

# Life story of Tunisian durum wheat landraces revealed by their genetic and phenotypic diversity

Safa Ben Krima, Amine Slim, Sandrine Gélisse, Hajer Kouki, Isabelle Nadaud, Pierre Sourdille, Amor Yahyaoui, Sarrah Ben M'barek, Frédéric Suffert, Thierry C. Marcel

# ► To cite this version:

Safa Ben Krima, Amine Slim, Sandrine Gélisse, Hajer Kouki, Isabelle Nadaud, et al.. Life story of Tunisian durum wheat landraces revealed by their genetic and phenotypic diversity. 2021. hal-03303679

# HAL Id: hal-03303679 https://hal.inrae.fr/hal-03303679v1

Preprint submitted on 28 Jul 2021

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

# Life story of Tunisian durum wheat landraces revealed by their genetic and phenotypic diversity

- 3 Safa BEN KRIMA<sup>1</sup>, Amine SLIM<sup>2</sup>, Sandrine GÉLISSE<sup>1</sup>, Hajer KOUKI<sup>3</sup>, Isabelle NADAUD<sup>4</sup>, Pierre
- 4 SOURDILLE<sup>4</sup>, Amor YAHYAOUI<sup>5</sup>, Sarrah BEN M'BAREK<sup>3</sup>, Frédéric SUFFERT<sup>1</sup>, Thierry C MARCEL<sup>1\*</sup>
- <sup>1</sup> University Paris-Saclay, INRAE, AgroParisTech, UMR BIOGER, 78850 Thiverval-Grignon, France;
- <sup>2</sup> National Gene Bank of Tunisia, Boulevard du Leader Yasser Arafat Z. I Charguia 1, Tunis 1080, Tunisia;
- <sup>3</sup> Regional Field Crop Research Center of Beja (CRRGC), BP 350, 9000 Beja, Tunisia;
- 8 <sup>4</sup> University Clermont-Auvergne, INRAE, UMR 1095 GDEC, 63000 Clermont-Ferrand, France;
- 9 <sup>5</sup> International Maize and Wheat Improvement Center (CIMMYT), Carretera Mexico-Veracruz Km. 45,
- 10 El Batán, 56237 Texcoco, México;
- 11
- 12 \*Corresponding author: thierry.marcel@inrae.fr
- 13

## 14 Abstract:

- 15 Durum wheat (*Triticum turgidum* L. subsp. *durum*) landraces represent a prominent genetic resource
- 16 for Mediterranean farming systems and breeding programs. Fourteen landraces sampled in Tunisia
- 17 were genotyped with 9 microsatellite markers and characterized with 15 morphological descriptors,
- 18 including resistance to the fungal disease *Septoria tritici* blotch (STB). The genetic diversity, nearly was
- as important within landraces populations (45%) than between populations (54%). It was structured in
- seven genetic groups and was only partly explained by the variety name or the locality of origin.
- 21 Populations were also greatly diversified phenotypically (Shannon-Weaver H'=0.54) with traits related
- to spike and awn colours being the most diversified. Resistance to STB was either qualitative in two populations or with varying degrees of quantitative resistance in the others. A P<sub>st</sub>-F<sub>st</sub> comparison
- indicate a local adaptation of the populations. Overall, the genetic structure of Tunisian durum wheat
- 25 landraces revealed a complex selection trajectory and seed exchanges between farmers.
- 26 **Keywords:** genetic resources, population genetics, phenotypic divergence, P<sub>st</sub>-F<sub>st</sub> comparison,
- 27 septoria leaf blotch.

#### 28 INTRODUCTION

29 The importance of diversity in plant genetic resources used in agriculture and the need for biodiversity 30 conservation is now widely recognized (Maxted & al., 2010). Intra- and interspecific diversity represent 31 a heritage value that is important to preserve because it is the basis for breeders sensu lato - from 32 traditional farmers to global breeding companies - to adapt crops to abiotic stresses (i.e. 33 heterogeneous and changing environments) and provide them with resistance to biotic stresses (i.e. 34 pests and diseases) (Bellon, 1996). The Green Revolution, which occurred between 1950 and the late 35 1960s, led to a loss of this diversity. Traditional varieties, thereafter called landraces, were threatened 36 by genetic extinction primarily due to their replacement by modern genetically uniform varieties (Villa 37 & al., 2007). Although humans have historically domesticated and cultivated more than 7,000 species, 38 few high-yielding modern varieties from a limited number of these species constitute nowadays most 39 of the world's food resources (Perrings, 2018). This evolution to modern production systems promotes 40 the cultivation of some varieties at the expense of traditional and local crops (De luca & al., 2018) and 41 consequently is responsible of a huge intraspecific genetic erosion (Perrings, 2018; Wallace & al, 2019). 42 Moreover, this steady decline of genetic diversity canalizes the evolution of pests and forces to adopt 43 management practices, *e.g.* use of pesticides that damage the agro-ecosystem (Conversa & al, 2020). 44 Wild relatives, weedy forms and traditional varieties are especially important for conservation 45 purposes. One of the most threatened components of agricultural plant genetic resources are 46 traditional varieties commonly referred as landraces (Brush, 1997), which constitute the bulk of 47 genetic diversity in domesticated species (Conversa & al., 2020; Poudel & Johnsen, 2009; Villa & al., 48 2007). Villa & al. (2007) define a landrace as being « a dynamic population of a cultivated plant that 49 has historical origin, distinct identity and lacks formal crop improvement, as well as often being 50 genetically diverse, locally adapted and associated with traditional farming systems ». Landraces are 51 essential heritage from farmer generations at the local and regional scale, as they are associated with 52 traditional farming systems and food trends (Negri, 2003). As a result, they are related to the biological, 53 historical, cultural and socio-economic contexts where they have been grown over generations 54 (Conversa & al, 2020), and are specifically well adapted to the environmental conditions of their 55 cultivation area (i.e. tolerance to biotic and abiotic stresses) (De Ron & al., 2018; A.C. Zeven, 1998). In 56 conclusion, there is a strong paradox between the loss of landraces being replaced by high yielding 57 modern varieties and the necessity for breeding programs to preserve intraspecific diversity to develop 58 innovative varieties and hybrids (De Ron & al., 2018; Govindaraj & al., 2015; Hammer & Diederichsen, 59 2009). This statement is particularly relevant to durum wheat.

60 Durum wheat (Triticum turgidum L. ssp. durum) is one of the most crucial crops in the Mediterranean countries. It's a selfing tetraploid species (2n = 4x = 28, AABB) that originated and diversified in the 61 62 Mediterranean basin (Martínez-Moreno & al., 2020), which is the largest durum wheat producing region worldwide, accounting for about 60% of the total growing area (Royo & al., 2017). This 63 64 traditional crop is the raw material for the fabrication of local dishes and products including couscous, 65 pasta, several kinds of bread and other semolina products such as bulgur and frike (Belaid 2000; Nazco 66 & al., 2014; Hammami & Sissons, 2020). Durum wheat originated in the Fertile Crescent around 10,000 67 years B.P. It spread to the western coast of the Mediterranean basin (MacKey, 2005) reaching North 68 Africa around 7,000 years B.P. (Feldman, 2001). During the migration, a combination of natural and 69 farmer's selection resulted in the development of local durum wheat landraces well adapted to their 70 region of origin and environment. But these durum wheat landraces were recently replaced by 71 improved, genetically uniform and more productive modern varieties also called 'elite cultivars' 72 (Soriano & al., 2016). In this context, the National Gene Bank of Tunisia (NGBT) has implemented an 73 ambitious program for the conservation of Tunisian durum wheat landraces both ex situ in gene banks 74 and in situ on farms. Prospecting activities carried out since 2012 by the NGBT revealed that durum 75 wheat landraces are still cultivated by some farmers in mountainous areas from the North and the 76 Center of Tunisia, under traditional farming systems. These landraces, transmitted by farmers from 77 one generation to the next, are designed by a variety name linked to a historical origin and specific 78 phenotypic characteristics. Previous studies have demonstrated that Tunisian durum wheat landraces 79 are genetically diversified (Medini & al., 2005; Robbana & al., 2019; Slim & al., 2019). Robbana & al 80 (2019) showed a variation of genetic diversity between six durum landraces using Diversity Arrays 81 Technology sequencing (DArTseq). They reported as well a higher level of genetic diversity between 82 landraces than within landraces. Slim & al. (2019) demonstrated for instance a great diversity of 41 83 Tunisian landraces using 16 molecular markers based on simple sequence repeats (SSRs), also called 84 microsatellites, with clear differentiation between landraces and elite cultivars. They detected also five 85 genetic clusters structuring landraces with a strong North-South stratification. SSRs have been 86 reported to be the most widely used markers to study the genetic diversity in wheat germplasm due 87 to their large distribution in the genome, codominant nature, high polymorphism, good reproducibility 88 and ease of application (Russell & al., 1997; Medini & al., 2005). These multiallelic markers allow to 89 capture higher variability than biallelic markers like Single Nucleotide Polymorphism (SNP), Amplified 90 Fragment Length Polymorphism (AFLP) or DArT markers (Semagn & al., 2014; Targońska & al., 2016; 91 Hurtado & al., 2008).

92 Several studies have shown that Tunisian durum wheat landraces are agro-morphologically diversified 93 and can be exploited for their large panel of technological properties (Ayed & al., 2010; Nazco & al., 94 2012; Ayadi & al., 2012; Chamekh & al. 2015; Babay & al. 2019; Bouacha & Rezgui, 2017; Yacoubi & al., 95 2020). Ayed & al. (2010) showed a huge phenotypic diversity of Tunisian durum wheat landraces for 96 six qualitative traits: seed colour, seed size, glume colour, glume pubescence, spike density and beak 97 length (Ayed & al., 2010). Nazco & al. (2012) highlighted that landraces from the western 98 Mediterranean countries such as Tunisia have heavier grains and higher grain-filling rates than those 99 from the eastern Mediterranean countries. Previous studies established that landraces overstep 100 improved genotypes for agronomic traits such as plant height, biomass, straw yield and also have high 101 grain yield, high nitrogen utilization efficiency, high nitrogen use efficiency, high NADH-dependent 102 glutamate synthase activity and high NADH-dependent glutamate dehydrogenase activity (Ayadi & al., 103 2012; Chamekh & al. 2015). Additional studies shown that landraces have a high protein content and 104 consequently high physico-chemical technological propriety of semolina and pasta (Babay & al., 2019, 105 Bouacha & Rezgui, 2017). Landraces are known to be sources of increased biomass and thousand 106 kernel weight, both being important traits for adaptation to drought and heat stresses (Yacoubi & al., 107 2019), but also of genetic resistances to pests, including fungal pathogens. Recently, Huhn & al. (2012) 108 screened a wheat collection including Tunisian durum wheat landraces and revealed that five lines are 109 moderately resistant to Fusarium head blight (Huhn & al., 2012). Ferjaoui & al. (2011, 2015) and Ouaja 110 & al. (2020) identified other lines with resistance to Septoria tritici blotch (STB). This foliar disease, 111 caused by the fungal pathogen Zymoseptoria tritici, can cause important yield losses in Tunisia 112 (Ferjaoui & al., 2015). Most of the Tunisian durum wheat landraces remain to be genetically and 113 phenotypically characterized, including for their level of resistance to STB disease.

114 The large majority of studies on durum wheat landraces examine collections of lines belonging to 115 different populations, with only a few or sometimes only one individual per population. It is advisable to extend the population-study to more individuals per population in order to better characterize 116 117 genetic and phenotypic diversity both between and within populations. To this end, we studied 14 118 Tunisian durum wheat landraces collected by the NGBT between 2015 and 2017. Some landraces had 119 the same name but were grown in different localities while others were grown by the same farmer but 120 had different names. We thus decided to investigate whether these landraces are different (at the 121 molecular and phenotypic levels) or not and what are their relationships. Concretely, these landraces 122 consisted here and in the whole manuscript as a 'population', *i.e.* a group of individuals collected at a

specific farmer's field and reported by each farmer to be a landrace. 16 to 51 individuals were randomly

selected from each population, which were characterized for: (i) their neutral genetic diversity and

structure with SSR markers; (ii) their phenotypic diversity based on agro-morphological characters and

126 their response to STB; and (iii) their phenotypic differentiation (P<sub>st</sub>) comparatively with neutral genetic

differentiation ( $F_{st}$ ) in order to determine whether phenotypic differences between populations were

- 128 due to selection.
- 129

#### 130 **RESULTS**

## 131 Identification of hexaploid lineages, historical contaminant of Tunisian durum wheat fields

A first genetic structure analysis of the durum wheat germplasm showed that the lineages from the 14 populations can be divided into 10 genetically distinct groups (strongest  $\Delta K$  of 14.93) (data not shown). Most genetic groups were composed of lineages coming from different "variety-locality" populations, designed by a "variety" name and its "locality" of origin. At k=10, 33 lineages corresponding to 23 unique multilocus genotypes (MLGs) belong exclusively to three genetic groups genetically close from each other. These lineages stood out for having a characteristic of spikes different from all the others,

- *i.e.* long cylindrical white spikes. These lineages were found in eight out of the 14 populations collected
  in the North or the Center of Tunisia, *i.e.* Roussia Joumine, Mahmoudi Amdoun, Mahmoudi Oued
  Sbaihia, Mahmoudi El Jouf, Chili El Jouf, Chili Lansarine, Aouija Msaken and Mahmoudi Msaken. They
  correspond to what is called 'mule's tail' or 'mare's tail' by Tunisian farmers. 'Mule's tails' are
  undesirable contaminants growing in Tunisian durum wheat fields, whose grains are too "soft" and
  become flour rather than semolina when milled. The karyotype analysis showed that all mule tails'
  lineages suspected to be hexaploid wheat species rather than tetraploid durum wheat carried indeed
- 145 42 chromosomes vs. 28 for 'Karim' and 'Mahmoudi-101' varieties (Figure 1). The highest proportion of
- 146 'mule's tail' was found on the two populations Mahmoudi Oued Sbaihia and Mahmoudi El Jouf, both
- 147 cultivated in the North East of Tunisia (Governorate Zaghouan). Therefore, 33 hexaploid lineages were
- eliminated from the dataset and further analyses were performed with tetraploid wheat lineages only.

# 149 Characterization using microsatellite markers

150 From the collection of 335 lineages of durum wheat, the polymorphic microsatellite (SSR) markers 151 amplified 61 different alleles (Table 1). Size ranges of alleles did not overlap between markers tagged 152 with the same dye. The number of alleles per locus ranged from three for the marker Xgpw2239 to 12 153 for the marker Xgwm372. In total, 23 private alleles were identified in eight out of the nine loci. The 154 low mean-H<sub> $\circ$ </sub> (0.003-0.014) and -H<sub>s</sub> (0.147-0.317) values revealed a low level of heterozygosity. All 155 markers were highly informative and polymorphic as evidenced from their high PIC value (ranging from 156 0.503 for Xgpw2239 to 0,966 for Xgwm413), and were characterized by a high Fixation index (0.961-157 0.986). An analysis of the SSR markers with the BayeScan program revealed no outlier loci. 158 Furthermore, genotype accumulation curves indicated that eight loci only are required to discriminate 159 between individuals in the studied populations (Figure-ESM1). Hence, this multiplex of nine SSR markers is a valuable tool for population genetics studies of durum wheat and genotyping results can 160 161 be further used for characterizing the diversity of the 14 durum wheat populations.

# 162 Distribution of genetic diversity

163 The AMOVA test determined that there is almost as much inter and intra-population genetic variation 164 (54% of genetic variability was explained by inter-populations variability vs. 45% by diversity within populations; Table 2). As the level of heterozygozity is very low (intra-individual variation around 1%),
the studied lineages can be considered fixed after one or two generations of selfing.

167 The genetic diversity indices (genotypic richness; Shannon, Stoddart-Taylor, Simpson and Eveness 168 indexes) for nine SSR markers calculated for each population are given in Table 3. The genotypic 169 richness ranges from 0.107 in the durum wheat population Chili El Jouf to 0.551 in Roussia Journine. 170 The Shannon index ranges from 0.545 in Chili El Jouf to 2.625 in Roussia Joumine. The Stoddart and 171 Taylor's index ranges from 1.33 for Chili El Jouf to 11.25 in Roussia Joumine. The Shannon Wiener index 172 is sensitive to genotypic richness in samples of uneven sizes (Grünwald & al., 2003). However, the high 173 positive correlation between the Shannon and Stoddart-Taylor indexes (r=0.86) reinforces the 174 conclusions that can be drawn. The Simpson's index, measuring the probability that two randomly 175 selected genotypes are different, varies from 0.25 in Chili El Jouf to 0.911 in Roussia Journine and was 176 also highly correlated with the Shannon index (r=0.96). The Eveness index ranges also from 0.25 in Chili 177 El Jouf to 0.801 in Roussia Joumine, indicating that the MLGs observed in Roussia Joumine are closer 178 to equal abundance than for the other populations. As estimates of genotypic diversity include 179 genotype richness and evenness of distribution of genotypes within the sample (Grünwald & al., 2003), 180 combining results of different evaluated parameters lead us to state that Roussia Joumine stands out 181 as being the most diversified population. Indeed, it has the highest genotypic richness, highest 182 genotypic diversity index and highest Eveness index. On the other hand, Chili El Jouf is the least 183 genetically diverse population.

Overall, 118 MLGs were identified from all populations, meaning that 64.8% of lineages are clonal. At
 equal population sizes, the eMLGs range from 2.6 for Chili El Jouf to 9.6 for Roussia Joumine. Roussia

186 Joumine stands out by its high number of MLGs.

#### 187 Genetic structure of Tunisian durum wheat populations

For a number of genetic groups (k) varying from k=3 to k=7, the analysis of the genetic structure of the 188 189 14 durum wheat populations highlighted that the strongest  $\Delta K$  was obtained for k=3 (with  $\Delta K$ =193.84), 190 k=5 (with  $\Delta$ K=77.46) and k=7 (with  $\Delta$ K=44.19) (Figure 2). The number of genetic groups k=7 was chosen 191 because it allows a better description of the genetic structuration of the 14 populations and it 192 maximises the genetic differentiation, *i.e.* F<sub>st</sub>, compared to the other partitioning of groups. The seven 193 populations Bidi Kasserine, Ajimi Kasserine, Chili Lansarine, Chili El Jouf, Mahmoudi Msaken, 194 Mahmoudi Sejnane and Roussia Joumine are made of individuals belonging mainly to one genetic 195 group, showing the homogeneity of individuals within these populations. The two populations from 196 Kasserine (collected from the same farmer), Bidi Kasserine and Ajimi Kasserine, are identical, 197 suggesting that they have a similar origin despite their different names. Similarly, individuals from the 198 populations Chili Lansarine and Chili El Jouf also belong to the same genetic group, implying that 199 populations with the same variety name but from different localities aroused from a common origin. 200 The structure of these populations is therefore explained by a combination of effects, including the 201 locality of origin and the variety name.

202 The minimum spanning network (MSN) visualizes relationships among MLGs and indicates the 203 existence of one MLG of high frequency in nine different populations (Figure 3). The populations 204 sharing this common MLG (MLG.66) are Mahmoudi Oued Sbaihia, Mahmoudi Joumine, Mahmoudi 205 Amdoun, Mahmoudi El Jouf, Richi El Jouf, Aouija Msaken, Beskri Msaken, Chili Lansarine and Chili El 206 Jouf. Other high frequency MLGs were found in two or three populations. Several MLGs detected a 207 single time were highly distant from other MLGs belonging to the same population. MLGs from the 208 population Roussia Journine were close to each other, and close to MLGs from the two populations 209 Mahmoudi Msaken and Mahmoudi Sejnane. These three populations were distant from the other 11 populations. MLGs belonging to Ajimi Kasserine and Bidi Kasserine were exceptionally close to eachother.

212 Genetic differentiation between populations was estimated with pairwise F<sub>st</sub> (F<sub>st</sub> values between pairs 213 of populations). The matrix of pairwise F<sub>st</sub> shows values ranging from 0.016 (between Mahmoudi El 214 Jouf and Mahmoudi Oued Sbaihia) to 0.887 (between Ajimi Kasserine and Mahmoudi Msaken), with 215 significant genetic differentiation between the majority of populations (Table 4). In some cases, the 216 genetic differentiation is not significant (5% threshold), for example between the two populations from 217 Kasserine (Bidi and Ajimi varieties), the two Mahmoudi populations from El Jouf and Oued Sbaihia, and 218 the two Mahmoudi populations from Oued Sbaihia and Joumine; while the two Mahmoudi populations 219 from El Jouf and Joumine were significantly different. Similarly, even if significant, the F<sub>st</sub> value 220 between the two Chili populations from El Jouf and Lansarine is low and very close to the significance 221 threshold. Similarities and differences between populations testify of the history of seed exchanges 222 that took place between farmers. Genetic differentiation between the seven genetic groups determined by STRUCTURE was also calculated using the F<sub>st</sub> index (Table-ESM1). An individual was 223 224 considered to belong to a specific genetic group when it has at least 50% affiliation to this group. The 225 matrix of pairwise F<sub>st</sub> shows values ranging from 0.36 to 0.64 between the seven genetic groups (Table-226 ESM1) indicating that they are significantly different from each other.

# Phylogenetic relationship between the 14 populations and selection trajectory of Tunisian landraces

229 Tunisian durum wheat landraces were collected and stored in genebanks since the studies realized by Félicien Boeuf (1875-1961) in the beginning of the 20<sup>th</sup> century. We studied the relationship between 230 231 the 14 durum wheat populations collected by the NGBT in 2015 and 2017 and 40 landraces collected 232 between 1911 and 1976 carrying the same variety name (seed samples received from NPGS and NGBT 233 genebanks) (Table-ESM2). The 14 populations, the 40 landraces and individuals from three seed lots 234 of the modern variety 'Karim' were genotyped with the nine microsatellite markers described earlier. 235 A neighbour-joining phylogenetic tree was built with microsatellite genotypes (Figure 4). The tree 236 revealed a complex relationship between historical landraces, as samples with the same variety name 237 did not necessarily clustered together. First, the historical landraces called Mahmoudi were mostly 238 divided into two clusters: 14 from the 24 Mahmoudi landraces clustered at the bottom of the tree and 239 were only admixed with the two Bidi landraces (PI534469, 1976) and (PI576736, 1976); 6 of the 240 remaining clustered at the top of the tree and were admixed with one Bidi (Cltr3811, 1912) and one 241 Chili (PI534336, 1976). In this last cluster, the Mahmoudi Msaken population was strongly related to 242 Chili (PI534336, 1976) (bootstrap support of 98.9), and the population Roussia Journine remains 243 relatively distant from other historical landraces and studied populations. This suggests that 244 Mahmoudi was submitted to at least two different environments, and that in each environment it was 245 confronted to different landraces with which some exchanges could have occurred through open 246 pollination. Second, most of the landraces called Chili were grouped in the same cluster in the middle 247 of the tree, together with few Mahmoudi and Bidi landraces. Among them, landraces Mahmoudi 248 (Cltr15501, 1972) and Bidi (PI157961, 1947) were identical to three Chili landraces. This identity could 249 not have arisen by chance or mutation and this strongly suggests that these two landraces belong to 250 the Chili group and that their names have been badly assigned. This raises the question about the 251 collection and maintenance of accessions in gene banks. Nine of the 14 studied populations also grouped in this cluster with Chili landraces: including Chili Lansarine and Chili El jouf, but also 252 253 Mahmoudi El Jouf, Mahmoudi Oued Sbaihia, Beskri Msaken, Mahmoudi Joumine, Aouija Msaken, Richi 254 El Jouf and Mahmoudi Amdoun; indicating a strong contribution of Chili historical landraces to the 255 genetic constitution of the studied populations. Third, the populations Ajimi Kasserine and Bidi 256 Kasserine grouped apart from other populations and landraces, and not surprisingly were strongly

- related to each other (bootstrap support of 100). Finally, in the NGBT and INAT seed lots of 'Karim',
- the same MLG was observed for all individuals with eight of the nine markers, leading us to conclude
- that these two seed lots are genetically homogeneous; variability of the ninth marker was attributed
- to a technical bias. For the CRP seed lot, three different MLGs were observed among which the MLG detected in the two other seed lots of 'Karim'. Thus, this third seed lot corresponds well to 'Karim' but
- is not genetically pure. More surprisingly, Mahmoudi Sejnane was strongly related to Karim (bootstrap
- support of 98.4), which may indicate that the farmer from Sejnane erroneously considered his seed lot
- as a landrace.

# 265 **Phenotypic diversity of the 14 populations**

# 266 -Shannon Index

267 The phenotypic diversity was assessed on 273 from the 335 genotyped lineages for the 15 evaluated 268 phenotypic traits (Table ESM3) using the Shannon diversity index (H') at the population and at the 269 genetic group levels (Table 5). At the population level, the mean H' for all characterized traits ranged 270 from H'=0.41 in Mahmoudi Msaken to H'=0.66 in Richi El Jouf, making Richi El Jouf the most 271 phenotypically diversified population. The spike colour (SC) was the most polymorphic trait over the 272 14 populations (mean H'=1.06), with spikes being mostly slightly coloured. The seed shape (SS) was 273 the least diversified trait over all populations (mean H'=0.08), with the majority of populations having 274 semi-elongated seeds.

- - At the genetic group level, the greatest diversity (H'=0.85) was found in genetic group 4 composed by individuals belonging to Richi El Jouf, Mahmoudi Msaken and Aouija Msaken. Six from the 15 evaluated
  - traits were highly polymorphic in this genetic group compared to others: SS (H'=0.87), spike density
  - (SD) (H'=0.93), length of spike without awns (LSWA) (H'=0.95), awn colour (AC) (H'=1.12), number of (LSWA)
  - spikelets per spike (NSS) (H'=1.28) and thousand grains weight (TGW) (H'=1.03). The genetic group 7
  - was the least phenotypically diversified (H'=0.28) but this group consists of only two lineages and is
  - therefore not representative. The phenotypic diversity observed within the genetic groups indicate
  - that they are not phenotypically homogeneous. Nevertheless, some phenotypic traits are clearly
  - distinct between them. For example, SS with parallel-edge spikes being dominant in genetic groups 2,
  - 3 and 7 (monomorphic H'=0) while less frequent in groups 1, 4 and 5. Genetic group 6 was
  - characterized by a particular stunted-spike shape, which is not listed in the official classification from
  - UPOV. Few individuals from genetic group 4 also had this particular stunted-spike shape. Moreover,
     SC was the most diverse phenotypic trait among all the genetic groups (mean H' = 1.00).

# 288 -Statistics on quantitative traits

- 289 Nine phenotypic traits were quantitatively evaluated and subjected to statistical analyses. Significant
- positive correlations (at p<0.01) were detected between heading date (HS) and plant height (HI) (0.71),
- and not surprisingly between length of awns (LA) and length of awns in relation to spike (LAS) (0.82)
- 292 (Figure-ESM2). Concerning *Septoria tritici* blotch (STB), a significant negative correlation (at *p*<0.01)
- was detected between area under disease progress curve (AUDPC) and HI (-0.43), and between AUDPC
- and HS (-0.57), corroborating that HI and HS can influence disease development. Early heading and short lineages tend to be more infected by STB. Finally, LSWA and SD were also significantly negatively
- 296 correlated (-0.50).
- 297 Phenotypes of all quantitative variables were not normally distributed. Only the trait NSS could be 298 normalized following a Box-Cox transformation (Box & Cox, 1964). A significant difference between 299 population means was found for 8 quantitative traits (at p<0.001); LAS was not significantly different

between population means. We observed a maximum of pairwise significant differences betweenpopulations for the trait NSS (35 pairs) and a minimum for the trait LA (five pairs) (Table 6).

The population Mahmoudi Sejnane cumulated the highest number of differences with other populations for quantitative phenotypic traits, except with the population Mahmoudi Msaken (Table 6). In total, 31 pairs of populations were not significantly different for any of the phenotypic traits studied, including the phylogenetically related (Figure 4) Richi El Jouf and Aouija Msaken, Mahmoudi Joumine and Beskri Msaken, Mahmoudi El Jouf and Mahmoudi Oued Sbaihia, Bidi Kasserine and Ajimi Kasserine, and even Mahmoudi Sejnane and Mahmoudi Msaken (Table 6).

#### 308 -Response of the 14 populations to STB infection

The 14 durum wheat populations were contrasted in their response to STB infection (Figure 5). The two populations Ajimi Kasserine and Roussia Joumine show a nearly complete and qualitative resistance level to the *Z. tritici* strain IPO91009, while Mahmoudi Sejnane appeared highly susceptible. The other 11 populations show varying degrees of resistance or susceptibility to STB. Within each population, a variation in response to STB was also observed between lineages.

## 314 -Factor Analysis of Mixed Data (FAMD)

315 A Factor Analysis of Mixed Data (FAMD) was performed on both quantitative and qualitative variables 316 (Figure 6). The first two principal components in the FAMD accounted for 16.5% and 9.8% of the total 317 variation, respectively, and together explained 26.3% of the total variation (Figure 6A). SS, HI, HS, NSS, 318 TGW, AUDPC and LA were the most important traits contributing to the first principal component 319 (Figure ESM3). Shape of grain (SG), SS, SD, AUDPC, TGW and LSWA contributed significantly to the 320 second principal component (Figure ESM3). FAMD allowed to discriminate between three groups of 321 populations. The first group includes Mahmoudi Sejnane and Mahmoudi Msaken (top left quadrant of 322 Figure 6A), which are differentiated from the others populations by their stunted spike shape (Figure 323 6B), earlier heading date, shorter plant height, higher susceptibility to STB, and lower number of 324 spikelets per spike (p < 0.05). The second group includes Aouija Msaken and Richi El Jouf (top right 325 quadrant of Figure 6A), which are distinguishable by their fusiform spike shape (Figure 6C), elongated 326 shape of grains (Figure 6D), and the highest TGW compared to other populations (p < 0.05). Finally, a 327 third group includes the other 10 populations, which are characterized by spikes with parallel edges 328 (Figure 6F) and semi-elongated grains (Figure 6E).

#### 329 -P<sub>st</sub>-F<sub>st</sub> comparaison and sensitive analysis

330 Comparisons of P<sub>st</sub> and F<sub>st</sub> show that most phenotypic traits are highly divergent reflecting therefore a 331 local adaptation rather than a genetic drift, because lower 95% CI for P<sub>st</sub> were higher than the F<sub>st</sub>=0.57 332 (Figure 7). Only for LAS, P<sub>st</sub> didn't differ from F<sub>st</sub> since the lower 95% CI=0.5066<F<sub>st</sub>. However, the 333 observed degree of differentiation is unlikely due to genetic drift for this trait (Leinonen & al., 2013). 334 Evidence for the robustness of the sensitive P<sub>st</sub>-F<sub>st</sub> analysis varied among the eight highly divergent 335 phenotypic traits but was exceptionally strong for some traits such as HI, HS and AUDPC, which had 336 critical c/h<sup>2</sup> around 0.1 (Table 7). Similarly, LSWA, NSS and TGW have a critical c/h<sup>2</sup> ranging from 0.16 337 to 0.21. Therefore, assuming that these traits are under selection and not under genetic drift is a very 338 robust inference since phenotypic variance across populations that is explained by additive genetic 339 effects would need to be at least six times lower than the phenotypic variation within populations to 340 be explained by genetic drift. For SD and LA, the higher critical  $c/h^2$  of 0.30 and 0.61, indicates that 341 phenotypic differentiation is lower and the conclusion of selection is clearly not conservative.

#### 343 **DISCUSSION**

344 Durum wheat landraces traditionally used by Tunisian farmers have been progressively replaced by 345 modern cultivars provided by international (i.e. CIMMYT) and Tunisian (i.e. INRAT) breeding programs. This led to a rapid increase in productivity of wheat and was accompanied by the multilateral trade 346 347 liberalization on its value chain as well as staple food affordability in both rural and urban Tunisian 348 areas (La Rovere & al., 2010). However, few farmers are still until now cultivating landraces in some 349 mountainous or dry areas of Tunisia characterized by low input farming systems, for their own 350 consumption or to be sold on local markets. Understanding the genetic and phenotypic diversity of 351 landraces can be used to legitimate the preservation of these important genetic resources in order to 352 improve the adaptability of durum wheat crop to marginal environments and threatening pathogens 353 (Pietrusińska & al., 2018).

#### **1-** Durum wheat landraces are genetically diverse and dynamic populations

#### 355 -Microsatellite markers

PIC values are a good indication of the informativeness of the multiplexed panel of the microsatellite (SSR) markers used in this study. According to the classification by Botstein & al. (1980), our nine SSRs were moderately (0.25<PIC<0.5) to highly (PIC>0.5) informative, therefore sufficient to discriminate between populations and useful for further genetic diversity studies and seed bank management.

The genetic diversity of the 14 populations assessed with these SSRs (He=0.57) was almost equivalent to the genetic diversity of several collections of durum wheat landraces from different Mediterranean countries such as Tunisia, Morocco or Italy with average gene diversity values between 0.44 and 0.69 (Nefzaoui & al., 2014; Slim & al., 2019; Sahri & al., 2014; Soriano & al., 2016). This high genetic diversity can explain why Tunisia has previously been considered as a centre of diversity for durum wheat (Ayed & al., 2010).

#### 366 -Repartition of genetic diversity

367 The Analysis of MOlecular Variance (AMOVA) evidenced the great genetic diversity present within the 368 durum wheat populations since variation was only slightly greater between populations (54%) than 369 within population (45%) (Table 2). It can be the result of outcrossing and fitness-relevant mutations. The global  $F_{st}$  value of 0.5 also supports a high differentiation between the 14 populations. Similar 370 371 results were obtained for Ethiopian durum wheat landraces with a slightly higher genetic variance 372 between populations (52%) than between individuals within genetic groups (48%) (Alemu & al., 2020). 373 However, several other studies rather report a greater genetic variance within than between 374 populations (Kabbaj & al., 2017; Soriano & al., 2016; Kyratzis & al., 2019; Asmamaw & al., 2019), 375 potentially explained by seed exchange or selection by farmers for similar traits.

#### 376 -Structuration of lineages

377 Based on a bayesian method, the 14 populations were stratified into seven genetic groups (Figure 2). 378 Only few lineages showed a low membership coefficient q-value suggesting a weak level of admixture 379 among the populations. This is probably due to the fact that wheat is mostly autogamous and to the 380 selection pressure exerted by farmers. Some genetic groups are composed of lineages belonging to a 381 single population while other genetic groups include lineages from different populations. Two genetic 382 groups (G2 and G6) are exclusive to a single population (Roussia Journine and Mahmoudi Sejnane, 383 respectively), as supported by the distant position of each population on the minimum spanning 384 network (Figure 3). Roussia Joumine is the only population with variety name 'Roussia', and this 385 landrace is likely to be genetically distinct from the others. Mahmoudi Sejnane seems unrelated to 386 other 'Mahmoudi' populations. This is also true for Mahmoudi Msaken composed only from lineages 387 belonging to a group (G4) which also includes individuals from the Richi El jouf and Aouija Msaken 388 populations. Another group (G5) is composed of lineages coming from eight populations, including 389 nearly all individuals from Chili El Jouf and Chili Lansarine. This suggest that both populations probably 390 have a common origin and thus highlights the pronounced impact of the variety name in the genetic 391 structuration. Furthermore, a group (G3) is composed of lineages coming from four populations, 392 including nearly all individuals from Ajimi Kasserine and Bidi Kasserine. This strong evidence of the 393 impact of locality on the genetic of Tunisian durum wheat landraces structuration, in addition to the 394 expected effect of the variety name, has been demonstrated in previous studies (Slim & al., 2019; 395 Soriano & al., 2016; Rufo & al., 2019). Finally, a group (G1) is composed of lineages coming from seven 396 populations having different origins and different names of variety, underlining the more complex 397 genetic structure of several populations. Genetic structuration of these 14 durum wheat populations 398 is therefore explained by a combination of effects, including the locality of origin and the variety name, 399 that likely results from the farmer-mediated selection and exchange trajectory of each population.

#### 400 -Agro-morphological diversity

401 Characterizing the phenotypic diversity of durum wheat landraces, known to be often agro-402 morphologically heterogeneous (Royo & al., 2017), is prominent towards an understanding of their life 403 history, in other words the selection trajectory and seed exchanges between farmers. Indeed, the 14 404 studied populations were agro-morphologically diversified with a mean Shannon-Wiener index of 405 H'=0.54, estimated with 15 phenotypic traits. In previous studies, even greater agro-morphological 406 diversities were observed with H'=0.77 within 30 Tunisian durum wheat landraces characterized for 407 11 phenotypic traits (Ayed & al., 2010) and H'=0.63 within a collection of 368 Tunisian durum wheat 408 accessions characterized for 9 phenotypic traits (Slim & al. 2011). The least diversified traits among the 409 14 populations were SG (shape of grain) (H'=0.08), SS (spike shape) (H'=0.14) and LAS (length of awns 410 in relation to spike) (H'=0.14). The most frequent shape of grains is semi-elongated, which is associated 411 with a spike shape with parallel edges. Thus, it suggests a selection pressure exerted by Tunisian 412 farmers on their landraces for the same spike shapes. However, grain and spike shapes appeared highly 413 diversified in several other studies (Belhadj & al., 2015; Bechere & al., 1996; Al Khanjari & al., 2008). 414 The most diversified traits among the 14 populations were SC (spike colour) (H'=1.06), PgA 415 (anthocyanin colouration of awns) (H'=0.82) and AC (awn colour) (H'=0.75) (Table 6). Spike and awn 416 colours are important for Tunisian farmers. For example, farmers associate the black colour of awns 417 with the Mahmoudi landrace distinguished for its higher quality for semolina and bread bakery. These 418 findings are consistent with previous studies showing that awn colour was often among the most 419 diversified traits in Tunisian and Omani landraces (Belhadj & al., 2015; Al Khanjari & al., 2008).

420 Population genetic groups did not appear congruent with the structure of agro-morphological diversity 421 while another study highlighted a reliable relationship between genetic and phenotypic population 422 structures, and the connection of both with the geographic origin of the landraces (Soriano & al., 2016). 423 The overall agro-morphological diversity measured with Shannon index does not differ significantly 424 between the 14 populations. A better discrimination between the populations could be obtained with 425 more informative traits and tighter descriptors. Traits related to spike and awn colours could be 426 described using a colour palette for better discrimination between shades, such as Roussia Joumine's 427 russet colour of awns. A broader list of spike shapes could be established including for example the 428 stunted form of spikes found in Mahmoudi Msaken and Mahmoudi Sejnane populations.

429

#### 431 **2- Assessing the origin of durum wheat landraces**

432 Characterization of the 14 populations revealed the complexity of the history of durum wheat
433 cultivation in Tunisia. Behind each population hides a particular life story that can only be understood
434 in light of converging evidences from genetic and agro-morphological descriptions of the populations,
435 and from their local context.

#### 436 -Roussia Joumine, standing-out redhead

437 The population Roussia Joumine, the only one with variety name 'Roussia', stood-out for the 438 homogeneous russet colour of spikes from all its characterized lineages, although this criterion is not 439 in the UPOV classification. It was also strongly genetically distinguishable from all the other studied 440 populations. In literature, Roussia 875 is a local landrace from the north of Tunisia identified in 1927 441 and registered from 1953 to 1973 (Ammar & al., 2011). Gharbi & El Felah (2013) reported that 442 'Roussia', keep its name from an original population that was cultivated extensively in field crops. Slim 443 & al. (2019) previously reported a dendrogram for the durum wheat collection of the NGBT on which 444 Roussia Joumine and Mahmoudi Amdoun clustered together, while in our study both populations were 445 highly differentiated (F<sub>st</sub>=0.6). Indeed, Roussia Journie showed higher values for all genetic indices 446 evaluated, and it was the most genetically diversified population while being phenotypically less 447 divergent. This landrace was as well remarkably resistant to STB and could be an interesting source of 448 resistances to this disease.

#### 449 -Chili El Jouf and Chili Lansarine, a common origin

450 The two populations Chili El Jouf and Chili Lansarine, were significantly weakly differentiated with SSR 451 markers (F<sub>st</sub>=0.03) and almost exclusively composed of lineages classified in the same genetic group 452 (Figure 2) and sharing an MLG (Figure 3). Both populations were grouped with 'Chili' landraces 453 collected by USDA in Tunisia in 1979 (Figure 4). In literature, the origin of 'Chili' landrace is not very 454 clear. According to Saade & al (1996) 'Chili' was selected from a commercial shipment imported from 455 Chile in the early 1960s and rapidly adopted by Tunisian farmers. Deghais & al. (2007) reported that, 456 in 1932, sir Racine, a French industrialist, sent from Marseille (France) 100 kg of Chili seeds to sir 457 Charles Fabre, a farmer in Bou Salem. In 1938, the Botanical Service of Tunis recovered a part of these 458 Chili seeds and incorporated them in regional field trials to evaluate its adaptative potential to regional 459 conditions. Propagation of Chili then took place as early as 1941/1942 and the Botanical Service of 460 Tunis selected the variety 'Chili-931', which was registered in the official Tunisian catalogue of varieties 461 in 1953. Since 1985, the cultivation of this variety has been largely abandoned with the exception of a 462 few niches and it was removed from the catalogue in 1993.

463 Chili El Jouf and Chili Lansarine were identical for most phenotypic traits studied, except for the NSS, 464 being higher in Chili El Jouf than in Chili Lansarine (Table 6). The difference between the two 465 populations could result from a selection by the farmers from El Jouf for a better yield, whose NSS is a 466 component. Moreover, as NSS is influenced by environmental conditions, such as temperature and 467 day length (Knezevic & al., 2007; Rawson, 1970), the difference between the two Chili populations 468 could also have resulted from a local adaptation to environmental conditions. Genetic analysis 469 demonstrated that Chili El Jouf had lower genetic diversity than all studied populations (Table 3).

#### 470 -Aouija Msaken and Richi El Jouf, distant cousins

The populations Aouija Msaken and Richi El Jouf stood out for their similar fusiform spike shape and
elongated grain shape (Figure 6). These populations are genetically heterogeneous and share lineages
belonging to the same genetic groups (Figure 2) but remain genetically differentiated from each other

474 (F<sub>st</sub>=0.27). Based on Neighbour joining clustering, Richi El Jouf and Aouija Msaken were genetically 475 closely related (bootstrap support of 85.2), with an F<sub>st</sub> index around 0.2 being the lowest between Richi 476 El Jouf and any other population. In addition, both populations share a common MLG (Figure 3) 477 suggesting that they have a common ancestor. Richi is a very old variety related to the species Triticum 478 polonicum (personal communication from NGBT). According to some studies, Aouija is also called 479 "Aouej", "Swabaa Aljia" or "Aouiji". Several forms were introduced in 1913 and 1916 by Boeuf from 480 Morocco where they were called "Zréa Laouaja" in relation to the hunchback shape of the grain 481 reminiscent of that of Triticum polonicum. Richi El Jouf was the most diversified population based on 482 all measured traits (H'=0.66), while Aouija Msaken had a lower phenotypic diversity (H'=0.50).

#### 483 -Ajimi Kasserine and Bidi Kasserine, from the same bag

484 The two populations collected in Kasserine, Bidi Kasserine and Ajimi Kasserine, were undifferentiated 485 from each other, genetically (F<sub>st</sub>=0.039) and phenotypically for quantitative traits. Both of them shared 486 three MLGs, had exclusively the same heading dates, same parallel edged spikes and same spike 487 shapes. Some lineages from Ajimi Kasserine were shorter, with shorter and more compact spikes, 488 longer awns and lower number of spikelets per spike, albeit statistically not significant. In addition, 489 some lineages of Ajimi Kasserine had white colour of seeds and spikes and high anthocyanin 490 pigmentation of awns that was not observed in Bidi Kasserine. The populations Ajimi Kasserine and 491 Bidi-Kasserine were genetically close based on neighbour joining clustering (bootstrap support of 100, 492 Figure 4) and on their F<sub>st</sub> index around 0.04. These results support the hypothesis of a common origin 493 for these two populations, considering that the NGBT collected seed samples from these two 494 populations from the same farmer in the region of Kasserine. Our findings are consistent with an 495 interview with the farmer suggesting a mishandling of seed bags that led to name two bags with 496 different variety names while they originated from the same landrace.

#### 497 -Majmoudi Sejnane, a fake identity

498 Mahmoudi Sejnane was highly differentiated from all the other 'Mahmoudi' populations ( $F_{st}$ : 0.53-0.8), 499 but was phenotypically identical to Mahmoudi Msaken. Both, Mahmoudi Sejnane and Mahmoudi 500 Msaken, were genetically homogeneous with lineages belonging to only one genetic group each, 501 respectively G4 and G6. These two populations were characterized by their stunted spike shape and 502 higher susceptibility to STB. The history of Mahmoudi landraces is unclear and several landraces may 503 have been wrongly attributed the variety name 'Mahmoudi' by farmers. In fact, it appears that the 504 population Mahmoudi Sejnane is genetically very close from the modern variety 'Karim' (bootstrap 505 88.4, Figure 4). A recent study by Slim & al. (2019), studying the same population Mahmoudi Sejnane 506 and the modern variety 'Karim' by neighbour joining cluster analysis, reached the same conclusion. 507 Thus, the population Mahmoudi Sejnane is suspected to be derived from the modern variety 'Karim'. 508 The short plant height of Mahmoudi Sejnane (and Mahmoudi Msaken) compared to other populations 509 and its high level of susceptibility to STB strongly looks alike the characteristics of the variety 'Karim', 510 which has been introduced in Tunisia more than 40 years ago by CYMMIT. It is by far the most cultivated variety in Tunisia. Its resistance to STB has undergone years of genetic erosion making it 511 512 today very susceptible to Z. tritici.

#### 513 -Mahmoudi Msaken, a probable 'Chili' origin

514 Mahmoudi Msaken was highly differentiated from all the other populations with variety name 515 'Mahmoudi'. The phylogenetic tree revealed that Mahmoudi Msaken is rather strongly related to a 516 landrace carrying the name 'Chili' (bootstrap support of 98.9), which do not group with other 'Chili' 517 landraces (Figure 4). However, the short length, stunted spike shape (Figure 6) and higher susceptibility 518 to STB looked alike the characteristics of Mahmoudi Sejnane, while these two populations were 519 genetically highly differentiated (F<sub>st</sub>=0.8). This suggests that Mahmoudi Msaken is inherited from the 520 landrace Chili (PI534336, 1976) but might have evolved through mutations and crosses with other 521 varieties, such as the widespread variety Karim. This hypothesis could be investigated further through

522 the phenotypic characterization of the historical landrace.

523 -Beskri Msaken, a 'Mahmoudi' on disguise

524 Beskri Msaken was composed of lineages belonging to the same genetic groups (i.e. G1 and G5) as 525 other populations with the variety name 'Mahmoudi' (Figure 2). Moreover, Beskri Msaken was weakly 526 differentiated from the populations Mahmoudi El Jouf and Mahmoudi Joumine, these populations 527 sharing two MLGs. The close relationship between 'Beskri' and 'Mahmoudi' landraces was previously 528 established (Robbana & al., 2019; Slim & al., 2019). Based on Neighbour joining clustering, the populations Beskri Msaken and Mahmoudi Joumine also appeared genetically close, even when the Fst 529 530 index was around 0.5. It suggests a common ancestor between Beskri Msaken and other Mahmoudi 531 populations.

#### 532 -'Mahmoudi', a popular variety with a complex history

533 The four populations Mahmoudi El Jouf, Mahmoudi Oued Sbaihia, Mahmoudi Joumine and Mahmoudi 534 Amdoun had a heterogeneous genetic structure but all have lineages belonging to G1 and G5 (Figure 535 2). In addition, Mahmoudi Oued Sbaihia and Mahmoudi Joumine had lineages belonging to G3. 536 Mahmoudi Amdoun was genetically differentiated from the three other populations with variety name 537 'Mahmoudi' (Fst 0.247-0.346) but remained identical phenotypically. The genetic differentiation between 'Mahmoudi' populations from El Jouf and Oued Sbaihia, and from El Jouf and Joumine, was 538 539 weak while they did differ phenotypically for quantitative traits. The complexity of the genetic 540 structure of 'Mahmoudi' populations emphasises the complexity to decipher and identify the 541 determinants of the life history, *i.e.* selection trajectory and seed exchanges between farmers, of those 542 landraces. In the literature, 'Mahmoudi' landraces were reported from Tunisia, Algeria, Libya or 543 Palestine (Deghais & al., 2007; Erroux, 1991; Boeuf, 1926; Gharbi & El Felah, 2013; Ammar & al., 2011). 544 In Tunisia, Boeuf (1926) reported seven indigenous botanical Mahmoudi varieties that were selected 545 and propagated by the Botanical Service of Tunis in the early 1900s. In the middle of the twentieth 546 century 'Mahmoudi' landraces became the most commonly cultivated landraces by traditional farmers 547 in Tunisia, even exclusively in certain locality such as Journie. Farmers questioned during a survey 548 appeared very attached to this landrace for its yield stability, and high quality for making couscous and 549 traditional bread. Based on pairwise F<sub>st</sub> analysis, Chili (PI534351, 1976) was almost genetically identical 550 to Mahmoudi Joumine (F<sub>st</sub>=0), Mahmoudi El Jouf (F<sub>st</sub>=0) and Mahmoudi Oued Sbaihia (F<sub>st</sub>=0.04). These 551 three varieties named 'Mahmoudi' were also weakly differentiated from Chili Lansarine and Chili El 552 Jouf, suggesting that the landrace 'Chili' has contributed to the complex structure of 'Mahmoudi' 553 landraces in Tunisia. On the phylogenetic tree with historical collections of Tunisian landraces, these three varieties named 'Mahmoudi' populations and Mahmoudi Amdoun, clustered with landraces 554 555 carrying the variety name 'Chili' rather than with landraces carrying the variety name 'Mahmoudi' 556 (Figure 4). It strengthens the hypothesis that seeds from 'Chili' have been introduced into 'Mahmoudi' 557 landraces. Also, the success of 'Mahmoudi' in Tunisia may explain why many seed lots (or populations) 558 have been called by traditional farmers with the name 'Mahmoudi' while having different origins, 559 resulting therefore in the complex genetic and phenotypic structure that we observed. Overall, our 560 findings emphasise the fact that the name of varieties is not sufficient to explain the origin and history 561 of Tunisian durum wheat landraces.

562

#### 564 **3- Challenges and opportunities in the conservation of durum wheat landraces**

#### 565 -Contamination of durum wheat landraces with a hexaploid wheat

566 Our study revealed the presence of hexaploid lineages in eight out of the 14 studied populations, easily 567 distinguishable in the farmers' fields because of their long cylindrical white spikes. They are considered 568 as impurities and result in a semolina with a flour-like texture with poorer quality. The origin of this 569 species in Tunisian durum wheat fields is not well understood. Some authors describe it as indigenous 570 and others as introduced by Europeans (Bœuf, 1926; Portères, 1958), while more recent studies 571 established that durum wheat fields grown with landraces are frequently contaminated with hexaploid 572 wheats (Zeven & Waninge, 1989; Figliuolo & al., 2007). In the middle of the twentieth century the 573 proportion of bread wheat in Tunisian durum wheat fields approached 50% (Bœuf, 1926; Portères, 574 1958), but this proportion is lower nowadays and was only 9% in our study. These bread wheats or 575 'mule's tail' existing as a mixture in the durum wheat fields are locally called « Baabous Bghal » 576 = « Babous el brel », « Dhil Bghal » or « Boujlida » = « Bou jelida » (in English: mule's tail), and Bœuf 577 and collaborators were unsuccessful in domesticating them through breeding (Bœuf, 1926; Gharbi & 578 Elfalah, 2013). Portères (1958) suggested that this form already existed before the arrival of Europeans, 579 and that durum wheat brought by Arabs in North Africa competed bread wheat due to its resistance 580 to drought and its better use for semolina. He further mentioned that the Romans, perhaps the 581 Phoenicians and Greeks, brought bread wheat to North Africa. Ducellier described in 1923 an Algerian wheat named 'Hachadi' (white, dense, very rough ear; strongly curved, inflated glumes; strongly 582 583 spreading ears; strong divergent beards; reddish or reddish-brown seeds), which might be synonymous 584 to Tunisian 'mule's tail' (Laumont & Erroux, 1962). 'Hachadi' is probably an old form of cultivated 585 wheat, which is nowadays never found in its pure state but always mixed with traditional durum or 586 bread wheats. Nevertheless, these ancient forms have no cultural value anymore and are not 587 interesting for breeders. Similarly, Laumont & Erroux (1962) reported the presence of a hexaploid 588 wheat form in Algerian durum wheat fields, known as 'Guelia' (root akli: to cook, because of the reddish colour of the grain which would have been scorched in the heat). This form of wheat has spread 589 590 in Algeria since 1950 as a result of the importation of wheat (Florence x Aurore) by Tunisian flour mills. 591 It looks like the « Baabous Bghal » or « Boujlida » (Laumont & Erroux, 1962), is morphologically 592 different from 'Hachadi' and characterised by rough, white ears, with a keeled husk similar to durum 593 wheat, and with a red grain often confused with durum wheat grains.

594 The presence of these undesirable hexaploid wheats in durum wheat fields obliges Tunisian farmers 595 to conduct a dynamic selection in order to eradicate 'mule's tail', unsuccessfully so far. Farmers who 596 are not undergoing this selection process are gradually losing the purity of their landrace, which can

lead to a loss of its diversity and sometimes to its complete loss due to resignation.

#### 598 -Divergence among landraces is explained by selection rather than genetic drift

599 The majority of evaluated quantitative traits were highly divergent ( $P_{st}$ > $F_{st}$ ), with a robust 600 approximation of Pst. Especially, HI, HS, AUDPC, NSS and LSWA were distinguishable with low critical 601  $c/h^2 < 0.20$ . Indeed, we can suppose that divergence in genes coding for these five traits exceeds what 602 is expected on the basis of genetic drift. It indicates that selection has resulted in different phenotypes 603 for these five traits between the studied populations, which is the footprint of local adaptation. 604 Selection made by farmers on their landraces should have contributed to this local adaptation. Indeed, 605 a local survey informed us that some farmers perform irregular selection based especially on spike 606 length, spike density and awn colour. Thus, the genetic variation present in the landraces for these 607 traits constitute an important resource to breed for improved varieties adapted to local constraints 608 and meeting Tunisian farmers' expectations.

#### 609 -Use of most resistant lineages to STB for conservation and breeding purposes

610 Landraces are a well-known source of resistance to diseases (Akem & al. 2000; Nazco & al., 2012; Xu 611 & al., 2018; Agnoun & al., 2019; Yao & al., 2019), such as durum wheat landraces and their resistance 612 to Septoria tritici blotch (STB) caused by Z. tritici. Previous studies by Ferjaoui & al. (2011, 2015) identified for instance new resistance genes to STB in 'Azizi27', 'Agili37', 'Agili39' and 'Derbessi12' 613 614 landraces. Ouaja & al. (2020) also identified valuable sources of resistance among a collection of 304 615 Tunisian durum wheat landraces. In our study, the 14 populations responded differently to STB (Figure 616 5). Roussia Journie and Ajimi Kasserine were the most resistant populations to the Z.tritici strain used 617 for inoculations, IPO91009. The resistance is qualitative (total resistance controlled by major genes) in 618 both populations suggesting the presence of highly effective resistance genes that could be useful for 619 breeding. Bidi Kasserine appeared slightly less resistant to STB than Ajimi Kasserine, although the two 620 populations are genetically and phenotypically identical. On the contrary Mahmoudi Sejnane was 621 highly susceptible to STB, suggesting that it does not carry any source of resistance to this disease or 622 that the resistance genes it may contain have been overcome by Z. tritici populations. Intermediary, 623 the remaining 10 populations expressed varying levels of resistance, probably quantitative (partial 624 resistance controlled by polygenes with moderate-to-small effects).

625 Intra-population variation of the level of resistance between lineages was observed as well (Figure 5), 626 reflecting the genetic diversity previously described within the populations. Few lineages from Roussia 627 Joumine and Ajimi Kasserine had some level of susceptibility. Inversely, several populations expressing 628 quantitative resistance contained few completely resistant lineages, such as Mahmoudi Amdoun and 629 Mahmoudi Oued Sbaihia. Moreover, all populations expressing quantitative resistance as a whole were 630 composed of lineages with varying levels of resistance, which was especially pronounced for 631 Mahmoudi Msaken. This intra-population diversity represents as well an important resource of genetic 632 variation, which should not be overseen.

633

#### 634 Conclusion

635 Our study showed a complex structuration of 14 durum wheat populations, only partly explained by 636 their geographic origin and name of variety. Among them, landraces called 'Chili' contributed a lot to 637 the history of Tunisian durum wheat landraces. These results highlight that a landrace is the outcome 638 of a complex selection trajectory, driven not only by the heritage of a biological material (seed lots) but also by traditional breeding practices (selection) deeply-rooted in a territory and subject to several 639 640 kinds of disruptions (exchange, mixing, loss, ingression of exogeneous material, misidentification, etc.). 641 These two components are an integral part of the "life history" of landraces. The dynamic nature of 642 this life story must be considered and characterized, even more thoroughly than we have done in this 643 study, for instance with sociologists, if one wants to better exploit the resources that landraces 644 represent. It is also crucial to enlarge prospection and collection efforts of actual cultivated landraces to retrace the history of these genetic resources and, particularly, distinguishing the "real" landraces 645 646 from the "false" by combining genetic and phenotypic approaches. The conservation of durum wheat 647 landraces is important for future breeding programs. And the role of gene banks is prominent to 648 develop appropriate and relevant in situ and ex situ conservation plans.

- 649
- 650
- 651

#### 652 Materials and Methods

#### 653 Plant material

654 Since 2012 the National Gene Bank of Tunisia (NGBT) conducts a program aiming at the conservation of landraces from traditional Tunisian crops, such as durum wheat. Farmers still cultivating those 655 656 landraces were identified and seed samples were collected for conservation purposes. Within this 657 framework, we focused on 14 populations, 10 being collected in 2015 from five localities and four 658 being collected in 2017 from three additional localities (Table 8). Each population was considered as 659 representative of a given durum wheat landrace and was designated by a variety name combined with 660 its locality of origin. Therefore, several of these "variety-locality" populations collected in different "localities" have the same "variety" name, and vice-versa. Seeds of each population were randomly 661 662 selected from the seed samples and sown under open field conditions in micro-plots (2 m x 1 m) at the experimental station of Thiverval-Grignon, France (48°50'21.3" N, 1°57'07.2" E). For each population, 663 664 32 plants were randomly selected and labelled from 1 to 32, except for the four populations collected 665 in 2017 for which 16, 23, 38 and 51 plants were selected, leading to a total of 448 plants. At the 666 inflorescence emergence stage (Zadoks' 50-59), the wheat spikes were bagged in permeable 667 cellophane wraps in cellophane to promote self-fertilization and prevent the expression of 668 heterozygosity (< 5% in cultivated wheat). When matured (Zadoks' 90-99), spikes from each plant were 669 collected and stored in separate bags before being threshed, spike by spike. This procedure was 670 repeated twice for the 10 populations collected in 2015 and only once for the four populations 671 collected in 2017. As each cycle of self-fertilization is expected to reduce natural intercrossing by 50%, 672 the residual heterozygosity was estimated less than 1.25%. Each lineage obtained from the 32 plants 673 originally selected and represented by seeds coming from a single spike were sown as a single head-674 row during the growing season 2017-2018 on the CRP-Wheat Septoria Phenotyping Platform from 675 CIMMYT-IRESA (Kodia Bou Salem, Tunisia). This trial was conducted as randomized incomplete block 676 design (12 blocks of 16 m x 1 m separated by the modern variety 'Karim'). Eight reference varieties 677 were included in each block: 'Karim', 'Khiar', 'Maali', 'Nasr', 'Salim', 'Agili-39' and 'Mahmoudi-101'. At 678 the end of the trial, six spikes per wheat lineage (i.e. head-row) were harvested, threshed, and stored 679 for further characterizations.

#### 680 Molecular analyses

#### 681 \*DNA isolation and genotyping

For each wheat lineage, one seed was sown and grown under controlled conditions (growth chamber) 682 with a 16h/24h light (300 photons  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup>) and a temperature of 18°C night/20°C day. Two weeks 683 after sowing, a 2-3 cm segment from the youngest leaf of each plant was sampled for DNA extraction 684 685 performed using the DNA plant MiniKit protocole (www.qiagen.com). Purity and concentration of the 686 extracted DNAs were estimated using a Nanodrop spectrophotometer (ND-1000). The DNA samples were normalized to a concentration of 20 ng. $\mu$ L<sup>-1</sup> and genotyped using nine SSR markers (Table 9). 687 688 Genotyping was outsourced at Eurofins (www.eurofins.fr). These SSR markers were assembled in a 689 single multiplex following the methodology described by Gautier & al. (2014). Briefly, the forward 690 primers were labelled with four fluorochromes (i.e. Ned, Fam, Pet, Vic) in a way that the same colour 691 was given only to markers with non-overlapping range of allele sizes. Candidate markers were 692 individually tested on eight reference durum wheat DNAs, before constitution and validation of the 693 multiplex. Chromatograms were visually inspected for all markers and for all individuals using Peak 694 scanner software version 1.0, before final assignment of SSR alleles. Individuals with missing data for 695 at least one marker were removed from our dataset not to influence the detection of unique multilocus 696 genotypes (MLGs) per population. Finally, genotypes of 335 lineages were obtained (13 to 32 lineages 697 per population), after the exclusion of hexaploid lineages (see below, section "Determination of the698 ploidy level in lineages").

699 \*Marker characterization

Neutrality of the SSR markers was preliminary confirmed using BayeScan v2.1 software, which allows
 identifying candidate loci under natural selection from genetic data using differences in allele
 frequencies between populations. The default settings were used (Foll & Gaggiotti, 2008). The plotting
 and identification of outliers was performed with R software using the output of the MCMC algorithm.

A genotype accumulation curve was drawn with the package poppr in Rstudio v3.5.2 to determine the minimum number of loci necessary to discriminate between individuals in a population. This function randomly sampled the loci without replacement and counted the number of multilocus genotypes observed.

- 708Estimates of number of alleles ( $N_a$ ), number of private alleles ( $N_{ap}$ ), mean observed heterozygosity ( $H_o$ ),709mean genetic diversity ( $H_s$ ) and estimate of  $F_{is}$  following Nei (1987) were obtained using the package710hierfstat of Rstudio v.5.2.
- 711 The Polymorphism Information Content (PIC) of each microsatellite locus was evaluated using the

formula of Botstein & al. (1980), with n corresponding to the number of alleles, p<sub>i</sub> to the frequency of

713 the  $i^{th}$  allele, and  $p_j$  to the frequency of the  $j^{th}$  allele:

714 
$$PIC = 1 - \sum_{i=1}^{n} p_i^2 - \sum_{i=1}^{n-1} \sum_{i=i+1}^{n} 2p_i^2 p_j^2$$

715 Markers with PIC value greater than 0.5 were considered highly informative (Botstein & al., 1980).

716 \*Population structure, molecular variance and diversity analyses

717 Population structure was analysed on MLGs with the Bayesian model-based clustering software 718 STRUCTURE v.2.3.4 (Pritchard & al., 2000). Simulations were performed under an admixture model 719 with independent allele frequencies. The analysis was carried out for a number of tested clusters (K) 720 ranging from 1 to 20, 10 runs, 100,000 burn-in period and 100,000 Markov Chain Monte Carlo (MCMC) 721 repetitions after burning. The most likely number of populations (K) was identified using the Delta K 722 method implemented in Structure Harvester. Structure results were summarized using Excel to obtain 723 the probability of each lineage to belong to each cluster. Lineages of the 14 populations were assigned 724 to one genetic cluster, or several genetic clusters when genotypes were admixed. A lineage was 725 assigned to a genetic group if > 50% of its genome fraction value was derived from that group.

- A Minimum Spanning Network on MLGs based on Nei distances with 1000 bootstraps and neighbor
   joining clustering method was performed using the package poppr (kamvar & al., 2014) of R Studio
   V3.5.2.
- To assess the partitioning of the total genetic variance within and among populations, an analysis of molecular variance (AMOVA) based on *F*-statistics with 999 permutations was performed using GenAlEx 6.5 (Peakall & Smouse, 2012).
- The genetic distance between populations was assessed by calculating pairwise F<sub>st</sub> using the software
   Arlequin v3.5.2.2 (Excoffier & Lischer, 2010).
- 734 Diversity measures incorporate both genotypic richness R (number of genotypes in a population; 735 Dorken & Eckert, 2001) and the Eveness index  $E_5$  (distribution of genotypes within a population;

Grünwald & al., 2003). Common diversity indices were calculated, i.e. Shannon Weaver (H), Stoddart
 and Taylor's index (G) and Simpson's index (lambda). Finally, several genetic diversity parameters,

- 738 including the number of alleles per population (Na), the observed heterozygosity (H<sub>o</sub>), the expected
- heterozygosity (H<sub>e</sub>), the number of multilocus genotypes (MLGs), the number of expected MLG at the
- smallest sample size ≥ 10 based on rarefaction (eMLGs), and Weir and Cockerham estimates of F
- statistics over all loci (F<sub>st</sub>), were estimated using the software Genclone v 2.0 (Arnaud-Haond & Belkhir,
- 2007), the packages poppr (Kamvar & al., 2014) and Hierfstat (Goudet, 2005) of Rstudio v3.5.2.
- 743 \*Phylogenetic analysis

744 Several international gene banks have accessions from durum wheat landraces collected through the 745 20<sup>th</sup> century in their collections. We ordered from the U.S. National Plant Germplasm System (NPGS) 746 of the USDA (https://npgsweb.ars-grin.gov/) seed samples corresponding to 39 accessions from durum 747 wheat landraces carrying the names 'Bidi', 'Chili' and 'Mahmoudi' from Tunisia, Algeria, Morocco and 748 Italy (Table-ESM2). A 40<sup>th</sup> accession, called 'Mahmoudi-101', was provided by the NGBT. Four 749 individuals of each of the 40 accessions were genotyped with the nine SSR markers as described 750 previously. In addition, three different seed lots were collected from the modern variety 'Karim' 751 provided by the NGBT, the CRP-Wheat Septoria Phenotyping Platform, and the National Agronomic 752 Institute of Tunisia (INAT). 30, 30 and 12 individuals from each respective lot were genotyped. Then, a 753 phylogenetic analysis was conducted integrating genotypes of the 14 studied populations, the 40 754 accessions from landraces and the three seed lots from 'Karim'. A phylogenetic tree based on 755 Neighbour Joining clustering method on Edward's distances, with 1000 bootstraps, was generated 756 using the package poppr of Rstudio V3.5.2.

## 757 Evaluation of phenotypic characters:

758 \*Determination of the ploidy level in lineages

759 Preliminary molecular and phenotypic analyses revealed lineages within eight populations belonging 760 to the same genetic groups and being phenotypically more similar to hexaploid wheat species rather 761 than tetraploid durum wheat. The ploidy level of 14 lineages, of the landrace Mahmoudi-101 and of 762 variety Karim was verified by establishing their karyotype. The seeds were sterilized 10 min with 763 sodium hypochlorite 5% and then rinsed with water. They were placed for 48 hours at 4°C in the dark 764 for germination on a petri dish containing a water-moistened filter paper and then moved at 22°C in 765 an oven for 48 hours. Roots of 2 cm were cut and placed in a glass tube containing a solution of ultra-766 pure water previously cooled for 24 hours on ice to stop mitosis. The roots were then fixed in an 767 ethanol/acetic acid solution (3:1) at 4°C for 24 hours. The roots were placed in 1 mL of acetocarmine 768 (carmin 10 g/L with 45% acetic acid) in a watch glass for 1 hour. The tip of a meristem is placed on a 769 slide with a drop of 45% acetic acid, covered with a coverslip and then taped gently with the tip of a 770 pencil to release the cells. The slide is lightly heated for a few seconds at 90 °C to remove the 771 cytoplasmic veil. The quality of the spreads chromosomes is observed under a conventional phase 772 contrast microscope. The photos are taken in bright-field at X40 magnification. Chromosome numbers 773 are counted for five cells to estimate the hexaploid/tetraploid nature of the plants.

- All hexaploid lineages were removed from further analyses on genetic and phenotypic diversity in the14 populations.
- 776 \*Agro-morphological characterization of landraces

Four representative spikes were randomly sampled at maturity from each lineage sown as a headrow

778 in the Kodia Bou Salem's field trial. Fifteen qualitative and quantitative agro-morphological traits were

measured on the plants in the field or on the sampled spikes: plant height (HI), heading date (HS), spike shape (SS), spike colour (SC), spike density (SD), length of awns in relation to spike (LAS), length of spike

without awns (LSWA), length of awns (LA), awn colour (AC), anthocyanin pigmentation of awns (PgA),

number of spikelets per spike (NSS), colour of grains (CG), shape of grains (SG), and thousand grain

- weight (TGW). Scoring was done according to the recommendations of the International Plant Genetic
   Resource Institute (IPGRI) and the International Union for the Protection of New Varieties of Plants
- 785 (UPOV) wheat descriptor lists (Table ESM3). Some lineages were lost during the field trial leaving us
- 786 with 273 lineages for which a complete genotypic and phenotypic data set was available.
- 787 \*Evaluation of landraces for resistance to Septoria tritici blotch

788 During the trial conducted on the CRP-Wheat Septoria Phenotyping Platform at Kodia Bou Salem, 789 lineages from landraces and reference varieties were evaluated for their resistance to Septoria tritici 790 blotch (STB). In the field, spreader rows of the modern variety 'Karim', highly deployed in Tunisia and 791 considered susceptible (Berraies & al., 2014), were sown in all blocks perpendicularly to head-rows. 792 Spreader rows and head-rows were spray-inoculated with the Zymoseptoria tritici strain IPO91009, 793 collected in Tunisia (Béja) in 1991 (Kema & al., 1996). The inoculation was performed twice on 15 and 794 29 January 2018, between Zadoks' 13 and (three fully unfolded leaves) and Zadoks' 26 (tillering ) stages 795 with a conidial suspension adjusted to the concentration 10<sup>6</sup> spores.mL<sup>-1</sup>. STB severity was assessed at two different dates following the "double digit" scoring method (Saari & Prescott, 1975). The area 796 797 under the disease progression curve (AUDPC) was calculated from the percentage of disease severity 798 (DS) at both observation dates (Sharma & Duveiller, 2007; Das & al., 1992) according to the formula:

799 
$$DS = \left(\frac{D1}{9}\right) \times \left(\frac{D2}{9}\right) \times 100$$

where, D1 is the first digit (vertical disease progress) and D2 is the second digit (severity of infection),and

802 
$$AUDPC = \sum_{i=1}^{n-1} ((DS_{i+1} + DS_i) \times (T_{i+1} - T_i))/2$$

803 where,  $DS_i$  is disease severity at the i<sup>th</sup> assessment,  $T_i$  is the time (number of days) on which the i<sup>th</sup> 804 assessment was performed, and n is the total number of assessments.

Statistical analyses were performed to assess differences between populations for their
 resistance/susceptibility to STB at p-value=0.05. Kruskall-Wallis tests were applied and Mann–Whitney
 tests were performed using Rstudio v 3.5.2.

808 \*Statistical analysis of quantitative traits

809 A matrix of correlations between the nine quantitative traits was calculated at confidence interval of 810 0.99 using the package corrplot of Rstudio v 3.5.2. One-way Anova and Kruskall-Wallis tests parametric and non-parametric tests, respectively - were applied depending on the trait's distribution 811 812 to test for phenotypic differences between populations. For the non-normally distributed traits, a first 813 normalization was performed using the boxcox function (Box & Cox, 1964) from the package MASS 814 (Venables & Ripley, 2002). In case a significant difference was detected for a trait, post-hoc tests were 815 performed using Rstudio v 3.5.2: a Tukey's Honest Significant Difference test (i.e. parametric) or a 816 Mann–Whitney test (*i.e.* non-parametric), at p-value=0.001.

817 \*Phenotypic diversity between populations and genetic groups

For qualitative phenotypic traits, we considered the most frequent phenotypic class among the four spikes of each lineage. For quantitative traits, the mean values of the four spikes were categorized into classes (Table ESM3). All qualitative and quantitative values grouped into classes were used to calculate the Shannon–Weaver diversity index (H) (Shannon & Weaver 1949; Jain & al. 1975) according to the formula:

 $H = -\sum_{i=1}^{n} p_i \log_2 p_i$ 

824 where p<sub>i</sub> is the frequency of individuals from the i<sup>th</sup> class and n is the number of classes for the 825 designated phenotypic trait.

826 A relative phenotypic diversity index (H') was calculated according to the formula:

827 
$$H' = H/H_{max}$$
 with  $H_{max} = \log_e (n)$ 

828

#### 829 \*Factor Analysis of Mixed Data (FAMD)

A Factor Analysis of Mixed Data (FAMD) was performed using the FactoMineR and factoextra packages in Rstudio v3.5.2, using both quantitative and qualitative variables to ensure a balance of their influence in the analysis. This method allows to reveal similarities between lineages and to explore associations between all variables.

#### 834 Relationship between genotypic and phenotypic diversity parameters

A P<sub>st</sub>-F<sub>st</sub> analysis was performed using the 273 lineages for which both phenotypic and genetic data
 were available to test whether phenotypic differences in nine quantitative traits between populations
 were due to selection or genetic drift. P<sub>st</sub> was used as an approximation of Q<sub>st</sub>, whose exact calculation
 requires common garden experiments to measure the additive genetic variances (Leinonen & al., 2008;
 Pujol & al., 2008; Brommer, 2011) and was not possible here.

The genetic differentiation  $F_{st}$  and the corresponding 95% confidence interval (CI) were calculated using the package hierfstat.  $P_{st}$  was estimated according to Brommer (2011) according to the formula: 842

843 
$$P_{st} = \frac{\frac{c}{h^2}\sigma_b^2}{\frac{c}{h^2}\sigma_b^2 + 2\sigma_w^2}$$

where  $\sigma_b^2$  and  $\sigma_w^2$  are the phenotypic variances between and within populations, respectively, c is an 844 estimate of the proportion of the total variance due to additive genetic effects across populations, and 845  $h^2$  is the heritability (the proportion of phenotypic variance due to additive genetic effects). When 846 Pst=Fst, the differentiation of quantitative traits may be the result of genetic drift, even if a contribution 847 of natural selection cannot be discarded nor estimated. If P<sub>st</sub>>F<sub>st</sub>, quantitative traits have a higher level 848 849 of differentiation, which could be an evidence for directional selection (or heterogeneous selection). 850 Finally, if P<sub>st</sub><F<sub>st</sub>, quantitative traits are less diversified than neutral differentiation, suggesting that 851 these traits have been under the influence of stabilizing selection (or homogeneous selection).

As the determination of c is difficult, a sensitive analysis was performed using the package Pstat of R studio v3.5.2 to infer the robustness of the estimation of  $c/h^2$  which is critical for P<sub>st</sub> to correctly approximate Q<sub>st</sub>. The lower critical  $c/h^2$  ratio is obtained when P<sub>st</sub> exceeds F<sub>st</sub> and at null assumption ( $c/h^2 = 1$ ) the proportion of phenotypic variance due to additive genetic effects is the same within and across populations. The trait is considered divergent at the point where P<sub>st</sub> exceeds F<sub>st</sub> ( $c<h^2$ ). P<sub>st</sub> values 857 were extracted at null assumption and P<sub>st</sub> with a 95% CI were estimated using Pstat package with 1000 858 permutations. If the lower 95% CI of P<sub>st</sub> is higher than F<sub>st</sub> then the quantitative trait is considered highly 859 divergent. For each trait with  $P_{st} > F_{st}$ , the robustness of the  $P_{st}$  as an approximation of  $F_{st}$ , indicating 860 local adaptation, was estimated by the critical  $c/h^2$  value. This ratio is referring to the value when the lower 95% CI of P<sub>st</sub> equal the upper 95% CI of F<sub>st</sub> as described in Brommer & al. (2011). As the critical 861  $c/h^2$  was here lower than 0.20 for some traits, comparing  $P_{st}$  and  $F_{st}$  allowed us to make robust 862 conclusions on the selection regime (Brommer, 2011). The critical c/h<sup>2</sup> ratio was calculated according 863 864 to the formula:

865

866 
$$\sigma_{w}^{2} (lower) = \frac{\left[\left[1 - P_{st}(lower)^{c} \right]}{2 \times P_{st}(lower)}\right]}{2 \times P_{st}(lower)}$$

867 Then :

868 
$$C/h^{2}(critical) = \frac{\left[2 \times \sigma_{w}^{2}(lower) F_{st}(upper)\right]}{1 - F_{st}(upper)}$$

869

870

## 871 Acknowledgments

872 This study was supported by grants from INRAE department Plant Health and Environment, for the 873 APÔGÉ Project and for the PhD thesis of Safa Ben Krima, covering the 2017-2020 period. We thank 874 Sonia Hamza (INAT, Tunisia) for assistance at the beginning of the project and for providing seeds of 875 the durum wheat cultivar Karim, Marie-Hélène Muller (INRAE AGAP) for her assistance with population 876 genetics analyses, and Anne-Lise Boixel and Florence Carpentier (INRAE BIOGER) for assistance with 877 statistical analyses. We are grateful with the genebanks from USDA ARS and from CIMMYT for 878 providing historical seed samples from Tunisian durum wheat landraces. And finally, we would like to 879 express our deep gratitude to Tunisian farmers who always welcomed us and allowed us to work with 880 their familial heritage, their durum wheat landraces.

881

# 882 Competing interests

883 The authors declare that they have no competing interests.

884

#### 885 References

Agnoun Y, Yelome I, Sié M, Albar L, Ghesquière A, Silue D. 2019. Resistance of selected *Oryza glaberrima* landraces and their intra-specific breeding lines to Beninese rice yellow mottle virus
 isolates. *Crop Protection* 119: 172–176. DOI: 10.1016/j.cropro.2019.01.022

- Akem C, Ceccarelli S, Erskine W, Lenné J. 2000. Using genetic diversity for disease resistance in
   agricultural production. *Outlook on Agriculture* 29: 25–25. DOI: 10.5367/00000000101293013
- Al Khanjari S, Filatenko A, Hammer K, Buerkert A. 2008. Morphological spike diversity of Omani wheat.
- 892 *Genetic Resources and Crop Evolution* **55**: 1185–1195. DOI: 10.1007/s10722-008-9319-9
- Alemu A, Feyissa T, Letta T, Abeyo B. 2020. Genetic diversity and population structure analysis based on the high-density SNP markers in Ethiopian durum wheat (*Triticum turgidum* ssp. *durum*). *BMC*
- 895 *Genetics* **21**: 18. DOI: 10.1186/s12863-020-0825-x

- Ammar K, Gharbi MS, Deghaies M. 2011. Wheat in Tunisia. In: Bonjean A, Angus WM & Van Ginkel M
  (Eds). *The world wheat book, A history of wheat breeding, Volume 2*. Lavoisier Publishing, Paris.
- Arnaud-Haond S, Belkhir K. 2007. GENCLONE: a computer program to analyse genotypic data, test for
   clonality and describe spatial clonal organization. *Molecular Ecology Notes* 7: 15–17. DOI:
   10.1111/j.1471-8286.2006.01522.x
- Asmamaw M, Keneni G, Kassahun T. 2019. Genetic diversity of Ethiopian durum wheat (*Triticum durum*Desf.) landrace collections as revealed by SSR Markers. *Advances in Crop Science and Technology* 7(1):
  1000413. DOI: 10.4172/2329-8863.1000413
- Ayadi S, Karmous C, Hammami Z, Tamani N, Trifa Y, Esposito S, Rezgui S. 2012. Genetic variability of
   nitrogen use efficiency components in Tunisian improved genotypes and landraces of durum wheat.
   *Aqricultural Science Research Journals* 2(11): 591–601. DOI: 10.13140/RG.2.1.4907.2401
- Ayed S, Karmous C, Trifa Y, Slama-Ayed O, Slim-Amara H. 2010. Phenotypic diversity of Tunisian durum
  wheat landraces. *African Crop Science Journal* 18(1): 35–42. DOI: 10.4314/acsj.v18i1.54197
- Babay E, Mnasri SR, Mzid R, Ben Naceur M, Hanana M. 2019. Quality selection and genetic diversity of
  Tunisian durum wheat varieties using SSR markers. *Bioscience Journal* 35(4): 1002–1012. DOI:
  10.14393/BJ-v35n4a2019-42301
- Bechere E, Belay G, Mitiku D, Merker A. 1996. Phenotypic diversity of tetraploid wheat landraces from
  northern and north-central regions of Ethiopia. *Hereditas* 124: 124–172. DOI: 10.1111/j.16015223.1996.00165.x
- Beharav A, Golan G, Levy A. 1997. Evaluation and variation in response to infection with *Puccinia striiformis* and *Puccinia recondita* of local wheat landraces. *Euphytica* 94: 287–293. DOI:
  10.1023/A:1002983824125
- Belaid A. Durum wheat in WANA: Production, trade, and gains from technological change. 2000. In:
  Royo C, Nachit M, Di Fonzo N, Araus JL (Eds). Seminar on durum wheat improvement in the
  Mediterranean region: New challenges. Zaragoza (Spain), CIHEAM, pp 35–49. URL:
  http://om.ciheam.org/om/pdf/a40/00600004.pdf
- 922 Belhadj H, Medini M, Bouhaouel I, Amara H. 2015. Analyse de la diversité phénotypique de quelques
- accessions autochtones de blé dur (*Triticum turgidum* ssp. *durum* Desf.) du sud tunisien. *Journal of new science* 24(5): 1115–1125. URL: https://www.jnsciences.org/agri-biotech/32-volume-24.html
- Bellon MR. 1996. The dynamics of crop infraspecific diversity: A conceptual framework at the farmer
  level. *Economic Botany* 50: 26–39. DOI: 10.1007/BF02862110
- Berraies S, Ammar K, Gharbi M, Yahyaoui A, Rezgui S. 2014. Quantitative inheritance of resistance to
  Septoria tritici blotch in durum wheat in Tunisia. *Chilean journal of agricultural research* 74: 35–40.
  DOI: 10.4067/S0718-58392014000100006
- Bœuf F. 1926. Amélioration de la culture du Blé en Tunisie. *Revue de botanique appliquée et d'agriculture coloniale* 63 (6<sup>ème</sup> année): 657–666. DOI: 10.3406/jatba.1926.4456
- 932 Botstein D, White RL, Skolnic M, Davis RW. 1980. Construction of a genetic linkage map in man using
- 933 restriction fragment length polymorphisms. *American journal of human genetics* 32(3): 314–331.
  934 PMID: 6247908
- Bouacha OD, Rezgui S. 2019. Spaghetti quality: Comparison between landraces and high yielding
  Tunisian durum wheat varieties. *Journal of New Sciences* 64(7): 4056–4060. E-ISSN: 2286-5314

- Box GEP, Cox DR. 1964. An analysis of transformations. *Journal of the Royal Statistical Society. Series B* (*Methodological*) 26(2): 211–252. URL: <u>https://www.jstor.org/stable/2984418</u>
- Brommer JE. 2011. Whither Pst? The approximation of Qst by Pst in evolutionary and conservation
  biology. *Journal of Evolutionary Biology* 24(6): 1160–1168. DOI: 10.1111/j.1420-9101.2011.02268.x
- Brush StB, Meng E. 1998. Farmers' valuation and conservation of crop genetic resources. *Genetic Resources and Crop Evolution* 45: 139–150. DOI: 10.1023/A:1008650819946
- 943 Chamekh Z, Karmous C, Ayadi S, Sahli A, Hammami Z, Fraj MB, Benaissa N, Trifa Y, Slim-Amara H. 2015.
  944 Stability analysis of yield component traits in 25 durum wheat (*Triticum durum* Desf.) genotypes under
  945 contrasting irrigation water salinity. *Agricultural Water Management* 152: 1–6. DOI:
  946 10.1016/j.agwat.2014.12.009
- 947 Conversa G, Lazzizera C, Bonasia A, Cifarelli S, Losavio F, Sonnante G, Elia A. 2020. Exploring on-farm
  948 agro-biodiversity: a study case of vegetable landraces from Puglia region (Italy). *Biodiversity and*949 *Conservation* 29: 747–770. DOI: 10.1007/s10531-019-01908-3
- Das MK, Rajaram S, Mundt CC, Kronstad WE. 1992. Inheritance of slow-rusting resistance to leaf rust
  in wheat. *Crop science* 32(6): 1452–1456. DOI: 10.2135/cropsci1992.0011183X003200060028x
- Deghais M, Kouki M, Gharbi MS, El Felah M. 2003. Les variétés de céréales cultivées en Tunisie: blé
  dur, blé tendre, orge et triticale. INRAT. 447p.
- Deghais M, Kouki M, Gharbi MS, El Felah M. 2007. Les variétés de céréales cultivées en Tunisie: blé
  dur, blé tendre, orge et triticale. INRAT. 445p.
- De Luca D, Cennamo P, Del Guacchio E, Di Novella R, Caputo P. 2018. Conservation and genetic
  characterisation of common bean landraces from Cilento region (southern Italy): high differentiation
  in spite of low genetic diversity. *Genetica* 146: 29–44. DOI: 10.1007/s10709-017-9994-6, PMID:
  29030763
- De Ron AM, Bebeli PJ, Negri V, Vaz Patto MC, Revilla P. 2018. Warm season grain legume landraces
   from the South of Europe for germplasm conservation and genetic improvement. *Frontiers in Plant Science* 9: 1524. DOI: 10.3389/fpls.2018.01524, PMID: 30405662
- Dorken ME, Eckert CG. 2001. Severely reduced sexual reproduction in northern populations of a clonal
  plant, *Decodon verticillatus* (Lythraceae). *Journal of Ecology* 89: 339–350. DOI: 10.1046/j.13652745.2001.00558.x
- 966 Ehdaie B, Waines JG, Hall AE. 1988. Differential responses of landrace and improved spring wheat
  967 genotypes to stress environments. *Crop Science* 28: 838–842. DOI:
  968 10.2135/cropsci1988.0011183X002800050024x
- 969 Erroux J. 1991. « Blé ». In : Encyclopédie berbère, 10, 1991, document B81, mise en ligne le 01 mai
  970 2013, consultée le 01 mars 2020. URL: <u>http://journals.openedition.org/encyclopedieberbere/1766</u>

971 Excoffier L, Lischer HEL. 2010. Arlequin suite ver 3.5: A new series of programs to perform population
972 genetics analyses under Linux and Windows. *Molecular Ecology Resources* 10(3): 564–567. DOI:
973 10.1111/j.1755-0998.2010.02847.x, PMID: 21565059

- 974 Feldman M. 2001. Origin of cultivated wheat. In: Bonjean AP, Angus WJ (Eds). The world wheat book,
- 975 *A history of wheat breeding*. Lavoisier Publishing, Paris, pp 3–56.

- Ferjaoui S, Sbei A, Aouadi N, Hamza S. 2011. Monogenic inheritance of resistance to Septoria tritici
  blotch in durum wheat 'Agili'. *International Journal of Plant Breeding* 5: 17–20.
- Ferjaoui S, M'Barek SB, Bahri B, Slimane RB, Hamza S. 2015. Identification of resistance sources
  to Septoria tritici blotch in old tunisian durum wheat germplasm applied for the analysis of
  the Zymoseptoria tritici-durum wheat interaction. Journal of Plant Pathology 97(3): 471–481. DOI:
  10.4454/JPP.V97I3.028
- Figliuolo G, Mazzeo M, Greco I. 2007. Temporal variation of diversity in Italian durum wheat
   germplasm. *Genetic Resources and Crop Evolution* 54(3): 615–626. DOI: 10.1007/s10722-006-0019-z
- Foll M, Gaggiotti O. 2008. A genome scan method to identify selected loci appropriate for both
   dominant and codominant markers: a Bayesian perspective. *Genetics* 180: 977–993. DOI:
   10.1534/genetics.108.092221, PMID: 18780740
- Gautier A, Marcel TC, Confais J, Crane C, Kema G, Suffert F, Walker A-S. 2014. Development of a rapid
   multiplex SSR genotyping method to study populations of the fungal plant pathogen *Zymoseptoria tritici. BMC Research Notes* **7**: 373. DOI: 10.1186/1756-0500-7-373
- 990 Gharbi MS, Elfelah M. 2013. Les céréales en Tunisie: plus d'un siècle de recherche variétale. Annales
   991 *de l'INRAT* 86 (Numéro Spécial Centenaire): 45–68. ISSN: 0365-4761
- Goudet J. 2005. Hierfstat, a package for R to compute and test variance components and F-statistics.
   *Molecular Ecology Notes* 5: 184–186. DOI: 10.1111/j.1471-8278.2004.00828.x
- Govindaraj M, Vetriventhan M, Srinivasan M. 2015. Importance of genetic diversity assessment in crop
   plants and its recent advances: an overview of its analytical perspectives. *Genetics Research International* 2015: Article ID 431487. DOI: 10.1155/2015/431487, PMID: 25874132
- 997 Grünwald NJ, Goodwin SB, Milgroom MG, Fry WE. 2003. Analysis of genotypic diversity data for
  998 populations of microorganisms. *Phytopathology* 93(6): 738–746. DOI: 10.1094/PHYTO.2003.93.6.738,
  999 PMID: 18943061
- Hammami R, Sissons M. 2020. Durum wheat products, couscous. In: Igrejas G, Ikeda T, Guzmán C (Eds). *Wheat quality for improving processing and human health*. Springer, Cham. DOI: 10.1007/978-3-03034163-3 15
- Hammer K, Diederichsen A. 2009. Evolution, status and perspectives for landraces in Europe. In:
   Vetelainen M, Negri V, Maxted N (Eds). *European landraces: on-farm conservation, management and use*. Bioversity Technical Bulletin No. 15, Bioversity International publisher, Rome, Italy, pp 23–43. URL:
   https://www.bioversityinternational.org/e-library/publications/detail/european-landraces-on-farm-
- 1007 conservation-management-and-use/
- 1008 Huhn MR, Elias EM, Ghavami F, Kianian SF, Chao S, Zhong S, Alamri MS, Yahyaoui A, Mergoum M. 2012.
- 1009 Tetraploid Tunisian wheat germplasm as a new source of Fusarium head blight resistance. Crop Science
- 1010 **52(1)**: 136–145. DOI: 10.2135/cropsci2011.05.0263
- 1011 Hurtado P, Olsen K, Buitrago C, Ospina C, Marin J, Duque M, Wongtiem P, Wenzel P, Killian A, Adeleke
- 1012 M, Fregene M. 2008. Comparison of simple sequence repeat (SSR) and diversity array technology
- 1013 (DArT) markers for assessing genetic diversity in cassava (*Manihot esculenta* Crantz). *Plant Genetic*
- 1014 *Resources* **6(3)**: 208–214. DOI: 10.1017/S1479262108994181

1015Jain SK, Qualset CO, Bhatt GM, Wu KK. 1975. Geographical patterns of phenotypic diversity in a word1016collectionofdurumwheats.CropScience15:700–704.DOI:101710.2135/cropsci1975.0011183X001500050026x

Kabbaj H, Sall AT, Al-Abdallat A, Geleta M, Amri A, Filali-Maltouf A, Belkadi B, Ortiz R, Bassi FM. (2017).
Genetic diversity within a global panel of durum wheat (*Triticum durum*) landraces and modern
germplasm reveals the history of alleles exchange. *Frontiers in Plant Science* 8: 1277. DOI:
10.3389/fpls.2017.01277, PMID: 28769970

- Kamvar ZN, Tabima JF, Grünwald NJ. 2014. Poppr: an R package for genetic analysis of populations with
   clonal, partially clonal, and/or sexual reproduction. *PeerJ* 2: e281. DOI: 10.7717/peerj.281, PMID:
   24688859
- Kema GHJ, Annone JG, Sayoud R, van Silfhout CH, van Ginkel M, de Bree J. 1996. Genetic variation for
  virulence and resistance in the wheat-*Mycosphaerella graminicola* pathosystem. I. Interactions
  between pathogen isolates and host cultivars. *Phytopathology* 86: 200–212. DOI: 10.1094/Phyto-86200
- Knezevic D, Paunovic A, Madic M, Djukic N. 2007. Genetic analysis of nitrogen accumulation in four
  wheat cultivars and their hybrids. *Cereal Research Communications* 35(2): 633–636. Proceedings of the
  VI. Alps-Adria Scientific Workshop, Obervellach, Austria, 30 April–5 May 2007. DOI:
  10.1556/CRC.35.2007.2.117
- Korzun V, Börner A, Worland AJ, Law CN, Röder MS. 1997. Application of microsatellite markers to
   distinguish inter-varietal chromosome substitution lines of wheat (*Triticum aestivum* L.). *Euphytica* 95:
   149–155. DOI: 10.1023/A:1002922706905
- Kuckuck H, Kobabe G, Wenzel G. 1991. Fundamentals of Plant Breeding. 1<sup>st</sup> Edition. Springer-Verlag
   Berlin Heidelberg, Germany, IX 236p. ISBN: 978-3-642-75394-7
- Kyratzis A, Nikoloudakis N, Katsiotis A. 2019. Genetic variability in landraces populations and the risk
  to lose genetic variation. The example of landrace 'Kyperounda' and its implications for *ex situ*conservation. *PLoS ONE* 14(10): e0224255. DOI: 10.1371/journal.pone.0224255
- 1041 La Rovere R, Thabet C, Ammar K, Sferi R. 2010. The Tunisian wheat sector in the new liberalization 1042 scenario. *New Medit* **9(1)**: 13–23.
- 1043 Laumont P, Erroux J. 1962. Les blés tendres cultivés en Algérie. *Annales de l'Institut national* 1044 *agronomique El Harrach* **3**: 1–60. URL: <u>https://www.asjp.cerist.dz/en/article/15083</u>
- Leinonen T, O'Hara RB, Cano JM, Merilä J. 2008. Comparative studies of quantitative trait and neutral
  marker divergence: a meta-analysis. *Journal of Evolutionary Biology* 21: 1–17. DOI: 10.1111/j.14209101.2007.01445.x, PMID: 18028355
- Leinonen T, McCairns R, O'Hara R, Merilä J. 2013. Q<sub>ST</sub>-F<sub>ST</sub> comparisons: evolutionary and ecological
   insights from genomic heterogeneity. *Nature Reviews Genetics* 14: 179–190. DOI: 10.1038/nrg3395
- Lovell DJ, Parker SR, Hunter T, Royle DJ, Coken RR. 1997. Influence of crop growth and structure on the
   risk of epidemics by *Mycosphaerella graminicola* (Septoria tritici) in winter wheat. *Plant pathology* 46:
   126–138. DOI: 10.1046/j.1365-3059.1997.d01-206.x
- MacKey J. 2005. Wheat, its concept, evolution andtaxonomy. In: Royo C, Nachit M, Di Fonzo N, ArausJL,
  Pfeiffer WH, Slafer GA (Eds). *Durum wheat breeding: current approaches and future strategies, Vol. I.*Howorth Press, New York, pp 3–62.

- Martínez-Moreno F, Solís I, Noguero D, Blanco A, Özberk I, Nsarellah N, Elias E, Mylonas I, Soriano JM.
  (2020). Durum wheat in the Mediterranean Rim: historical evolution and genetic resources. *Genetic*
- 1058 *Resources and Crop Evolution* **67**: 1415–1436. DOI: 10.1007/s10722-020-00913-8

Maxted N, Kell S, Toledo Á, Dulloo E, Heywood V, Hodgkin T, Hunter D, Guarino L, Jarvis A, Ford-Lloyd
B. 2010. A global approach to crop wild relative conservation: Securing the gene pool for food and
agriculture. *Kew Bulletin* 65: 561–576. DOI: 10.1007/s12225-011-9253-4

Medini M, Hamza S, Rebai A, Baum M. 2005. Analysis of genetic diversity in Tunisian durum wheat
 cultivars and related wild species by SSR and AFLP markers. *Genetic Resources and Crop Evolution* 52:
 21–31. DOI: 10.1007/s10722-005-0225-0

1065 Morris EK, Caruso T, Busco F, Fischer M, Hancock C, Maier TS, Meiners T, Müller C, Obermaier E, Prati 1066 D, Socher SA, Sonnemann I, Wäschke N, Wubet T, Wurst S, Rillig MC. 2014. Choosing and using diversity 1067 indices: insights for ecological applications from the German biodiversity exploratories. *Ecology and* 1068 *evolution* **4(18)**: 3514–3524. DOI: 10.1002/ece3.1155, PMID: 25478144

Nazco R, Villegas D, Ammar K, Peña RJ, Moragues M, Royo C. 2012. Can Mediterranean durum wheat
 landraces contribute to improved grain quality attributes in modern cultivars? *Euphytica* 185: 1–17.
 DOI: 10.1007/s10681-011-0588-6

Nazco R, Peña RJ, Ammar K, Villegas D, Crossa J, Moragues M, Royo C. 2014. Variability in glutenin
subunit composition of Mediterranean durum wheat germplasm and its relationship with gluten
strength. *The Journal of Agricultural Science* **152(3)**: 379–393. DOI: 10.1017/S0021859613000117,
PMID: 24791017

1076 Nefzaoui M, Udupa S, Gharbi M, Bouhadida M, Iraqi D. 2014. Molecular diversity in Tunisian durum
1077 wheat accessions based on microsatellite markers analysis. *Romanian Agricultural Research* **31**: 33–
1078 39. ISSN: 2067-5720

Negri V. 2003. Landraces in central Italy: where and why they are conserved and perspectives for their
 on-farm conservation. *Genetic Resources and Crop Evolution* 50: 871–885; DOI:
 10.1023/A:1025933613279

1082 Nei M. 1987. Molecular evolutionary genetics. Columbia University Press, New York.

Ouaja M, Aouini L, Bahri B, Ferjaoui S, Medini M, Marcel TC, Hamza S. 2020. Identification of valuable
 sources of resistance to *Zymoseptoria tritici* in the Tunisian durum wheat landraces. *European Journal of Plant Pathology* 156: 647–661. DOI: 10.1007/s10658-019-01914-9

- Peakall R, Smouse PE. 2012. GenAlEx 6.5: genetic analysis in Excel. Population genetic software for
  teaching and research--an update. *Bioinformatics* (Oxford, England) 28(19): 2537–2539. DOI:
  10.1093/bioinformatics/bts460
- Perrings C. 2018. Conservation beyond protected areas: the challenge of landraces and crop wild
   relatives. In: Dayal V, Duraiappah A, Nawn N (Eds). *Ecology, Economy and Society*. Springer, Singapore.
   DOI: 10.1007/978-981-10-5675-8\_8
- 1092 Pielou EC. 1975. Ecological diversity. John Wiley and Sons. New York, pp 165.

1093 Pietrusińska A, Monika Ż, Piechota U, Słowacki P, Moskal K. 2018. Searching for diseases resistance
 1094 sources in old cultivars, landraces and wild relatives of cereals. A review. *Annales UMCS, Agricultura*

1095 LXXIII(4): 45–60. DOI: 10.24326/asx.2018.4.5

- 1096 Portères R. 1958. Les appellations des céréales en Afrique (suite). *Journal d'agriculture tropicale et de* 1097 *botanique appliquée* **5(4-5)**: 311–364. DOI: 10.3406/jatba.1958.2469
- Poudel D, Johnsen FH. 2009. Valuation of crop genetic resources in Kaski, Nepal: Farmers' willingness
  to pay for rice landraces conservation. *Journal of Environmental Management* **90(1)**: 483–491. DOI:
  10.1016/j.jenvman.2007.12.020, PMID: 18359142
- Pritchard JK, Stephens M, Donnelly P. 2000. Inference of population structure using multilocus
   genotype data. *Genetics* 155: 945–959. PMCID: PMC1461096
- Pujol B, Wilson AJ, Ross RIC, Pannell JR. 2008. Are Q (ST) -F-ST comparisons for natural populations
  meaningful? *Molecular Ecology* 17: 4782– 4785. DOI: 10.1111/j.1365-294X.2008.03958.x, PMID:
  19140971
- 1106 Rawson HM. 1970. Spikelet number, its control and relation to yield per ear in wheat. *Australian* 1107 *Journal of Biological Sciences* 23: 1–16. DOI : 10.1071/BI9700001
- Robbana C, Kehel Z, Ben Naceur M, Sansaloni C, Bassi F, Amri A. 2019. Genome-wide genetic diversity
  and population structure of Tunisian durum wheat landraces based on DArTseq technology. *International Journal of Molecular Sciences* 20(6): 1352. DOI: 10.3390/ijms20061352, PMID: 30889809
- 1111 Röder MS, Korzun V, Wendehake K, Plaschke J, Tixier MH, Leroy P, Ganal MW. 1998. A microsatellite 1112 map of wheat. *Genetics* **149(4)**: 2007–2023. PMID: 9691054
- Rosenberg NA, Mahajan S, Ramachandran S, Zhao C, Pritchard JK, Feldman MW. 2005. Clines, clusters,
  and the effect of study design on the inference of human population structure. *PLoS Genetics* 1(6):
  e70. DOI: 10.1371/journal.pgen.0010070
- Royo C, Soriano JM, Alvaro F. 2017. Wheat: a crop in the bottom of the Mediterranean diet pyramid. *In Mediterranean Identities Environment, Society, Culture.* InTech. DOI: 10.5772/intechopen.69184
- Rufo R, Alvaro F, Royo C, Soriano JM. 2019. From landraces to improved cultivars: assessment of
  genetic diversity and population structure of Mediterranean wheat using SNP markers. *PLoS ONE* 14
  (7): e0219867. DOI: 10.1371/journal.pone.0219867
- Russell J, Fuller J, Macaulay M, Hatz BG, Jahoor A, Powell W, Waugh R. 1997. Direct comparison of
  levels of genetic variation among barley accessions detected by RFLPs, AFLPs, SSRs and RAPDs. *Theoretical and Applied Genetics* 95: 714–722. DOI: 10.1007/s001220050617
- Saade ME. 1996. Adoption and impact of high yielding wheat varieties in Northern Tunisia. *CIMMYT Economics Working Paper*. 96-03. Mexico, D.F. ISSN: 0258-8587
- Saari EE, Prescott JM. 1975. A scale for appraising the foliar intensity of wheat disease. *Plant Disease Reporter* 59: 377–380.
- Sahri A, Chentoufi L, Arbaoui M, Ardisson M, Belqadi L, Birouk A, Roumet P, Muller MH. 2014. Towards
  a comprehensive characterization of durum wheat landraces in Moroccan traditional agrosystems:
  analysing genetic diversity in the light of geography, farmers' taxonomy and tetraploid wheat
  domestication history. *BMC evolutionary biology* 14: 264. DOI: 10.1186/s12862-014-0264-2
- Semagn K, Babu R, Hearne S, Olsen M. 2014. Single nucleotide polymorphism genotyping using
  Kompetitive Allele Specific PCR (KASP): overview of the technology and its application in crop
  improvement. *Molecular Breeding* 33: 1–14. DOI: 10.1007/s11032-013-9917-x

- Shannon CE, Weaver W. 1949. The mathematical theory of communication. The University of Illinois.
  Urbana, Chicago, London. pp. 3–24.
- Shannon CE. 2001. A mathematical theory of communication. *ACM SIGMOBILE Mobile Computing and Communications Review* 5(1): 3–55. DOI: 10.1145/584091.584093
- 1139 Sharma RC, Duveiller E. 2007. Advancement toward new Septoria Blotch resistance wheats in south 1140 Asia. *Crop Science* **47**: 961–968. DOI: 10.2135/cropsci2006.03.0201
- 1141 Simpson EH. 1949. Measurement of diversity. *Nature* **163**: 688. DOI: 10.1038/163688a0
- Slim A, Ayed S, Slama-Ayed O, Robbana C, Jaime AT, Slim-Amara H. 2011. Morphological diversity of
  some qualitative traits in tetraploid wheat landrace populations collected in the South of Tunisia. *International Journal of Plant Breeding* 5(1): 67–70.
- 1145 Slim A, Piarulli L, Chennaoui Kourda H, Rouaissi M, Robbana C, Chaabane R, Pignone D, Montemurro
- 1146 C, Mangini G. 2019. Genetic structure analysis of a collection of Tunisian durum wheat germplasm.
- 1147 International Journal of Molecular Sciences **20**: 3362. DOI: 10.3390/ijms20133362, PMID: 31323925
- Spitze K. 1993. Population structure in *Dahpnia obtusa*: quantitative genetic and allozymic variation. *Genetics* 135(2): 367–374. PMID: 8244001
- 1150 Somers DJ, Isaac P, Edwards K. 2004. A high-density microsatellite consensus map for bread wheat
- (*Triticum aestivum* L.). *Theoretical and Applied Genetics* **109**: 1105–1114. DOI: 10.1007/s00122-0041740-7, PMID: 15490101
- Soriano JM, Villegas D, Aranzana MJ, García del Moral LF, Royo C. 2016. Genetic structure of modern
   durum wheat cultivars and Mediterranean landraces matches with their agronomic performance. *PLoS ONE* 11: e0160983. DOI: 10.1371/journal.pone.0160983, PMID: 27513751
- Sourdille P, Singh S, Cadalen T, Brown-Guedira GL, Gay G, Qi L, Gill BS, Dufour P, Murigneux A, Bernard
  M. 2004. Microsatellite-based deletion bin system for the establishment of genetic-physical map
  relationships in wheat (*Triticum aestivum* L.). *Functional & Integrative Genomics* 4: 12–25. DOI:
  10.1007/s10142-004-0106-1, PMID: 15004738
- Stoddart JA, Taylor JF. 1988. Genotypic diversity: Estimation and prediction in samples. *Genetics* 118:
  705–711. PMID: 17246421
- 1162 Targońska M, Bolibok-Brągoszewska H, Rakoczy-Trojanowska M. 2016. Assessment of genetic diversity
  1163 in secale cereale based on SSR markers. *Plant Molecular Biology Reporter* 34: 37–51. DOI:
  1164 10.1007/s11105-015-0896-4
- Venables WN, Ripley BD. 2002. Modern Applied Statistics with S. Fourth Edition, Springer, New York.
   ISBN 0-387-95457-0, URL: <u>http://www.stats.ox.ac.uk/pub/MASS4</u>
- Villa TCC, Maxted N, Scholten M, Ford-Lloyd B. 2007. Defining and identifying crop landraces. *Plant Genetic Resources* 3(3): 373–384. DOI: 10.1079/PGR200591
- 1169 Wallace M, Bonhomme V, Russell J, Stillman E, George TS, Ramsay L, Wishart J, Timpany S, Bull H,
- 1170 Booth A, Martin P. 2019. Searching for the origins of bere barley: a geometric morphometric approach
- 1171 to cereal landrace recognition in archaeology. Journal of Archaeological Method and Theory 26: 1125–
- 1172 1142. DOI: 10.1007/s10816-018-9402-2
- Weir BS, Cockerham CC. 1984. Estimating F-statistics for the analysis of population structure. *Evolution* **38(6)**: 1358–1370. DOI: 10.2307/2408641, PMID: 28563791

1175 Xu XD, Feng J, Fan JR, Liu ZY, Li Q, Zhou YL, Ma ZH. 2018. Identification of the resistance gene to 1176 powdery mildew in Chinese wheat landrace Baiyouyantiao. *Journal of Integrative Agriculture* **17**: 37– 1177 45. DOI: 10.1016/S2095-3119(16)61610-6

Yacoubi I, Nigro D, Sayar R, Masmoudi K, Seo YW, Brini F, Giove SL, Mangini G, Giancaspro A, Marcotuli
I, Colasuonno P, Gadaleta A. 2020. New insight into the North-African durum wheat biodiversity:
phenotypic variations for adaptive and agronomic traits. *Genetic Resources and Crop Evolution* 67:
445–455. DOI: 10.1007/s10722-019-00807-4

Yao F, Zhang X, Ye X, Li J, Long L, Yu C, Li J, Wang Y, Wu Y, Wang J, Jiang Q, Li W, Ma J, Wei Y, Zheng Y,
Chen G. 2019. Characterization of molecular diversity and genome-wide association study of stripe rust
resistance at the adult plant stage in Northern Chinese wheat landraces. *BMC Genetics* 20: Article
number 38. DOI: 10.1186/s12863-019-0736-x, PMID: 30914040

1286 Zeven AC, Waninge J. 1989. The presence of three groups of Scalavatis and other hexaploid bread
1287 wheat plants contaminating durum wheat fields in Cyprus. *Euphytica* 43: 117–124. DOI:
1288 10.1007/BF00037904

1189 Zeven AC. 1998. Landraces: a review of definitions and classifications. *Euphytica* 104: 127–139. DOI:
1190 10.1023/A:1018683119237

- 1191 Website: <u>https://graingenes.org/GG3/</u>
- 1192
- 1193

#### 1194 Figures

#### 1195



**Figure 1. Karyotypes** of A. a 'mule's tail' lineage carrying 42 chromosomes (hexaploid), and B. the durum wheat cultivar Karim carrying 28 chromosomes (tetraploid).



1198

1199Figure 2. Admixture proportions of the 14 durum wheat populations estimated with STRUCTURE1200(K=3 to K=7) leading to the identification of different genetic groups (G1 to G7 on the left of the1201figure). Each vertical bar represents an individual. The colour proportion within each bar represents1202the posterior probability of assignment of each individual to one of the groups of genetic similarity.

1203 The range of assignment probability varied from 0 to 100%.

bioRxiv preprint doi: https://doi.org/10.1101/2020.08.14.251157. this version posted August 14, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



#### 1205

1206 Figure 3. Minimum spanning network between multilocus genotypes (MLGs) of the 14 durum wheat

1207 populations. Each node represents a different MLG with the size proportional to the frequency of the MLG. Colours represent the population of origin of each MLG. Edges represent minimum genetic 1208 1209 distances between MLGs based on Nei distances (1000 bootstraps) and Neighbor-Joining clustering 1210 method. Nodes that are more closely related have thicker and darker edges, whereas nodes that are more distantly related have lighter and thinner edges. 1211

bioRxiv preprint doi: https://doi.org/10.1101/2020.08.14.251157. this version posted August 14, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



1214 Figure 4. Phylogenetic tree with the 14 studied durum wheat populations (in violet), the 40 landraces

1215 collected from the USDA and the NGBT (in black) and three seed lots of the modern cultivar Karim

1216 (in green), resulting from Neighbor-Joining cluster analysis based on Edward's distances (1000

1217 **bootstraps).** Bootstrap support values expressed in percentages are indicated on the nodes only if

1218 there are > 50%.

1219



1221 Figure 5. Septoria tritici blotch (STB) AUDPC (area under the disease progress curve) for the 14 durum

1222 wheat populations obtained after their inoculation in field conditions at Kodia Bou Salem with the

**Z.** *tritici* strain IPO91009. Means are represented by a red point. Populations with significantly different means are indicated by different letters after Kruskal-Wallis and Mann–Whitney tests at

1225 *p*=0.05.



Pictures of corresponding spike and grain shapes.



**Figure 7.** Output of the Factor Analysis of Mixed Data (FAMD). A. Categorical variable factor map projects the classes of qualitative variables in the plane of 1 and 2 dimensions underlining their contribution. Classes of Spike Shape: 1SS, 2SS, 2SSB, 3SS, 5SS; classes of Spike Colour: 1SC, 2SC, 3SC; classes of Awn Colour: 1AC, 2AC, 3AC, 4AC; classes of Anthocyanin colouration of Awns: 1PgA, 3PgA, 5PgA, 7PgA, 9PgA; classes of Colour of Grain: 1CG, 2CG, 3CG, 4CG; classes of Shape of Grain: 1SS, 2SS, 3SS. **B. Correlation circle represents the projection of quantitative variables on the 1 and 2 dimensions underlining their contribution.** HI: plant height; HS: heading date; AUDPC: area under disease progress curve; SD: spike density; LAS: length of awns in relation to spike; LSWA: length of spike without awns; LA: length of awns; NSS: number of spikelets per spike; TGW: thousand grains weight.

bioRxiv preprint doi: https://doi.org/10.1101/2020.08.14.251157. this version posted August 14, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

1231



1232

1233Figure 8. Comparison of phenotypic differentiation of the 14 durum wheat populations ( $P_{st}$ : red solid1234line; dotted red lines represents the 95% Cl). The neutral genetic differentiation ( $F_{st}$ - dotted green line1235for  $F_{st}$ =0.57098 was calculated using the package hierfstat with Rstudio version 3.5.2), while the ratio1236c/h² ranged from 0 to 3 where c represents the proportion of the total variance and h² the heritability.

1237 The lowest value of  $c/h^2$  for which  $P_{st}$  exceeds  $F_{st}$  = critical value of  $c/h^2$  can be considered an indication 1238 of the robustness of using  $P_{st}$  as an alternative for  $Q_{st}$  (Brommer, 2011).

bioRxiv preprint doi: https://doi.org/10.1101/2020.08.14.251157. this version posted August 14, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

#### 1240 Tables

1241

# 1242 Table 1: Polymorphism of the 9 SSR (microsatellite) markers used to characterize the 14 durum wheat populations: number

of alleles (N<sub>a</sub>), number of private alleles (N<sub>ap</sub>), mean observed heterozygosity (H<sub>o</sub>) mean expected heterozygosity (H<sub>s</sub>), fixation
 index (F<sub>is</sub>) following Nei (1987) expected heterozygosity over all (H<sub>e</sub>), and Polymorphism Information Content (PIC) values.

	Xgpw2103	Xgpw2239	Xgpw4082	Xgpw7148	Xgwm193	Xgwm285	Xgwm372	Xgwm4004	Xgwm413
Na	4	3	6	6	9	8	12	5	8
N <sub>ap</sub>	2	0	3	2	5	3	4	1	3
H₀	0.005	0.005	0.003	0.008	0.006	0.014	0.007	0.005	0.012
Hs	0.147	0.224	0.198	0.267	0.192	0.353	0.323	0.313	0.317
Fis	0.965	0.977	0.986	0.969	0.971	0.961	0.977	0.984	0.962
He	0.396	0.561	0.524	0.413	0.528	0.691	0.643	0.665	0.738
PIC	0.322	0.423	0.466	0.387	0.492	0.639	0.618	0.600	0.704

1245

# 1247 Table 2: Analysis of Molecular Variance (AMOVA) based on SSR (microsatellite) markers and using the F<sub>st</sub>

#### 1248 measure, for 335 individuals belonging to 14 durum wheat populations.

Source	df	SS	MS	Estimated variance	Variance (%)
Between populations	13	934.965	71.920	1.459	54%
Within populations	321	782.721	2.438	1.204	45%
Within individuals	335	10.500	0.031	0.031	1%
Total	669	1728.187		2.694	100%

df: degree of freedom; SS: sums of squares; MS: mean square; variance (%): percentage of total variance contributed by each component.

	biof
	χ, ×į
~	þ
Nh:	epri
ich	nt c
was	<u>.</u>
no	đ.
t ce	s://
nifi	<u>0</u> .
dbe	οīά
d Ac	10.
ieer	1
ſe	1/2
riew	22
) is	08
the	4
au	25
thor	15
-Ť	2
=	÷
Inde	his
inder. A	his vers
ınder. All rig	his version
inder. All rights	his version pos
ınder. All rights res	his version postec
inder. All rights reserv	his version posted Au
inder. All rights reserved.	his version posted Augu:
inder. All rights reserved. No	his version posted August 1
inder. All rights reserved. No reu	his version posted August 14, 2
Inder. All rights reserved. No reuse	his version posted August 14, 2020
inder. All rights reserved. No reuse allo	his version posted August 14, 2020. TI
inder. All rights reserved. No reuse allowed	his version posted August 14, 2020. The c
inder. All rights reserved. No reuse allowed with	his version posted August 14, 2020. The copy
inder. All rights reserved. No reuse allowed witho	his version posted August 14, 2020. The copyrig
inder. All rights reserved. No reuse allowed without p	his version posted August 14, 2020. The copyright h
inder. All rights reserved. No reuse allowed without perm	his version posted August 14, 2020. The copyright holds
inder. All rights reserved. No reuse allowed without permissi	his version posted August 14, 2020. The copyright holder fo
inder. All rights reserved. No reuse allowed without permission.	his version posted August 14, 2020. The copyright holder for the
inder. All rights reserved. No reuse allowed without permission.	his version posted August 14, 2020. The copyright holder for this p
inder. All rights reserved. No reuse allowed without permission.	his version posted August 14, 2020. The copyright holder for this prep

Populations	Number of lineages <sup>1</sup>	Na <sup>2</sup>	H₅³	H₀⁴	Number of MLG⁵	Number of eMLG <sup>6</sup>	R <sup>7</sup>	Shannon index H <sup>8</sup>	Stoddart and Taylor's index (G) <sup>9</sup>	Simpson's index LAMDA <sup>10</sup>	Eveness E5 <sup>11</sup>
Ajimi Kasserine	32	2.333	0.138	0.010	10	5.19	0.29	1.466	2.60	0.615	0.480
Aouija Msaken	28	2.556	0.159	0	5	2.86	0.148	0.608	1.35	0.260	0.420
Beskri Msaken	26	2.333	0.382	0	9	5.48	0.320	1.528	2.96	0.663	0.545
Bidi kasserine	25	2.667	0.268	0.036	12	7.36	0.458	1.977	4.37	0.771	0.542
Chili El jouf	29	2.111	0.096	0	4	2.60	0.107	0.545	1.33	0.250	0.460
Chili Lansarine	28	2.444	0.185	0	11	6.93	0.370	1.894	4.04	0.753	0.539
Mahmoudi Amdoun	21	2.444	0.367	0	7	5.30	0.300	1.402	2.74	0.635	0.568
Mahmoudi El Jouf	13	2.444	0.332	0.009	6	6.00	0.416	1.285	2.45	0.592	0.554
Mahmoudi Joumine	19	2.333	0.285	0	8	6.25	0.388	1.587	3.19	0.687	0.565
Mahmoudi Msaken	15	1.333	0.057	0	4	3.72	0.214	0.857	1.77	0.436	0.569
Mahmoudi Oued Sbaihia	22	2.889	0.407	0.015	11	7.26	0.476	1.895	4.10	0.756	0.549
Mahmoudi Sejnane	16	2.111	0.267	0	7	6.22	0.400	1.667	4.27	0.766	0.760
Richi El jouf	31	3.000	0.348	0.022	7	4.10	0.200	1.283	2.80	0.643	0.691
Roussia Joumine	30	3.111	0.342	0	17	9.61	0.551	2.625	11.25	0.911	0.801
					12	Fst=0.572					

1250 Table 3: Genetic diversity of the 14 durum wheat populations as evaluated from 9 SSR (microsatellite) markers

1: Number of genotyped lineages by population after eliminating hexaploid individuals and genotypes containing missing value for at least one marker. 2: Mean average of alleles. 3: Mean expected heterozygosity. 4: Mean observed heterozygosity. 5: Number of MultiLocus Genotypes. 6: Number of expected MLG at the smallest sample size based on rarefaction. 7: Genotypic Richness (Dorken & Eckert, 2001). 8: Shannon-Wiener Index of MLG diversity (Arnaud-Haond & al., 2007; Shannon 2001). 9: Stoddart and Taylor's index of MLG diversity (Stoddart & Taylor, 1988).10: Simpson's Index (Simpson, 1949). 11: Evenness, E<sub>5</sub> (Pielou, 1975; Grünwald & al., 2003). 12: F<sub>st</sub> over all loci (Weir & Cockrham, 1984).

Table 4: Matrix of F<sub>st</sub> between the 14 durum wheat populations

	Ajimi Kasserine	Aouija Msaken	Beskri Msaken	Bidi Kasserine	Chili El Jouf	Chili Lansarine	Mahmoudi Amdoun	Mahmoudi El Jouf	Mahmoudi Joumine	Mahmoudi Msaken	Mahmoudi Oued Sbaihia	Mahmoudi Sejnane	Richi El Jouf
Aouija Msaken	0.808*												
Beskri Msaken	0.688*	0.570*											
Bidi kasserine	0.039	0.733*	0.579*										
Chili El Jouf	0.845*	0.778*	0.236*	0.765*									
Chili Lansarine	0.788*	0.705*	0.192*	0.699*	0.035*	]							
Mahmoudi Amdoun	0.679*	0.590*	0.343*	0.576*	0.489*	0.409*							
Mahmoudi El Jouf	0.722*	0.618*	0.047	0.591*	0.149*	0.088*	0.315*						
Mahmoudi Joumine	0.758*	0.656*	0.053	0.650*	0.139*	0.111	0.346*	0.033					
Mahmoudi Msaken	0.887*	0.757*	0.579*	0.804*	0.862*	0.768*	0.703*	0.698*	0.704*				
Mahmoudi Oued Sbaihia	0.614*	0.541*	0.112*	0.490*	0.208*	0.140*	0.247*	0.016	0.108	0.615*			
Mahmoudi Sejnane	0.737*	0.731*	0.559*	0.638*	0.772*	0.689*	0.587*	0.586*	0.641*	²0.800*	0.527*		
Richi El Jouf	0.680*	0.270*	0.233*	0.591*	0.354*	0.287*	0.315*	0.188*	0.253*	0.575*	0.178*	0.570*	
Roussia Joumine	0.707*	0.680*	0.506*	0.606*	0.724*	0.667*	0.612*	0.559*	0.577*	0.645*	0.525*	0.647*	0.569*

\*Significance of the F<sub>st</sub> at a threshold of 5%.

1253

		HI	HS	AUDPC	SS	SC	SD	LAS	LSWA	LA	AC	PgA	NSS	CG	SG	TGW	H' mean ±SE
	H'																
	Bidi Kasserine	0.75	0.00	0.17	0.00	0.57	0.69	0.00	0.72	0.37	0.69	1.13	0.83	0.40	0.00	0.49	0.45±0.09
	Richi El Jouf	0.48	0.00	0.83	0.50	1.02	0.81	0.21	0.71	0.74	0.75	1.11	0.71	0.17	0.91	0.90	0.66± 0.08
	Aouija Msaken	0.17	0.00	0.62	0.44	0.84	0.70	0.00	0.57	0.61	1.09	0.78	0.84	0.00	0.25	0.65	0.50± 0.09
	Mahmoudi El Jouf	0.63	0.48	0.30	0.00	1.24	0.53	0.00	0.68	0.68	0.66	0.92	0.82	0.00	0.00	0.74	0.51± 0.10
	Mahmoudi Oued Sbaihia	0.70	0.00	0.66	0.17	1.34	0.61	0.00	0.73	0.46	1.25	0.79	0.68	0.55	0.00	0.91	0.59± 0.11
ons	Chili El Jouf	0.57	0.00	0.82	0.14	0.90	0.47	0.00	0.47	0.79	0.74	0.72	0.41	0.70	0.00	0.69	0.50± 0.08
ulati	Ajimi Kasserine	0.74	0.00	0.09	0.00	1.15	0.26	0.00	0.41	0.79	0.84	0.88	0.58	0.54	0.00	0.29	0.44± 0.10
Рор	Beskri Msaken	0.75	0.00	0.46	0.72	1.13	0.56	0.00	0.62	0.76	0.76	0.78	1.06	0.68	0.00	0.75	0.60±0.09
	Chili Lansarine	0.62	0.00	0.66	0.00	1.25	0.35	0.00	0.43	0.65	1.27	0.71	0.80	0.65	0.00	0.76	0.54 <u>+</u> 0.11
	Mahmoudi Msaken	0.52	0.94	0.77	0.00	0.84	0.26	0.00	0.00	0.96	0.32	0.26	0.57	0.32	0.00	0.45	0.41±0.09
	Mahmoudi Sejnane	0.53	1.38	1.08	0.00	0.82	0.61	0.49	0.61	0.78	0.39	0.87	0.21	0.00	0.00	0.35	0.54 <u>+</u> 0.10
	Mahmoudi Amdoun	0.84	0.00	0.53	0.00	1.35	0.71	0.28	0.00	0.83	0.74	0.93	0.92	1.13	0.00	0.99	0.62±0.12
	Roussia Joumine	0.90	0.00	0.14	0.00	1.08	0.51	0.35	0.74	0.86	0.00	0.84	0.88	1.13	0.00	0.71	0.54 <u>+</u> 0.11
	Mahmoudi Joumine	0.65	0.00	0.85	0.00	1.30	0.39	0.57	0.98	0.92	0.93	0.79	0.46	0.98	0.00	0.85	0.64 <u>+</u> 0.10
	mean H'	0.63	0.20	0.57	0.14	1.06	0.53	0.14	0.55	0.73	0.75	0.82	0.70	0.52	0.08	0.68	
	H'																
$S^1$	G1	0.60	0.00	0.60	0.36	1.43	0.70	0.48	0.89	0.89	1.07	0.72	0.92	1.06	0.22	0.58	0.70±0.09
idno	G2	0.93	0.00	0.14	0.00	1.08	0.51	0.38	0.74	0.97	0.00	0.84	0.88	1.13	0.00	0.29	0.53±0.11
ic gr	G3	0.82	0.00	0.15	0.00	1.07	0.42	0.00	0.60	0.65	1.12	1.14	0.86	0.39	0.00	0.58	0.52±0.11
enet	G4	0.80	0.49	0.93	0.87	0.99	0.93	0.14	0.95	0.93	1.12	0.95	1.28	0.55	0.81	1.03	0.85±0.07
U	G5	0.80	0.05	0.75	0.27	1.13	0.72	0.00	0.44	0.72	1.06	1.00	0.97	0.80	0.00	0.97	0.64±0.10
	G6	0.37	0.58	1.14	0.00	0.86	0.62	0.54	0.58	0.81	0.43	0.88	0.23	0.00	0.00	0.00	0.47±0.09

Table 5. Estimates of Shannon–Weaver Diversity Index H', mean H' and standard error (±SE) of each of the 14 populations and seven genetic groups<sup>1</sup> based on the evaluation of lineages with 15 phenotypic traits.

G7	0.00	0.00	0.31	0.00	0.46	0.62	0.00	0.62	0.62	0.00	0.00	0.00	0.00	0.91	0.62	0.28±0.09
mean H'	0.62	0.16	0.58	0.21	1.00	0.65	0.22	0.69	0.80	0.68	0.79	0.73	0.56	0.28	0.58	

<sup>1</sup> Genetic groups defined by STRUCTURE software at K=7.

	Bidi Kasserine	Richi El Jouf	Aouija Msaken	Mahmoudi El Jouf	Mahmoudi Oued Sbaihia	Chili El Jouf	Ajimi Kasserine	Beskri Msaken	Chili Lansarine	Mahmoudi Msaken	Mahmoudi Sejnane	Mahmoudi Amdoun	Roussia Joumine
Richi El Jouf	SD, LSWA		·	•	•	•	·			•	•	•	
Aouija Msaken	SD, LSWA		]										
Mahmoudi El Jouf			SD										
Mahmoudi	NSS	SD	SD, LA,										
Oued Sbaihia			LSWA			-							
Chili El Jouf	NSS	SD	SD, LSWA				7						
Ajimi Kasserine		AUDPC, SD, LSWA, TGW	AUDPC, SD LSWA, TGW			AUDPC		_					
Beskri Msaken	NSS						AUDPC, SD, LSWA		_				
Chili Lansarine	NSS	NSS, SD	NSS, SD			NSS	NSS, AUDPC						
Mahmoudi	NSS, HS	NSS, HI,	NSS, HI,	NSS	NSS, HI, HS	NSS,	NSS, HS,	NSS, HS	NSS				
Msaken		HS, LSWA	HS, LSWA			HI, HS	AUDPC, SD				1		
Mahmoudi	NSS, HS	NSS, HI,	NSS, HI,	NSS, HI	NSS, HI, HS	NSS,	NSS, HI, HS,	NSS, HI,	HI, HS				
Sejnane	AUDPC,	HS, TGW	HS, LA,	AUDPC	AUDPC, SD	HI, HS,	AUDPC, SD	HS,					
	SD		LSWA, TGW			SD		AUDPC, TGW					
Mahmoudi	NSS	TGW	NSS, HI, LA,			ні	NSS, SD			HS, NSS	HS, AUDPC	]	
Amdoun			TGW										
Roussia	NSS, SD,	AUDPC,	HI, AUDPC		SD, LSWA	AUDPC,	SD, LSWA		AUDPC	NSS, HS,	NSS, HS		
Joumine	LSWA	TGW	TGW			SD,				AUDPC,	AUDPC,		
						LSWA				LSWA	LSWA		
Mahmoudi	NSS		LA, LSWA				AUDPC, SD			NSS, HS	NSS, HS,		AUDPC,
Joumine											TGW		LA,
													LSWA

Colour gradient: cells are getting darker as the number of phenotypic traits significantly different between the two populations is increasing. <sup>1</sup> Abbreviations: Table ESM3 1256

# Table 7. P<sub>st</sub>-F<sub>st</sub> comparison for quantitative traits measured in 273 lineages belonging to the 14 durum wheat populations. Lower values of the critical c/h<sup>2</sup> indicate a more robust inference of local adaptation.

Traits <sup>µ</sup>	P <sub>st</sub> at c/h <sup>2</sup> =1 <sup>\$</sup> (null asymption)	F <sub>st</sub>	95% CI lower	95% Cl upper	Critical c/ h <sup>2 f</sup>
	(	0.57098	0.5218	0.6180	
HI	0.9580		0.9353	0.9760	0.1118
HS	0.9692		0.9417	0.9881	0.1001
AUDPC	0.9506		0.9319	0.9680	0.1182
SD	0.9073		0.8432	0.9625	0.3008
LAS	0.5784		0.5066	0.7535	
LSWA	0.9123		0.8904	0.9399	0.1991
LA	0.7732		0.7252	0.8585	0.6129
NSS	0.9299		0.9064	0.9537	0.1670
TGW	0.9220		0.8813	0.9551	0.2179

 $^{\mu}$  Abbreviations: Table ESM3

 $c/h^2 = 1$  is the null assumption meaning that the proportion of phenotypic variance caused by additive genetic effects is the same for between-population and within-population variances (Brommer, 2011) f critical c/h<sup>2</sup> ratio was calculated according to the formula (Brommer, 2011).

1260

#### 1263 Table 8. List of the 14 durum wheat populations: locality, suspected origin/year and year of sampling.

Populations	Locality-Gouvernorate	Suspected origin/year of	Year of
		landrace*	sampling
Ajimi Kasserine	Kasserine-Kasserine	Unavailable	2015
Aouija Msaken	Msaken -Sousse	North Africa, 1909	2015
Beskri Msaken	Msaken-Sousse	Algeria, 1909	2015
Bidi Kasserine	Kasserine-Kasserine	North Africa, 1908	2015
Chili El Jouf	El Jouf -Zaghouan	France, 1932	2015
Chili Lansarine	Lansarine-Manouba	France, 1932	2015
Mahmoudi Amdoun	Amdoun-Beja	Tunisia, 1983	2017
Mahmoudi El Jouf	El Jouf -Zaghouan	Tunisia, 1983	2015
Mahmoudi Joumine	Joumine-Bizerte	Tunisia, 1983	2017
Mahmoudi Msaken	Msaken-Sousse	Tunisia, 1983	2015
Mahmoudi Oued Sbaihia	Oued Sbaihia -Zaghouan	Tunisia, 1983	2015
Mahmoudi Sejnane	Sejnane-Bizerte	Tunisia, 1983	2017
Richi El Jouf	El Jouf-Zaghouan	Tibar, Tunisia 1908/1909	2015
Roussia Joumine	Joumine-Bizerte	Bizerte, Tunisia, 1927	2017

\*Deghaïs & al. 2003, 2007; Ammar & al. 2011

Markers	Chromosomes	Size Range (bp) <sup>1</sup>	Dye colour	Motif	Primers F	Primers R	Tm <sup>2</sup>	Reference
Xgwm413	2B	88-110	Ned	(GA) <sub>20</sub>	TGCTTGTCTAGATTGCTTGGG	GATCGTCTCGTCCTTGGCA	60°C	Röder & al., 1998
Xgpw7148	3B	87-100	Fam	(AG) <sub>21</sub>	GCACACAACGACACTTGCTT	GCTTAGCTGCTTGCTTTGTG	60°C	Sourdille & al., 2004
Xgwm193	6BS	155-192	Ned	(CT) <sub>21imp</sub> (CA) <sub>8</sub>	CTTTGTGCACCTCTCTCCC	AATTGTGTTGATGATTTGGGG	60°C	Korzun & al., 1997
Xgpw2239	4AS	191-197	Vic	(CT) <sub>19</sub>	CAACCATATGCCCAGGAGAC	TGTTGCTGTCTGAAACAGGG	60°C	Sourdille & al., 2004
Xgwm285	3B	209-235	Ned	(GA) <sub>19</sub>	ATGACCCTTCTGCCAAACAC	ATCGACCGGGATCTAGCC	60°C	Somers & al., 2004
Xgpw4082	4B	211-235	Fam	(TG) <sub>23</sub> (GA) <sub>16.5</sub>	CTTTCTTTCCCCTCCTGTCC	ATCATCACAAATGCAGCGAG	60°C	Sourdille & al., 2004
Xgpw4004	5A	209-256	Pet	Unavailable	CGCCTCGGATTCTATTCTTG	CTTACTGCGGCCTTGAGTTG	60°C	GrainGenes
Xgpw2103	7AL (7B)	230-308	Vic	(TC) <sub>16</sub>	CGTATGCAGCATGGCATC	GCTATGTTGTGTGGCATTGG	60°C	Sourdille & al., 2004
Xgwm372	2A	296-331	Fam	(GA) <sub>26</sub>	AATAGAGCCCTGGGACTGGG	GAAGGACGACATTCCACCTG	60°C	Röder & al., 1998

1267 9. Description of the panel of 9 SSR (microsatellite) markers used for population genetics.

<sup>1</sup>Size range (bp: base pairs) for each marker corresponds to the minimum and maximum size identified in all the durum wheat populations; <sup>2</sup>Tm: melting temperature.