Grain yield, agronomic traits, and protein content of two- and six-row barley genotypes under terminal drought conditions

Vesna Kandic1*, Dejan Dodig1, Mile Secanski1, Slaven Prodanovic2, Gordana Brankovic2, and Primoz Titan3

1Maize Research Institute “Zemun Polje”, Slobodana Bajića 1, 11080 Belgrade, Serbia.
2Corresponding author (vesna.kandic@gmail.com).
3University of Belgrade, Faculty of Agriculture, Nemanjina 6, 11080 Belgrade, Serbia.
4RGA Research, Genetics and Agrochemistry Ltd., Brodarska ulica 27, Krog, 9000 Murska Sobota, Slovenia.

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ABSTRACT

Small grain cereals in lower latitude areas usually mature under terminal drought conditions that affect their agronomic performance. An experiment was conducted to compare agronomic traits, grain yield, and protein content under control and terminal drought conditions of 15 two-row and 10 six-row barley (Hordeum vulgare L.) genotypes. The experiment was set up at two locations for two growing seasons (2011, 2012) and two treatments. One treatment was terminal drought (D) simulated by the mechanical removal of all leaf blades 7 d after the heading of each genotype, and a control (C) treatment in which plants were left intact. On average, defoliation caused a greater reduction in grain yield and protein content of the six-row genotypes (37.6% and 12.3%, respectively) than the two-row genotypes (28.8% and 7.1%, respectively). On the other hand, test weight of six-row genotypes showed better tolerance to terminal drought. According to the multivariate function analysis, the genotypes of both types of barley with a high test weight, a longer uppermost internode, and a longer grain filling period had high protein content under terminal drought stress. In contrast to six-row genotypes, it is possible to breed two-row genotypes that not only have high grain yield but also high protein content under both optimal and drought stress conditions.

Key words: Abiotic stress, barley quality, grain filling, grain yield, Hordeum vulgare.

INTRODUCTION

Barley (Hordeum vulgare L.) has three main uses: feed, food, and malt production (Newton et al., 2011). Barley grain has lower protein and energy levels than wheat, but it is the only cereal and the only staple food resource in arid and semi-arid regions of some developing countries. Climate change will be a significant challenge for delivering grain with consistent yield and quality in the future due to the complex effects of atmospheric CO₂ and changing temperature and rainfall patterns on barley development (Nuttall et al., 2017). This is the case in Mediterranean and Southeastern European environments where small grain cereals usually mature under terminal drought conditions. Studies showed that yield components and protein content (PC) decreased and increased, respectively, when barley was subjected to water stress after anthesis (Samarah et al., 2009; Wu et al., 2017). Dry matter for grain filling is a result of a post-anthesis photosynthetic activity and remobilization of storage reserves deposited prior to anthesis. If climatic conditions (predominantly water scarcity and enhanced temperatures) are limiting, the contribution of current photosynthesis to grain filling declines and the remobilization of stored carbohydrates from senescence tissues increases (Merah et al., 2017). During the vegetative and early reproductive stages of cereal development, assimilated C is temporarily stored as carbohydrate in vegetative
sink tissues, such as the stem and leaf sheaths, and those reserves are subsequently remobilized for the transport to reproductive sink tissues and filling the grain during the later stages of plant development (Scofield et al., 2009). Arisnabarreta and Miralles (2008) and Nuttall et al. (2017) reported that barley yield (carbohydrate deposition) in a wide range of environments was considered to be limited by the number and capacity of grains to store DM (sink-limitation) rather than the availability to assimilate for grain filling (source-limitation). In contrast, protein deposition is largely a limited source and relies on N reserves accumulated in the leaves and stems in the pre-anthesis phase (Kohl, 2015). At the end of the grain filling process, mature barley grain consists of starch, proteins, β-glucan, and minor components. Starch is a major storage component in the starchy endosperm and increased starch content usually results in larger grains. Test weight (TW) is grain density or weight per unit of volume of a grain at a standardized moisture level. Grains with a higher TW have a higher percentage of large, plump kernels with a greater proportion of starch-rich endosperm and lower PC. This is the reason why TW, in addition to PC, is an important indicator of barley quality.

Based on spike morphology, that is, floret fertility, barley is divided in two types. In two-row barley genotypes, only the central spikelet is fertile, while the remaining two lateral spikelets are either sterile or reduced. On the other hand, in six-row barley genotypes, in addition to the central spikelet, two lateral spikelets are also fertile and can develop grain. These two types generally differ in their end use: six-row barley is mainly used as feed due to its higher grain PC and less uniform grain size and weight compared with two-row barley (Lang et al., 2013; Zwirek et al., 2019), while two-row barley is more often used as a malting material in beer production, producing high malt extract, lighter color, and less enzyme content than the six-row type (Gupta et al., 2010). The aim of this study was to evaluate the effect of terminal drought conditions simulated by defoliation on grain yield, agronomic traits, and protein content of 25 barley genotypes of both barley types and to examine a correlation of these traits under optimal and stress conditions.

MATERIALS AND METHODS

Plant material and growing conditions

The experiment consisted of 15 two-row and 10 six-row winter barley genotypes (Kandic et al., 2018) and was conducted at two locations (5 km apart) in northern Serbia, Zemun Polje (ZP) (44°86'N, 20°33'E) and Skolsko dobro (SD) (44°86’N, 0°28'E) during the 2010-2011 and 2011-2012 growing seasons. The experiment used a randomized block design with two replicates and two treatments. Each plot consisted of five 1 m long rows, 20 cm spacing, and a typical seeding rate of 350 seeds m⁻². The first treatment was the control (C) with intact plants, while all leaf blades were manually removed 7 d after the heading of each genotype in the second treatment (D). Drought conditions during grain filling were simulated by inhibiting current photosynthesis (as a result of defoliation) and plants were forced to rely on mobilizing stem reserves to maintain grain filling (Dodig et al., 2016; 2018).

Soil at the ZP site is a slightly calcareous Chernozem (CaCO₃ 1.5%) with ~ 3.0% humus content, pH (KCl) 7.1, soil available P and K content of 57 and 35 mg 100 g⁻¹, respectively, and 0.19% total N content. Soil at the SD site is also a Chernozem with higher Ca (CaCO₃ 6.4%), lower humus (2.5%), soil pH (KCl) 7.4, soil available P and K content of 26 and 22 mg 100 g⁻¹, respectively, and 0.15% total N content. Soil characteristics at the sites correspond to the 60 cm layer. Fertilizer rates were determined based on the soil chemical characteristics and available amounts of N, P, and K in the soil. A total of 40 kg N ha⁻¹, 40 kg P ha⁻¹, and 40 kg K ha⁻¹ and a total of 50 kg N ha⁻¹, 50 kg P ha⁻¹, and 50 kg K ha⁻¹ were applied prior to sowing in the ZP and SK sites, respectively. The top dressing was applied at 30 kg N ha⁻¹ at both sites in March or April. Standard cropping practices were applied to keep plots pest and disease free.

Genotypes were sown in late October and harvested in late June of the following year. To prevent the reduction of the accumulated reserve and storage capacity in stems, plots were manually irrigated from early March (beginning of tillering) to the end of April (beginning of heading) when water in the top 0.75 m soil layer had decreased below 50% field capacity. Soil moisture content was determined weekly by the gravimetric method from samples collected at three depths (0.25, 0.50, and 0.75 m) and at a frequency depending on the rainfall pattern. Two irrigation events of 50 mm added water and another irrigation event of 30 mm added water were applied in the 2010-2011 and 2011-2012 seasons, respectively.

The meteorological data for the winter barley crop cycle in spring (March-June) were recorded at the nearest weather station of the Republic Hydrometeorological Service of Serbia, approximately 5 km away from both experimental sites. Averaged daily mean temperatures during grain filling (May and June) were 19.9 °C (2011) and 20.8 °C (2012), while
the number of days with the maximum temperature above 30 °C in the May-June period were 7 and 15 d, respectively (Kandic et al., 2018). Higher temperatures in 2012 were combined with a lower amount of precipitation during grain filling (98 mm in 2012 vs. 118 mm in 2011). To obtain more precise temperature information during grain filling, average temperatures (Ta) for every genotype were calculated using daily mean temperatures during the whole grain filling period of a specific genotype.

Data collection
Spike sampling was initiated at the start of defoliation and continued at a 5-d interval until full maturity (a total of nine samples were drawn for each plot). Five uniform plants per plot were collected from the center of the plot at each sampling and used to obtain the number of sterile kernels (SK, excluding lateral florets), number of fertile kernels (KN), and length of the uppermost internode (LI, length between the last developed node and the collar, cm). Grain yield (YLD, kg ha⁻¹), test weight (TW, kg hL⁻¹), and total protein content (PC, %) were calculated at full maturity for each plot based on the 13% moisture level. Protein content was determined by near infrared spectroscopy (Chopin Technologies). The grain filling duration (GFD) was calculated as accumulated growing degree days (GDD) from heading. The GDD was computed with 0 °C as a base temperature (Grigorieva et al., 2010; Ibrahim et al., 2018) on the basis of a daily mean temperature by the following formula: 

$$T_n = \frac{(T_{\text{max}} + T_{\text{min}})}{2} - T_b$$

where $T_n$ is the daily degree-day, $T_{\text{max}}$ is the maximum daily temperature, $T_{\text{min}}$ is the minimum daily temperature, and $T_b$ is the base temperature.

Data analysis
To evaluate significant differences among genotypes, treatments, and environments (year-location combination), a three-way ANOVA was performed. The significance of differences between treatments C and D and between two- and six-row barley genotypes was estimated by the t-test with letter groupings generated by 5% and 1% levels of significance. The principal component analysis (PCA) described by Yan and Rajcan (2002) was used to visually display relationships between observed traits. Separate biplots were constructed for two treatments and barley types. A positive correlation between two traits was represented by an acute angle between them, while an obtuse angle represented a negative correlation. All calculations were made with the Minitab (trial version 17) and Excel software packages.

RESULTS AND DISCUSSION
The greatest average reduction of studied traits due to defoliation across years and sites for both barley types was recorded for YLD (37.6% and 28.8% in the groups of six-row and two-row genotypes, respectively) (Figure 1). On the other hand, KN was almost at the same level in both treatments (Figure 1). Yield reductions were similar to results indicated by Gonzalez et al. (2007), who reported that the average grain yield reduction for six- and two-row barley after applying terminal drought stress was 38.9% and 27.2%, respectively. Stress caused by defoliation increased SK of six-row genotypes by 16.7% and prolonged GFD by 1.09%. In two-row barley, SK increased by 7.4%, while GFD after defoliation was slightly shorter (0.45%). Samarah et al. (2009) reported that GFD for barley genotypes (two- and six-row) grown under water stress treatments was shorter in both types than in the well-watered treatment. In the stress treatment, shortening

Figure 1. Percentage average decrease/increase of investigated traits of two-row (black bars) and six-row (white bars) barley genotypes in defoliation vs. control treatment.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Percentage Change</th>
</tr>
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<tbody>
<tr>
<td>YLD</td>
<td>37.6%</td>
</tr>
<tr>
<td>KN</td>
<td>0%</td>
</tr>
<tr>
<td>LI</td>
<td>0%</td>
</tr>
<tr>
<td>TW</td>
<td>0%</td>
</tr>
<tr>
<td>PC</td>
<td>0%</td>
</tr>
<tr>
<td>GFD</td>
<td>0%</td>
</tr>
</tbody>
</table>

PC: Protein content; GFD: grain filling duration; TW: test weight; YLD: grain yield; LI: length of uppermost internode; KN: number of fertile kernels; SK: number of sterile kernels.
of the uppermost internode was greater in six-row barley (2.0%) than in two-row barley (0.6%), as well as TW reduction (2.3% and 1.1%, respectively). Moreover, TW decreased more in six-row barley (2.3%) than in two-row barley (1.1%).

The PC decrease was slightly higher in the group of six-row genotypes (12.3%) than in two-row genotypes (11.9%). Other authors generally reported that PC was higher under (terminal) drought conditions (Afshari-Behbahanizadeh et al.,

**Figure 2.** Mean values of investigated traits across years (2011, 2012) and treatments (control C, defoliation D), and averaged (C aver and D aver) for both years.

Uppercase letters indicate a difference between two-row (black bars) and six-row (white bars) barley genotypes separately for each treatment. Lowercase letters indicate a difference between the two treatments separately for each barley type. The same lowercase or uppercase letters are not significantly different at P < 0.05 according to the t-test.
in the C treatment (71.9 and 73.6 kg hL\(^{-1}\), respectively); this suggests that two-row types were less sensitive/susceptible

73.6 kg hL\(^{-1}\), respectively), while six-row genotypes had significantly (P < 0.05) lower values in the D treatment than

treatments C and D, but there was a nonsignificant difference between the two types in the same treatment

2012, there was a significant difference (P < 0.05) within the same barley type (either two- and/

Averaged across years, defoliation did not significantly influence LI of the barley types. The LI values established in

the highest direct effects on grain yield in both six-row and two-row barley genotypes under terminal drought conditions.

Given that protein deposition relies on N reserves accumulated in leaves and stems in the pre-anthesis phase, we

2016). However, since protein deposition mostly relies on N reserves accumulated in leaves and stems (Distelfeld et al.,

A comparative analysis of two barley types under different conditions was performed. Figure 2 shows mean values of

investigated traits over 2 yr (2011, 2012) and treatments (control, defoliation) and averaged for both years. Significant (P < 0.05) differences between the two barley types were obtained for most traits. The average number of SK in both years and both treatments was significantly higher in the group of six-row genotypes than in the group of two-row genotypes. Several authors (Arisnabarreta and Miralles, 2006) reported that spikelet abortion was generally higher in six-row barley types because they have more fertile spikelet primordia per spike (Koppolu et al., 2013; Alqudah and Schnurbusch, 2014). In 2011, a significant difference was recorded in the group of six-row genotypes between treatments C (5.5) and D (7.4) for SK, while the difference between treatments C and D was nonsignificant (1.3 vs. 1.4) in the group of two-row genotypes. Both barley types had higher SK in 2012 than in 2011 when twice as many days with temperatures above 30 °C were recorded. Two-row barley had higher SK by 18.8% and 17.6% in 2012 than in 2011 in treatments C and D, respectively. On the other hand, SK of six-row barley was higher by 25.6% and 3.9% in 2012 than in 2011 in treatments C and D, respectively. In 2012, the difference between the same barley types under different treatments was nonsignificant.

The two barley types were significantly (P < 0.01) different regarding the number of fertile kernels (KN) per spike in both treatments. On average, higher KN values were recorded across years in six-row genotypes (53.6 and 53.1 in C and D, respectively) than in two-row genotypes (26.0 and 26.2 in C and D, respectively). Barley row type, that is, KN is controlled by at least five loci: Six-rowed spike 1 (Vrs1 [syn = HvHox1]), Vrs2, Vrs3, Vrs4, and Intermediate spike-c (Int-c [syn = Vrs5]), and all five genes have been identified (Sakuma et al., 2017). The barley spike is formed during the vegetative stage and depends on the environmental conditions under which the morphogenesis of the generative organs takes place during the orthogenetic process. Arisnabarreta and Miralles (2008) considered that the critical period for the grain number of six-row types was 30 d before flowering and even earlier for two-row types (between 40 and 10 d before flowering). The number of fertile spikelets was affected by the number of assimilates available to the spike in the early stages of its development (Arisnabarreta and Miralles, 2008). In the present study, defoliation on the seventh day after heading did not reduce KN because the final number of grains had already been formed. According to Alqudah and Schnurbusch (2014), actual anthesis/fertilization events in winter barley occur before flowering (when anther extrudes), for example, after the awn tipping stage or heading (according to the Zadoks scale GS49; Zadoks et al., 1974).

The TW is grain density or weight per unit of volume of a grain at a standardized moisture level. On average, across years and treatments, higher values for TW were recorded for two-row genotypes (74.0 kg hL\(^{-1}\)) than for six-row genotypes (72.8 kg hL\(^{-1}\)). In 2011, there was a nonsignificant difference between the two types, as well as within the same type in treatments C and D. In 2012, there was a significant difference (P < 0.05) within the same barley type (either two- and/or six-row) in treatments C and D, but there was a nonsignificant difference between the two types in the same treatment (either C and/or D). Overall for both years, two-row genotypes had similar TW values in treatments C and D (74.5 and 73.6 kg hL\(^{-1}\), respectively), while six-row genotypes had significantly (P < 0.05) lower values in the D treatment than in the C treatment (71.9 and 73.6 kg hL\(^{-1}\), respectively); this suggests that two-row types were less sensitive/susceptible to drought stress during grain filling. The higher TW values for two-row types in both treatments and both years are in accordance with the results obtained by other authors, who reported that two-row types generally produced larger seeds with a higher TW (Frégeau-Reid et al., 2001; Djekic et al., 2017).

Given that protein deposition relies on N reserves accumulated in leaves and stems in the pre-anthesis phase, we compared barley types for LI) in both treatments. Previous studies performed on wheat genotypes (Dodig et al., 2016; 2017) showed that one of the most important traits for grain weight per spike was peduncle length in both terminal drought and control treatments. Other authors (Nikkhah et al., 2010; Shobbar et al., 2017) reported that peduncle length had one of the highest direct effects on grain yield in both six-row and two-row barley genotypes under terminal drought conditions. Averaged across years, defoliation did not significantly influence LI of the barley types. The LI values established in two-row genotypes were 24.2 and 24.1 cm in treatments C and D, respectively, while the corresponding values in six-row genotypes were 23.9 and 23.4 cm. The difference between two types in the same treatment was also nonsignificant. Averaged across treatments and barley types, the LI of the studied genotypes was shorter by 25.5% in 2012 when higher temperatures were combined with lower precipitation during grain filling compared with 2011. Modarresi et al. (2010)
reported that wheat peduncle length was highly sensitive to heat stress during grain filling and decreased by 20.2%. The only significant influence of defoliation (P < 0.05) was recorded in D12 when six-row genotypes had shorter LI than two-row genotypes (18.8 vs. 20.7 cm).

The combined PCs for 2 yr were significantly different (P < 0.01) for the same barley type between treatments C and D. On average, two-row barley PC was not significantly higher (12.9%) than 6-row barley (12%). The average PC values obtained by other authors for both barley types/genotypes varied from 12% to 14% (Ingvorsen et al., 2016). Frégeau-Reid et al. (2001) reported that PC for two-row barley was greater than for six-row barley, while in the study by Lahouar et al. (2017), the value was higher in six-row genotypes than in two-row barley genotypes. Elia et al. (2010) carried out a study of cross-breeding European two-row barley with American six-row barley and found that the high protein allele was carried by the two-row parent. Averaged for both years and types, PCs were lower in treatment D than in treatment C (12% and 13.6%, respectively). In 2011, there was a significant (P < 0.01) difference between the two barley types within the same treatment in favor of the two-row barley type, while in 2012, this difference was nonsignificant, and six-row genotypes had a slightly higher PC.

Averaged across years, there were nonsignificant differences between barley types for GFD in both treatments in 2011. In 2012, two-row genotypes had a significantly (P < 0.05) longer grain filling period in treatment C than in treatment D, but the group of six-row genotypes maintained grain filling at an approximately similar level in both treatments (D and C). Dias and Lidon (2009) reported that drought after anthesis shortened the grain filling period; however, according to our results, this happened only in 2012 when drought stress was accompanied with higher mean temperatures.

The highest mean YLD (8239 kg ha⁻¹) was recorded in two-row genotypes in C11, and the lowest (3204 kg ha⁻¹) in six-row genotypes in D12. Averaged for the 2 yr, there were significant differences between two- and six-row types in treatments C (P < 0.05) and D (P < 0.01). In treatment D, two-row genotypes had 25.1% higher yield than six-row genotypes (5279 and 3954 kg ha⁻¹, respectively). On the other hand, yields recorded in treatment C were 7419 and 6336 kg ha⁻¹, respectively, which means that the yield established in two-row genotypes was higher by 14.6%. The higher yield of two-row genotypes is in accordance with results reported by Bratkovic (2014), but contrary to results reported by Samarah et al. (2009). In 2011, two- and six-row genotypes had a significantly (P < 0.01) lower yield in treatment D than in treatment C, while there was nonsignificant difference for yield when the two types were compared within a treatment. In 2012, both barley types had a significantly (P < 0.05) lower yield in treatment D than in treatment C. In 2012, six-row genotypes had a significantly lower yield than two-row genotypes in both treatments (C: 5241 kg ha⁻¹ vs. 6599 kg ha⁻¹, respectively, and D: 3204 kg ha⁻¹ vs. 4853 kg ha⁻¹, respectively).

The results of three separate factorial ANOVA for the two barley types (Table 1) indicate that the environment factor (on average 36.9% and 36.0% for two- and six-row genotypes, respectively) had the most significant influence on the variability of all the traits, followed by the genotype factor (on average 13.3% and 19.8% for two- and six-row genotypes, respectively) and the G × E interaction (11.5% and 9.4%, respectively). The great effect of environment on most traits was probably due to the contrasting seasons in which the experiments were set up. The influence of the G × T interaction on most of the traits was nonsignificant and this indicates that there was nonsignificant difference among the investigated genotypes in their response to stress caused by defoliation. These results are different from those obtained when two- and six-row genotypes were observed together and in which the most significant influence on variability was the genotype factor and the treatment factor had a greater influence than the interactions (Kandic et al., 2018). Most of the YLD variation in the two-row group was associated with the treatment (31.5%) and in the six-row group with the environment (35.3%). The environment had the strongest influence on PC as well as on TW and LI for both barley types. Furthermore, the environment factor had the highest impact on KN of two-row genotypes (71.7%), while the variability of KN in six-row genotypes was mostly influenced by the genotype (54.3%). The variability of SK in both types was mostly influenced by genotypic differences. The effect of the genotype was significant for all measured traits except for GFD, indicating a high degree of genetic variation that could be exploited in the breeding program of two- and six-row barley.

**Multivariate analysis**

A PCA was performed to determine the interrelationship between YLD, agronomic traits, and PC under control and stress conditions for each barley type (Figure 3). In the two-row barley group in treatment C, the strongest correlations were between GFD and LI (positive) and between PC and TW (negative). The YLD had a positive correlation with KN,
SK, and PC. In treatment D, the strongest positive correlations were between LI and SK, and the strongest negative between TW, PC, and KN. The YLD was positively associated with TW, PC, and GFD. These results suggest that two-row genotypes could have both high yields and PCs under both optimal and drought stress conditions. Under defoliation, LI had a greater impact on PC compared with optimal conditions, probably due to the fact that after the removal of the leaf blades, protein deposition, as a limited source, only relied on N reserves accumulated in the stems. Besides genotypes with longer LI, those which could prolong the grain filling period under terminal drought stress also had a higher PC and YLD. When testing the population of malting barley in environments suffering severe drought, Moody et al. (2001) reported that grain size, as a primary contributor to final YLD, had been enhanced by longer peduncles, less kernels per spike, and earlier maturity.

For the six-row genotypes in the control treatment, the strongest positive correlation was detected between KN and LI, while the strongest negative correlations were established between PC, TW, and YLD. Although no difference was found between the two barley types for GFD, a strong association between YLD and GFD indicates that high yielding genotypes (in both barley types) under optimal conditions would tend to have a longer grain filling period. Under defoliation, the strongest positive correlation was between KN and SK, as well as between GFD and TW. The YLD had the strongest negative association with KN and SK. Moreover, YLD and PC were also negatively correlated, but not so strongly as in treatment C. On the other hand, PC was positively associated with all traits, except YLD. In contrast to two-row barley, this means that it would be very difficult to develop six-row barley genotypes with both high yield and PC.

Across two- and six-row genotypes, a positive correlation between PC and Ta under defoliation is in accordance with the results indicated by Reinhardt et al. (2013), who reported that high temperatures during the grain filling period prior to harvest increased the PC of barley genotypes. Our results suggest that genotypes of both barley types with high TW, longer LI, and ability to prolong the grain filling period under stress conditions, would have a higher PC. High mean temperatures under terminal drought stress influenced YLD of both barley types to a greater extent than under optimal growing conditions.

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CONCLUSIONS

Terminal stress simulated by the removal of all leaf blades 7 d after heading caused a significant (P < 0.05) reduction of protein content (PC) and grain yield (YLD) in two-row genotypes, and PC, YLD, and test weight (TW) in six-row barley genotypes. The reduction of YLD and PC was greater in the group of six-row genotypes, suggesting that this type is more sensitive to terminal drought stress than two-row barley genotypes. Our results showed that two-row genotypes had both high YLD and PC under both optimal and drought stress conditions. On the other hand, this was not possible in the group of six-row genotypes due to a strong negative correlation between these traits. Generally, to select genotypes of both barley types with high PC under stress conditions, the priority should be for those genotypes with high TW, longer uppermost internode, and the longest grain filling period.

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