## Yield performance of chromosomally engineered durum wheat-*Thinopyrum*

# ponticum recombinant lines in a range of contrasting rain-fed environments

## across three countries

- 4 Ljiljana Kuzmanović a\*, Roberto Ruggeri a, Jason A. Able b, Filippo M. Bassi c, Marco Maccaferri d,
- 5 Roberto Tuberosa <sup>d</sup>, Pasquale De Vita <sup>e</sup>, Francesco Rossini <sup>a</sup>, Carla Ceoloni <sup>a\*</sup>
- 6 <sup>a</sup> Department of Agricultural and Forest Sciences (DAFNE), University of Tuscia, Viterbo, Italy
- 7 b School of Agriculture, Food and Wine, Waite Research Institute, The University of Adelaide, Glen Osmond, Australia
- 8 °ICARDA, Biodiversity and Integrated Gene Management, Rabat Institutes, Rabat, Morocco
- 9 d Department of Agricultural Sciences, University of Bologna, Bologna, Italy
- CREA-CI Cereal Research Centre for Cereal and Industrial Crops, Foggia, Italy
- \*\*Corresponding authors:
- 13 <u>ljiljanakuzmanovic@gmail.com</u> (L. Kuzmanović)
- 14 <u>ceoloni@unitus.it</u> (C. Ceoloni)
  - Abstract

15

16

1

2

- 17 Introgressions of *Thinopyrum ponticum* 7AgL chromosome segments, spanning 23%, 28% and 40%
- of the distal end of durum wheat 7AL arm, were previously shown to contain multiple beneficial
- 19 gene(s)/QTL for yield-related traits, in addition to effective disease resistance (Lr19, Sr25) and
- quality (Yp) genes. In the present study, durum wheat near isogenic recombinant lines (NIRLs),
- harbouring each of the three introgressions, were included for the first time in multi-location field
- trials, to evaluate general and environment-specific effects of the alien chromatin on 26 yield-related
- traits. The results from nine different trials across contrasting environments of Italy, Morocco and
- South Australia over four years revealed that the overall impact of 7AgL introgressions into the
- 25 tetraploid wheat background did not imply, except in one environment, major yield penalty. The
- 25 tetrapion wheat background and not impry, except in one environment, major yield penaity. The
- comprehensive effect of the three 7AgL segments on individual yield-contributing traits, resulted in significant increases of biomass m<sup>-2</sup> (+9%), spike number m<sup>-2</sup> (+13%), grain number m<sup>-2</sup> (+11%) and
- spikelet<sup>-1</sup> (+8%), but also in a general, significant decrease of grain weight (-8%). When the separate
- 28 spirited (1070), but also in a general, significant decrease of grain weight (-070). When the separate
- NIRLs were analysed, each of the three 7AgL segments turned out to be associated with variation of
- specific yield components. The effects of the 40%-long segment proved to be the most stably
- expressed across environments and involved significant increases of spike and grain number m<sup>-2</sup> (13%
- and 15%, respectively), grain number spike<sup>-1</sup> (10%) and spike fertility index (46%), though
- accompanied by a significant decrease in thousand grain weight (-23%). In spite of this trade-off
- 34 between grain number and grain weight, their interplay was such that in four trials, including dryer
- environments, a grain yield advantage was observed. This evidence, and comparison with the two
- other NIRLs, substantiates the hypothesized existence of major gene(s)/QTL for grain number in the
- 37 most proximal 28-40% 7AgL region, exclusive to the 40%-long 7AgL introgression. The present
- study represents an important validation of the use of chromosomally engineered genetic stocks for
- durum wheat improvement, targeting not only disease resistance and quality traits but also relevant
- 40 yield components.
- 41 **Keywords**: Alien introgression, Chromosome engineering, Grain number, Tiller number, Wheat
- 42 breeding

## Introduction

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

Durum wheat (*Triticum durum* var. *durum*, 2n = 4x = 28, genomes AB) is cultivated on approximately 8% of the world's wheat area, and is economically important in the Mediterranean basin, North America's Great Plains and Mexico, as well as Australia, Russia, Kazakhstan, India, Ethiopia and Argentina (Bassi and Sanchez-Garcia, 2017; Royo et al., 2014). Due to the socio-economic challenges emerging from the rapid human population increase across the world, breeding for continual improvement in yield potential (among other traits) is a priority for durum wheat breeders. Given that wheat provides calories for about 20% of human nutrition (FAO, 2013), current yield increases of not more than 1% year-1 for both durum and bread wheat, will be insufficient to meet the food demand in the imminent future (Fischer and Edmaedas, 2010; Ray et al., 2013). Further, with ongoing climate changes, especially temperature extremes and water shortages, already determining a progressive displacement of durum wheat cultivation from traditional areas (Ceoloni et al., 2014a; Habash et al., 2009), the goal of significantly increasing yield potential is even more challenging. An approximate 6% loss in global wheat yield was recently predicted for each degree-Celsius increase in temperature (Liu et al., 2016; Zhao et al., 2017). Yield reduction will depend on the interaction of temperature with other limiting factors, such as rainfall, CO<sub>2</sub> emissions, and nitrogen supply, thus requiring environment-specific breeding strategies to select for highly adapted genotypes (Elía et al., 2018; Fleury et al., 2010; Luo et al., 2005; Tricker et al., 2018; Zhao et al., 2017). Consequently, for a particularly complex trait such as yield, understanding and dissecting the multi-layered genotype × environment interaction in multi-environment trials is a crucial step towards closing the gap between the actual and attainable yields, particularly when several different constraints are present (Araus et al., 2003a; Bassi and Sanchez-Garcia, 2017; Dodig et al., 2012; Maccaferri et al., 2011; Marti and Slafer, 2014; Parent et al., 2017; Slafer et al., 2014; Tardieu and Tuberosa, 2010; Zaim et al. 2017). To develop durum genotypes with improved yield and adaptability to more frequent incidence of drought and heat stress and/or altered rainfall distribution (Habash et al., 2009; Tadesse et al., 2016), enhancing the genetic background through targeted introgressions is a powerful strategy, given that cultivated germplasm represents only a very small fraction of the variability present in nature (Royo et al., 2009 and references therein; Zaim et al. 2017). Wheat-alien introgression experiments conducted in the past proved to be a valid approach to harness the genetic diversity of alien, mostly wild, segments of wheat-related gene pools (Ceoloni et al., 2014b, 2017a; Dempewolf et al., 2017; **Prohens** et al., 2017; al., 2017). To achieve Zhang et this, addition/substitution/translocation/recombinant lines, harbouring parts of alien genomes, can be used to facilitate the introduction of desired genes into stable and adapted genotypes (e.g. Ceoloni et al.,

2015). Targeted and precise exploitation of useful genes from these materials is readily possible through efficient sexual means, foremost the cytogenetic methodologies of "chromosome engineering" (Ceoloni et al., 2005, 2014a, 2014b). This approach, integrated with continuously developing techniques of genome and chromosome analysis (e.g. marker-assisted selection, association mapping, next generation sequencing, *in situ* hybridization), represents a unique platform for creation of novel and breeder-friendly genetic stocks.

The use of wild relatives for yield improvement in wheat has so far been sporadic, as their productivity is poor, and conspicuous effects on wheat yield rarely observed (Ceoloni et al., 2015; Dempewolf et al., 2017; Zhang et al., 2017). Noteworthy examples regard mostly the hexaploid bread wheat, more widely cultivated, and benefiting from a higher ploidy level with respect to durum wheat, hence a higher buffering ability toward chromosome manipulations (reviewed in Ceoloni et al., 2014a, 2015; Mondal et al., 2016). One of the most notable and documented cases of alien introgression with positive effects on wheat yield, is the transfer of a portion from the group 7 chromosome arm (= 7AgL or 7el<sub>1</sub>L) of the decaploid perennial species *Thinopyrum ponticum* (Popd.) Barkworth & D. R. Dewey (2n = 10x = 70, genomes  $E^eE^eE^x$ StSt, see Ceoloni et al., 2014b) onto the 7DL and 7AL arm of bread and durum wheat, respectively. In bread wheat, the sizeable 7AgL translocation named T4 (~70% of the recipient 7DL arm, harbouring Lr19+Sr25+Yp genes) led to increased grain yield, biomass and grain number (10-35%) across a number of non-moisture stress environments, and in different backgrounds of CIMMYT germplasm (Monneveux et al., 2003; Reynolds et al., 2001; Singh et al., 1998; Tripathi et al., 2005; Miralles et al., 2007). Under water stress, however, yields for CIMMYT T4 derivatives turned out to be lower than control lines (Monneveux et al., 2003; Singh et al., 1998), as was the case for T4 derivatives developed in Australian adapted genetic backgrounds, when tested in high- and low-yielding environments (Rosewarne et al., 2015).

In durum wheat, three fractions of the same 7AgL chromatin, spanning 23%, 28% and 40% of the recipient 7AL arm of cv. Simeto, and all containing the *Lr19+Sr25+Yp* genes (Ceoloni et al., 2005), were separately introgressed into near-isogenic recombinant lines (NIRLs), and observed across four years in one rain-fed locality of Central Italy (Kuzmanović et al., 2014; 2016). The range of increases in grain yield, biomass and grain number was 3-39%, depending on the recombinant type, season and experimental procedure (spaced plants in Kuzmanović et al., 2014, plot trials in Kuzmanović et al., 2016). In addition, and in contrast to the bread wheat (T4) studies, characterization of the three durum wheat-*Th. ponticum* recombinants comprised more traits, including detailed phenology, spike fertility and flag leaf attributes, and revealed 19 enhanced traits in association with the presence of specific 7AgL portions. This enabled a structural-functional dissection of the 7AgL

chromatin incorporated onto the durum 7AL, with consequent assignment of yield-contributing genes (previously associated to the entire T4 segment) to defined 7AgL sub-regions (Kuzmanović et al., 2014, 2016). The increase of several yield-related traits was recorded in each of the three durum wheat-*Th. ponticum* NIRLs. However, the one carrying the 28%-long 7AgL segment was identified as the best performing line, with a high number of yield-related traits (tiller/spike number, flag leaf dimensions and chlorophyll content, grain yield, biomass, duration of stem elongation phase) being evidently enhanced by genetic factor(s) located within the 23-28% chromosomal stretch of its 7AgL segment.

With no information from other environments on the expression of 7AgL and its effects on yield in durum wheat, the aim of the present work was to assess the yield performance of the same three durum wheat-*Th. ponticum* recombinants across an array of rain-fed environments located in three different countries, and to evaluate possible environment/segment-specific associations with final yield and individual yield contributing traits, in view of using these recombinants across environments or in site-directed breeding programs.

## Materials and methods

125 Plant materials

110

111

112

113

114

115

116

117

118

119

120

121

122

123

- Three durum wheat-*Th. ponticum* NIRLs, named R5-2-10, R112-4 and R23-1 (hereafter referred to
- as R5, R112 and R23, respectively), developed in the background of cv. Simeto by repeated
- backcrossing (BC) (Ceoloni et al., 2005), were used across four years and three countries. Simeto
- 129 (pedigree: selection from Capeiti 8 x Valnova) is a variety released in 1988, well adapted to the Italian
- growing conditions. The NIRLs have portions of *Th. ponticum* 7AgL chromosome arm replacing
- 131 23%, 28% and 40% of their distal 7AL arm, respectively, and all three lines include the
- 132 Lr19+Yp+Sr25 genes in the sub-telomeric region. Similarly to the plant material described in
- Kuzmanović et al. (2016), each of the genotypes analysed, corresponding here to BC<sub>5</sub>F<sub>5-9</sub> (R5 and
- R112) and BC<sub>4</sub>F<sub>5-9</sub> (R23) progenies, was represented by either being a homozygous carrier ("+") or
- non-carrier ("-") of the given 7AgL segment. Each "+" and "-" NIRL included two families
- originating from sister lines.
- 137 Field experiments
- A total of nine rain-fed field trials were carried out over four years and four locations where durum
- is typically cultivated (two in Italy, one in Morocco and in one Australia) and used for the multi-

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

environment yield assessment. Details on all trials are reported in Table 1. Years and locations were combined and hereafter referred to as environments, with a specific acronym assigned in Table 1. Two of the nine trials have been described previously in Kuzmanović et al. (2016) (VT12 and VT13), from where a subset of traits was considered in the present analysis. In ten environments, all three NIRLs with respective controls were used, while in AUS14, only R5 and R112 were analysed. Sowing densities applied were those commonly used in each of the experimental locations (Table 1). In all field experiments, complete randomized block designs with three replicates for each sister line was used, resulting in a total of 24 plots in AUS14 and 36 plots in the other eight environments (2 per each sister line, i.e. 6 per each +/- NIRL). Meteorological data during the growing seasons for daily temperatures (minimum, mean and maximum) and rainfall (Table 2) were retrieved from meteorological stations at experimental sites, except for MOR14, for which the data were downloaded from NASA's (National Aeronautics and Space Administration, USA) site for Prediction of Worldwide Energy Resource (http://power.larc.nasa.gov). All trials were managed according to standard local practices including fertilization, weed, pest and disease control, in order to avoid, in particular, leaf rust spreading on Lr19 non-carrier plants (- NIRLs), hence to eliminate the indirect yield-contributing factor of *Lr19*-carriers (+ NIRLs).

Environment acronym	Location	Latitude	Longitude	Altitude (m)	Season	Total rainfall (mm)	Mean temperature at heading (C°)	NIRLs tested	Sowing date	Crop cycle length (days)	Sowing density (seed/m²)	Plot size (m²)
VT12	Experimental farm of the University of Tuscia, Viterbo (Central Italy)	42° 25' N	12° 4' E	301	2011/12	248	13.6	R5, R112, R23	17/11/2011	226	350	2.3
VT13	"	"	"	"	2012/13	534	17.2	"	20/12/2012	192	350	2.3
VT14	и	"	"	"	2013/14	676	12.7	"	29/11/2013	213	350	4.5
VT15	п	"	"	"	2014/15	337	16.5	"	15/12/2014	197	350	4.5
BO14	Experimental farm of the University	44° 33' N	11° 24' E	33	2013/14	598	13.1	"	07/11/2013	236	350	4.2
MOR14	of Bologna, Cadriano (North Italy) Marchouch (Morocco)	33° 36' N	-6° 43' W	440	2013/14	229	13.0	"	03/12/2013	189	300	6
MOR15	п	"	"	"	2014/15	349	14.1	"	18/11/2014	204	300	6
AUS13	Waite campus, University of Adelaide, Urrbrae (South Australia)	34° 58' S	138° 38' E	48	2013	305	16.9	"	19/06/2013	179	150	1.6
AUS14	"	"	"	"	2014	296	14.3	R5, R112	15/05/2014	193	150	5

Table 1. Description of locations and field experiments analysed in this study (NIRL, Near Isogenic Recombinant Line).

Environment	Sowing to hea	ding			Heading to harvest						
	Rainfall (mm)	T <sub>min</sub> (°C)	T <sub>mean</sub> (°C)	$T_{max}$ (°C)	Rainfall (mm)	T <sub>min</sub> (°C)	T <sub>mean</sub> (°C)	T <sub>max</sub> (°C)			
VT12	155	2.7	7.9	13.8	93	11.4	18.6	25.8			
VT13	413	3.8	8.4	13.7	121	11.4	17.3	23.7			
VT14	486	4.1	9.0	14.3	190	11.3	17.5	24.0			
VT15	312	4.0	8.4	13.5	25	12.0	18.8	25.8			
BO14	422	4.0	8.5	12.9	176	13.3	19.5	25.8			
MOR14	180	5.8	10.7	17.5	49	11.7	18.6	27.2			
MOR15	342	6.6	10.9	16.4	7	13.2	19.5	27.2			
AUS13	266	10.0	14.2	18.4	39	12.7	19.6	24.3			
AUS14	260	9.1	13.3	17.5	36	12.8	18.9	25.1			

**Table 2.** Weather conditions for growing seasons in nine environments analysed, as retrieved from meteorological stations at experimental sites, or in the case of MOR14 only, downloaded from NASA's site for Prediction of Worldwide Energy Resource (<a href="http://power.larc.nasa.gov">http://power.larc.nasa.gov</a>). Environment acronyms are as described in Table 1.

## *Measurements of yield-related traits*

157

158

159

160

161

162

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

A list of traits and details on environments in which the materials were analysed, as well as on the sample type and number for each replicated plot, are reported in Table 3. Measurements of all traits were performed as described in Kuzmanović et al. (2016) with some modifications. HD was considered as number of days from sowing to heading. Samples of culms and flag leaves were randomly chosen within each plot. All dry weights of culms, spikes, chaff and total aboveground biomass were recorded after 48 h oven drying at 65°C. SIA was calculated as SDWA/TDWA ratio, SFI as the ratio between GNS and CHAFF at maturity (Isidro et al., 2011). SFI is a spike fertility parameter positively correlated with fruiting efficiency (Abbate et al., 2013; Martino et al., 2015), calculated as the ratio between GNS and SDWA (e.g., Slafer et al., 2015; Terrile et al., 2017). FLA was determined using the formula  $FLW \times FLL \times 0.75$  (Dodig et al., 2010). Chlorophyll content was measured by using a hand-held meter SPAD 502 (Konica-Minolta, Japan) at medium milk (Zadoks 75; Zadoks et al., 1974), late milk (Zadoks 77) and very late milk (Zadoks 79) developmental stages. HI was determined as GYM2/BM2 in MOR14, MOR15, AUS13 and AUS14, where GYM2 and BM2 were obtained directly from the total harvested area. In other environments, HI was determined on 25-culm samples plot<sup>-1</sup>, and at harvest used for BM2 estimation, once the total plot area was trashed and weighed (=GYM2/HI). TGW was obtained from weighing two 100-seed samples plot<sup>-1</sup> and then used to evaluate GNM2 (= $GYM2 \times 1000/TGW$ ).

Trait	Acronym	Environ	Environment									No. sample
		VT12	VT13	VT14	VT15	BO14	MOR14	MOR15	AUS13	AUS14	type/plot	plot <sup>-1</sup>
1 Harvest index	HI	X	X	X	X	X	X	X	X	X	TH/25C	1
2 Spike No. m <sup>-2</sup>	SNM2	X	X	X	X	X	X	X	X	X	TH	1
3 Biomass m <sup>-2</sup> (g)	BM2	X	X	X	X	X	X	X	X	X	TH	1
4 Grain yield m <sup>-2</sup> (g)	GYM2	X	X	X	X	X	X	X	X	X	TH	1
5 Grain No. m <sup>-2</sup>	GNM2	X	X	X	X	X	X	X	X	X	TH	1
6 1000 grain weight (g)	TGW	X	X	X	X	X	X	X	X	X	G	2
7 Grain yield spike-1 (g)	GYS	X	X	X	X	X	X	X	X	X	IC	6
8 Grain No. spike <sup>-1</sup>	GNS	X	X	X	X	X	X	X	X	X	IC	6
9 Grain No. spikelet <sup>-1</sup>	GNSP	X	X	X	X	X	X	X	X	X	IC	6
10 Spikelet No. spike-1	SPN	X	X	X	X	X	X	X	X	X	IC	6
11 Plant height (cm)	PH	X	X	X	X	X	X	X	X	X	IC	10
12 Days to heading	HD	X	X	X	X	X	X	X	X	X	day count	1
13 Spike dry weight at anthesis (g)	SDWA	X	X	X	X	X			X	X	IC	6
14 Tiller dry weight at anthesis (g)	TDWA	X	X	X	X	X			X	X	IC	6
15 Spike index at anthesis	SIA	X	X	X	X	X			X	X	IC	6
16 Spike dry weight at harvest (g)	SDW	X	X	X	X	X			X	X	IC	6
17 Tiller dry weight at harvest (g)	TDW	X	X	X	X	X			X	X	IC	6
18 Chaff dry weight at harvest (g)	CHAFF	X	X	X	X	X			X	X	IC	6
19 Spike length (cm)	SL	X	X	X	X	X			X	X	IC	6
20 Spike fertility index	SFI	X	X	X	X	X			X	X	IC	6
21 Flag leaf width (cm)	FLW	X	X	X	X	X			X	X	FL	10
22 Flag leaf length (cm)	FLL	X	X	X	X	X			X	X	FL	10
23 Flag leaf area (cm²)	FLA	X	X	X	X	X			X	X	FL	10
24 Chlorophyll content at Zadoks 75	CHLZ75	X	X	X	X	X				X	FL	10
25 Chlorophyll content at Zadoks 77	CHLZ77	X	X	X	X	X			X		FL	10
26 Chlorophyll content at Zadoks 79	CHLZ79									X	FL	10

Table 3. Traits and sample details assessed across nine environments analysed in this study [TH, total plot harvest; 25C, 25 culms; G, grain sample (1 or 2); IC, individual culms (5-10); FL, individual flag leaves (5-10)]. Environment acronyms are as described in Table 1.

## Statistical analyses

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

All analyses were performed by SYSTAT12 software (Systat Software Incorporated, San Jose, CA, USA). To investigate the effects of genetic and environmental factors and interactions between them on recorded traits, an analysis of variance (ANOVA) was performed, applying to datasets a general linear model (GLM) as a mixed effect model. Three such models were employed (= GLM1-3), depending on the data subset taken into consideration, as not all of the 25 traits and not all of the three NIRLs were analysed in all nine environments. In GLM1 and GLM2, datasets comprised environments where all three NIRLs were tested, while in GLM3, only the AUS14 dataset was analysed. GLM1 included traits No. 1-12 listed in Table 3, GLM2 traits No. 1-25, while GLM3 comprised traits No. 1-24 and 26. GLM1 was applied for the analysis of the overall effect of presence/absence of alien segments on main yield-related traits across environments while GLM2 and GLM3 were applied for the analysis of individual 7AgL segment effects. Each variable (i.e. trait measured) was entered as a 'dependent' factor against 'independent' factors. The latter were: genotype background (G), i.e. background genetic information from the recurrent variety, environment (E), presence/absence of the 7AgL segment [7AgL alone in GLM1 or 7AgL(G), i.e. nested in the background, in other GLMs], and replicate [R alone in GLM3 and environment-nested, R(E), in other GLMs]. The latter factor was used in the models as the error. First order  $[E \times G; R(E); E \times 7AgL; 7AgL(G)]$ , and second order  $[E \times G; R(E); E \times 7AgL; 7AgL(G)]$ 7AgL(G)] interactions between the above factors were analysed as well. In all analyses three levels of significance were considered, corresponding to P < 0.05, P < 0.01 and P < 0.001. When significant factors and/or interactions between them (F values) were observed, a pairwise analysis was carried out by the Tukey Honestly-Significant-Difference test at the 0.95 confidence level.

A correlation matrix was built for a subset of traits recorded in all environments. Each pair of variables was correlated by calculating Pearson's correlation coefficients (r value), while the significance levels were obtained using the Bonferroni method. Simple linear regression analysis was carried out by applying the least squares method for data fitting a 0.95 confidence level.

In order to analyse any genotype-specific response for grain yield (GYM2) across environments, the environmental index was calculated (= average value of all participating genotypes) in each environment and combined with the environmental mean of each genotype in a simple linear regression (b coefficient statistics). GYM2 values were transformed in logarithmic to attain linearity and homogeneity of errors (Finlay and Wilkinson 1963) and to observe the relative ("intrinsic") variability of interest (Becker and Leon, 1988).

#### Results

215

216

217

218

219

220

221

222

223

224

225

226

227

228

229

230

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

**Environments** 

The field trials described here cover a wide range of environmental conditions, as usually observed in areas of durum wheat cultivation. Overall, environments were comparable for mean temperatures (Table 2), with a little more variation between them in the period from sowing to heading (6.3°C) than from heading to maturity (2.3°C). By contrast, the environments were considerably different for the rainfall amount received (Table 1) and its distribution during the growing season (Table 2). Large differences in rainfall input were recorded particularly from heading to maturity, being in the range from 7 to 190 mm. The two seasons in South Australia were virtually identical for the precipitation received, nearly approaching the average values of the agricultural southeast areas of the State, while being the warmest years on record (since 1910), particularly through the second half of the season (www.bom.gov.au). The two trials in Morocco were also characterized by very warm seasons, with average daily maximum temperatures from May until harvest, often exceeding the site's average daily maximum values (22-26°C) by up to 16°C (www.weatherspark.com). Although less abundant, precipitation events were better distributed in MOR14 than in MOR15 (Table 2). As for the Italian trial sites, season 2012 was more typical and very favourable for durum wheat cultivation when compared to the 2013 to 2015 seasons. The latter three seasons were characterized by exceptional (and numerous) precipitation events and an overall increase in temperature during the entire crop cycle (www.informatoreagrario.it). VT12 had rainfall distribution and temperature trends that were conducive for good crop growth during the entire life cycle (Table 2; Kuzmanović et al., 2016). Still, in VT12, drought stress was present with precipitation amounts significantly below the site's mean, higher maximum temperatures during the second part of the and growth cycle (www.informatoreagrario.it). Conversely, seasons 2013 to 2015 in Italy had unusually rainy and mild winters, with a full soil moisture profile. Particularly heavy and prolonged rain periods prior to sowing and during the grain filling period occurred in VT13, VT14 and BO14, while VT15 resulted in very dry and hot conditions from heading onwards with respect to the former three environments and to the site's mean values.

## Yield response across environments

According to the observed highly significant R<sup>2</sup> values of linear regression (Fig. 1a), all NIRLs (both + and –) positively responded to better thermo-pluviometric patterns and higher environmental indices across environments. The b coefficient values around 1 for R5 and R112 genotypes were indicative of

an average yield stability (Finlay and Wilkinson, 1963), and the consistently higher grain yield with respect to the site's mean indicated their general adaptability. In contrast, the R23 NIRL pair, despite b coefficient values around 1 (i.e. average stability indicator), had poorer grain yield than the site's mean in all environments, probably due to the lower adaptability of the genetic background of its representative families, less isogenic compared to the recurrent cv. Simeto parent than the two other NIRLs (see § 2.1).

With an average grain yield across environments between 1.03 and 7.05 t/ha (Fig. 1b), environments were arbitrarily classified as: low-yielding (grain yield lower than 2 t/ha), medium-yielding (grain yield between 2 and 5 t/ha), and high-yielding (grain yield higher than 5 t/ha). The productivity of R5 and R112 NIRLs was typically higher than the environmental means, as shown from their yield gains ranging 3-30% and 2-27%, respectively. Yield of R23 NIRLs (both "+" and "-") was always under the site's mean (-1 to -47%). Yet, in four out of eight environments (all but AUS14, Table 1), the presence of the R23 7AgL segment evidently had a positive effect, reducing the background-dependent yield disadvantage (Fig. 1b). The two more productive 7AgL-carrier lines (R5+ and R112+), displayed yield gains with respect to their control NIRLs in 5 and 3, respectively, out of the nine environments. The highest gain of R5+ (and of all 7AgL+ lines) was observed in low-yielding AUS13 (+30%), a similar value to that recorded in medium-yielding MOR15 (+26%). In the same MOR15, R112+ showed the highest gain (+21%). By contrast, a notable yield penalty was observed for R112+ in low-yielding AUS14 (-18%), and for R23+ in medium-yielding MOR15 (-47%).

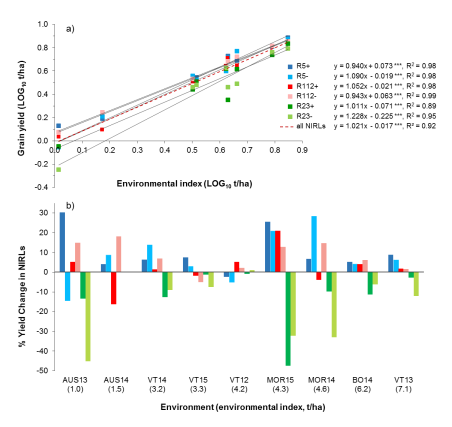


Fig. 1. Regression lines showing the relationship of individual grain yields of the six NIRLs and nine

environments analysed (a), and the percentage yield change of the six NIRLs with respect to the site's

mean (b). Environment acronyms are as described in Table 1.

7AgL-associated effects on yield and yield-related traits

267

268

269

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

Across all eight environments where all three NIRLs were tested (all but AUS14, Table 1), when data

from the three "+" recombinant or three "-" control lines were pooled, yield parameters turned out to

be increased to a variable extent in 7AgL+ vs. 7AgL- genotypes (Table 4 and Supplementary Table

1). The most significant positive 7AgL effects were observed for SNM2 (+13%), BM2 (+9%), GNM2

(+11%), and GNSP (+8%) accompanied though by a significant decrease in TGW and GYS. However,

HI of 7AgL+ lines remained unchanged (Table 4). Although the environment factor alone (E,

Supplementary Table 1) was highly significant for all traits, confirming that the environments tested

were different, the association between E and 7AgL turned out to be not significant for any of the traits

(as observed from the Tukey test for E × 7AgL, data not shown). ANOVA performed for each of the

nine environments separately, in view of highlighting the 7AgL effects in the various contrasting

conditions, revealed specific associations between positive 7AgL-linked effects and sites (Table 4).

SNM2, GNM2 and GNS were significantly increased in 7AgL+ vs. 7AgL- NIRLs in several

environments, particularly in the lowest- (AUS13) and highest-yielding (BO14, VT13), as well as in

the medium-yielding VT12. AUS13 turned out to be the only environment in which GYM2 together

with BM2 were significantly increased (+18% and +14%, respectively) in 7AgL+ vs. 7AgL- NIRLs.

This may indicate potential for adaptation to stressed conditions conferred by the 7AgL introgressions.

Trait	NIRLs	Environm	ent								
		ALL	AUS13	AUS14	VT14	VT15	VT12	MOR15	MOR14	BO14	VT13
HI	7AgL+	0.359	0.234	0.375	0.424	0.453	0.406	0.262	0.297	0.442	0.482
	7AgL-	0.360	0.224	0.420	0.410	0.466	0.412	0.289	0.306	0.447	0.483
	<i>P</i> -value	0.516	0.420	0.056	0.351	0.578	0.484	0.533	0.666	0.790	0.943
	7AgL effect (%)		4	-11	3	-3	-1	-9	-3	-1	0
SNM2	7AgL+	262.6	121.3	54.9	188.6	179.8	284.9	371.8	338.5	372.4	370.5
	7AgL-	233.3	91.6	52.9	187.0	164.9	259.2	356.9	354.0	343.0	342.2
	<i>P</i> -value	0.009**	0.000***	0.865	0.880	0.265	0.036*	0.684	0.643	0.041*	0.028*
	7AgL effect (%)	13	32	4	1	9	10	4	-4	9	8
BM2	7AgL+	1064.3	474.1	376.3	726.3	739.1	1041.8	1681.3	1570.7	1397.0	1512.2
	7AgL-	979.6	416.2	402.5	794.3	679.4	1012.5	1487.5	1543.0	1399.6	1443.2
	<i>P</i> -value	0.018*	0.001**	0.583	0.218	0.254	0.471	0.065	0.636	0.973	0.120
	7AgL effect (%)	9	14	-7	-9	9	3	13	2	0	5
GYM2	7AgL+	384.8	110.9	136.8	311.1	332.6	421.5	424.9	467.0	612.2	721.2
	7AgL-	372.6	94.3	169.0	330.2	316.8	416.4	428.3	475.4	625.4	688.6
	<i>P</i> -value	0.782	0.029*	0.115	0.433	0.304	0.752	0.959	0.844	0.685	0.281
	7AgL effect (%)	3	18	-19	-6	5	1	-1	-2	-2	5
GNM2	7AgL+	7871.2	1132.5	3058.1	5681.6	5703.0	15589.9	9349.4	10297.3	10234.4	12297.9
	7AgL-	7100.5	813.1	3323.2	5544.7	5101.9	13635.0	8488.2	9188.1	9526.1	10425.8
	<i>P</i> -value	0.000***	0.000***		0.903	0.172	0.038*	0.353	0.110	0.172	0.001**
	7AgL effect (%)	11	39	-8	2	12	14	10	12	7	18
TGW	7AgL+	50.8	34.8	46.6	55.4	59.2	60.5	46.4	45.6	60.1	60.3
	7AgL-	55.3	38.1	51.0	59.9	62.3	66.8	49.7	51.1	65.9	66.3
	<i>P</i> -value	0.000***	0.006**	0.148	0.097	0.417	0.032*	0.274	0.048*	0.021*	0.005**
	7AgL effect (%)	-8	-9	-9	-8	-5	-9	-7	-11	-9	-9
GYS	7AgL+	2.3	1.0	2.7	3.1	2.9	2.6	3.1	2.2	2.3	2.9
	7AgL-	2.4	1.1	3.2	3.0	3.2	2.8	3.5	2.3	2.6	3.0
	<i>P</i> -value	0.000***	0.168	0.022*	0.595	0.130	0.041*	0.071	0.368	0.124	0.269
	7AgL effect (%)	-4	-8	-18	3	-9	-6	-13	-6	-11	-4
GNS	7AgL+	43.1	24.6	59.4	54.8	49.8	43.4	52.6	46.9	42.3	47.7
	7AgL-	39.7	21.8	66.3	47.7	51.3	41.4	56.6	44.9	43.9	43.7
	<i>P</i> -value	0.057	0.028*	0.023*	0.004**	0.460	0.280	0.142	0.358	0.532	0.006**
	7AgL effect (%)	9	13	-10	15	-3	5	-7	5	-3	9
GNSP	7AgL+	2.4	1.2	2.4	3.0	2.9	2.5	2.9	2.4	2.2	3.0
	7AgL-	2.2	1.1	2.2	2.6	2.9	2.4	2.8	2.3	2.3	2.8
	<i>P</i> -value	0.005**	0.078	0.072	0.007**	0.814	0.185	0.462	0.547	0.594	0.021*
	7AgL effect (%)	8	11	7	13	-1	4	4	3	-3	7
SPN	7AgL+	18.7	20.9	18.8	18.5	17.2	17.2	18.4	19.7	19.2	15.9
	7AgL-	18.7	20.9	18.9	18.1	17.7	17.1	20.7	19.7	19.4	15.7
	P-value	0.145	0.910	0.904	0.512	0.233	0.703	0.044*	0.965	0.613	0.485
	7AgL effect (%)		0	0	2	-3	1	-11	0	-1	2
PH	7AgL+	82.4	74.7	82.6	91.8	75.1	87.1	85.0	84.7	86.2	82.4
	7AgL-	81.4	72.3	81.7	90.9	76.2	87.0	89.4	83.9	86.4	81.3
	<i>P</i> -value	0.965	0.214	0.490	0.862	0.777	0.984	0.357	0.776	0.960	0.773
	7AgL effect (%)		3	1	1	-1	0	-5	1	0	1
HD	7AgL+	130.0	99.4	129.5	142.2	139.3	153.6	131.7	105.8	166.4	132.0
	7AgL-	127.0	95.3	126.6	141.0	138.7	153.2	130.4	105.7	165.6	130.6
	P-value	0.011*	0.000***		0.499	0.334	0.575	0.609	0.904	0.562	0.011*
	7AgL effect (%)	2	4	2	1	0	0	1	0	1	1

**Table 4.** Main yield-related traits unit area<sup>-1</sup> and spike<sup>-1</sup> of 7AgL-carrier vs. 7AgL-non carrier durum wheat-*Th. ponticum* NIRLs across environments (HI, harvest index; SNM2, spike number m<sup>-2</sup>; BM2, biomass m<sup>-2</sup>; GYM2, grain yield m<sup>-2</sup>; GNM2, grain number m<sup>-2</sup>; TGW, thousand grain weight; GYS, grain yield spike<sup>-1</sup>; GNS, grain number spike<sup>-1</sup>; GNSP, grain No. spikelet<sup>-1</sup>; SPN, spikelet No. spike<sup>-1</sup>; PH, plant height; HD, heading date). In eight out of the nine environments, all three recombinants and their respective controls were analysed (ALL); in AUS14, only R5 and R112 NIRLs were included. Positive and negative 7AgL effects are highlighted in *green* and *red*, respectively. \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001. Environments are ordered from the lowest- to the highest- yielding one, and their acronyms are as described in Table 1.

The more extensive and detailed GLM2 model (Supplementary Table 2), used to examine individual effects of each of the three 7AgL segments (Table 5), showed that E alone was highly significant for all traits, but also the majority of interactions including it. Across all environments, the influence of the individual 7AgL segments on the various traits [7AgL(G), Supplementary Table 2] was different, as revealed by analysis of the three "+" vs. "-" NIRL pairs. Taken the eight environments where all three recombinants were analysed (all but AUS14, Table 1) as a whole, the R5+ recombinant did not show any significant trait change due to its 7AgL segment (Table 5), although in the AUS13 environment alone several significant differences emerged (see § 3.3.1). Also the comparison between R112+ and its R112- control across all environments showed only minor differences for HI (-3%), TGW (-4%), HD (+4%) and FLL (-9%) (Table 5).

On the other hand, R23+ was the recombinant for which the highest number of significant differences with respect to its R23- control were observed irrespective of the E factor (Table 5). While a number of yield-related parameters were depressed by the presence of the 40%-long 7AgL segment [TGW, GYS, SPN, PH, dry weight at anthesis and maturity (SDWA, TDWA, SDW, TDW, CHAFF), flag leaf dimensions (FLL, FLA)], the same 7AgL portion was clearly associated with significant increases of several other parameters directly contributing to grain yield, i.e. SNM2 (+13%), GNM2 (+15%), GNS (+10%), GNSP (+10%) and SFI (+46%) (Table 5).

Trait	R5+			R5-			R112+			R112-			R23+			R23-		
	Mean	SE		Mean	SE		Mean	SE		Mean	SE		Mean	SE		Mean	SE	
HI	0.39	0.01	a	0.38	0.02	a	0.37	0.01	b	0.38	0.02	a	0.31	0.02	c	0.32	0.02	с
SNM2	251.1	16.0	AB	228.8	19.6	AB	263.0	15.9	AB	230.4	18.9	AB	273.5	15.8	A	241.8	18.6	В
BM2	1041.3	62.1	ns	932.1	76.4	ns	1067.1	73.7	ns	938.5	73.8	ns	1084.1	65.8	ns	1079.1	76.9	ns
GYM2	418.3	29.4	ns	382.6	39.7	ns	396.0	28.7	ns	373.0	35.9	ns	340.9	27.1	ns	360.5	34.3	ns
GNM2	7376.0	591.9	В	6929.3	775.9	В	7660.7	656.7	В	6956.7	807.2	В	8576.8	803.4	A	7458.8	849.2	C
TGW	57.5	1.5	A	56.0	2.1	A	53.2	1.7	В	55.6	2.0	A	41.6	1.1	C	54.2	2.0	В
GYS	2.6	0.1	A	2.3	0.2	A	2.4	0.1	A	2.4	0.2	A	2.0	0.1	В	2.5	0.2	A
GNS	42.5	1.4	ab	38.8	1.9	b	42.5	1.6	ab	40.1	2.0	ab	44.1	1.9	a	40.2	2.6	b
GNSP	2.4	0.1	A	2.2	0.1	A	2.3	0.1	A	2.3	0.1	A	2.3	0.1	A	2.1	0.1	В
SPN	18.2	0.3	C	18.3	0.4	C	18.6	0.3	BC	18.1	0.3	C	19.2	0.3	В	20.0	0.5	A
PH	77.9	0.9	c	75.8	1.0	cd	76.1	0.8	d	73.2	0.9	d	93.2	1.2	b	97.8	1.4	a
HD	127.2	3.1	C	123.6	4.0	C	129.2	3.2	В	123.7	3.9	C	133.6	3.1	A	135.0	3.7	A
SDWA	0.76	0.03	AB	0.79	0.04	AB	0.74	0.03	AB	0.78	0.03	A	0.55	0.02	C	0.66	0.02	В
<b>TDWA</b>	4.46	0.14	A	4.82	0.20	A	4.39	0.14	A	4.51	0.15	A	3.83	0.14	В	4.65	0.22	A
SIA	0.17	0.01	ns	0.16	0.01	ns	0.17	0.00	ns	0.17	0.00	ns	0.15	0.00	ns	0.15	0.01	ns
SDW	3.26	0.12	A	3.17	0.12	A	3.20	0.12	A	3.17	0.12	A	2.57	0.11	В	3.02	0.16	Α
TDW	5.26	0.21	b	4.99	0.23	b	5.09	0.23	b	4.92	0.25	b	4.75	0.20	c	6.19	0.18	a
CHAFF	0.87	0.04	ab	1.02	0.10	ab	0.94	0.05	a	0.93	0.05	ab	0.74	0.04	b	0.90	0.04	a
SL	6.5	0.1	ns	6.7	0.2	ns	6.4	0.1	ns	6.3	0.1	ns	7.2	0.1	ns	7.3	0.2	ns
SFI	50.5	2.7	В	48.1	4.3	В	51.9	3.1	В	48.3	4.0	В	68.5	4.6	A	46.9	3.9	В
FLW	1.8	0.03	A	1.7	0.04	AB	1.9	0.03	AB	1.8	0.03	В	1.7	0.02	В	1.7	0.04	C
FLL	20.3	0.59	ab	21.7	0.90	ab	19.9	0.59	bc	21.8	0.90	a	18.8	0.57	c	20.9	0.79	a
FLA	31.1	1.67	A	30.0	1.80	A	31.1	1.70	A	32.0	1.80	A	26.3	1.39	В	30.0	2.11	A
CHLZ75	49.8	1.6	ns	53.1	1.6	ns	49.5	1.5	ns	51.0	2.0	ns	41.8	1.3	ns	45.1	1.4	ns
CHLZ77	42.8	1.2	A	43.1	1.5	A	44.4	1.3	A	42.6	1.5	AB	39.3	1.2	В	43.4	1.4	AB

**Table 5**. Mean values and standard errors (SE) of yield-related traits of the three durum wheat-*Th. ponticum* NIRLs (+, 7AgL carriers; –, 7AgL non-carriers) across eight environments where all three recombinants and their respective controls were analysed (HI, harvest index; SNM2, spike number  $m^{-2}$ ; BM2, biomass  $m^{-2}$ ; GYM2, grain yield  $m^{-2}$ ; GNM2, grain number  $m^{-2}$ ; TGW, thousand grain weight; GYS, grain yield spike<sup>-1</sup>; GNS, grain number spike<sup>-1</sup>; GNSP, grain number spike<sup>-1</sup>; PH, plant height; HD, heading date; SDWA, spike dry weight at anthesis; TDWA, tiller dry weight at anthesis; SIA, spike index at anthesis; SDW, spike dry weight at maturity; TDW, tiller dry weight at maturity; CHAFF, chaff dry weight; SL, spike length; SFI, spike fertility index; FLW, flag leaf width; FLL, flag leaf length; FLA, flag leaf area; CHLZ75, flag leaf chlorophyll content at Zadoks 75; CHLZ77, flag leaf chlorophyll content at Zadoks 77). Letters in each row correspond to the ranking of the Tukey test at P < 0.01 (capital) and P < 0.05 (lower case) levels.

 $7AgL \times E$  interaction

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

340

341

342

343

344

345

346

347

348

349

350

351

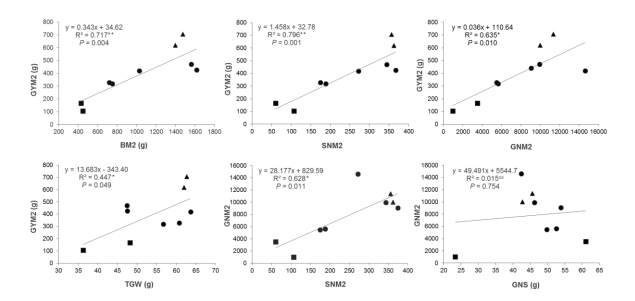
352

353

Individual alien segment effects were significant to a variable extent in particular environments, and for specific traits, as revealed by the Tukey test (data not shown) for significant  $E \times 7AgL(G)$ interactions (GLM2, Supplementary Tables 2 and 3). From this analysis across environments, significant differences between R5+ and R5- control were observed in the low-yielding AUS13 environment. These included increase of HI (+27%) and FLW (+15%), as well as decrease of FLL (-13%) and CHAFF (-38%). Considering the AUS13 dataset alone, additional significant differences in favour of R5+ could be highlighted, namely a 52% increase in grain yield (GYM2), paralleled by significantly higher values of GYS (+30%), GNM2 (+45%), GNS (+21%), SFI (+77%) and BM2 (+21%) (see Supplementary Table 3). As for the R112+ recombinant, the multi-environment dataset showed TGW (-13%), FLL (-15%) and FLW (-17%) to be significantly depressed compared to the R112- control in AUS13, while biomass production. (BM2) to be boosted in MOR15 (+24%) Data from AUS14, in which R5 and R112 NIRLs only were grown, revealed that presence of both 7AgL segments was not particularly advantageous for the main yield traits (traits No. 1-8 from Table 3; Supplementary Table 3). The GLM3 model identified only a few significant differences associated with the presence of the 7AgL segment ([7Ag(G)]; Supplementary Table 4): incremental, for FLW of both recombinants (+37% for R5+ and +25% for R112+), FLA of R5+ (+42%) and HD of R112+ (+6%); detrimental, for GNSP of R112+ (-14%) and for CHL75 of R5+ (-7%) (Supplementary Table 3). On the other hand, the R23+ NIRL was highly responsive to different environments for several traits. The significant enhancing effect of its 7AgL segment for GNM2 and SFI was particularly pronounced in VT12-VT13 (+30% and +39%, respectively) and MOR14 (+56%) for the former trait, while in VT13-VT14 (+35% and +59%, respectively) and BO14 (+95%) for the latter (Supplementary Table 3). Some of the identified depressing effects of the 40%-long 7AgL segment were significantly associated with certain, even contrasting environments: TGW in VT12-15, BO14 and AUS13 (ranging -18 to -27%); GYS in MOR15 (-43%); dry weight at maturity (SDW, TDW) in BO14 (-27% and -28%, respectively); FLA in BO14 and VT14 (-22% and -33%, respectively). Similarly to the R5 case, analysis of the AUS13 dataset alone showed significant increases of GYM2 (+58%), GNM2 (over 3 fold), SNM2 (+128%), GNS (+49%), GNSP (+67%) and BM2 (+33%) to be also associated to R23+ vs. R23-, as well as significant reduction of TGW (-19%), GYS (-30%), TDWA (-19%) and TDW (-43%) (Supplementary Table 3).

## Correlations of yield traits

Scatter plots of regression analysis ( $R^2$ ) for the pairs of main yield traits are shown in Fig. 2, and coefficients of correlation (r) are reported in Table 6. For all environments, final grain yield per area (GYM2) was mostly dependent on SNM2, BM2 and GNM2 ( $R^2 = 0.80, 0.72, \text{ and } 0.64, \text{ respectively}$ ), while the contribution of TGW was lower ( $R^2 = 0.45$ ). Nevertheless, as observed from significant r coefficients, the involvement of given traits to GYM2 formation in each environment was different. In line with linear regression analysis, the strongest and the most transversal positive correlation of GYM2 was observed with grain number and biomass, shown by a significant r value in eight out of the total nine environments tested (58-93% for GNM2 and 61-95% for BM2). In three low- and medium-yielding environments, GYM2 was positively influenced also by SNM2, GYS and GNSP (r = 50-79%). TGW confirmed to be less important for final grain yield, the two parameters being significantly correlated in only two medium-yielding environments (r = 60-82%). Finally, GNM2, the key trait for wheat grain yield increases, as expected, was primarily influenced by SNM2 (r = 62-96%, significant in five environments), grain number spike<sup>-1</sup> (GNS) and spikelet<sup>-1</sup> (GNSP) (r = 66-96%, significant in three and four environments, respectively), particularly in low- and medium-to-low environments.



**Fig. 2**. Scatterplots of the means of grain yield and grain number  $m^{-2}$  vs. main grain yield components from the three durum wheat-*Th. ponticum* NIRLs evaluated across nine environments (GYM2, grain yield  $m^{-2}$ ; BM2, biomass  $m^{-2}$ ; SNM2, spike number  $m^{-2}$ ; GNM2, grain number  $m^{-2}$ ; TGW, thousand-grain weight; GNS, grain number spike<sup>-1</sup>). \*P < 0.05, \*\*P < 0.01, "ns" not significant. Low- (GY < 2 t ha<sup>-1</sup>), medium- (GY 2-5 t ha<sup>-1</sup>) and high- (GY > 5 t ha<sup>-1</sup>) yielding environments are represented by *square*, *circle* and *triangle* symbols, respectively.

377	Pairwise traits	Environme	ent			63       *       0.58       *       0.93       ***       0.72       **       0.63       *       0.56         03       0.02       0.82       ***       0.60       *       0.41       0.26         02       0.54       *       0.55       -0.29       -0.21       0.53         04       *       0.81       ***       -0.23       0.64       *       0.61       *       0.64         9       0.09       0.22       0.73       **       0.45       0.16         05       0.07       -0.14       0.49       0.37       0.23         06       0.24       0.04       0.73       **       0.47       0.55         0       0.01       -0.62       *       -0.56       -0.49       -0.45         49       -0.26       -0.75       **       -0.61       *       -0.46       -0.44				
		AUS13	AUS14	VT14	VT15	VT12	MOR15	MOR14	BO14	VT13
378	GYM2 - GNM2	0.85 ***	0.87 **	0.90 ***	0.63 *	0.58 *	0.93 ***	0.72 **	0.63 *	0.56
	GYM2 - TGW	0.35	0.36	0.26	0.03	0.02	0.82 ***	0.60 *	0.41	0.26
	GYM2 - SNM2	0.50 **	0.74	0.83 ***	0.22	0.54 *	0.55	-0.29	-0.21	0.53
379	GYM2 - BM2	0.68 ***	0.87 **	0.95 ***	0.64 *	0.81 ***	-0.23	0.64 *	0.61 *	0.64 **
	GYM2 - GYS	0.62 ***	0.17	0.73 **	0.19	0.09	0.22	0.73 **	0.45	0.16
	GYM2 - GNS	0.82 ***	-0.09	0.58	0.25	0.07	-0.14	0.49	0.37	0.23
380	GYM2 - GNSP	0.79 ***	0.19	0.65 *	0.26	0.24	0.04	0.73 **	0.47	0.55
	GYM2 - PH	-0.48 **	0.07	0.05	0.10	0.01	-0.62 *	-0.56	-0.49	-0.45
381	GYM2 - HD	-0.33	-0.38	-0.35	-0.49	-0.26	-0.75 **	-0.61 *	-0.46	-0.44
201	GNM2 - SNM2	0.81 ***	0.96 ***	0.76 ***	0.61	0.62 **	0.56	-0.07	-0.11	0.71 **
	GNM2 - GNS	0.76 ***	0.96 ***	0.70 **	0.35	0.66 **	-0.22	0.49	0.29	0.54
382	GNM2 - GNSP	0.75 ***	-0.20	0.70 **	0.13	0.75 ***	0.02	0.73 **	0.33	0.50

**Table 6**. Pearson's correlation coefficients between pairs of main yield-related traits involved in coarse and fine regulation of final grain yield across nine environments tested (GYM2, grain yield m<sup>-2</sup>; GNM2, grain number m<sup>-2</sup>; TGW, thousand grain weight; SNM2, spike number m<sup>-2</sup>; BM2, biomass m<sup>-2</sup>; GYS, grain yield spike<sup>-1</sup>; GNS, grain number spike<sup>-1</sup>; GNSP, grin number spikelet<sup>-1</sup>; PH, plant height; HD, heading date). Correlations are underlined using a heat colour map of green (positive correlations; dark green P < 0.001, medium green P < 0.01, light green P < 0.05) and red (negative correlations; medium red P < 0.01, light red P < 0.05) shades. Environment acronyms are as described in Table 1.

#### Discussion

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

412

413

414

415

416

417

418

419

420

To evaluate, for the first time, the effect of different environmental conditions on yield and yieldcontributing traits of three durum wheat-Th. ponticum recombinant lines, having 23%, 28% and 40% of 7AgL chromatin on their 7AL distal end, nine rain-fed field trials in four locations worldwide over four years were undertaken. The present analysis showed that the overall incidence of the 7AgL introgressions into the tetraploid wheat background on the trade-off between the various yield components, did not result, except for one environment (AUS14, see ahead), in major yield penalty, nor in any alteration of harvest index. In fact, taking all three 7AgL segments (all "+" vs. all "-" NIRLs comparison) and the nine environments together, an average, albeit not significant, 3% increase in grain yield m<sup>-2</sup> was detected for the +NIRLs (Table 4). Given the high heterogeneity among environments, this minor difference resulted from four incremental cases (ranging from +1% to 18%) and five others exhibiting a yield reduction (1-2% in three of them). The same analysis of the effects of 7AgL segments as a whole but on individual yield-contributing traits highlighted that yield performances associated to the alien introductions were mainly due to significant increases of biomass m<sup>-2</sup>, spike/tiller number m<sup>-2</sup>, related grain number m<sup>-2</sup> and spikelet<sup>-1</sup>, and to a general, significant decrease of grain weight (Table 4). This evidence confirms previous observations in one location only (Kuzmanović et al., 2014, 2016) and indicates stable expression of such traits across environments. The 7AgL case thus appears to support the consolidated evidence that in both bread and durum wheat, coarse regulation of yield is based primarily on grain number and biomass (Marti et al., 2016; Pedro et al., 2011; Slafer et al., 2014).

Grain yield of all NIRLs was generally higher in environments where more rainfall was recorded, particularly from heading onwards (e.g., Central and Northern Italy; Tables 1 and 2), which is in line with what observed for durum wheat grown under Mediterranean rain-fed conditions (Araus et al., 2003b). Similarly, previous studies indicate that bread wheat T4 derivatives, whose sizeable 7AgL segment includes those of the durum wheat recombinants described here, benefit from higher water availability (Singh et al., 1998; Monneveaux et al., 2003; Rosewarne et al., 2015). Nonetheless, the results from the environment-by-environment ANOVA (Table 4) revealed the potential of durum wheat 7AgL+ lines, taken as a whole, for yield increase also under heat and drought stress conditions, such as those of AUS13.

Considering the separate NIRL pairs across the range of environments, a number of 7AgL segment-associated effects emerged. R5+ exhibited an average 9% increase (albeit not significant) in grain yield (GYM2) vs. its control 7AgL— line (Table 5), as well as the highest grain yield gains with respect to the sites' mean in eight out of the nine trials (Fig. 1b). While grain yield increase of R5+

422

423

424

425

426

427

428

429

430

431

432

433

434

435

436

437

438

439

440

441

442

443

444

445

446

447

448

449

450

451

452

453

454

vs. R5– plants was slight in high- and medium-yielding environments (1-4% in VT12, VT13, VT15, BO14, MOR15), in low-yielding AUS13 it amounted to a significant 52%, paralleled by a significant increase in several other yield parameters (Supplementary Table 3). These figures, however, were not mimicked by the AUS14 environment, which was characterized by similar meteorological conditions to AUS13 (Tables 1 and 2) but a longer growth cycle (earlier sowing) and a remarkable increase in flag leaf size of R5+ vs. R5- plants (Supplementary Table 3). Flag leaf size is known to be positively correlated with photosynthetic activity and yield in wheat under drought (Foulkes et al., 2007; Habash et al., 2007; Quarrie et al., 2006). The longer AUS14 growing season likely favoured vegetative plant growth, as also evidenced by greater biomass at anthesis (SDWA and TDWA) besides that bigger flag leaf area (Supplementary Table 3). However, the latter was probably not advantageous for R5+ (and so for R112+, see ahead), likely leading to high transpiration and dehydration rates, both negatively affecting final yield (e.g., Izanloo et al., 2008; Tardieu 2005).

As to the R112+ recombinant, average values of all trials showed increased values vs. R112for spike (+14%) and grain number (+10%) m<sup>-2</sup>, as well as 6% higher grain yield (Table 5), in addition to a consistent tendency for wider flag leaf (significant in AUS14, VT12, VT13 and VT15, Supplementary Table 3) and higher chlorophyll content at late grain filling stages in several environments (Supplementary Table 3). These observations confirm that yield formation in R112+ depends mostly on tiller number development and potentially increased photosynthetic activity of leaves, both contributing to grain number formation (see also Kuzmanović et al., 2014, 2016). Tiller number is greatly influenced by the environment and determines wheat adaptive ability under rainfed conditions (Elhani et al., 2007; Zhang et al., 2010); thus, it is not surprising that higher grain yield of R112+ vs. control plants (+3-7%) was observed in only three of the experimental sites, where more favourable conditions for tillering were evidently met (VT12, VT15, MOR15, Supplementary Table 3). At high-yielding sites, R112+ produced essentially the same as the controls (BO14, VT13), while a major yield penalty was recorded in low-yielding AUS14 (-29%, Supplementary Table 3), concomitantly with a significant, environment-specific decrease in spike fertility (-14% of grains spikelet<sup>-1</sup>). Similarly to the R5+ case (see above), the significantly larger flag leaf size (+25% FLW), accompanied by higher chlorophyll content (+23%, though not significant) at late grain filling in the R112+ case, seemed to be at a disadvantage for R112+ grain yield in AUS14 (Supplementary Table 3). Capacity of a wheat plant to maintain flag leaf greenness generally enhances photoassimilation (Foulkes et al., 2007; Peremarti et al., 2014); yet, in environments such as South Australia, where high irradiance often comes along with drought stress, increased temperatures and high wind (Izanloo et al., 2008; Fleury et al., 2010), high chlorophyll content could provoke oxidative damage and thus reduce photosynthetic capacity and, ultimately, yield potential (Long et al., 2015; Parry et al., 2011;

456

457

458

459

460

461

462

463

464

465

466

467

468

469

470

471

472

473

474

475

476

477

478

479

480

481

482

483

484

485

486

487

488

Tricker et al., 2018; Zhu et al., 2010). Another possible reason for the reduced yield of R112+ in the Australian environment might reside in the interaction of its root system with the soil features at this site. Previous simulations determined that a shallow root system is more favourable in this type of environment (reviewed in Izanloo et al., 2008), which contrasts with the root system architecture (RSA) of R112+, found to be characterized by increased seminal root angle, total root length and root dry weight (Virili et al., 2015). Under drought conditions, a widened root angle and deeper roots were found to be crucial adaptive mechanisms associated to yield increases in rice (Uga et al., 2013; 2015; Ahmadi et al., 2014) and wheat (Lopes and Reynolds, 2010; Manchadi et al., 2006; Slack et al., 2018). However, the shallow, clay-limestone soils, typical of most Adelaide Plain's and of our testing site, might have hindered the potential of the R112+ root system, limiting its access to water and nutrients at deeper layers. In fact, it was reported that root impedance reduces leaf elongation and the number of tillers in wheat (Jin et al., 2015). Therefore, the R112+ recombinant looks to be unsuitable for dry environments with combined stress factors, such as heat, drought and hostile soil structure. Instead, to exploit at best its 7AgL-linked positive effects on tiller/spike number, flag leaf photosynthetic activity (see also Kuzmanović et al., 2014, 2016), and RSA characteristics (Virili et al., 2015), it could be profitably employed in breeding directed to environments with optimal thermo-pluviometric patterns and soil characteristics (as VT12, VT15 and MOR15).

Finally, yield performance of the R23+ NIRL represents an intriguing case. In fact, while the background genotype, common to the R23- NIRL, was probably responsible for generally lower adaptability and consequent productivity throughout environments compared to the other two NIRL pairs (see § 3.2, Figure 1 and Supplementary Table 3), specific, both positive and negative, effects linked to its 40%-long 7AgL segment were consistently expressed in the multi-environment analysis (Table 5). The most important incremental effects validated throughout all trials were on spike fertility index, grain and spike number m<sup>-2</sup> (Table 5 and Supplementary Table 3). Enhancement of the first two traits strongly supports the suggested existence of a large effect QTL for grain number within the 28-40% 7AgL portion specific to the R23+ introgression (Kuzmanović et al., 2014; 2016). Interestingly, a couple of important QTL for spike fertility traits (No. of spikelets and grains spikelet 1) were recently identified in the homoeologous 7AL region of durum wheat (Giunta et al., 2018). Furthermore, the results showed that R23+, similarly to R112+, and so in the shared 23-28% 7AgL segment, harbours a gene/QTL for tiller/spike number m<sup>-2</sup>, evidently a major factor of its yield potential, particularly in the dryer southern Australian (AUS13) and Moroccan (MOR14) environments (Supplementary Table 3; see also Kuzmanović et al., 2014; 2016). However, in contrast to the R112+ case, in which higher tiller number is likely to have primarily contributed to a more efficient plant source (i.e. leaf area, chlorophyll content), in R23+ the increased tiller number in

490

491

492

493

494

495

496

497

498

499

500

501

502

503

504

505

506

507

508509

510

511

512

513

514

515

516

517

518

519

520

521

522

combination with remarkably higher spike fertility had probably a more prominent effect on the genotype's sink (i.e. grain number m<sup>-2</sup>). Confirming a different yield formation from that hypothesized for R112+, flag-leaf related traits were instead significantly depressed in R23+. Moreover, the same genotype also had reduced biomass at anthesis and maturity when compared to its R23- control (about -18%, Table 5). Both traits appear to be highly correlated with grain number (e.g. Fischer, 2008; González et al., 2011; Slafer et al., 2015); hence, it was unexpected that R23+ could consistently sustain its large gain in grain number m<sup>-2</sup> and in spike fertility index (+15% and +46%, respectively) across eight environments (all but AUS14, where R23 was not tested; Table 5). Mechanisms involved in the control of fertile florets survival rate, rather than their absolute number, are likely to be responsible for this (Kuzmanović et al., 2016 and references therein). Recent work by Marti et al. (2016) established that growth and N partitioning in the two weeks before anthesis are key factors in determining differences in wheat yield performance, particularly for grain number. The same authors suggest that this growth period is not relevant for total biomass production at anthesis, but is crucial for determining the relative grain number. As the nature of this relationship is still uncertain (Slafer et al., 2015; Terrile et al., 2017 and references therein), it would be worthwhile investigating the modes of how assimilates are accumulated and translocated in R23+ during spike development just prior to anthesis. Guo et al. (2017) suggested a critical role of carbohydrate metabolism and phytohormones in regulating the floret primordia survival (FPS) and final grain number in wheat. Floret primordia formation (FPF) and their survival are known to be proportional to the availability of assimilates allocated for their development during spike growth before anthesis (Ferrante et al., 2013; González et al., 2011; Slafer et al., 2015). In R23+ plants, the number of fertile florets at anthesis (mirroring the FPF) showed to be similar to its R23- control (Kuzmanovic et al., 2016), while the number of seeds at maturity was enhanced, as a consequence of higher FPS (Table 5 and Supplementary Table 3; see also Kuzmanovic et al., 2014, 2016). Therefore, translocation of assimilates to grains is likely to be affected rather than their accumulation in spike tissues. Particularly noteworthy is the occurrence of increased spike fertility of R23+ vs. R23– not only in environments characterized by profuse rainfall throughout the crop cycle, such as VT14 and BO14, but also in AUS13 and MOR14, where these favourable conditions were not met (Tables 1 and 2; Supplementary Table 3). Despite the consistently lower thousand-grain weight of R23+ compared to R23- plants, higher grain number of R23+ was likely responsible for increments in grain yield (+7-58%) observed in four out of the eight environments (Table 5 and Supplementary Table 3). Due to the grain number vs. grain weight trade-off, increased fruiting efficiency is considered to be generally irrelevant for actual yield increases (Slafer et al., 2015). Nonetheless, this negative relationship may not be "constitutive", i.e. the factor(s) increasing fruiting efficiency may be independent of the size of florets, i.e. potential grains (Slafer et al. 2015; Ferrante et al. 2015). Also, resource partitioning to developing florets may be increased and mortality of distal florets reduced, thus ensuring yield gains (Slafer et al., 2015, and references therein). Similar mechanisms may apply to the R23+ case. Interestingly, the highest grain yield increases of R23+ were observed in dry environments of AUS13 and MOR14 (+58% and +34%, respectively), indicating that R23+ maintains its higher seed set even under drought and/or heat stress conditions. This finding is of great relevance, as grain number is recognized as the most susceptible yield component to drought (Richards et al., 2010)., Furthermore, spike fertility was recently shown to be the key trait contributing to yield performance of durum wheat grown under severe and protracted heat stress (Sall et al., 2018).

The possibility of fully unlocking the R23+ yield potential for practical exploitation remains, however, somewhat challenging, because of the linkage between the gene(s)/QTL for grain number and spike fertility, and a gene(s) causing segregation distortion (*Sd*) and possibly correlated depression of a number of morpho-physiological traits, including grain development (Ceoloni et al., 2014b; Kuzmanović et al., 2016). Nonetheless, this linkage drag could be broken, e.g. via induced mutations or further homoeologous recombination, or possibly countered by transferring the R23 7AgL segment to different recipient cultivars, particularly choosing those exhibiting high grain weight, the trait mostly depressed in R23+ vs. R23- across environments (Table 5). In this view, previous evidence that, similar to the bread wheat T4 case, the interaction of the R23 40%-long 7AgL segment with varying durum wheat genetic backgrounds results in improved transmission of the recombinant chromosome, with little or no effect on plant phenotype (Ceoloni et al., 2014b), is encouraging.

Apart from genotype-dependent "stabilization" that the R23 7AgL segment may require, all segments can be readily moved into more site-directed (R112) or across environment (mainly R5 and R23) breeding pipelines for exploitation of their Lr19+Sr25+Yp genes, as well as for additional traits contributing to yield increase and stability. Moreover, all three recombinants can be further enriched with other useful alien genes. Recently, several new recombinants in bread and durum wheat have been obtained by chromosome engineering, in which highly effective gene(s)/QTL for resistance to *Fusarium* head blight and crown rot, originating from *Thinopyrum* species, were pyramided onto the most telomeric portions of the 7AgL segments described here (Ceoloni et al., 2017b; Forte et al., 2014; Kuzmanović et al., 2017). Preliminary results have shown normal fertility of the new recombinants and even higher yields when compared to control plants. This evidence contributes to strengthen the validity of targeted exploitation of alien variation to enhance yield potential in wheat species (Ceoloni et al., 2014a; Mondal et al., 2016; Zaïm et al., 2017, and references therein).

## **Conclusions**

Analysis of the three 7AgL introgression lines into durum wheat across a range of variable environments has validated their potential as donors of yield-contributing traits, besides that of additional beneficial attributes originating from the wild *Th. ponticum* donor. Being developed in a cultivar best adapted to Italian growing conditions (Simeto), the three recombinants exhibited variable performances in stressed environments, like South Australia and Morocco. In general, under no significant water stress, 7AgL lines responded well, without significant yield losses, mainly through contribution of grain and spike number, and, to some extent, flag leaf size and function at late grain filling stages. Also the observed significant increases in grain yield and grain number for 7AgL+ vs. 7AgL- NIRLs in more heat and drought-stressed environments, indicate that the 7AgL yield-related gene/QTL content may also be beneficial under adverse growing conditions. The analysed set of NIRLs represent a valuable toolkit for deciphering physiological mechanisms and identifying genes involved in grain yield regulation, as they display significantly different phenotypes for a number of traits associated with specific 7AgL segments.

# Acknowledgements

- 572 Financial support from MIUR (Italian Ministry of Education, University and Research), grant PRIN
- 573 (Progetti di Ricerca scientifica di rilevante Interesse Nazionale) 2010-11 on 'Identification and
- 574 characterization of yield- and sustainability-related genes in durum wheat', as well as from Lazio
- 575 region—FILAS project "MIGLIORA", are gratefully acknowledged. The authors would like to
- acknowledge Poh Chong and Alessandra Bitti for technical assistance with the Australian and Italian
- 577 trials, respectively.

## References

- 580 Abbate, P.E., Pontaroli, C., Lázaro, L., Gutheim, F. 2013. A method of screening for spike fertility 581 in wheat. J. Agr. Sci. 151, 322–330, doi:10.1017/S0021859612000068.
- Ahmadi, N., Audebert, A., Bennett, M.J., Bishopp, A., de Oliveira, A.C., Courtois, B., Diedhiou, A.,
- Diévart, A., Gantet, P., Ghesquière, A., Guiderdoni, E., Henry, A., Inukai, Y., Kochian, L., Laplaze, L., Lucas, M, Luu, D.T., Manneh, B., Mo, X., Muthurajan, R., Périn, C., Price, A.,
- Eaplaze, L., Lucas, M, Luu, D.T., Mannen, B., Mo, A., Muthurajan, R., Perin, C., Price, A., Robin, S., Sentenac, H., Sine, B., Uga, Y., Véry, A.A., Wissuwa, M., Wu, P., Xu, J. 2014. The
- roots of future rice harvests. Rice, 7, 29.
- Araus, J.L., Bort, J., Steduto, P., Villegas, D., Royo, C. 2003a. Breeding cereals for Mediterranean conditions: ecophysiological clues for biotechnology application. Ann. Appl. Biol. 142, 129-141.
- Araus, J.L., Villegas, D., Aparicio, N., García del Moral, L.F., El Hani, S., Rharrabti, Y., Ferrio, J.P., Royo, C. 2003b. Environmental factors determining carbon isotope discrimination and yield in durum wheat under Mediterranean conditions. Crop Sci. 43, 170-180.
- Bassi, F.M., Sanchez-Garcia, M. 2017. Adaptation and stability analysis of ICARDA durum wheat elites across 18 Countries. Crop Sci. 57, 2419–2430, doi: 10.2135/cropsci2016.11.0916.
- Becker, H.C., and Leon, J. 1988. Stability analysis in plant breeding. Plant breeding, 101, 1-23.
- Ceoloni, C., Forte, P., Gennaro, A., Micali, S., Carozza, R., Bitti, A. 2005. Recent developments in durum wheat chromosome engineering. Cytogenet. Genome Res. 109, 328–334, http://dx.doi.org/10.1159/000082416.
- Ceoloni, C., Kuzmanović, L., Forte, P., Gennaro, A., Bitti, A., 2014a. Targeted exploitation of gene pools of alien Triticeae species for sustainable and multi-faceted improvement of the durum wheat crop. Crop Pasture Sci. 65, 96–111.
- Ceoloni, C., Kuzmanović, L., Gennaro, A., Forte, P., Giorgi, D., Grossi, M.R., Bitti, A. 2014b.
   Genomes, chromosomes and genes of perennial Triticeae of the genus *Thinopyrum*: the value of their transfer into wheat for gains in cytogenomic knowledge and 'precision' breeding. In:
   Tuberosa, R., Graner, A., Frison, E. (Eds.), Advances in Genomics of Plant Genetic Resources.
   Springer, Dordrecht, The Netherlands, pp. 333–358.
- Ceoloni, C., Kuzmanović, L., Forte, P., Virili, M.E., Bitti, A. 2015. Wheat-perennial Triticeae introgressions: major achievements and prospects. In: Molnár-Láng M, Ceoloni C, Doležel J, eds. Alien introgression in wheat Cytogenetics, Molecular Biology, and Genomics. pp 273-313; Springer. ISBN: 978-3-319-23493-9, doi: 10.1007/978-3-319-23494-6.
- 611 Ceoloni, C., Kuzmanović, L., Ruggeri, R., Rossini, F., Forte, P., Cuccurullo, A., Bitti, A. 2017a.

  Harnessing genetic diversity of wild gene pools to enhance wheat crop production and sustainability: challenges and opportunities. Diversity, 9, 55, doi:10.3390/d9040055.
- Ceoloni, C., Forte, P., Kuzmanović, L., Tundo, S., Moscetti, I., De Vita, P., Virili, M.E., D'Ovidio, R. 2017b. Cytogenetic mapping of a major locus for resistance to Fusarium head blight and crown rot of wheat on *Thinopyrum elongatum* 7EL and its pyramiding with valuable genes from a *Th. ponticum* homoeologous arm onto bread wheat 7DL. Theor. Appl. Genet. 130, 2005-2024, doi:10.1007/s00122-017-2939-8.
- Dempewolf, H., Baute, G., Anderson, J., Kilian, B., Smith, C., Guarino, L. 2017. Past and future use of wild relatives in crop breeding. Crop Sci. 57, 1070–1082, doi: 10.2135/cropsci2016.10.0885.

- Dodig, D., Zoric, M., Kobiljski, B., Savic, J., Kandic, V., Quarrie, S., Barnes, J. 2012. Genetic and association mapping study of wheat agronomic traits under contrasting water regimes. Int. J. Mol. Sci. 13, 6167-6188, doi:10.3390/ijms13056167.
- Dodig, D., Zorić, M., Kobiljski, B., Šurlan-Momirović, G., Quarrie, S.A., 2010. Assessing drought tolerance and regional patterns of genetic diversity among spring and winter bread wheat using simple sequence repeats and phenotypic data. Crop Pasture Sci. 61, 812–824.
- Elía, M., Slafer, G.A., Savin, R. 2018. Yield and grain weight responses to post-anthesis increases in maximum temperature under field grown wheat as modified by nitrogen supply. Field Crops Res. 221, 228–237, https://doi.org/10.1016/j.fcr.2018.02.030.
- Elhani, S., Martos, V., Rharrabti, Y., Royo, C., García del Moral, L.F., 2007. Contribution of main stem and tillers to durum wheat (*Triticum turgidum* L.var. durum) grain yield and its components grown in Mediterranean environments. Field Crops Res. 103, 25–35.
- FAO. 2013. FAOSTAT Statistics Database, available at <a href="http://www.fao.org/faostat/en/#home">http://www.fao.org/faostat/en/#home</a>. Food and Agriculture Organization of the United Nations, Rome, Italy. (verified on 2/11/2017).
- Ferrante, A., Savin, R., Slafer, G.A. 2013. Floret development and grain setting differences between modern durum wheats under contrasting nitrogen availability. J. Exp. Bot. 64, 169–184.
- Ferrante, A., Savin, R., Slafer, G.A. 2015. Relationship between fruiting efficiency and grain weight in durum wheat. Field Crops Res. 177, 109–116, http://dx.doi.org/10.1016/j.fcr.2015.03.009.
- Finlay, K.W., Wilkinson, G.N. 1963. The analysis of adaptation in a plant-breeding programme. Aust.
   J. Agric. Res. 14, 742-754.
- Fischer, R.A., 2008. The importance of grain or kernel number in wheat: a reply to Sinclair and Jamieson. Field Crops Res. 105, 15–21.
- Fischer, R.A., Edmeades, G.O. 2010. Breeding and cereal yield progress. Crop Sci. 50, S85–S98.
- Fleury, D., Jefferies, S., Kuchel, H., Langridge, P. 2010. Genetic and genomic tools to improve drought tolerance in wheat. J. Exp. Bot. 61, 3211–3222, doi:10.1093/jxb/erq152.
- Forte, P., Virili, M.E., Kuzmanović, L., Moscetti, I., Gennaro, A., D'Ovidio, R., Ceoloni, C. 2014. A novel assembly of *Thinopyrum ponticum* genes into the durum wheat genome: pyramiding Fusarium head blight resistance onto recombinant lines previously engineered for other beneficial traits from the same alien species. Mol. Breed. 34, 1701–1716.
- Foulkes, M.J., Sylvester-Bradley, R., Weightman, R., Snape, J. W. 2007. Identifying physiological traits associated with improved drought resistance in winter wheat. Field Crops Res. 103, 11-24.
- 653 Giunta, F., De Vita, P., Mastrangelo, A. M., Sanna, G., Motzo, R. 2018. Environmental and genetic 654 variation for yield-related traits of durum wheat as affected by development. Front. Plant Sci., 655 9, 8, doi: 10.3389/fpls.2018.00008.
- 656 González, F.G., Miralles, D.J., Slafer, G.A. 2011. Wheat floret survival as related to pre-anthesis 657 spike growth. J. Exp. Bot. 62, 4889–4901.
- 658 Grossi, M.R., Gennaro, A., Forte, P., Bitti, A., Ceoloni, C. 2009. Segregation distortion following 659 introgression of *Thinopyrum ponticum* DNA into durum wheat. Proceedings of the 53<sup>rd</sup> Italian 660 Society of Agricultural Genetics Annual Congress, Torino, Italy, 16-19 September 2009,
- http://www.geneticagraria.it/attachment/SIGA\_2009/7\_20.pdf

- Guo Z., Chen, D., Alqudah, A.M., Röder, M.S., Ganal, M.W., Schnurbusch, T. 2017. Genome-wide
   association analyses of 54 traits identified multiple loci for the determination of floret fertility
   in wheat. New Phytologist 214, 257–270.
- Habash, D.Z., Bernard, S., Schondelmaier, J., Weyen, J., Quarrie, S.A., 2007. The genetics of nitrogen use in hexaploid wheat: N utilization, development and yield. Theor. Appl. Genet. 114, 403–419.
- Habash, D.Z., Kehel, Z., Nachit, M. 2009. Genomic approaches for designing durum wheat ready for climate change with a focus on drought. J. Exp. Bot. 60, 2805–2815, doi:10.1093/jxb/erp211.
- Isidro, J., Álvaro, F., Royo, C., Villegas, D., Miralles, D.J., García del Moral, L.F. 2011. Changes in duration of developmental phases of durum wheat caused by breeding in Spain and Italy during the 20<sup>th</sup> century and its impact on yield. Ann. Bot. -Lond. 107, 1355–1366.
- Izanloo, A., Condon, A.G., Langridge, P., Tester, M., Schnurbusch, T. 2008. Different mechanisms of adaptation to cyclic water stress in two South Australian bread wheat cultivars. J. Exp. Bot 59, 3327–3346, doi:10.1093/jxb/ern199.
- Jin K., Shen, J., Ashton, R.W., White, R.P., Dodd, I.C., Phillips, A.L., Parry, M.A.J., Whalley, W.R. 2015. The effect of impedance to root growth on plant architecture in wheat. Plant Soil 392, 323–332, doi: 10.1007/s11104-015-2462-0.
- Kuzmanović, L., Gennaro, A., Benedettelli, S., Dodd, I.C., Quarrie, S.A., Ceoloni. C. 2014. Structural-functional dissection and characterization of yield-contributing traits originating from a group 7 chromosome of the wheatgrass species *Thinopyrum ponticum* after transfer into durum wheat. J. Exp. Bot. 65, 509-525. doi:10.1093/jxb/ert393.
- Kuzmanović, L., Ruggeri, R., Virili, M.E., Rossini, F., Ceoloni, C. 2016. Effects of *Thinopyrum* ponticum chromosome segments transferred into durum wheat on yield components and related morpho-physiological traits in Mediterranean rain-fed conditions. Field Crops Res. 186, 86–98, http://dx.doi.org/10.1016/j.fcr.2015.11.007.
- Kuzmanović, L., Tundo, S., Ciorba R., Forte P., Bitti A., D'Ovidio R., Ceoloni, C. 2017. Transfer of a major QTL for resistance to Fusarium head blight from *Thinopyrum elongatum* onto durum wheat 7AL chromosome arm and its pyramiding with other useful genes from *Th. ponticum*. Proceedings of the Joint Meeting SIBV-SIGA, Pisa, Italy – September, 19-22<sup>nd</sup>, 2017. ISBN 978-88-904570-7-4, <a href="http://www.geneticagraria.it/attachment/SIGA">http://www.geneticagraria.it/attachment/SIGA</a> 2017/3 22.pdf.
- Liu, B., Asseng, S., Müller, C., Ewert, F., Elliott, J., Lobell, D.B., Martre, P., Ruane, A.C., Wallach, 692 D., Jones, J.W., Rosenzweig, C., Aggarwal, P.K., Alderman, P.D., Anothai, J., Basso, B., 693 Biernath, C., Cammarano, D., Challinor, A., Deryng, D., De Sanctis, G., Doltra, J., Fereres, E., 694 Folberth, C., Garcia-Vila, M., Gayler, S., Hoogenboom, G., Hunt, L.A., Izaurralde, R.C., 695 Jabloun, M., Jones, C.D., Kersebaum, K.C., Kimball, B.A., Koehler, A.-K., Kumar, S.N., 696 Nendel, C., O'Leary, G.J., Olesen, J.E., Ottman, M.J., Palosuo, T., Prasad, P.V.V., Priesack, 697 E., Pugh, T.A.M., Reynolds, M., Rezaei, E.E., Rötter, R.P., Schmid, E., Semenov, M.A., 698 Shcherbak, I., Stehfest, E., Stöckle, C.O., Stratonovitch, P., Streck, T., Supit, I., Tao, F., 699 Thorburn, P., Waha, K., Wall, G.W., Wang, E., White, J.W., Wolf, J., Zhao, Z., Zhu, Y. 2016. 700 Similar estimates of temperature impacts on global wheat yield by three independent methods. 701 Nat. Clim. Change, 6, 1130-1136. 702
- Long, S.P., Marshall-Colon, A., Zhu, X.-G. 2015. Meeting the global food demand of the future by engineering crop photosynthesis and yield potential. Cell 161, 56-66, http://dx.doi.org/10.1016/j.cell.2015.03.019.

- Lopes, M.S., Reynolds, M.P. 2010. Partitioning of assimilates to deeper roots is associated with 706 cooler canopies and increased yield under drought in wheat. Funct. Plant Biol. 37, 147–156. 707
- Luo, Q., Bellotti, W., Williams, M., Bryan, B. 2005. Potential impact of climate change on wheat 708 yield in South Australia. Agric. For. Meteorol. 132, 273-285. 709
- Maccaferri, M., Sanguineti, M.C., Demontis, A., El-Ahmed, A., del Moral, L.G., Maalouf, F., Nachit, 710
- M., Nserallah, N., Ouabbou, H., Rhouma, S., Royo, C., Villegas, D., Tuberosa, R. 2011. 711
- Association mapping in durum wheat grown across a broad range of water regimes, J. Exp. Bot. 712
- 62, 409–438, https://doi.org/10.1093/jxb/erq287. 713
- Manschadi, A.M., Christopher, J., de Voil, P., Hammer, G.L. 2006. The role of root architectural 714 traits in adaptation of wheat to water-limited environments. Funct. Plant Biol. 33, 823-837, 715
- 716 https://doi.org/10.1071/FP06055.
- Marti, J., Slafer, G.A. 2014. Bread and durum wheat yields under a wide range of environmental 717 conditions. Field Crops Res. 156, 258-271. 718
- Marti, J., Araus, J.L., Slafer. G.A. 2016. Sink-strength determines differences in performance 719 720 between breadand durum wheat. Field Crops Res. 198, 101-111, http://dx.doi.org/10.1016/j.fcr.2016.08.020. 721
- Martino, D.L., Abbate, P.E., Cendoya, M.G., Gutheim, F., Mirabella, N.E., Pontaroli, A.C. 2015. 722 Wheat spike fertility: inheritance and relationship with spike yield components in early 723 generations. Plant Breeding 134, 264-270, doi:10.1111/pbr.12262. 724
- Miralles, D.J., Resnicoff, E., Carretero, R. 2007. Yield improvement with Lr19 translocation in 725 wheat: which plant attributes are modified? In: Spiertz, J.H.J., Struik, P.C., Laar, H.H. van 726 727 (Eds.), Scale and complexity in plant systems research: gene-plant-crop relations. Frontis, 21, 169-176. 728
- Mondal, S., Rutkoski, J.E., Velu, G., Singh, P.K., Crespo-Herrera, L.A., Guzmán, C., Bhavani, S., 729 Lan, C., He X., Singh, R.P. 2016. Harnessing diversity in wheat to enhance grain yield, climate 730 resilience, disease and insect pest resistance and nutrition through conventional and modern 731 breeding approaches. Front. Plant Sci. 7, 991, doi: 10.3389/fpls.2016.00991. 732
- Monneveux, P., Reynolds, M.P., Gonzalez Aguilar, J., Singh, R.P. 2003. Effects of the 7DL.7Ag 733 734 translocation from Lophopyrum elongatum on wheat yield and related morphophysiological traits under different environments. Plant Breeding 122, 379–384. doi:10.1046/j.1439-735 0523.2003.00856.x 736
- Parent, B., Bonneau, J., Maphosa, L., Kovalchuk, A., Langridge, P., Fleury, D. 2017. Quantifying 737 wheat sensitivities to environmental constraints to dissect G x E in the field. Plant Physiol., pp-738 00372. 739
- Parry M.A.J., Reynold, M., Salvucci, M.E., Raines, C., Andralojc, P.J., Zhu, X.-G., Price, G.D., 740 741 Condon, A.G., Furbank, R.T. 2011. Raising yield potential of wheat. II. Increasing photosynthetic capacity and efficiency. J. Exp. Bot. 62, 453–467, doi:10.1093/jxb/erq304. 742
- Pedro, A., Savin, R., Habash, D.Z., Slafer, G.A., 2011. Physiological attributes associated with yield 743 and stability in selected lines of a durum wheat population. Euphytica 180, 195–208. 744
- Peremarti, A., Marè, C., Aprile, A., Roncaglia, E., Cattivelli, L., Villegas, D., Royo, C. 2014. 745
- 746 Transcriptomic and proteomic analyses of a pale-green durum wheat mutant shows variations
- 747 in photosystem components and metabolic deficiencies under drought stress. BMC Genomics 748 15, 125.

- Prohens, J., Gramazio, P., Plazas, M., Dempewolf, H., Kilian, B., Díez, M.J., Fita, A., Herraiz, F.J.,
- Rodríguez-Burruezo, A., Soler, S., Knapp, S., Vilanova, S. 2017. Introgressiomics: a new
- approach for using crop wild relatives in breeding for adaptation to climate change. Euphytica
- 752 213, 158, https://doi.org/10.1007/s10681-017-1938-9.
- Quarrie, S.A., Pekic-Quarrie, S., Radosevic, R., Rancic, D., Kaminska, A., Barnes, J.D., Leverington,
- M., Ceoloni, C., Dodig, D., 2006. Dissecting a wheat QTL for yield present in a range of
- environments: from the QTL to candidate genes. J. Exp.Bot. 57, 2627–2637.
- Ray, D.K., Mueller, N.D., West, P.C., Foley, J.A. 2013. Yield Trends Are Insufficient to Double Global Crop Production by 2050. PLOS One 8, e66428.
- Reynolds, M.P., Calderini, D.F., Condon, A.G., Rajaram, S. 2001. Physiological basis of yield gains in wheat associated with the LR19 translocation from *Agropyron elongatum*. Euphytica 119, 137–141.
- Richards, R.A., Rebetzke, G.J., Watt, M., Condon, A.G. (Tony), Spielmeyer. W., Dolferus, R. 2010.

  Breeding for improved water productivity in temperate cereals: phenotyping, quantitative trait loci, markers and the selection environment. Funct. Plant Biol. 37, 85–97
- Rosewarne, G., Bonnett, D., Rebetzke, G., Lonergan, P., Larkin, P.J. 2015. The potential of *Lr19* and
   *Bdv2* translocations to improve yield and disease resistance in the high rainfall wheat zones of
   Australia. Agronomy 5, 55-70.
- Royo, C., Elias, E.M., Manthey, F.A., 2009. Durum wheat breeding. In: Prohens, J., Nuez, F., Carena,
   M.J. (Eds.), Handbook of Plant Breeding, Vol. 3. Springer Science+Business Media, LLC,
   Cereals, pp. 199–226.
- Royo, C., Nazco, R., Villegas, D. 2014. The climate of the zone of origin of Mediterranean durum
   wheat (*Triticum durum* Desf.) landraces affects their agronomic performance. Genet. Resour.
   Crop Evol. 61, 1345–1358, doi 10.1007/s10722-014-0116-3.
- Sibikeev, S.N., Voronina, S.A., Krupnov, V.A. 2000. Effects from 7DL-7Ae# 1 translocation on resistance to environmental factors and grain quality of bread wheat. In: EWAC newsletter: proceedings of the 11<sup>th</sup> EWAC conference, 24-28 July 2000, Novosibirsk, Russia, pp. 169-170.
- Sall, A.T., Cisse, M., Gueye, H., Kabbaj, H., Ndoyel, I., Filali-Maltouf, A., Belkadi, B., El-Mourid, M., Ortiz, R., Bassi, F.M. 2018. Heat tolerance of durum wheat (*Tritcum durum* Desf.) elite germplasm tested along the Senegal River. J Agr Sci. 10, 217-233, https://doi.org/10.5539/jas.v10n2p217.
- Singh, R.P., Huerta-Espino, J., Rajaram, S., Crossa, J. 1998. Agronomic effects from chromosome translocations 7DL.7Ag and 1BL.1RS in spring wheat. Crop Sci. 38, 27–33.
- Slack, S., York, L.M., Roghazai, Y., Lynch, J., Bennett, M., Foulkes, J. 2018. Wheat shovelomics II:
  Revealing relationships between root crown traits and crop growth. BioRxiv 280917; doi: https://doi.org/10.1101/280917.
- Slafer, G.A., Savin, R., Sadras, V.O. 2014. Coarse and fine regulation of wheat yield components in response to genotype and environment. Field Crops Res. 157, 71–83, http://dx.doi.org/10.1016/j.fcr.2013.12.004
- Slafer, G.A., Elia, M., Savin, R., García, G.A., Terrile, I.I., Ferrante, A., Miralles, D.J., González, F.G. 2015. Fruiting efficiency: an alternative trait to further rise wheat yield. Food Energy Secur 4, 92–109, doi: 10.1002/fes3.59.

- 791 Tadesse W., Amri, A., Sanchez-Garcia, M., El-Bouhssini, M., Karrou, M., Patil, S., Bassi F., Baum,
- M., Oweis, T. 2016. Improving wheat production in the Central and West Asia and North Africa
- 793 (CWANA) region. In: Langridge, P. (Ed.), Achieving sustainable cultivation in wheat, Vol. 2,
- Burleigh Dodds Science Publishing Limited, Cambridge, UK, http://dx.doi.org/10.19103/AS.2016.0004.39.
- Tardieu, F. 2005. Plant tolerance to water deficit: physical limits and possibilities for progress. C. R. Geoscience 33, 57–67, doi:10.1016/j.crte.2004.09.015.
- Tardieu, F., Tuberosa, R. 2010. Dissection and modelling of abiotic stress tolerance in plants. Curr.
   Opin. Plant Biol. 13, 206-212.
- Terrile, I.I., Miralles, D.J., González, F.G. 2017. Fruiting efficiency in wheat (*Triticum aestivum* L.): 800 801 Trait response to different growing conditions and its relation to spike dry weight at anthesis Field and grain weight harvest. Crops Res. 201, 86-96, 802 803 http://dx.doi.org/10.1016/j.fcr.2016.09.026.
- Tricker, P.J., El Habti, A., Schmidt, J., Fleury, D. 2018. The physiological and genetic basis of combined drought and heat tolerance in wheat. J Exp. Bot., doi.org/10.1093/jxb/ery081.
- Tripathi, S.C., Sayre, K.D., Kaul, J.N. 2005. Planting systems on lodging behavior, yield components, and yield of irrigated spring bread wheat. Crop Sci. 45, 1448–1455.
- Uga, Y., Sugimoto, K., Ogawa, S., Rane, J., Ishitani, M., Hara, N., Kitomi, Y., Inukai, Y., Ono, K., Kanno, N., Inoue, H., Takehisa, H., Motoyama, R., Nagamura, Y., Wu, J., Matsumoto, T.,
- Takai, T., Okuno, K., Yano, M. 2013. Control of root system architecture by DEEPER
- ROOTING 1 increases rice yield under drought conditions. Nat. Genet. 45, 1097-1102, doi:10.1038/ng.2725.
- Uga, Y., Kitomi, Y., Ishikawa, S., Yano, M. 2015. Genetic improvement for root growth angle to enhance crop production. Breeding Sci. 65, 111–119, doi:10.1270/jsbbs.65.111.
- Virili, M.E-, Kuzmanović, L., Bitti, A., Salvi, S., Tuberosa, R., Ceoloni, C. 2015. Analysis of seminal root architecture in durum wheat-*Thinopyrum ponticum* recombinant lines. Proceedings of the Joint Meeting SIBV-SIGA, Milan, Italy 8-11<sup>th</sup> September, 2015. ISBN 978-88-904570-5-0, http://www.geneticagraria.it/attachment/SIGA\_2015/2\_08.pdf.
- Zadoks, J.C., Chang, T.T., Konzak, C.F. 1974. A decimal code for the growth stages of cereals. Weed Res. 14, 415–421.
- Zhang, H., Turner, N.C., Simpson, N., Poole, M.L. 2010. Growing-season rainfall, ear number and the water-limited potential yield of wheat in south-western Australia. Crop Pasture Sci. 61, 296–303.
- Zhang H., Mittal, N., Leamy, L.J., Barazani, O., Song, B.-H. 2017. Back into the wild—Apply untapped genetic diversity of wild relatives for crop improvement. Evol. Appl. 10, 5-24, doi: 10.1111/eva.12434.
- Zaïm, M., El Hassouni, K., Gamba, F., Filali-Maltouf, A., Belkadi, B., Sourour, A., Amri, A., Nachit, M., Taghouti, M., Bassi, F. M. 2017. Wide crosses of durum wheat (*Triticum durum* Desf.)
- reveal good disease resistance, yield stability, and industrial quality across Mediterranean sites.
- Field Crops Res. 214, 219-227, <a href="http://dx.doi.org/10.1016/j.fcr.2017.09.007">http://dx.doi.org/10.1016/j.fcr.2017.09.007</a>.
- Zhao, C., Liu, B., Piao, S., Wang, X., Lobell, D.B., Huang, Y., Huang, M., Yao, Y., Bassu, S., Ciais,
- P., Durand, J.-L., Elliott, J., Ewert, F., Janssens, I.A., Li, T., Lin, E., Liu, Q., Martre, P., Müller,
- 833 C., Peng, S., Peñuelas, J., Ruane, A.C., Wallach, D., Wang, T., Wu, D., Liu, Z., Zhu, Y., Zhu,

Z., Asseng, S. 2017. Temperature increase reduces global yields of major crops in four independent estimates. PNAS 114, 9326–9331.

836 837 Zhu, X.-G., Long, S.P., Ort, D.R. 2010. Improving photosynthetic efficiency for greater yield. Annu. Rev. Plant Biol. 61, 235–61.