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Published Version:

Environment-oriented selection criteria to overcome controversies in breeding for drought resistance in wheat / Poggi, Giovanni Maria; Corneti, Simona; Aloisi, Iris; Ventura, Francesca. - In: JOURNAL OF PLANT PHYSIOLOGY. - ISSN 0176-1617. - ELETTRONICO. - 280:(2023), pp. 153895-153909. [10.1016/j.jplph.2022.153895]

Availability:

This version is available at: https://hdl.handle.net/11585/909906 since: 2022-12-19

Published:

DOI: http://doi.org/10.1016/j.jplph.2022.153895

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Poggi, Giovanni Maria; Corneti, Simona; Aloisi, Iris; Ventura, Francesca: Environment-oriented selection criteria to overcome controversies in breeding for drought resistance in wheat

JOURNAL OF PLANT PHYSIOLOGY VOL. 280 ISSN: 0176-1617

DOI: 10.1016/j.jplph.2022.153895

The final published version is available online at: https://dx.doi.org/10.1016/j.jplph.2022.153895

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Environment-oriented selection criteria to overcome controversies in breeding for drought resistance in wheat

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15 Abstract

Wheat is one of the most important cereal crops, representing a fundamental source of calories and 16 protein for the global human population. Drought stress (DS) is a widespread phenomenon, already 17 affecting large wheat-growing areas worldwide, and a major threat for cereal productivity, resulting 18 in consistent losses in average grain yield (GY). Climate change is projected to exacerbate DS 19 incidence and severity by increasing temperatures and changing rainfall patterns. Estimating that 20 wheat production has to substantially increase to guarantee food security to a demographically 21 expanding human population, the need for breeding programs focused on improving wheat drought 22 resistance is manifest. Drought occurrence, in terms of time of appearance, duration, frequency, and 23 severity, along the plant's life cycle varies significantly among different environments and different 24 agricultural years, making it difficult to identify reliable phenological, morphological, and functional 25 traits to be used as effective breeding tools. The situation is further complicated by the presence of 26 confounding factors, e.g., other concomitant abiotic stresses, in an open-field context. Consequently, 27 the relationship between morpho-functional traits and GY under water deficit is often contradictory; 28 moreover, controversies have emerged not only on which traits are to be preferred, but also on how 29 one specific trait should be desired. In this review, we attempt to identify the possible causes of these 30 disputes and propose the most suitable selection criteria in different target environments and, thus, 31 32 the best trait combinations for breeders in different drought contexts. In fact, an environment-oriented approach could be a valuable solution to overcome controversies in identifying the proper selection 33 criteria for improving wheat drought resistance. 34

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Keywords: Climate change, phenotyping, morphological and functional traits, target environment,grain yield

1 List of abbreviations

- 2 Chlorophyll-a (Chl a)
- 3 Chlorophyll-a singlet state (1Chl*)
- 4 Chlorophyll-a triplet state (3Chl*)
- 5 Critical Temperature (Tcrit)
- 6 Drought Stress (DS)
- 7 Drought Susceptibility Index (DSI)
- 8 Dry Weight (DW)
- 9 Fresh Weight (FW)
- 10 Full Turgor Weight (TW)
- 11 Genome Wide Association Study (GWAS)
- 12 Genotype x Environment (GxE)
- 13 Geometric Mean Productivity (GMP)
- 14 Gibberellic Acid (GA)
- 15 Grain Filling (GF)
- 16 Grain Yield (GY)
- 17 Harmonic Mean (HARM)
- 18 Harvest Index (HI)
- 19 Heat Stress (HS)
- 20 Internal CO₂ Concentration (Ci)
- 21 Leaf Area Index (LAI)
- 22 Leaf Rolling (LR)
- 23 Marker Assisted Selection (MAS)
- 24 Maximum quantum efficiency of photosystem II (Fv/Fm)
- 25 Minimal Chl a fluorescence (F₀)
- 26 Mean Productivity (MP)
- 27 Net Photosynthesis (Pn)
- 28 Non-Photochemical Quenching (NPQ)
- 29 Peduncle Length (PL)
- 30 PhotoSystem II (PSII)
- 31 Plant Height (PH)
- 32 Polyethylene Glycol (PEG)
- 33 Quantitative Trait Loci (QTLs)
- 34 Reactive Oxygen Species (ROS)
- 35 Relative Water Content (RWC)
- 36 Ribulose 1,5-BisPhosphate (RuBP)
- 37 Stomatal Conductance (g_s)
- 38 Stress Susceptibility Index (SSI)
- 39 Stress Tolerance Index (STI)
- 40 Thousand Kernel Weight (TKW)
- 41 Tolerance Index (TOL)
- 42 Transpiration (E)
- 43 Vapour Pressure Deficit (VPD)
- 44 Water Use Efficiency (WUE)
- 45 Yield Index (YI)
- 46 Yield Stability Index (YSI)
- 47

1 Wheat worldwide relevance and drought threat

2 Wheat is one of the most important cereal crops worldwide. It ranks first for global harvested area (219 Mha in 2020), and second in terms of global production (761 Mt in 2020) behind corn 3 (FAOSTAT dedicated website, visited 04/26/2022), thereby representing a fundamental agricultural 4 source of calories and protein for the human population, and an essential source of income for millions 5 of farmers (Pena-Bautista et al., 2017). Drought stress (DS) is one of the major and widespread threats 6 for cereal productivity, resulting in consistent losses in average grain yield (GY), affecting large areas 7 of Europe, Africa, Asia, Australia, South America, Central America, and North America (Daryanto 8 et al., 2016; Hazaymeh et Hassan, 2016, Zampieri et al., 2017). Climate change is projected to 9 exacerbate DS incidence and severity by increasing temperatures and changing rainfall patterns. 10 Extreme meteorological events, e.g. heat waves and drought spells, which have already increased in 11 many regions, are projected to intensify, both in intensity and duration, in most parts of the world, 12 including many wheat-growing regions that are naturally prone to drought, such as Mediterranean 13 climatic zones and semi-arid tropics (Berger et al., 2016; Jia et al., 2019). Estimating that wheat 14 production has to increase by about 70% by 2050 to guarantee food security to a demographically 15 16 expanding human population (CIMMYT, 2014; Mylonas et al., 2020), the need for breeding programs which focus on the development of wheat cultivars with improved drought resistance is 17 unquestionable (Merchuk-Ovnat et al., 2016). Increasing wheat drought resistance is thus a key goal 18 and involves not only bread wheat (Triticum aestivum L.), but also durum wheat (Triticum turgidum 19 L. ssp. durum). The latter, despite representing only 8% of total wheat production (Sukumaran et al., 20 21 2018; Igrejas et al., 2020), is typically grown in semiarid countries of the Mediterranean basin (Southern Europe, North Africa, and the Middle East) (Del Moral et al., 2003), where it is a 22 fundamental staple food. This area, where about 60% of global durum wheat is cultivated, mainly as 23 a rainfed crop (Giraldo et al., 2016), is already prone to low and erratic precipitation profiles and is 24 severely threatened by climate change, as climatic projections foresee a rise in temperatures and a 25 simultaneous reduction in precipitation for this region, with important yield losses expected (Flato et 26 27 al., 2013).

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29 Difficulties in breeding for drought resistance

Drought resistance can be defined as the capacity of a crop to successfully produce a satisfactory 30 yield in water shortage conditions (Luo et al., 2010). It is a complex quantitative trait, resulting from 31 numerous mechanisms adopted by plants to cope with water scarcity, comprising a broad spectrum 32 of morphological, physiological, biochemical and molecular processes activated at different 33 phenological stages (Rampino et al., 2006; Nezhadahmadi et al., 2013). All these adaptations can 34 confer drought resistance through drought escape (ability to complete the life cycle before stressful 35 36 conditions appears), avoidance (ability to maintain a good water status in a drought context) or tolerance (ability to withstand dehydration) (Ludlow, 1989 - cited by Kooyers, 2015). The highly 37 significant Genotype x Environment (GxE) interactions consistently reduce yield heritability in 38 drought contexts (Farooq et al., 2014), making genetic improvement very challenging for breeders. 39 Differences in drought occurrence along the wheat life cycle, in terms of time of appearance, duration, 40 frequency and severity, are highly significant among different environments (Reynolds et al., 2005). 41 Moreover, in rainfed conditions, weather fluctuations in terms of temperature and precipitation 42 distribution in the cropping season produce a consistent genotype x year interaction, further 43 complicating the direct or indirect selection for yield in an open-field environment (Ahmadizadeh et 44 45 al., 2012; Budak et al., 2013; Mohammadi et al., 2019). The concomitancy with other abiotic stresses

must also be taken into consideration (see section 2.5). In this review, we report first on the direct 1 selection for yield and highlight strengths and weaknesses of this approach. Phenological, 2 morphological, and functional traits associated with GY in drought conditions, commonly used in the 3 indirect selection for yield, are then dissected, highlighting the controversies that have emerged 4 regarding their reliability as effective breeding tools. In fact, morpho-functional traits related to wheat 5 drought resistance are well-known and documented, as reviewed by Khadka et al. (2020), but 6 evidence is still contradictory regarding the reliability of such traits as indirect selection criteria for 7 final GY. Disputes remain over which of these traits are most desirable (e.g., tall vs short genotypes), 8 as a consequence of the high variability of drought scenarios in terms of duration, timing and severity. 9 In an attempt to overcome these controversies, here we analyze the available literature, identify the 10 possible causes of these disputes and, finally, offer a proposal for the most suitable selection criteria, 11 and thus the best trait combinations, for breeders in different drought contexts. 12

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14 **1. Direct selection for yield**

Despite all the above-mentioned difficulties, the selection for yield (commonly referred to as "direct 15 selection") is still considered a valuable strategy, precisely because of the general lack of 16 understanding of physiological and molecular processes underlying DS response in wheat (Dugasa et 17 al., 2019; Patel et al., 2019). In an attempt to overcome the discrepancies arising from differences in 18 drought frequency, duration, and/or severity, several researchers have suggested selection criteria 19 based only on yield in non-stress conditions, where the low GxE interaction should permit the full 20 expression of a genotype's yield potential (Rajaram et Van Ginkle, 2001 - cited by Patel et al. 2019). 21 The main limitation of this approach is that genotypes with high potential yield under non-stress 22 23 conditions may not necessarily result in high yield under drought (Patel et al. 2019). Similarly, genotypes with satisfactory yield under stress conditions may not be the best choice in favorable 24 25 agronomic years because of a low yield potential (Mohammadi et al., 2010). A possible solution is to prefer genotypes with high yield potential in mild-stress environments, and to prefer highly resistant 26 ones with low yield potential in severe stress contexts (Voltas et al., 1999 - cited by Mohammadi et 27 al., 2010). Instead of this dichotomous selection, authors currently prefer a mid-point solution that 28 considers yield in both stress and non-stress conditions, aiming to a more exhaustive, comprehensive 29 and efficient selection, especially under unpredictable yearly drought scenarios in rainfed 30 environments (Mohammadi et al., 2010; Patel et al., 2019). For this purpose, multiple indices have 31 been proposed in the past (Fischer & Maurer, 1978; Rosielle and Hamblin, 1981; Bouslama and 32 Schapaugh, 1984; Fernandez, 1992; Schneider et al., 1997; Gavuzzi et al., 1997), as reported in Table 33 34 1.

35 Several authors have conducted experiments to identify the most suitable indices for successful breeding (Golabadi et al., 2006; Mohammadi et al., 2010; Mohammadi et a., 2011; Ahmadizadeh et 36 al., 2012; Patel et al., 2019). Based on their findings, it is advisable to conduct the selection process 37 under mild stress conditions. This is due to the fact that severe stress causes massive yield reduction 38 for most genotypes, thus reducing phenotypic variability. Moreover, different indices provide 39 different information. SSI, TOL, and YSI allow identifying genotypes with satisfactory yield in 40 stressful environments, but with low yield potential in favorable ones. On the contrary, STI, GMP, 41 MP, YI, and HARM identify high-yielding genotypes both in stress and non-stress contexts. 42 Therefore, the most suitable index for screening of drought-resistant genotypes mainly depends on 43 the environment, in particular by taking into account how frequently and severely drought years occur 44 compared to favorable ones (Table 1). 45

1 Direct selection, in general, allows breeders to bypass the understanding of the complex morpho-

2 functional and biochemical processes regulating the overall phenomenon of drought resistance.

3 However, this approach is very expensive in terms of time and resources, and is largely affected by

4 the inevitable fluctuation of agricultural years.

5 Modern Marker Assisted Selection (MAS) strategies can represent a valid solution to improve the efficiency of direct selection. Wheat landraces and wild progenitors carry an interesting allelic 6 repertoire for abiotic stresses resistance (Budak et al., 2013; Lopes et al., 2015). Typically, these 7 genotypes are characterized by satisfactory stable yields in harsh environments, but low yields in 8 high-input unstressed conditions. Differently, elite modern wheat cultivars are highly responsive in 9 terms of GY in high-input management, but in case of stress occurrence and adverse conditions their 10 yield benefit is not so obvious (Rebetzke et al., 2011). MAS can be used to transfer from wild relatives 11 to modern cultivars genes/Quantitative Trait Loci (QTLs) conferring tolerance to several biotic and 12 abiotic stresses (Merchuk-Ovnat et al., 2016). Molero et al. (2022) performed a Genome Wide 13 Association Study (GWAS) on 149 spring wheat lines, identifying promising exotic alleles for heat 14 stress (HS) tolerance in Aegilops tauschii (the ancestral donor of the D genome). Similarly, Merchuk-15 Ovnat et al. (2016), used a recombinant inbred line population derived from a cross between durum 16 wheat and wild emmer, to map QTLs conferring drought resistance. These studies demonstrate that 17 breeding programs based on the introgression of QTL alleles from wheat wild progenitors could 18 represent a valuable strategy to produce high yielding modern wheat cultivars, resistant to the effects 19 of abiotic stresses (including drought), but at the same time with no yield penalties under favorable 20 conditions. 21

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23 **2. Indirect selection for yield**

GY is the result of complex interactions with the environment and is determined not only by drought 24 conditions, but also by a wide range of other biotic and abiotic factors (Reynolds et al., 2005). For 25 this reason, a different approach is the so-called "physiological breeding", meaning an indirect 26 selection for yield, based on various traits associated with GY (Sukumaran et al., 2018). This 27 comprehensive approach is still evolving in knowledge and its applicability is often hindered by 28 contradictory information regarding several traits. For example, the relative importance of a given 29 trait in determining GY is often unclear, making it difficult to identify the most desirable single trait 30 31 associated with drought resistance. As a consequence, there is a lack of simple and reliable 32 combinations of selection criteria to be used as screening tools and the issue is still debated (Merah et al., 2001; Habash et al., 2009; Sukumaran et al., 2018). 33

In this review, we report on controversies that emerge from the literature regarding the main parameters used in screening methods, based on phenology, morphology, and physiology, so as to offer a state-of-the-art overview and to propose selection criteria combinations to be adopted in different target environments.

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39 2.1 Phenology

DS during different plant phenophases impacts GY in different ways: in the early phases, going from
germination to the double ridge stage, compromises correct crop establishment and Leaf Area Index
(LAI); from double ridge to anthesis impacts negatively on spikelet number and number of kernels

per spike; at anthesis, it reduces spikelet number and fertility; from anthesis to maturity, drought 1 compromises Grain Filling (GF), thereby lowering Thousand Kernel Weight (TKW) (Giunta et al., 2 1993; Acevedo et al., 2002; Del Moral et al., 2003). The relative importance of each of these phases 3 in terms of final yield is not absolute, but depends strongly on the environment and, therefore, on 4 when, how severely, and for how long the stress occurs. Consequently, conflicting information 5 regarding the most sensitive phases to DS can be found in the literature. For example, with respect to 6 GY, Del Moral et al. (2003), citing several authors, reports that the most sensitive period to drought 7 in durum wheat goes from the double ridge stage to anthesis, as it has negative effects on spikelet 8 number and kernels per spike. On the contrary, DS from anthesis to maturity might not have such 9 negative effects on final yield, since the duration and rate of GF can be maintained relatively stable 10 thanks to the remobilization of stored pre-flowering assimilates. On the contrary, Mohammadi et al. 11 (2019), while recognizing the possible yield loss linked to early drought because of disturbance in 12 spike development, states that durum wheat yield reduction during GF is typically more strongly 13 affected, since most of wheat grain carbohydrates derive from post-anthesis photosynthesis. This 14 apparent contradiction could be largely explained by the severity of the stress, other than obviously 15 its timing in the target environment. In fact, it is widely understood that in case of severe stress, durum 16 wheat yield reduction is mainly due to a drop in spikelet number and number of kernels per spike; 17 conversely, in the case of mild stress, yield loss is mainly due to lower TKW (Giunta et al. 1993). 18 Phenology is a plastic and flexible trait, meaning that it can be hastened or slowed down (Van Andel 19 and Jager, 1981). Figure 1 summarizes phenological alterations due to drought observed in wheat. 20 Simane et al. (1993) found that early DS (at tillering) increases the time necessary for durum wheat 21 22 to reach both anthesis and maturity. They also found that mid (flowering) and late (GF) DS did not 23 affect time to flowering, but significantly shortened GF duration. Ihsan et al. (2016), studying drought 24 effects on phenology in bread wheat, observed different results. Imposing the stress immediately after crop establishment (2 weeks after sowing), DS accelerated bread wheat phenological development 25 already from tillering, pushing all the genotypes involved in the experimentation to early maturity. In 26 particular, days to complete 50% heading and crop physiological maturity were the most reduced 27 compared to the other phenological stages. So, they concluded that both booting and GF are 28 significantly accelerated by early drought. Qaseem et al. (2019) found that also stress imposed from 29 heading reduces bread wheat days to anthesis and days to maturity. 30

31 The different genetic background of the cultivars examined might, at least in part, explain the apparently contradictory results. Different genotypes can in fact undertake different strategies to cope 32 33 with water scarcity, either drought escape or drought avoidance (Shavrukov et al., 2017). It is quite well established that terminal drought reduces GF duration, as a consequence of accelerated leaf 34 senescence, reduced photosynthesis, downregulation of enzyme activities, and sink limitation 35 (Farooq et al., 2014; Ihsan et al., 2016; Pour-Aboughadareh et al., 2020a), with a negative impact on 36 TKW. Terminal drought is typical in Mediterranean-climate regions, comprising the Mediterranean 37 Basin (North Africa, southern Europe, and the Middle East), California, central Chile, the Cape region 38 in South Africa, and SW Australia, where hot and dry summers follow wet and temperate winters 39 (Acevedo et al, 1999; Shavrukov et al., 2017; Del Pozo et al., 2019). In these areas, wheat is rainfed, 40 so DS typically emerges in spring, around anthesis, and progressively increases throughout GF 41 (Simane et al., 1993; del Moral et al., 2003; Mohammadi et al., 2019). The semi-arid tropics, where 42 wheat cropping relies on stored soil moisture, also experience terminal drought, thereby limiting GY 43 during the grain development stages (Berger et al., 2016). In contexts like these, early heading and 44 early maturity genotypes are preferable, since they avoid exposure to terminal drought by escape 45 strategies (Simane et al., 1993; Mohammadi et al, 2019; Yashavanthakumar et al., 2021). Early 46 heading and maturity should be accompanied in resistant genotypes by a longer GF period, in the 47

attempt to limit stress effects on grain starch accumulation (Yashavanthakumar et al., 2021; Ihsan et 1 al., 2016; Simane et al., 1993). Therefore, in Mediterranean-type climate areas and in semi-arid 2 tropical zones, genotypes characterized by early maturity, i.e. short pre-heading phase and increased 3 GF duration, should be preferred. This combination of traits minimizes exposure to terminal drought, 4 while assuring grain starch accumulation via prolonged photosynthesis (retarded leaf senescence) and 5 remobilization of prestored assimilates (strategy 1 in Table 2). An alternative solution to preserve 6 7 high GY by avoiding terminal drought is to prefer genotypes that can reach early maturity by combining a longer pre-heading phase, and a short (but highly efficient) GF period; this maximizes 8 earlier-forming yield components (such as tiller number or spike number per plant) and 9 photosynthetic production, while GF influences only TKW (Al-Karaki 2012; Singh et al. 2014). This 10 solution (strategy 2 in Table 2) derives from evidence highlighting that a reduced pre-heading phase 11 (and therefore early flowering) could reduce yield potential; it has, however, been experimentally 12 demonstrated that it is possible to obtain genotypes with early flowering, yet having a high yield 13 potential (Shavrukov et al. 2017). As flowering time has a strong genetic component, this trait has 14 been widely exploited in the last century by breeders operating in terminal drought contexts. The 15 predominant breeding strategy is, therefore, to allow wheat to escape terminal DS by selecting for 16 "short-cycle" genotypes, with early flowering and early maturity, while favoring GF duration 17 (Shavrukov et al., 2017; Mohammadi et al., 2019) (strategy 1 in Table 2). In such genotypes, drought 18 19 escape requires that the crop complete its short life cycle before the stress appears by a rapid and active metabolism. Therefore, the "short cycle" trait does not per se imply that the genotype is 20 sensitive or tolerant to drought. Hence, phenological traits which are desirable for one environment, 21 22 may be inappropriate for another (Simane et al., 1993).

23

24 **2.2 Morphological traits**

DS is known to cause morphological changes in wheat (Nezhadahmadi et al., 2013). The simplest, 25 and most widely used morphological traits to phenotype wheat on a large scale, are plant height (PH), 26 peduncle length (PL), and leaf rolling (LR). Reduction in PH and PL as a consequence of drought is 27 well documented (Ihsan et al., 2016; Kamrani, 2015; Nezhadahmadi et al., 2013), mostly due to the 28 dehydration of the protoplasm, leading to turgor loss and reduced cell expansion and division 29 (Saleem, 2003). Leaf rolling (LR) instead is caused by loss of turgor and poor osmotic adjustment in 30 leaf tissues (Monneveux et al., 2004). The attention of breeders has focused on these traits in order to 31 understand if, in addition to their simple application, they could represent a reliable screening 32 strategy. Recent evidence and controversies regarding these morphological traits are reported. 33

34 2.2.1 Plant height (PH)

Reduction in wheat PH has been an essential trait for breeding in the last century, and contributed to 35 36 the huge yield increase gained during the so-called Green Revolution. Reduction in plant size led in fact to new varieties less prone to lodging, and with a boosted resource portioning directed to the 37 38 spike, thus maximizing the Harvest Index (HI) (Matthews et al., 2006; De Vita et al., 2007; Rebetzke et al., 2011; Monneveux et al., 2012). The selection of semi-dwarf high performing wheats was 39 accelerated by the identification of specific dwarfing alleles (*Rht-B1b* and *Rht-D1b*), so that already 40 at the end of the 1990s, about 70% of the varieties on the market contained at least one dwarfing allele 41 (Hedden, 2003; Evans, 1998). This process of intense genetic selection has however led to a notable 42 genetic erosion in the new cultivated varieties, losing many alleles of interest, related also to tolerance 43 to abiotic stresses. A source of allelic repertoire of drought resistance is represented by tall-size wheat 44

landraces and wild emmer wheat, left behind by the bottleneck produced by twentieth century 1 breeding (Budak et al., 2013; Lopes et al., 2015; Merchuk-Ovnat et al., 2016). In general, semi-dwarf 2 modern wheats are considered to be highly performing in high-input well-managed environments, 3 while in harsh environments, characterized by stress occurrence and low-input management, yield 4 benefits are not so obvious, in comparison with tall-size landraces, which display better yield stability 5 in adverse conditions (Rebetzke et al., 2011; Blum et Sullivan, 1997). There are, however, 6 contradictory indications. If some authors indicate a positive relationship between PH and drought 7 resistance (Acevedo et al, 2002; Gao et al., 2020), or a positive correlation between PH and yield 8 parameters in drought contexts, such as fertile spikes and TKW (Bennani et al., 2016; El-Rawy et al., 9 2014), other authors find opposite relationships, and recommend the selection of small-size plants as 10 a breeding strategy for increasing drought resistance (Mohammadi et al., 2019). Finally, some authors 11 did not observe any correlation between PH and drought tolerance indices in rainfed conditions 12 (Kamrani, 2015). Again, these controversies are ascribable to differences in DS intensity, duration, 13 and moment of appearance in the different experimental conditions. Jatavev et al. (2020), 14 summarizing many research results obtained all over the world in recent years, reports that in a 15 context of strong and prolonged DS, such as in the very dry areas of Pakistan, China or Iran, breeders 16 always found a positive correlation between yield and PH, favoring tall wheats with wild-haplotypes 17 (i.e., *Rht-B1a-RhtD1a*). In the case of severe and lasting drought, stress appears already during crop 18 establishment and vegetative growth stage, and then exacerbates during the season. In such 19 conditions, semi-dwarf genotypes are clearly adversely affected. In fact, semi-dwarf Rht-B1b and 20 *Rht-D1b* wheats have reduced coleoptile length and seedling vigor, requesting a shallow sowing, 21 exposing young seedlings to higher risk of DS in case of low moisture of the very first centimeters of 22 23 soil, resulting in poor early growth and seedling emergence (Rebetzke et al., 2001a; Rebetzke et al., 24 2011; Wang et al., 2015a). Moreover, early drought exacerbates the phenotype of semi-dwarf wheats, further reducing their size, resulting in smaller biomass and LAI as well as fewer tillers and spikes, 25 ultimately causing reduced yield via shrunken and small grains (Jatayev et al., 2020). On the contrary, 26 in these contexts, tall wheats can faster produce more biomass, LAI and tillers, passing favorably to 27 the reproductive stage, and having more assimilates to be re-mobilized successively to the spike 28 (Butler et al., 2005; Jatayev et al., 2020). In the case of terminal drought, typical of the Mediterranean 29 climate, with short exposure to stress, semi-dwarf wheats have an advantage over tall genotypes, 30 being able to express their high yield potential without being thwarted by prolonged stress, especially 31 if dwarfing alleles are accompanied by early heading and early maturity traits, enabling drought 32 escape (Pour-Aboughadareh et al., 2020a; Mohammadi et al., 2019; Jatayev et al., 2020). 33

34 In summary, traditional tall-size wheats show a productive advantage in case of early, severe and prolonged DS. On the contrary, semi-dwarf wheats are preferable in contexts of mild stress and / or 35 terminal stress, which allows them to express their high yield potential. Several *Rht* genes responsive 36 to gibberellic acid (GA) have been identified, different from Rht-B1b and Rht-D1b that are GA-37 insensitive; these (e.g., Rht8 and Rht13) are responsible for greater coleoptile growth and seedling 38 vigor, thus favoring crop establishment in water deficit conditions. In fact, greater coleoptile length 39 and seedling vigor allow deeper sowing, assuring germinating seeds to reach soil moisture in deeper 40 layers, without compromising a good emergence (Rebetzke et al., 2007). Furthermore, some Rht 41 genotypes show a less pronounced reduction in stature (such as Rht-B1bE529K, Rht24 and Rht 25). 42 These *Rht* genes could pave the way towards new wheat ideotypes for drought contexts, combining 43 the productive advantages of semi-dwarf genotypes in mild-stress and well-watered environments, 44 and a lower susceptibility in case of early and severe stress (Rebetzke et al., 2011; Wang et al., 2014; 45 Wang et al., 2015a; Jatayev et al., 2020; Mo et al., 2018a; Mo et al., 2018b; Tian et al., 2017). 46

1 2.2.2 Peduncle length (PL)

Like PH, also PL, typically shortened by water deficiency, has received great attention. The peduncle 2 is the last internode of the main stem, located immediately below the initial tip of the spike. The 3 importance of the peduncle in supporting the GF process, via remobilization of pre-stored 4 assimilates, especially in contexts of DS in which the contribution of late photosynthesis to GF is 5 reduced, is well known (Yang and Zang, 2006; Vosoghi Rad et al., 2022). Some authors have 6 suggested that the peduncle could also play an active non-negligible role in late photosynthesis by 7 having anatomical and physiological advantages over the flag leaf, e.g., higher stomatal density 8 (Kong et al., 2010). Nevertheless, the relationship between PL and yield under DS conditions remains 9 controversial, and depends on the different environments in which the studies are conducted. Authors 10 observing the highest PL reduction, and the highest positive correlation between PL and GY in 11 drought conditions, impose water shortage quite early in the life cycle (Bogale et al., 2011; Soares et 12 al., 2020; Ahmad et al., 2020). When water shortage appears later, no significant correlation emerges 13 between PL and GY (Villegas et al., 2007; Pour-Aboughadareh et al., 2020a; Vosoghi Rad et al., 14 2022), and significant PL reduction due to DS is not always observed, as the peduncle reaches its 15 maximum length within 5 - 10 days after anthesis (Vosoghi Rad et al., 2022). No significant 16 correlation between PL and tolerance indexes emerges comparing neither rainfed nor irrigated wheat 17 in Iran (Kamrani et al. 2015). Peduncle dry matter translocation to the spike, and its efficiency, under 18 DS conditions, might not be related to PL, but to its specific weight (given by the ratio between 19 peduncle weight and length). Furthermore, the lower internodes seem to provide the most significant 20 contribution in terms of dry matter remobilization to the spike (Vosoghi Rad et al. 2022). While 21 Vosoghi Rad et al. (2022) reported a positive correlation between peduncle specific weight and yield 22 in drought conditions, Villegas et al. (2007) found the opposite relationship. They identified a positive 23 correlation between peduncle weight and drought susceptibility index (DSI), suggesting that a lighter 24 25 peduncle could be an indicator of drought resistance, insofar as a heavier peduncle would be characterized by a higher proportion of structural carbohydrates over water-soluble ones that can be 26 translocated to the spike. A further complexity is given by the fact that PL reduction is also an effect 27 of *Rht* genes (Rebetzke et al., 2011), and in general PL gives the highest contribution to PH (Vosoghi 28 29 Rad et al., 2022), whose complicated relationship with yield in arid and semi-arid contexts has been treated above. 30

31 2.2.3 Leaf rolling (LR)

Another morphological trait, commonly taken into consideration in cereals for the selection of 32 drought resistant genotypes, is LR, phenomenon that allows wheat to reduce the leaf surface exposed 33 to solar radiation by about 41%-48% (Clarke, 1986), consequently reducing leaf temperature and 34 35 transpiration (E). Furthermore, rolling creates a microclimatic condition of greater humidity around the leaf surface, allowing the stomata to remain open, thus favoring photosynthesis (Kadioglu and 36 Terzi, 2007), as well as enabling the interception of atmospheric water via foliar water uptake (Ali et 37 al., 2022). While LR is recognized as an important drought avoidance mechanism (O'Toole et al. 38 1979; Clarke, 1986; Kadioglu and Terzi, 2007), its contribution to yield is not much investigated in 39 40 wheat, and the results are not always univocal. Bogale et al. (2011) found a positive and significant correlation between LR and GY in durum wheat grown in pots under water deficit during anthesis. 41 On the contrary, Monneveux et al. (2004) highlights no significant impact of LR on GY, when bread 42 wheat in open-field experiments is subjected to mild long-lasting DS after anthesis. Moreover, several 43 authors report that the degree of LR poorly correlates with leaf water potential and leaf water loss 44 rate in both bread and durum wheat (Clarke et al., 1986; Yang et al., 1991), with LR starting only 45 after consistent water loss, probably due to a good capacity for osmotic adjustment, which delays the 46

phenomenon (Clarke et al., 1986; Schonfeld et al., 1988). The contradictory reliability of LR as an 1 effective breeding tool is confirmed by genomic prediction analysis that produced an estimated 2 breeding value of 0.16 for the trait, which is much lower, for example, compared to the estimated 3 value obtained for PL (0.72) (Alemu et al., 2021). However, some genotypic variations in LR have 4 been reported (Rebetzke et al., 2001b - cited by Sirault et al., 2015), and QTLs are starting to emerge 5 (Peleg et al., 2009; Alemu et al., 2021). LR is usually phenotyped with a visual score, which assigns 6 to each leaf a value on a scale of 1-3 or 1-5 (Bogale et al., 2011; Olivares-Villegas et al., 2007). 7 Therefore, the operator's subjectivity can strongly influence the assigned score, making objective 8 phenotyping difficult; the task is further complicated by the possibility of a strong intra-genotypic 9 variability in open-field plots (Clarke et al., 1986). Recently, new objective, quantitative, and 10 repeatable protocols to compare genotypic differences in LR have been proposed (Sirault et al., 2015), 11 based on computer vision analysis of transverse leaf sections. Up to date, LR, which occurs only after 12 strong dehydration when other strategies have failed (e.g., osmotic adjustment), does not appear as a 13 solid breeding tool when stress appears suddenly, and/or is mild and long-lasting (Clarke et al. 1986). 14 Nevertheless, LR could be an interesting selection criterion in case of severe progressive drought. Its 15 genetic basis still remains to be investigated, but there is the possibility that the importance of this 16 trait may grow in the future, given the opportunities supplied by new objective phenotyping 17 approaches. 18

19

20 2.3 Early vigor parameters

Early drought is critical as it can jeopardize proper crop establishment, the first essential component 21 in the expression of yield potential. Satisfactory seed germination and seedling growth, ensuring a 22 successful crop establishment, define together the early vigor. Seed water absorption, germination 23 percentage, seedling vigor index (calculated as the product of shoot length and germination 24 25 percentage, divided by 100), root length, and coleoptile length are the main early vigor traits that can be inhibited under water deficit conditions (Acevedo et al., 2002; Almaghrabi et al., 2012; Ahmad et 26 al., 2018). All these parameters, which are quick, easy, and cheap to phenotype, have attracted the 27 attention of breeders as tools for improving wheat drought resistance. Since the 1960s, selection for 28 these traits is generally carried out in germinating seeds using polyethylene glycol (PEG) (Parmar et 29 Moore, 1966). PEG simulates drought by inducing a uniform and controlled osmotic stress, with no 30 direct physiological damage as it is inert, non-ionic, and impermeable to cell membranes, thus 31 influencing osmotic potential without entering the apoplast (Hohl and Schopfer, 1991). In order for 32 PEG to mimic drought without altering plant hydraulic properties (therefore without being directly 33 absorbed by the plant) it is essential to use a high-molecular-weight (> 6000) PEG (Kaufmann et 34 Eckard, 1971). For these reasons, the use of PEG-6000 to simulate water deficit has been sustained 35 by several authors (Almaghrabi et al., 2012; Van den Berg et al., 2006; Tuberosa, 2012). Significant 36 differences among wheat genotypes for the aforementioned traits emerged in different studies, 37 suggesting the existence of genetic variability to be exploited for obtaining breeding gains (Pour-38 Aboughadareh et al., 2020b; Almaghrabi et al., 2012; Moayedi et al., 2009; Dhanda et al., 2004). 39 Despite representing an interesting tool for targeting early vigorous genotypes, the main drawback of 40 many PEG-6000 screenings is that they stop at the seedling stage, without considering the rest of the 41 life cycle. Consequently, promising genotypes identified as "drought resistant", and recommended as 42 suitable for cultivation under water deficit conditions, may not show such features if drought appears 43 later in the life cycle. Even though some evidence of an interesting correlation between seedling 44 45 growth under DS and adult plant drought resistance is starting to emerge (Dodig et al., 2015), future

research requires further validation of preliminary seedling screening results via pot or open-field 1 experiments. At any rate, early vigor remains a trait of interest to ensure successful crop establishment 2 and early drought resistance, and preserve final yield. In addition, regardless of early drought 3 conditions, a quick canopy cover reduces water loss from the ground, thereby preserving soil water 4 content for the future crop, i.e., improving water use efficiency in the crop cycle (Blum 2009; 5 Rebetzke et al., 2011; Tuberosa, 2012). 6

7

8 2.3.1 Number of tillers

9 Early-season drought reduces the number of productive tillers (Acevedo et al., 2002; Sarto et al., 2017), thus reducing the spike number and, ultimately, GY (Khadka et al., 2020). In the case of early 10 drought, selecting for this trait is a valuable strategy, as it allows to preserve the above-mentioned 11 early-forming yield components. However, to select for a high tiller number may not always be 12 advantageous. In fact, this approach might not be advisable under terminal drought scenarios as the 13 total number of non-productive tillers could be higher. This would increase crop water consumption 14 before anthesis, limiting the amount of stored available water in the soil during terminal drought 15 (Elhani et al., 2007; Ribot et al., 2012). Having fewer tillers is also a typical characteristic of semi-16 dwarf high-yield genotypes (Jatayev et al., 2020), as limiting the number of tillers allows the plant to 17 invest fewer resources on structural carbohydrates, maximizing the HI. As mentioned before, the 18 high-yield potential of these genotypes is maintained in the event of mild drought, but is typically 19 compromised in case of severe stress. In these scenarios of strong drought, taller genotypes with a 20 higher number of tillers have an advantage, preserving the ability to successfully produce GY. In fact, 21 Wasaya et al. (2021), evaluating 14 wheat genotypes, observed no correlation between PH and GY, 22 and between tiller number and GY, in the case of mild drought; on the contrary, both these 23 correlations were positive and highly significant under severe drought. Consequently, the number of 24 tillers could represent a valuable breeding tool in early, long-lasting, and severe water deficit 25 conditions. 26

2.4 Physiological traits 27

Functional and biochemical responses to DS are well documented in wheat and mainly involve 28 stomata closure to prevent excessive water loss. As a consequence, internal CO₂ concentration (Ci) 29 drops causing net photosynthesis (Pn) inhibition and production of Reactive Oxygen Species (ROS) 30 leading to oxidative damage, so that plant growth and productive performance are strongly 31 compromised (Nezhadahmadi et al., 2013; Ahmad et al., 2018). A valuable breeding target is, 32 therefore, to select for varieties able to maintain a good water status in drought contexts. 33

2.4.1 Relative Water Content (RWC) 34

Different authors have proposed leaf Relative Water Content (RWC) as a better indicator of leaf water 35 status compared to leaf water potential for wheat. RWC is easy to measure and calculate, by simply 36 weighing fresh leaf tissue after excision (FW), at full turgor (TW) and after drying (DW), following 37 the formula (Merah et al., 2001): 38

 $RWC(\%) = [(FW - DW)/(TW - DW)] \times 100$ 39

By closely reflecting the balance between leaf water supply and E rate, it highlights significant genetic 40

- variation with high heritability, and is informative at different stages of the plant's life cycle. 41 Moreover, it has proven to positively correlate with E and photosynthetic rate, HI, and GY in both
- 42

durum and bread wheat (Merah et al., 2001; Moayedi et al., 2011; Mohammadi et al., 2019; Ahmad
et al., 2018). RWC is considered a reliable tool for cereal breeding (Teulat et al., 2003; Rampino et
al., 2006). In fact, as long as the crop can maintain a high leaf RWC, plant physiology is unaffected
by stressful environmental conditions, allowing good productive performances (Beltrano et al., 2006).
On the contrary, when leaf RWC drops, plant water balance is disrupted, affecting normal physiology
and, ultimately, yield (Ahmad et al., 2018). If RWC drops, wheat implements a set of biochemical
and functional responses; some of them are extensively treated in the literature, but their relative

- 8 importance as breeding tools is still unclear.
- 9

10 2.4.2 Osmotic adjustment

Osmotic adjustment is a typical mechanism implemented by wheat under DS conditions, in the 11 attempt to avoid dehydration effects (Nezhadahmadi et al., 2013). The amino acid proline is the main 12 osmoregulator accumulated, especially in leaves (Ahmad et al., 2018). It stabilizes membranes, 13 prevents enzyme inactivation and represents a source of rapidly available nitrogen after stress relief 14 (Dib et al., 1994; Rampino et al., 2006; Ahmad et al., 2018). Thus, high proline levels are generally 15 considered an indicator of drought tolerance (Nezhadahmadi et al., 2013). Nevertheless, the effective 16 role of proline accumulation in enhancing DS resistance and its reliability as an effective screening 17 tool in wheat breeding programs, is still unclear and controversial (Moayedi et al., 2011; Hong-Bo et 18 al., 2006; Rampino et al., 2006). For example, imposing DS to 10-days-old Triticum durum and 19 Aegilops seedlings, Rampino et al. (2006) concluded that increasing proline parallels RWC decrease, 20 as a consequence of dehydration perception by the plant. Accordingly, proline accumulation was 21 higher and faster in sensitive genotypes, which had lower RWC. On the contrary, several authors 22 found a positive correlation between the ability of a genotype to maintain a high RWC and proline 23 accumulation, both at seedling and later growth stages, concluding that osmotic adjustment mediated 24 by proline has a key role in reducing cell water loss in drought conditions. These data suggest that 25 targeting high-proline drought-resistant genotypes is a valuable breeding tool (Hong-Bo et al., 2006; 26 Bayoumi et al., 2008; Moayedi et al., 2011). Saeedipour et al. (2013), comparing tolerant vs sensitive 27 genotypes, imposed stress after anthesis and observed higher and faster proline accumulation in the 28 29 tolerant ones, thus reaching opposite conclusions as compared to Rampino et al. (2006). Moreover, Chandrasekar et al. (2000), comparing durum and bread wheat genotypes, observed lower RWC 30 reduction under DS in durum wheat, but higher proline accumulation in bread wheat, concluding that 31 proline accumulation does not contribute to the higher level of drought tolerance of tetraploid relative 32 to hexaploid wheat. Finally, if some authors indicate a significant positive correlation between proline 33 accumulation and GY in stress conditions (Dib et al., 1994; Bayoumi et al., 2008), other authors find 34 35 that this correlation is weak and non-significant (Mwadzingeni et al., 2016a). These contradictory indications are due to the fact that many studies on proline are carried out in early stages of the life 36 cycle, without taking into consideration repeated measurements in the most critical phases of 37 38 phenological development and often without taking into account final GY. Furthermore, there is a lack of comprehensive studies based on systematic screening of a large germplasm pool. As a result, 39 40 information about proline's relationship with yield and other stress-related physiological parameters in critical growth stages is scarce and controversial. Mwadzigeni et al. (2016a) aimed to solve this 41 problem by conducting a study on 96 wheat genotypes, both in the greenhouse and open field, in two 42 agronomic seasons and by imposing terminal DS (after 50% heading). They found that proline content 43 had a non-significant correlation with GY, yield components (except for a negative correlation with 44 TKW), and other agronomic traits. Thus, using proline content as a biochemical marker of breeding 45 interest is not supported. However, while free proline accumulation does not represent a valuable 46

breeding tool *per se*, its contribution to drought resistance ought to be considered in conjunction with
 other fundamental traits of functional adaptation, which could depend on the genotype, the presence

- 3 of other stressing factors, and the phase of the life cycle.
- 4 2.4.3 Stomatal conductance, transpiration and Water Use Efficiency (WUE).

Since the flag leaf is traditionally recognized as a main contributor to GY formation (Evans et al., 5 1970; Sylvester-Bradley et al., 1990), water deficit effects on its physiology and biochemistry have 6 aroused great interest over time, in the attempt to find reliable breeding tools. DS is known to cause 7 stomata closure, leading to a drop in stomatal conductance (gs) thereby reducing both E and Pn 8 (Subrahmanyam et al., 2006; Nezhadahmadi et al., 2013; Ahmad et al., 2018). The decline in g_s limits 9 leaf tissue water loss, but at the same time inhibits CO₂ uptake, hence photosynthesis and eventually 10 dry matter accumulation (Sallam et al., 2019). Thus, while reduced g_s helps to preserve leaf water 11 status via reduced E (Farooq et al., 2009), on the other hand high gs positively correlates with GY 12 (Bahar et al., 2009). Therefore, drought-resistant genotypes must be able to maintain high gs in 13 drought conditions, while maintaining high Pn and, ultimately, yield (Saeidi et al., 2015; Wasaya et 14 al., 2021). Once again, the solution to this dilemma relies on the environment. Stomatal closure is, in 15 fact, a drought avoidance mechanism; as such, it is more closely linked to a plant's ability to survive, 16 than to produce, thus limiting yield potential. Li et al. (2021), conducting a meta-analysis on wheat 17 drought adaptation mechanisms, concluded that in severely water stressed environments, drought 18 avoidance mechanisms (such as stomatal closure) confer an advantage in terms of GY production, 19 while avoidance strategies are not effective in the case of moderate or mild stress. Thus, gs inhibition 20 may be a target trait in harsh environments to ensure plant survival and preserve GY, while genotypes 21 able to maintain high g_s may have higher yield potential in mildly stressful contexts. 22

Also, gs is strictly connected to the concept of Water Use Efficiency (WUE), namely the ratio between 23 plant total biomass and volume of consumed water. In fact, reduced g_s means reduced E, hence lower 24 water consumption by the plant. For decades, breeders have focused on improving WUE, considering 25 it synonymous of high yield performance with lower water use in drought stressed environments 26 (Blum, 2009). However, concerns about this strategy have emerged, mainly because WUE genotypic 27 variability is driven more by differences in water consumption than biomass production (Blum, 2005; 28 29 Blum, 2009). Consequently, excluding severe DS conditions in which stomatal closure is a necessary drought avoidance strategy for plant survival, selecting for genotypes with higher WUE means 30 selecting for genotypes that are affected by low soil moisture level, leading to stomatal closure and 31 reduced dry matter accumulation, rather than genotypes able to extract more water from the soil 32 (Blum, 2009; Tuberosa, 2012). 33

Further complexity derives from those plant water losses not linked to CO₂ fixation, i.e., non-stomatal 34 E and nocturnal g_s. Non-stomatal E refers to water loss via evaporation through the leaf cuticle, which 35 accounts for up to 30% of leaf water loss in stressful conditions (Hasanuzzaman et al., 2018). 36 Nocturnal E is also not negligible, representing 14-55% of daytime water losses for wheat (Schoppach 37 et al., 2014; Schoppach et al., 2020; McAusland et al., 2021). In conclusion, gs is a trait of interest to 38 improve wheat drought resistance. Selection should be oriented to genotypes with low gs only in a 39 context of severe water shortage. In moderate to mild drought scenarios, genotypes able to extract 40 more water from the soil and thus able to maintain high gs to sustain CO₂ fixation and GY should be 41 preferred. In parallel, attention should be shifted from WUE to the maximization of stomatal E while 42 minimizing water losses not related to CO₂ assimilation (non-stomatal E and nocturnal g_s). Non-43 44 stomatal water loss reduction can be achieved by increasing leaf epicuticular waxiness (Richards et al., 1996; Acevedo et al., 2002), and evidence of genotypic variation to exploit nocturnal stomatal E
in wheat is starting to emerge (Schoppach et al., 2020; McAusland et al., 2021).

Finally, the use of g_s as a breeding tool in open-field or greenhouse conditions is complicated by the 3 fact that environmental conditions interacting with stomatal opening/closure can change rapidly and 4 several times during the day (depending on cloud cover, solar radiation, and relative humidity), 5 making it difficult to measure this parameter reliably and, therefore, to perform unbiased 6 comparisons on a large number of genotypes and in multiple replicate (Tuberosa, 2012; Monneveux 7 et al., 2012). In fact, gs does not depend exclusively on drought but is regulated also in response to 8 Vapour Pressure Deficit (VPD), defined as the difference between the saturation (i.e. the maximum 9 amount of water vapor that the air can hold) and actual vapor pressure in the atmosphere (Grossiord 10 et al., 2020). Higher VPD determines a higher evapo-transpirative demand of the atmosphere, 11 increasing E. So, as VPD increases, E increases till the plant is unable to satisfy the request, and 12 closes the stomata (Franks and Farquhar, 1999). 13

So, independently of water availability in the root zone, if leaf -to-air VPD increases too much (for 14 example in case of high temperature or wind speed), plants respond by closing the stomata to prevent 15 excessive water loss (limiting E). This happens when atmospheric evapotranspirative demand is too 16 high and cannot be counterbalanced by water absorption and transport towards the leaf. This limit 17 value varies from species to species, and depends mainly on the hydraulic conductivity characteristics 18 of the xylem (Franks and Farquhar, 1999). There is no unanimous consensus on the specific 19 mechanisms driving stomatal closure in response to changes in VPD. It could involve active sensing 20 of the water status in the leaf or in the stomatal guard-cells, likely mediated by hormonal signals like 21 abscisic acid (Grossiord et al., 2020). When there is no soil water shortage, but still the crop 22 experiences stomatal closure in response to high VPD, decreased stomatal sensitivity to VPD has 23 been proposed as a possible contributor to the relationship between gs and yield in grain crops 24 (Richards, 2000). 25

Schoppach et al. (2012) revealed significant phenotypic diversity across wheat genotypes for g_s and 26 E sensitivity to VPD. The researchers concluded that genotypes with early stomata closure in response 27 to both atmospheric or edaphic induced water stress can store more water in the soil and utilize it 28 more sparingly during drought episodes. On the contrary, the drought tolerance of gs insensitive 29 genotypes to soil drying and/or increasing VPD may represent a different strategy, favoring CO₂ 30 fixations over water conservation. Those findings are consistent with the idea that drought resistance 31 is highly dependent on drought scenarios, with conservative and insensitive genotypes preferable in 32 case of severe or mild-moderate stress, respectively. Therefore, as noted by Medrano et al. (2002), gs 33 is responsive to all factors related to leaf water status, thus representing an integrative breeding trait for 34 the overall effects of plant water stress. 35

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- 37

38 2.4.4 Flag leaf net photosynthesis (Pn)

Flag leaf Pn is considered a major contributor to GY, mostly during the GF stage, when other leaves start to senesce (Evans et al., 1970; Sylvester-Bradley et al., 1990; Loss et Siddique, 1994). As drought causes a drop in RWC and g_s, Pn is inhibited. Selection for genotypes able to maintain high Pn rates under water deficit is thus a clear target for wheat breeders, especially in post-anthesis (Inoue et al., 2004). If decreased Pn depended solely on decreased g_s, selecting and phenotyping for both

traits would be redundant. However, in several cases, no significant correlation was found between 1 gs and Pn in drought-stressed wheat, and reduced gs was not considered the main cause of Pn decrease 2 for both bread and durum wheat exposed to drought (Bogale et al., 2011; Siddique et al., 1999). Other 3 authors instead indicate a strong correlation between Pn and gs drop in wheat under drought (Mu et 4 al., 2021; Sikder et al., 2015; Wang et al., 2015b). This is due to the fact that in C3 plants drought can 5 cause non-stomatal Pn limitations, related to impaired ATP synthesis and ribulose 1,5-bisphosphate 6 (RuBP) regeneration, altered leaf photochemistry, decreased Rubisco activity, and permanent 7 photoinhibition, as reported in Medrano et al. (2002) and Flexas et Medrano (2002). The prevalence 8 of stomatal or non-stomatal limitations to Pn depend on stress severity and duration, as well as on the 9 contextual presence of other stress sources. For example, in a Mediterranean climate, terminal 10 drought is typically accompanied by HS (Li et al., 2013). Terminal HS in wheat is known to inhibit 11 Pn due to increased photorespiration, as Rubisco kinetics is affected (Farooq et al., 2011). Some 12 studies suggest that Pn inhibition is mainly driven by stomatal closure in mild stress - even if 13 impairment in ATP synthesis and RuBP regeneration have been demonstrated too - while 14 photosynthetic limitation to Pn is predominant under severe drought (Ashraf et al., 2013; Athar, 2005; 15 Flexas et Medrano, 2002). Regardless of stomatal or non-stomatal limitation, the fact that some 16 authors find a significant correlation between flag leaf Pn and GY under drought, or between flag leaf 17 Pn and stress resistance (Zheng et al., 2011; Inoue et al., 2004) while others do not (Bogale et al., 18 19 2011; Guóth et al., 2009) raises some doubts about the use of Pn as a breeding target. One must consider that GY, as well as Pn and other gas exchange parameters, are highly integrative traits, 20 strongly dependent on micro-climatic conditions during sampling. Moreover, the contribution of flag 21 22 leaf Pn to GY may vary among genotypes and during the life cycle, depending on other assimilates 23 sources (e.g., remobilization from reserves, ear photosynthesis, etc.) (Asseng et Van Herwaarden, 24 2003; Li et al., 2017; Ding et al., 2018). Maintaining high photosynthetic capacity in the flag leaf is clearly an interesting and desirable trait to select for. Nevertheless, it is not easy to phenotype (given 25 26 the high plasticity of the trait during the day and/or over time) and should be selected in combination with other favorable traits to maximize the positive impact on GY. 27

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29 2.4.5 Chlorophyll performance parameters

When Chlorophyll-a (Chl a) is excited to the singlet state (¹Chl*) as a result of light absorption, under 30 CO₂-limiting conditions (e.g. stomata closure) the amount of harvested light energy and generated 31 reducing power can easily exceed the rate of its consumption by the photosynthetic machinery. 32 Therefore, the plant can employ several mechanisms to dissipate the excess of excitation energy, i.e. 33 emission of fluorescence and heat. Both thermal dissipation and photochemistry mechanisms reduce 34 the amount of emitted fluorescence, and non-photochemical processes that quench chlorophyll 35 fluorescence are collectively called Non-Photochemical Quenching (NPQ). Otherwise, energy can be 36 dissipated via the triplet state (³Chl*), a significant valve for excess excitation that can however 37 generate ROS, extremely damaging to the photosynthetic apparatus (Müller et al., 2001; Grieco et 38 39 al., 2020). Altered leaf photochemistry and energy flow in Photosystem II (PSII), the most vulnerable part of the photosynthetic apparatus (Sherstneva et al., 2022), have received great attention, thanks to 40 the possibility to easily assess the downregulation of photosynthesis in living plants, measuring 41 chlorophyll fluorescence parameters by specialized equipment and easy-to-use instruments. In 42 particular, Fv/Fm, i.e., maximum quantum efficiency PSII, is indicated by several authors as a 43 selection criterion to improve wheat drought tolerance and/or GY in drought contexts (Flagella et al., 44 1995; Araus et al., 1998; Almeselmani et al., 2011; Farshadfar et al., 2014; Mohammadi et al., 2019). 45

This is based on evidence that a decrease in Fv/Fm measured on dark-adapted leaves is related to 1 down- regulation of photosynthesis and photoinhibition (Maxwell and Johnson, 2000). Nevertheless, 2 some authors argue that PSII primary photochemistry, and thus Fv/Fm, is quite insensitive to DS (Lu 3 et Zhang, 1999; Subrahmanyam et al. 2006). This is supported by studies that do not find a significant 4 correlation between chlorophyll fluorescence parameters and GY in drought-stressed wheat, 5 concluding that these are not proper tools to phenotype wheat genotypes for drought resistance 6 (Bogale et al., 2011; Pour-Aboughadareh et al., 2020a). These contradictory conclusions may be 7 explained by differences in stress severity and by the presence of other stressing factors that inhibit 8 wheat PSII maximum quantum efficiency, e.g. HS and high radiation intensity (Sharma et al., 2012; 9 Sharma et al., 2015). In fact, Fv/Fm seems to be compromised by DS only in cases of severe 10 dehydration (Živčák et al., 2008). Thus, when assessing its suitability as a screening tool for wheat 11 drought tolerance, misleading results may be due to the concomitant presence of confounding factors, 12 such as intense solar radiation and high temperature. On the other hand, field conditions of terminal 13 drought accompanied by high temperature and solar radiation are typical of the Mediterranean 14 climate, making Fv/Fm an interesting breeding tool for this type of context. 15

Another widely used Chl a fluorescence metrics to measure PSII damage in case of HS, is the critical 16 temperature (T_{crit}) above which minimal Chl a fluorescence (F₀) rises rapidly, indicating incipient 17 damage to PSII (Hűve et al., 2011). Being *T*crit associated with increased thylakoid membrane fluidity 18 and disruption of the light-harvesting antennae (Geange et al., 2021), it has been used to examine the 19 vulnerability to HS across plant species (Sharma et al., 2020). So, as for Fv/Fm, T_{crit} can represent an 20 interesting breeding tool for wheat, in those contexts where drought is accompanied by high 21 temperature and high solar radiation, leading to excessive leaf temperature. Posch et al. (2022) 22 combining data from both field trials and controlled-environment, and meta-analysis on previously 23 published data, highlighted that leaf T_{crit} varies widely among wheat species and genotypes (by up to 24 20 °C). Moreover, T_{crit} also shows a significant genotype by phenology interaction, mainly due to an 25 increasing trend in T_{crit} as plants progressed from heading to anthesis and GF. Genotypes with 26 enhanced thermal safety margins, thus increased T_{crit} , could be particularly important in heat stressed 27 water-limited environments, considering that the concomitant presence of both stresses highly limit 28 29 g_s and transpirational cooling, resulting in increased leaf temperature (refer to 2.5 section).

Exploring more complex photosynthetic performance parameters than fluorescence alone, such as 30 NPQ, may allow to better discriminate drought resistant wheat genotypes. However, most of the 31 studies on NPQ variations in response to drought were carried out in controlled environments, on a 32 33 small number of genotypes. According to the experimental conditions, i.e. phenological phase of stress onset and drought duration and severity, NPQ can increase (Subrahmanyam et al., 2006; Zlatev 34 2009; Guóth et al., 2009; Zivcak et al., 2014; Yaghoubi Khanghahi et al., 2020; Zhu et al., 2020; 35 36 Grieco et al., 2020), decrease (Shangguan et al., 2000; Sherstneva et al., 2021) or not vary (Yudina et 37 al., 2020), making it difficult at the state of the art to determine the potentiality of this trait for highthroughput screening. Moreover, NPQ might respond differently to DS and HS (Zhu et al., 2020), 38 complicating the analysis in those environments where drought is frequently accompanied by heat 39 waves. Recently, Grieco et al. (2020) provided a base for developing wheat phenotyping for DS 40 41 tolerance based on NPQ. This study represents a breakpoint, since the majority of previous studies emphasize photodamage as the only cause of the downregulation of photosynthesis in case of harsh 42 DS conditions, while Grieco et al. (2020) monitored NPQ kinetics in slowly increasing levels of DS 43 in wheat, as it usually occurs in the field. Authors noted that NPQ values started to increase 13-17 44 days after increasing drought imposition, when soil substrate reached 30% of field capacity, with no 45

- 1 detectable change in Fv/Fm. Subsequently, after 18-24 days of increasing DS (substrate at 20% of
- 2 field capacity), both NPQ and Fv/Fm increased, indicating PSII damage. However, Grieco et al.
- 3 (2020) limited their investigation on DS at early phenological stages (BBCH 23 33), thus at the
- 4 moment not giving information on NPQ suitability as a breeding tool in case of terminal drought.
- 5
- 6

7 2.5 Combined heat and drought stress

8 Several concomitant environmental factors may have a non-negligible role on the opportunity to use a specific indicator as a proxy to effectively address DS resistance. In the climate change scenario, 9 HS is the main environmental constraint that typically affects wheat simultaneously with drought, 10 especially during reproductive and GF stages (Barnabas et al., 2008; Li et al., 2013; Hlaváčová et al., 11 2018). GF is affected by short periods (few days) of temperatures higher than 34 ± 2 °C (the so-called 12 heat-shock), or by prolonged periods of daily maximum temperature up to 32°C (chronic heat stress) 13 (Al-Khatib and Paulsen, 1984; Yang et al., 2002; Farooq et al., 2011). So, heat shock conditions are 14 those typically occurring during heat waves (3-7 days), while chronic HS is produced by the general 15 rise in temperatures during reproductive and GF stages. 16

17 Heat and drought impacts on wheat physiology are largely overlapping. In fact, HS induces higher evapotranspiration, RWC decreases, and stomata closure, leading to reduced gs, E and Pn (Faroog et 18 al., 2011). HS causes a reduction in Pn also as a consequence of increased Rubisco oxygenase activity 19 at high temperatures (photorespiration). Moreover, Pn drop due to high temperatures is attributed also 20 to NPQ, disruption in the structure of chloroplasts, and proteins/enzymes degradation/inactivation 21 due to oxidative damages (Farooq et al., 2011). So, both HS and DS affect Pn directly (stomata 22 closure) and indirectly (downregulation of photosynthetic metabolism), as well as both stress lead to 23 oxidative damage, i.e. lipid peroxidation, membrane disruption, protein degradation and enzymes 24 inactivation, reducing the functionality of leaves (Hlaváčová et al., 2018). 25

Studying the effects of single and combined DS and heat-shock on wheat physiology at anthesis, Wang et al. (2010) observed that the stress combination resulted in a greater decrease in Pn, g_s and E of flag leaves than drought or heat-shock alone. Similarly, Hlaváčová et al. (2018) observed that heatshock during anthesis and GF significantly increased the negative impact of drought on CO₂ assimilation rate. Combined terminal drought and chronic HS resulted in more severe consequences on Pn and g_s , considerably reducing growth and yield traits (Shah and Paulsen, 2003; Prasad et al., 2011).

33 Therefore, the aforementioned physiological parameters remain valid breeding tools for drought resistance even in case of concomitant HS. Indeed, in mild terminal drought conditions, genotypes 34 able to extract more water from the soil, thus maintaining higher gs to sustain Pn, should be preferred 35 also if drought is accompanied by HS, as this strategy would ensure more CO₂ fixation, higher E 36 (leading to tissue temperature regulation), less oxidative damage (linked to lower ROS formation), 37 thus limiting also the negative effects of HS. Similarly, the indications about phenological breeding 38 strategies remain valid if DS is accompanied by HS, as they both increase ovule and pollen sterility, 39 reduce spikelet number, accelerate plant senescence, and reduce GF duration, enhancing the reduction 40 in TKW (Barnabás et al., 2008; Ji et al., 2010; Farooq et al., 2011; Prasad et al., 2011). So, in the 41 42 context of terminal drought, even when accompanied by terminal HS, early heading and early 1 maturity genotypes can be considered preferable, avoiding exposure to terminal stressful conditions

by escape strategies. As exposed in section 2.1, a short cycle could be obtained by the combination
of a short pre-heading phase with an increased GF duration, or by a longer pre-heading phase, and a

4 short (but highly efficient) GF period.

5 On the contrary, proline accumulation and PSII inhibition could become more relevant breeding tools when drought is accompanied by HS, due to the synergic effect of DS and HS on proline accumulation 6 and PSII inhibition via increased thylakoid membrane fluidity and loss of electron-transport 7 dependent integrity. Wang et al. (2010), studying the effects induced by drought, heat and their 8 combination in wheat flag leaves, observed the sharpest increase in proline accumulation and osmotic 9 adjustment in case of stress combination, while Prasad et al. (2011), observed a synergism of the 10 combined stresses on chlorophyll content decline. Also Urban et al. (2018), imposing single and 11 combined DS and HS for two weeks after anthesis, observed that Fv/Fm showed a significant 12 interactions with temperature and water availability, resulting in an amplified decline of this 13 parameter under the combined influence of drought and heat (temperature above 35 °C). So, 14 chlorophyll fluorescence parameters and osmotic adjustment are indicators of HS tolerance in wheat, 15 with strong correlation with GY (Ullah et al., 2022), thus representing a possible criterion for future 16 breeding under combined HS and DS. 17

In conclusion, breeding strategies based on improving drought resistance, could be useful also in case 18 of simultaneous HS. However, despite it is known that a large number of physiological, biochemical 19 and growth processes are affected by interactions between DS and HS, there is still a limited 20 knowledge in terms of a precise quantification of the effects of these combined stresses on production 21 parameters, that would be crucial for reliable predictions of climate change impacts on wheat 22 (Barnabás et al., 2008; Hlaváčová et al., 2018). Acknowledging the complexity in understanding the 23 contributions of each stress, especially under field conditions (Farooq et al., 2011; Prasad et al., 2011), 24 precise information on combined heat and drought would be precious, not only for addressing 25 successful breeding programs, but also for the improvement of crop models, assessing simultaneous 26 stresses impacts on growth and yield (Hlaváčová et al., 2018). 27

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29 2.6 Concluding remarks

30 Breeding improvements for drought resistance in both bread and durum wheat can be obtained via direct or indirect selection for yield. Direct selection for yield is hampered by its low heritability, GY 31 being a highly integrative trait, dependent on a wide range of biotic and abiotic interactions. For this 32 reason, physiological breeding, i.e., the selection of morpho-functional traits associated with yield 33 performance in water-stressed environments, attracts most attention. The main obstacle to 34 physiological breeding is deciding which traits should be preferred as a reliable set of selection 35 criteria, given the broad spectrum of drought affected environments in which timing, duration, and 36 severity of the stress vary greatly. In fact, literature shows many conflicting results regarding the 37 correlation between various morpho-functional traits and GY under water-limiting conditions. In this 38 review, we offer an insight into the reasons for these controversies, while discussing the ease of use 39 of each trait. Moreover, we provide an interpretation of how selection criteria should be combined 40 for selection in different target environments, characterized by different drought scenarios. Table 3 41 summarizes our conclusions. Thus, this review does not intend to be an exhaustive treatment of all 42 phenotypic traits associated with DS, nor of the modern high-throughput phenotyping strategies and 43 44 genomic approaches available. For example, root architecture traits are extremely important for DS

resistance (Manschadi et al., 2006), as well as imaging spectroscopy can be applied in remote sensing to evaluate numerous traits associated with abiotic stress tolerance (Jangra et al., 2021). New rising high-throughput phenotyping platforms, providing precise measurements of desired traits among thousands of field-grown plants, associated with genomic technologies, will accelerate breeding programs (Mwadzingeni et al., 2016b; Jangra et al., 2021). The purpose of this review is to highlight the contradictions found in the literature for different morpho-physiological and phenological traits, and to provide an environment-oriented perspective to overcome them.

8 For example, in the Mediterranean climate type contexts, characterized by mild terminal drought, selection should be oriented to semi-dwarf early maturing genotypes, able to rapidly complete the life 9 cycle, minimizing the temporal exposure to drought, expressing their high yield potential, with 10 prompted g_s and Pn, as to maximize assimilates accumulation, increasing the efficiency of GF 11 12 process, hampered by the terminal stress. On the other hand, where DS appears early in the life cycle, and progressively increase its severity, early vigorous, tall genotypes, with high tillers number should 13 be preferred, able to successfully germinate in stressful condition, quickly covering bare soil 14 (reducing water loss from the ground) and producing more photosynthetic biomass, meaning more 15 assimilates to be re-mobilized successively to the spike. Moreover, in contexts like these, drought 16 avoidance features, such as LR, could be considered. 17

18 In conclusion, climate change makes the risk of water stress more frequent, jeopardizing wheat yields more often in areas where it is grown in rainfed conditions. For this reason, it is essential to work on 19 the selection of new varieties of wheat, both bread and durum, capable of giving good yields even in 20 the presence of periods of drought. This is a rather complicated challenge, as wheat can be affected 21 by drought at various stages of its life cycle. This makes the combination of phenology, water stress 22 and physiological characteristics a complex system. It is therefore essential that the selection criteria 23 are environment-oriented, in order to overcome controversies in breeding for drought resistance in 24 25 wheat.

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Acknowledgments

We acknowledge Giulia Feduzi for the artwork herein presented and Prof. Stefania Biondi for offering
help in reviewing and revising the manuscript.

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Table 1. Most common stress susceptibility/tolerance indices, considering yield in both stress and non-stress conditions. Index name and corresponding equations are reported, together with environment classification and citation reports.

equations are rep		ortea, together with		labbiliteation and en	anon repond.
Index name	Equation	Reference author(s)	Most suitable selection environment	Web of Science citations (April 7 th 2022)	Scholar citations (April 7 th 2022)
Stress susceptibility index (SSI)	$ \begin{array}{c} [1-(Ys \ / \ Yp) \ / \ 1-(\hat{Y}s \\ / \ \hat{Y}p)] \end{array} $	Fischer and Maurer, 1978 [33]	А	1145	3332
Tolerance index (TOL)	(Yp – Ys)	Rosielle and Hamblin, 1981 [34]	А	611	1870
Yield Stability Index (YSI)	Ys / Yp	Bouslama and Schapaugh, 1984 [35]	А	345	1011
Mean productivity (MP)	(Ys + Yp) / 2	Rosielle and Hamblin, 1981 [34]	В	611	1870
Harmonic mean (HARM)	2 (Yp×Ys) / (Yp+Ys)	Schneider et al., 1997 [37]	В	129	460
Geometric mean productivity (GMP)	$(\mathbf{Yp} \mathbf{x} \mathbf{Ys})^{\frac{1}{2}}$	Fernandez, 1992 [36]	В	94	2003
Stress Tolerance index (STI)	$(Yp x Ys) / (\hat{Y}p)^2$	Fernandez, 1992 [36]	В	94	2003
Yield Index (YI)	Ys / Ŷs	Gavuzzi et al., 1997 [38]	В	172	602

Note: in the above equations Yp and Ys are respectively the yields of a given genotype under optimum (potential) and stressed conditions; $\hat{Y}p$ and $\hat{Y}s$ are respectively the mean yields of all genotypes under study under optimum and stressed conditions

- A = to be adopted only in systematically drought stressed environment

- B = to be adopted in yearly dependent drought interested environment

(A or B classification has been implemented combining the results of Golabadi et al., 2006 [39]; Mohammadi et al., 2010 [31]; Mohammadi et al., 2011 [40]; Ahmadizadeh et al., 2012 [23]; Patel et al., 2019 [29])

Table 2. Developmental stage trait selection to obtain early maturity for terminal drought escape based on two strategies.

	STRATEGY 1	STRATEGY 2
	Yashavanthakumar et al., 2021 [53]; Ihsan et al., 2016 [47] ; Simane et al., 1993 [46]; Shavrukov et al., 2017 [49]; Mohammadi et al, 2019 [25].	Al-Karaki 2012 [54]; Singh et al. 2014 [55].
DEVELOPMENTAL STAGE		
Pre-heading period (BBCH 1 – 4)	short	long
Heading (BBCH 5)	early	Not early
Flowering (BBCH 6)	early	Not early
Grain filling period (BBCH 7-8)	long	Short but high rate

Table 3: Suitable trait combinations	for different drought scenarios	
Trait	Early-season drought	Terminal drought (post-anthesis)
	(pre-anthesis)	
Early vigor		
Peduncle Length		
RWC		\checkmark
LAI		
High number of tillers		
Low number of tillers		
Tall size	\checkmark	
Semi-dwarf		${\bf \boxtimes}$
Early flowering and maturity		\checkmark
Prolonged - or short but high rate		$\overline{\checkmark}$
- grain filling		
Flag leaf Pn		\checkmark
Fv/Fm - T _{crit}		\checkmark
Trait	Mild-moderate drought	Severe drought
High g _s	\checkmark	
Reduced g _s		\checkmark
Epicuticular waxiness	${\bf \bigtriangledown}$	
Reduced nocturnal stomatal		
transpiration		
High number of tillers		
Low number of tillers	$\overline{\checkmark}$	
Leaf rolling		
Flag leaf Pn	\checkmark	
Fv/Fm - T _{crit}		
Reduced NPQ		
RWC	☑	
Tall-size		$\overline{\checkmark}$
Semi-dwarf		



