions. For both  
 324 conditions, an analysis of variance (Table 3) reveals that the differences in  $AGW_{spike}$  are explained  
 325 mainly by the *a priori* classification of the cultivars into HCv or LCv with 61.3% of total variance  
 326 explained for the well-watered and 47.9% for the water-deficit conditions. Although highly  
 327 significant, the other effects never explained more than 4.2% of the variance in  $AGW_{spike}$  (Table 3).  
 328 In particular, the interaction term between  $NGS_{spike}$  and the group of cultivars ( $NGS \times G$ ) explained  
 329 only 0.4 and 0.2% under well-watered and water-deficit conditions, respectively. Taken together,  
 330 these results indicate that the cultivar differences in  $AGW_{spike}$  were not due to populational effects at  
 331 spike level.

332 **Table 3.** Analysis of variance (ANOVA) followed by the statistics of Wald test for AGW<sub>spike</sub> (mg grain<sup>-1</sup>) under well-watered and water-deficit conditions.

Condition	Source of variation <sup>1</sup>	Degrees of freedom	Percentage of variance explained	Wald statistic	P-value <sup>2</sup>
Well-watered	Number of grains per spike (NGS)	1	0.7	35.74	***
	Block (B)	3	0.8	40.31	***
	Group (G) of cultivars (Cv)	1	61.3	2992.91	***
	Cv nested in G	2	3.0	147.32	***
	NGS × G	1	0.4	17.44	***
	NGS × B	3	0.2	7.14	ns
	G × B	3	0.5	22.59	***
	G × B × NGS	3	0.1	2.79	ns
	Residual	1618	33.1		
Water-deficit	NGS	1	0.9	34.49	***
	B	3	4.2	155.77	***
	G	1	47.9	1758.29	***
	Cv nested in G	2	1.1	41.77	***
	NGS × G	1	0.2	6.89	**
	NGS × B	3	0.3	9.32	*
	G × B	3	0.8	28.07	***
	G × B × NGS	3	0.2	7.03	ns
	Residual	1631	44.4		

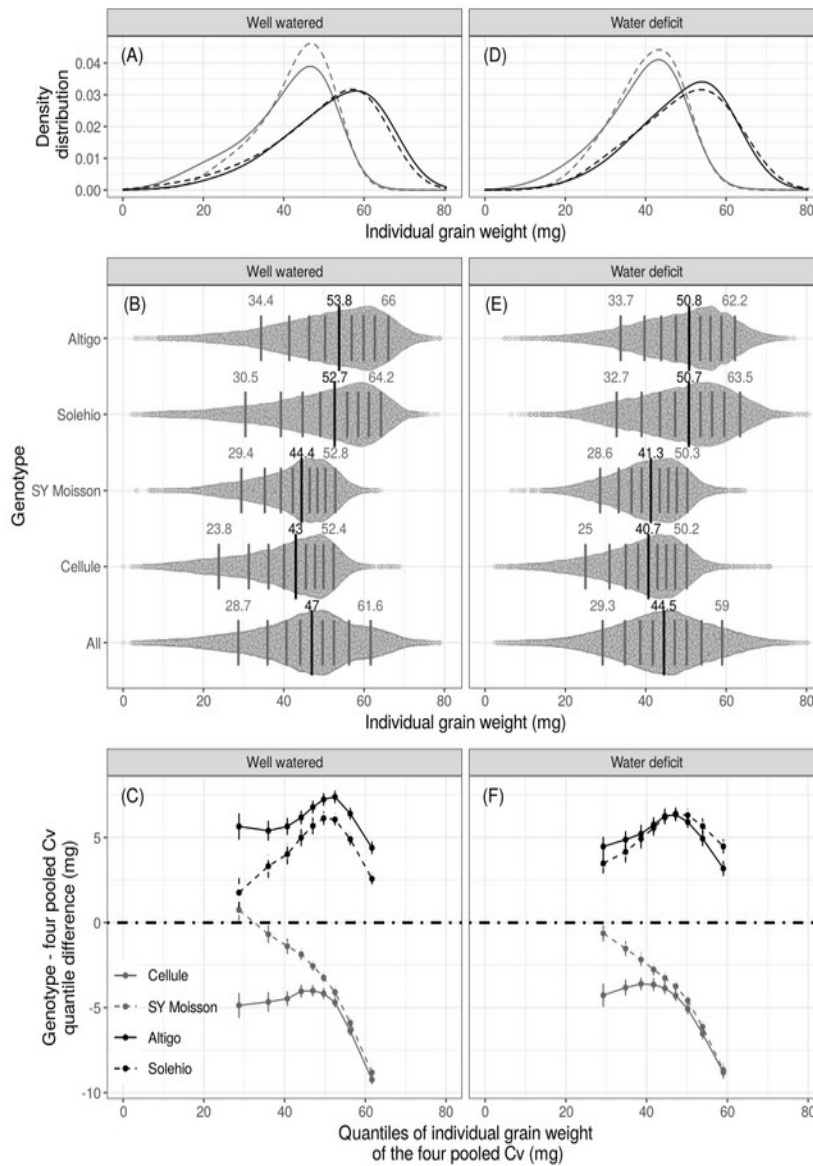
333 <sup>1</sup>: G: HCv = High AGW cvs (Altigo and Solehio) ; LCv = Low AGW cvs (Cellule and SY Moisson)

334 <sup>2</sup>: ns : P> 0.05; \* : P<0.05; \*\* : P<0.01 ; \*\*\* : P<0.001

### 335 3.3. Cultivar differences in individual grain weight were observed, irrespective of grain decile

336 The second populational effect proposed as a possible driver of cultivar differences in  
337  $AGW_{plot}$ , occurs within the spike, at the individual grain level. Under this assumption, cultivars with  
338 larger spikes (higher NGS) will present higher proportions of small grains within spikes and this  
339 will negatively affect  $AGW_{plot}$ . Assessing individual grain weight distributions per spike is made  
340 difficult due to (i) the limited number of grains within a spike and (ii) the large variation in the  
341 numbers of grains between spikes. Given the absence of clear a relationship between  $NGS_{spike}$  and  
342  $AGW_{spike}$ , the individual grain weight distribution was assessed only at the whole population level.

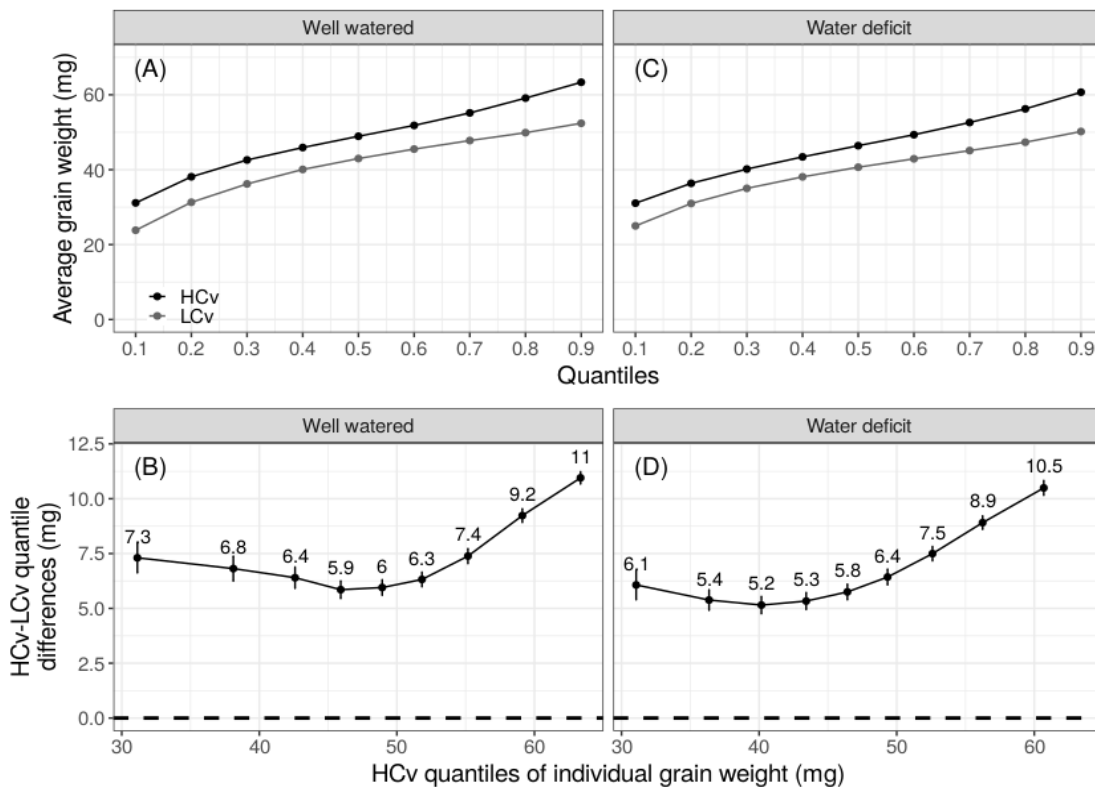
343 As expected, given that the choice of the four cultivars was based on their *a priori* differences in  
344  $AGW$ , clear differences in individual grain weight could be observed between Altigo and Solehio  
345 (HCv cultivars) on the one hand and SY Moisson and Cellule (LCv cultivars) on the other hand  
346 under both environmental conditions (Fig. 4A and 4D). All distributions were negatively skewed,  
347 with an excess of small grains. The shift function allowed comparison of the individual grain weight  
348 decile values, on the one hand of the whole-grain population made by pooling the four cultivars  
349 and, on the other hand, with each cultivar independently. This representation allows to visualisation  
350 of both the strong differences between the HCv and the LCv cultivars and of the relative  
351 homogeneity of the cultivars within each of these groupings, in spite of the divergent behaviours of  
352 the first four deciles (Fig. 4C and 4F) that relate to small skewness differences among cultivars.



353

354 **Fig. 4:** Distributions of the individual grain weight (mg) of the four cultivars under well-watered (A, B, C) and water-  
 355 deficit (D, E, F) conditions. Top panels (A, D) represent the density distribution of individual grain weight per cultivar.  
 356 Distribution parameters (variance, skewness) for each condition and each cultivar: (A) Alligo (mean= 51.79, variance=  
 357 156.91, skewness= -0.69); Solehio (49.91, 178.58, -0.85); Cellule (40.60, 122.71, -0.73); SY Moisson (42.78, 83.04, -  
 358 0.68); (D) Alligo (mean= 49.16, variance= 125.47, skewness= -0.56); Solehio (49.34, 136.68, -0.40); Cellule (38.96,  
 359 98.21, -0.65); SY Moisson (40.38, 70.07, -0.44). Middle panels (B, E) show the deciles values of individual grain  
 360 weight distributions for the four cultivars and the whole grain population mixing the four cultivars (All). Numbers are  
 361 given for the first decile, median and last decile. Bottom panels represent the shift function i.e., the differences between  
 362 the deciles of each cultivar from the deciles of whole population.

363 Individual grain weight differences between the HCv and LCv cultivars are even more striking  
 364 when averaged per group of cultivars. Under both conditions, individual grain weight from all  
 365 deciles is higher for HCv than for LCv, regardless of the decile considered (Fig. 5A and 5C).  
 366 Remarkably, these differences between cultivar groups tended to increase with grain weight (Fig.  
 367 5B and 5D) and these relationships were very similar under both environmental conditions.



368

369 **Figure 5:** Mean individual grain weight per decile for two groups of cultivars HCv and LCv under well-watered (A, B)  
 370 and water-deficit (C, D) conditions. HCv are Altigo and Solehio; LCv are Cellule and SY Moisson. Top panels (A, C)  
 371 represent the mean individual grain weight per decile per group. Bottom panels represent the shift function i.e., the  
 372 differences between the deciles of the HCv cultivars and the deciles of the LCv cultivars as a function of the HCv  
 373 deciles. The numbers indicate the differences (mg) for each decile.

374 The conclusions drawn from the graphical analysis of the shift function are supported by an  
 375 analysis of variance (Table 4). Indeed, apart from the large effect of grain decile, the main factor  
 376 affecting individual grain weight was the group of cultivars (14.2 for the well-watered conditions  
 377 and 16.9% for the water-deficit conditions). This result is independent of grain weight decile, as  
 378 indicated by the small interaction term (<1% of the variance explained in both environmental  
 379 conditions). In other words, HCvs had higher AGW than LCvs due to their having constitutively  
 380 heavier grains.

381 **Table 4.** Analysis of variance (ANOVA) followed by the statistics of Wald test for individual grain weight (mg) under well-watered and water-deficit conditions.

Condition	Source of variation <sup>1</sup>	Degrees of freedom	Percentage of variance explained	Wald statistic	P-value <sup>2</sup>
Well-watered	Block (B)	3	0.16	704.47	***
	Grain Decile (GrD)	9	70.69	316262.95	***
	Group (G) of Cultivars (Cv)	1	14.17	63398.26	***
	Cv nested in G	2	0.69	3101.77	***
	G × B	3	0.09	410.19	***
	G × GrD	9	0.86	3832.66	***
	B × GrD	27	0.01	56.03	***
	G × B × GrD	37	0.02	69.84	***
	Residual	59582	13.32		
Water-deficit	B	3	1.68	3903.87	***
	GrD	9	54.70	127044.86	***
	G	1	16.94	39343.56	***
	Cv nested in G	2	0.21	482.45	***
	G × B	3	0.35	819.85	***
	G × GrD	9	0.38	891.54	***
	B × GrD	27	0.06	133.76	***
	G × B × GrD	37	0.02	40.47	**
	Residual	59582	25.66		

382 <sup>1</sup>: Grain deciles (GrD) ; G: HCv = High AGW cvs (Altigo and Solehio) ; LCv = Low AGW cvs (Cellule and SY Moisson)

383 <sup>2</sup>: \*\*: P<0.01 ; \*\*\* : P<0.001.

## 384 4. Discussion

### 385 4.1. Relevance of the selected cultivars to exploring the grain number/AGW relationship

386 Most studies comparing wheat cultivars and showing differences in grain number and AGW  
387 have focused on the dichotomy between old cultivars and new ones, or between tall cultivars and  
388 semi-dwarf ones (Chen et al., 2019; Ferrante et al., 2017; Miralles and Slafer, 1995b), or they have  
389 explored cultivars showing wide-ranging yield potentials (e.g., Peltonen-Sainio et al., 2007;  
390 Quintero et al., 2018). All these studies, together with an abundance of studies that have focused on  
391 yield responses to different management practices (e.g., Acreche and Slafer, 2006; Feng et al.,  
392 2019; Fischer, 1993; Li et al., 2016) have concluded that yield variations are mainly related to  
393 variations in GNM2 (e.g., Slafer et al., 2014 and references therein). Given the negative relationship  
394 that commonly exists between GNM2 and AGW (Slafer et al., 2014; Slafer and Savin, 1994),  
395 studies that rely on a high variability in GNM2, only highlight the dominant effect of GNM2 on  
396 AGW. Therefore, this limits their ability to determine the component of AGW genetic variation,  
397 that can be attributed to constitutive genetic differences.

398 One way to resolve this problem is to drastically restrict the range of variation in GNM2, so  
399 as to allow a focus on AGW. In the studies cited above (Ferrante et al., 2017; Peltonen-Sainio et al.,  
400 2007; Quintero et al., 2018), the range of values of GNM2 varied by more than ten-fold across the  
401 cultivars and/or conditions they tested. For example, Ferrante et al. (2017) observed GNM2 values  
402 that varied between 2 500 and 25 000, Peltonen-Sainio et al. (2007) between 2 300 and 19 000 and  
403 Quintero et al. (2018) between 11 000 and 25 000 grains m<sup>-2</sup>. In our study, the range in GNM2 was  
404 much narrower i.e., between 13 000 and 17 000 grains m<sup>-2</sup> under well-watered conditions and  
405 between 10 000 and 13 000 grains m<sup>-2</sup> under water-deficit conditions (Table 2). Nevertheless,  
406 significant differences in AGW were still evident (Table 2). This suggests the observed differences  
407 in AGW were not driven solely by GNM2 differences. The four modern, high-yielding winter bread  
408 wheat cultivars we used, expressed consistently high (Altigo and Solehio) and low (Cellule and SY  
409 Moisson) values of AGW. However, these strong differences in yield components are also  
410 accompanied by earliness differences between LCv and HCv cultivars (Table 2). Earliness  
411 differences between cultivars are suspected to have a strong confounding effect on the  
412 establishment of yield components, in particular when developmental differences generate  
413 differences in exposure to climatic stresses. In our experiment, no acute uncontrolled abiotic stress  
414 event occurred during the crop cycle (Fig. 1, Fig. S1). Therefore, we do not believe that the limited



415 precocity differences observed will have caused significant differences in yield component  
416 establishment.

417 We thus conclude that the four cultivars we study here will allow us to determine if a  
418 proportion of the determination of the AGW differences could be constitutive. In addition, the two  
419 contrasting growing conditions (well-watered and water-deficit) will allow us to determine if our  
420 conclusions are reasonably independent of the growing environment. In this case it will allow a  
421 degree of generalisation for our results.

422

## 423 **4.2. AGW cultivar differences are poorly explained by populational effects reflecting their** 424 **constitutive nature**

425 The results of numerous studies published in the last few years support the idea that a large  
426 portion of the trade-off between GNM2 and AGW originates from populational effects. In other  
427 words, an increase in GNM2 will be associated with an increase in the relative contributions of  
428 grains having a lower weight potential than the rest of the grain population (Acreche and Slafer,  
429 2006; Miralles and Slafer, 1995a) and this will directly and negatively affect AGW.

430 From the putative populational effects suggested above, the one acting at the spike  
431 population level has often been mentioned but rarely studied directly. Xu et al. (2015) showed that  
432 tillers of higher order, produce spikes with lower grain numbers per spike and lower AGW. In  
433 particular, spikes from the first primary tillers have (depending on genotype) between 4 and 12%  
434 less grains per spike than spikes from the main stem. This effect is even stronger for tillers of higher  
435 order. The same observation was highlighted by Thorne and Wood (1987) with a much higher  
436 (28%) reduction in NGS in tillers compared to main stems. Such results imply that any increase in  
437 SNM2 made through an increased proportion of tillers (of any order) would induce a higher  
438 proportion of smaller spikes in the population. In the present study, the number of grains per spike  
439 ( $NGS_{\text{spike}}$ ) of individual spikes was recorded but information on the tiller order that produced these  
440 spikes was not. Nevertheless, given that the plant density at sowing under our conditions (250  
441 plants  $m^{-2}$ ) and the observed SNM2, are within the range 344-485 spikes  $m^{-2}$  (Table 2), our spike  
442 populations would have been composed mainly of spikes originating from the main stem with the  
443 calculated number of spikes per plant within the narrow range of 1.4 to 1.9 spikes per plant,  
444 depending on genotype and condition. The low SNM2 observed in our experiment probably  
445 explains the absence of clear genotypic differences in the NGS distribution (Fig. 2). This absence of  
446 populational effect observed at the spike level is likely to explain the absence of a relationship

447 between NGS and  $AGW_{spike}$  in spite of the large range of  $NGS_{spike}$  observed under both conditions  
448 and in all genotypes (Fig. 3). This implies that the hypothesis that spike populational effects on  
449 AGW probably relies on the effects of tiller order on AGW, rather than on a direct effect of  $NGS_{spike}$   
450 on AGW. To fully validate this proposition, the characterisation of both the individual NGS (such  
451 as in the present experiment) and the order of the tillers carrying each spike (as made by Xu et al.,  
452 2015 and Thorne and Wood, 1987) would require additional experimental work.

453 The second populational effect suggested in the literature having a putative effect on AGW,  
454 operates at the within-spike level. From this point of view, increasing grain number per  $m^2$  through  
455 an increase in NGS is accompanied by a higher proportion of small grains positioned in basal or  
456 apical spikelets within the spike (e.g., Feng et al., 2019; Ferrante et al., 2017; Miralles and Slafer,  
457 1995a). It is well established that, within spikelets, the grains in distal florets are smaller than in  
458 basal florets (e.g., Baillot et al., 2018; Miralles and Slafer, 1995a; Rawson and Evans, 1970). Taken  
459 together both relationships lead to a decrease in  $AGW_{spike}$  (Acreche and Slafer, 2006).

460 A problem is the estimation of the proportion of small grains within a population of grains is  
461 not straightforward, as the weight threshold criterion for ‘small’ is necessarily arbitrary. An  
462 alternative used by Acreche and Slafer (2006) is to define the proportion of small grains as the ratio  
463 between the number of grains located in distal positions as a proportion of the total number of  
464 grains in the spike. However, this method does not account for actual individual grain weights and  
465 can be misleading as an additional grain in a distal position on a central spikelet is not necessarily  
466 smaller than a proximal grain in a more basal or apical spikelet, as suggested by the results of Feng  
467 et al. (2018; Fig. 4).

468 Therefore, we used a method based on an analysis of individual grain weight deciles to  
469 compare cultivars. Within this analytical framework, the hypothesis of cultivar AGW differences  
470 driven by populational effects should result in higher weight differences between cultivars for the  
471 first deciles than for the last deciles. Our results clearly don’t support this hypothesis as cultivar  
472 differences in individual grain weight tend to increase with deciles in a similar fashion under the  
473 two environmental conditions (Fig. 5, Table 4). In other words, the LCv cultivars do not have lower  
474 AGWs than the HCv cultivars due to an increased proportion of small grains but because all grains  
475 are smaller, in particular the larger ones.

476 In conclusion, we note that our data did not expose either of the two populational effects  
477 previously suggested as the main determinants of AGW difference between cultivars. This does not  
478 mean that such populational effects never occur (they would likely dominate the trade-off between

479 GNM2 and AGW when large changes in GNM2 are produced) but that a component of the AGW  
480 differences observed between cultivars has a constitutive nature, as already suggested by Quintero  
481 et al. (2018). However, these authors' conclusions were based exclusively on the observation that  
482 both the proximal and distal grains (G2 and G4) of the two central spikelets were larger in the high  
483 AGW cultivars than in the low AGW cultivars. Thus, their analyses did not account for the diversity  
484 observed either in the spike population (variation in spike grain number distribution) or in the grain  
485 population caused by variation in spikelet number per spike. Our study explicitly evaluated the  
486 populational effects and showed that these were extremely weak and thus not sufficient to explain  
487 the cultivar differences in AGW. The constitutive nature of grain size difference has thus been  
488 highlighted for all grains (Fig. 5), independently of their weights and independently of the spike  
489 from which they originated. In agreement with Calderini et al. (2021), our conclusion implies that  
490 AGW can to some extent be manipulated independently of grain number.

491

#### 492 **4.3. Towards new targets for grain-weight breeding**

493 It is widely accepted that the grain yield increases achieved in recent decades have come  
494 about mainly through increases in grain number per m<sup>2</sup>. Meanwhile, AGW has been increased only  
495 marginally over the same period (e.g., Brancourt-Hulmel et al., 2003; Philipp et al., 2018; Shearman  
496 et al., 2005) and has been selected principally for its effects on grain quality, rather than on grain  
497 yield (Philipp et al., 2018; Xie et al., 2018). The existence of a significant part of AGW genetic  
498 variation that can be attributed to constitutive cues (i.e., independent of variations in grain number)  
499 offers new avenues for manipulating yield. First, gene manipulation (in particular, expansin gene  
500 manipulation) has been proposed as a way to increase GW (Lizana et al., 2010), without having a  
501 negative effect on GN, and so increase GY (Calderini et al., 2021). Second, this approach could rely  
502 on the QTLs already identified for grain weight (e.g., Yan et al., 2017), and especially on those that  
503 have been reported to be independent of the trade-off between AGW and grain number (Griffiths et  
504 al., 2015). Nevertheless, such QTLs should be used with care since the AGW quantification on  
505 which they rely were potentially compromised by populational effects that negatively influences the  
506 mean AGW. Using a methodology that provides access to single grain weights (as does the present  
507 study) should allow application of GWAS approaches, focusing specifically on the mean weights of  
508 the larger grains (defined, for example, as the 25% of larger grains). Our results suggest that this  
509 particular grain sub-population is unaffected by putative populational effects. This approach should  
510 be validated, independently of the target environment.

511 **Declaration of Competing Interest**

512 The authors declare they have no known competing financial interests or personal relationships that  
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514

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526

527 **Supplementary material**

528

529

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