

1 **Applied phenomics and genomics for improving barley yellow dwarf resistance in winter**
2 **wheat**

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28

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31 Selection (GS)

32

33 **Abstract**

34 Barley yellow dwarf (BYD) is one of the major viral diseases of cereals. Phenotyping BYD in
35 wheat is extremely challenging due to similarities to other biotic and abiotic stresses. Breeding
36 for resistance is additionally challenging as the wheat primary germplasm pool lacks genetic
37 resistance, with most of the few resistance genes named to date originating from a wild relative
38 species. The objectives of this study were to, i) evaluate the use of high-throughput phenotyping
39 (HTP) from unmanned aerial systems to improve BYD assessment and selection, ii) identify
40 genomic regions associated with BYD resistance, and iii) evaluate genomic prediction models
41 ability to predict BYD resistance. Up to 107 wheat lines were phenotyped during each of five
42 field seasons under both insecticide treated and untreated plots. Across all seasons, BYD
43 severity was lower with the insecticide treatment and plant height (PHTM) and grain yield
44 (GY) showed increased values relative to untreated entries. Only 9.2% of the lines were positive
45 for the presence of the translocated segment carrying resistance gene *Bdv2* on chromosome 7DL.
46 Despite the low frequency, this region was identified through association mapping. Furthermore,
47 we mapped a potentially novel genomic region for resistance on chromosome 5AS. Given the
48 variable heritability of the trait (0.211 – 0.806), we obtained relatively good predictive ability for
49 BYD severity ranging between 0.06 – 0.26. Including *Bdv2* on the predictive model had a large
50 effect for predicting BYD but almost no effect for PHTM and GY. This study was the first
51 attempt to characterize BYD using field-HTP and apply GS to predict the disease severity.
52 These methods have the potential to improve BYD characterization and identifying new sources
53 of resistance will be crucial for delivering BYD resistant germplasm.

54 **Introduction**

55 Wheat (*Triticum aestivum* L.) is one of the most essential food crops in the world and is
56 constantly threatened by several biotic stresses (Savary *et al.* 2019). Among the most important
57 viral stresses is barley yellow dwarf (BYD). This disease is widespread across the world, caused
58 by viruses and transmitted by aphids (Shah *et al.* 2012), and can cause significant yield
59 reductions in susceptible cultivars. In Kansas, BYD is the fourth most significant wheat disease
60 in terms of average estimated yield losses with an average yield loss of approximately 1%
61 estimated over the past 20 years (Hollandbeck *et al.* 2019), equivalent to a loss of more than \$10
62 million per year. However, yield losses are highly variable ranging from 5% to 80% in a single
63 field depending on the environment, management practices, the host, and the genetic
64 background, (Miller and Rasochová 1997; Perry *et al.* 2000; Gaunce and Bockus 2015).
65 Moreover, the wide host range and the complex lifestyle of its vectors make BYD extremely
66 difficult to manage, and different management strategies (e.g., planting date and control of vector
67 populations) are inconsistent depending on climate and location (Bockus *et al.* 2016). Thus, in
68 many production environments, particularly in the Central and Eastern regions of Kansas, BYD
69 is often the most economically impactful disease.

70

71 Barley yellow dwarf disease symptoms are highly variable depending on the crop, variety, time,
72 and developmental stage when the infection occurs, aphid pressure, and environmental
73 conditions (Shah *et al.* 2012; Choudhury *et al.* 2019b). BYD characterization in the field is
74 extremely challenging as the symptoms can easily be confused with other viral disease
75 symptoms such as wheat streak mosaic virus symptoms, nutrient deficiencies, or environmental
76 stresses like waterlogging (Shah *et al.* 2012). Typical BYD symptoms can be observed at all
77 levels of plant organization – leaf, roots, and flowers. Leaf discoloration in shades of yellow,
78 red, or purple, specifically starting at the tip of the leaf and spreading from the margins toward
79 the base is common as well as a reduction in chlorophyll content (Jensen and Van Sambeek
80 1972; D’arcy 1995). Often the entire plant visually appears stunted or dwarfed from a reduction
81 in biomass by reducing tiller numbers. Spike grain yield is decreased through a reduction in
82 kernels per spike and kernel weight which also affects grain quality (Riedell *et al.* 2003;
83 Choudhury *et al.* 2019b). Quality can be further reduced by a reduction in starch content (Peiris

84 *et al.* 2019). Below ground effects of BYD have also been reported including reduced root
85 growth (Riedell *et al.* 2003).

86
87 Currently, there is no simple solution to control BYD (Walls *et al.* 2019), however, the use of
88 genetic resistance and tolerance is the most appealing and cost-effective option to control this
89 disease (Comeau and Haber 2002; Choudhury *et al.* 2017; 2019b). Resistance and tolerance
90 could be different genetic mechanisms, namely stopping virus replication and minimizing
91 disease symptoms respectively, but within this paper all mention of resistance includes both
92 genetic resistance and tolerance. Breeding strategies involving genetic resistance can target
93 either the aphids or the virus. Resistance to aphids can be achieved by three different strategies,
94 antixenosis, antibiosis, or tolerance (Girvin *et al.* 2017). To date, most breeding efforts have
95 been directed to the identification of viral tolerance, also known as ‘field resistance’, that refers
96 to the ability of the plant to yield under BYD infection and is associated with a reduction of
97 symptoms of infection independent of the virus titer (Foresman *et al.* 2016). Field resistance has
98 been reported to be polygenic, falling under the quantitative resistance class, where several genes
99 with very small effects control the resistance response (Qualset *et al.* 1973, Cisar *et al.* 1982;
100 Ayala *et al.* 2002; Choudhury *et al.* 2019a; c).

101
102 Presently, no major gene conferring immunity or a strong resistant phenotype to BYD has been
103 identified in bread wheat, and only four resistance genes have been described for BYD. Located
104 on chromosome 7DS, *Bdv1* is the only gene described from the primary pool of wheat and was
105 originally identified in the wheat cultivar ‘Anza’ (Qualset *et al.* 1984; Singh *et al.* 1993). This
106 gene provides resistance to some but not all the viruses that cause BYD (Ayala-Navarrete and
107 Larkin 2011). The other three named genes were all introduced into wheat through wide
108 crossing from intermediate wheatgrass (*Thinopyrum intermedium*) (Ayala *et al.* 2001; Zhang *et*
109 *al.* 2009). *Bdv2* and *Bdv3* are both located on a translocation segment on wheat chromosome
110 7DL (Brettell *et al.* 1988; Sharma *et al.* 1995), while *Bdv4* is located on a translocation segment
111 on chromosome 2D (Larkin *et al.* 1995; Lin *et al.* 2007). *Bdv2* was the first gene successfully
112 introgressed in wheat breeding programs from the tertiary gene pool for BYD resistance (Banks
113 *et al.* 1995) and deployed into varieties.

114

115 In addition to the four known resistance genes, other genomic regions associated with BYD
116 resistance have been identified through genetic mapping. These regions have been described on
117 nearly all wheat chromosomes but have not been genetically characterized (Ayala *et al.* 2002;
118 Jarošová *et al.* 2016; Choudhury *et al.* 2019a; b; c). Moreover, two recent studies have reported
119 that some of these new genomic regions display additive effects (Choudhury *et al.* 2019a; b).
120 Additive genetic effects had already been reported in lines combining *Bdv2* and *Bdv4* (Jahier *et*
121 *al.* 2009).

122
123 Taken together, research indicates that resistance genes to BYD in wheat are rare. With a lack of
124 major genes and difficulty to characterize resistance in the wheat pool likely due to the polygenic
125 nature of many small effect loci, identifying resistance has been limited. Nevertheless, breeding
126 programs have devoted large efforts for breeding BYD resistance due to the economic
127 importance of this disease, with some of the greatest success coming from wide crosses to the
128 tertiary gene pool.

129
130 Breeding for BYD resistance can be improved by applying strategies for more effective
131 evaluation and utilization of the identified resistance. To get a better understanding of BYD and
132 its quantitative nature, consistent and high-throughput methods are needed for the identification
133 of resistant wheat lines for large-scale selection in breeding programs (Aradottir and Crespo-
134 Herrera 2021). Effective selection on the quantitative resistance with low heritability can be
135 aided by the high-throughput genotyping, high-throughput phenotyping (HTP), or a combination
136 of both.

137
138 Access to high-density genetic markers at a very low-cost, owing to the rapid developments in
139 DNA sequencing, have enabled breeding programs to apply molecular breeding for quantitative
140 traits. Genomic selection (GS) is a powerful tool to breed for quantitative traits with complex
141 genetic architecture and low heritability (e.g., yield, quality, and diseases such as Fusarium head
142 blight), because it has greater power to capture loci with small effect compared with other
143 marker-assisted selection strategies (Meuwissen *et al.* 2001; Poland and Rutkoski 2016). In
144 addition to molecular data, HTP using unmanned aerial systems (UAS), or ground-based sensors
145 is providing high density phenotypic data that can be incorporated into breeding programs to

146 increase genetic gain (Haghighattalab *et al.* 2016; Crain *et al.* 2018; Wang *et al.* 2020). Using
147 precision phenotyping for disease scoring can improve the capacity for rapid and non-biased
148 evaluation of large field-scale numbers of entries (Poland and Nelson 2011). Taken together
149 improvements in genomics and phenomics have the potential to aid breeding progress for BYD
150 resistance.

151
152 In an effort to accelerate the development of resistant lines, we combined high throughput
153 genotyping and phenotyping to assess BYD severity in a large panel of elite wheat lines. We
154 evaluated the potential of HTP data to accurately assess BYD severity as well as identify genetic
155 regions associated with BYD resistance and inform whole genome prediction to identify resistant
156 lines.

157

158 **Materials and Methods**

159

160 **Plant Material**

161 A total of 381 different wheat genotypes were characterized for BYD resistance, including 30
162 wheat cultivars and 351 advanced breeding lines in field nurseries over five years (Table S1). In
163 each nursery, an unbalanced set of 52 – 107 wheat entries were evaluated including both
164 cultivars and breeding lines (Table 1). The BYD susceptible cultivar ‘Art’ and BYD resistant
165 cultivar ‘Everest’ were included in all the nurseries (seasons) as checks.

166

167 **Field Experiments**

168 Nurseries for BYD field-screening were conducted during five consecutive wheat seasons (2015
169 – 2016 to 2019 – 2020) (Table 1). Seasons 2015 – 16 and 2016 – 17 were conducted at Kansas
170 State University (KSU) Rocky Ford experimental station (39°13'45.60" N, 96°34'41.21" W),
171 while the 2017 – 18, 2018 – 19, and 2019 – 20 nurseries were planted at KSU Ashland Bottoms
172 experimental station (39°07'53.76" N, 96°37'05.20" W). The nurseries were established for
173 natural infections by planting about three weeks earlier than the normal planting window in mid-
174 September. The susceptible cultivar ‘Art’ was planted as a spreader plot in the borders and as a
175 control check plot also with the resistant cultivar ‘Everest’. The experimental unit was 1.5m ×
176 2.4m with a six-row plot on 20cm row spacing.

177

178 A split-plot field design with two or three replications was used where the main plot was
179 insecticide treatment, and the split plot was the wheat genotype. Three replications were used
180 for proof of concept during the first two seasons but then two replications were chosen as a
181 balance of space and number of entries for the following seasons. For the treated replications the
182 seed were treated at planting with Gaucho XT (combination of insecticide and fungicide) at a
183 rate of 0.22 ml/100g of seed, followed with foliar insecticide applications starting from
184 approximately 2 – 3 weeks after planting through heading. Depending on field conditions, spray
185 treatments were conducted every 14 – 21 days if average air temperatures remained above 10°C.
186 Foliar insecticides were applied to the treated replications in a spray volume of 280.5 L/ha using
187 a Bowman MudMaster plot sprayer equipped with TeeJet Turbo TwinJet tips. Insecticide
188 applications consisted of a rotation of Warrior II, Lorsban, and Mustang Max at rates of
189 0.14L/ha, 1.17L/ha, and 0.29L/ha, respectively. For the control insecticide treatment (untreated),
190 the seed were treated with Raxil MD (fungicide) at a rate of 0.28ml/100g of seed, and no foliar
191 insecticide applications were applied. Foliar fungicide Nexicor was applied to the whole
192 experiment at a rate of 0.73L/ha, at both planting and heading, to control all other diseases so the
193 main disease pressure was focused on BYD.

194

195 **Phenotypic Data**

196 Individual plots were assessed for i) BYD severity characterized as the typical visual symptoms
197 of yellowing or purpling on leaves using a 0 – 100% visual scale, determined directly after spike
198 emergence by recording the proportion of the plot exhibiting the symptoms (Table 1), ii) manual
199 plant height (PTHT_M, meters), and iii) grain yield (GY, tons/ha). Experimental plots were
200 harvested using a Kincaid 8XP plot combine (Kincaid Manufacturing., Haven, KS, USA). Grain
201 weight, grain moisture and test weight measurements for each plot was recorded using a Harvest
202 Master Classic GrainGage and Mirus harvest software (Juniper Systems, Logan, UT, USA).
203 Visual phenotypic assessment was recorded using the Field Book phenoapp (Rife and Poland
204 2014).

205

206 **High-Throughput Phenotyping**

207 To compliment the manually recorded phenotypic data, we applied HTP using a ground-based
208 proximal sensing platform or an UAS (Table 2). Seasons 2015 – 16 and 2016 – 17 were
209 characterized by the ground platform as described in Barker *et al.* (2016) and Wang *et al.* (2018).
210 For the other three seasons, we used a quadcopter DJI Matrice 100 (DJI, Shenzhen, China)
211 carrying a MicaSense RedEdge-M multispectral camera (MicaSense Inc., United States). The
212 HTP data was collected on multiple dates throughout the growth cycle from stem elongation to
213 ripening (GS 30 – 90; Zadoks *et al.* 1974) (Table 2). Flight plans were created using CSIRO
214 mission planner application and missions were executed using the Litchi Mobile App (VC
215 Technology Ltd., UK, <https://uavmissionplanner.netlify.app/>) for DJI Matrice100. The aerial
216 image overlap rate between two geospatially adjacent images was set to 80% both sequentially
217 and laterally to ensure optimal orthomosaic photo stitching quality. All UAS flights were set at
218 20m above ground level at 2m/s and conducted within two hours of solar noon. To improve the
219 geospatial accuracy of orthomosaic images, white square tiles with a dimension of 0.30m ×
220 0.30m were used as ground control points and uniformly distributed in the field experiment
221 before image acquisition and surveyed to cm-level resolution using the Emlid REACH RS+
222 Real-Time Kinematic Global Navigation Satellite System unit (Emlid Ltd., HongKong, China).

223
224 An automated image processing pipeline (Wang *et al.* 2020) was used to generate the
225 orthomosaics and extract plot-level plant height (PTHT_D (m), Singh *et al.* 2019) and the
226 normalized difference vegetation index (NDVI) (Rouse *et al.* 1974), calculated as:

227
228
$$\text{NDVI} = \frac{\text{NIR}-\text{Red}}{\text{NIR}+\text{Red}} \quad [\text{Eq. 1}]$$

229
230 where NIR and Red are the near-infrared and red bands of the multispectral images and NDVI is
231 the output image. Both traits were selected based on potential BYD characterization where the
232 most typical BYD symptoms include chlorosis and stunting of the plants, thus, influencing
233 NDVI and PTHT.

234

235 **Statistical Data Analyses**

236 First, the adjusted mean best linear unbiased estimator (BLUE) was calculated for each entry for
237 all the different traits for each season (Table S1), using the following model:

238

$$239 \quad y_{ijklm} = \mu + G_i + T_j + GT_{ij} + R_{k(j)} + B_{l(kj)} + C_{m(kj)} + e_{ijklm} \quad [\text{Eq. 2}]$$

240

241 where y_{ijklm} is the phenotype for the trait of interest, μ is the overall mean, G_i is the fixed effect
242 of the i^{th} entry (genotype), T_j is the fixed effect of the j^{th} insecticide treatment, GT_{ij} is the fixed
243 effect of the interaction between the i^{th} entry and the j^{th} insecticide treatment (genotype by
244 treatment effect), $R_{k(j)}$ is the random effect of the k^{th} replication nested within the j^{th}
245 insecticide treatment and distributed as iid $R_{k(j)} \sim N(0, \sigma_R^2)$, $B_{l(kj)}$ is the random effect of the
246 l^{th} row nested within the k^{th} replication and j^{th} treatment distributed as iid $B_{l(kj)} \sim N(0, \sigma_B^2)$,
247 $C_{m(kj)}$ is the random effect of the m^{th} column nested within the k^{th} replication and j^{th} treatment
248 and assumed distributed as iid $C_{m(kj)} \sim N(0, \sigma_C^2)$, and e_{ijklm} is the residual for the $ijklm^{\text{th}}$ plot
249 and distributed as iid $e_{ijklm} \sim N(0, \sigma_e^2)$. The ‘lme4’ R package (Bates *et al.* 2014) was used for
250 fitting the models.

251

252 The BLUEs were used to inspect trait distributions and to calculate Pearson correlations between
253 all traits. In addition, BLUE values were used to calculate the reduction in GY for each entry as
254 the difference of GY between the untreated and insecticide treated main plots. This variable
255 reflects the level of BYD resistance of each entry, and it was used to perform GWAS and GS
256 analyses.

257

258 For NDVI and PTHT_D, the plot-level observed values extracted for the different phenotypic dates
259 were fitted to a logistic non-linear regression model (Fox and Weisberg 2011) as,

260

$$261 \quad y = \frac{\theta_1}{1 + e^{-(\theta_2 + \theta_3 x)}} + \varepsilon \quad [\text{Eq. 3}]$$

262

263 where is y the phenotype for the trait of interest at the time-point x measured as days after
264 January 1, θ_1 is the maximum value (upper asymptote) represented by the final PTHT or
265 maximum achieved NDVI, θ_2 is the inflection point that represents the greatest rate of change in
266 the growth curve, either senescence for NDVI or height of growth, θ_3 is the lag phase or onset of
267 senescence or growth rate from time x where x is the calendar day of the year since January 1,
268 and ε is the residual error (Figure S1). The “nlme” R package was used for model fitting
269 (Pinheiro *et al.* 2015). The model parameters obtained for each trait (θ_{1NDVI} , θ_{2NDVI} , θ_{3NDVI} ,
270 θ_{1PTHT_D} , θ_{2PTHT_D} , and θ_{3PTHT_D}) were used in addition to the other phenotypic traits to calculate
271 BLUEs, distributions, correlations, and BLUPs.

272
273 Secondly, we used a mixed linear model to calculate the best linear unbiased predictors (BLUPs)
274 for each entry in each nursery (season) (Table S1), using the same model as described in
275 equation 2 but defining G_i , T_j , and GT_{ij} as random effects. BLUPs were used because of the
276 unbalanced nature of the data (not all lines were evaluated in all the seasons). The BLUPs
277 calculated for each season were then combined for GWAS and GS. Furthermore, we calculated
278 broad-sense heritability on a line-mean basis by splitting the data based on whole plot treatment
279 for insecticide treatments as:

$$281 \quad H^2 = \frac{\sigma_G^2}{\sigma_G^2 + \frac{\sigma_e^2}{r}} \quad [\text{Eq. 4}]$$

282
283 where σ_G^2 is the genotypic variance, σ_e^2 is the residual error variance, and r is the number of
284 replications.

285 286 **Genotypic Data**

287 A total of 346 wheat entries were genotyped using genotyping-by-sequencing (GBS) (Poland *et al.*
288 *et al.* 2012) and sequenced on an Illumina Hi Seq2000. Single nucleotide polymorphisms (SNPs)
289 were called using Tassel GBSv2 pipeline (Glaubitz *et al.* 2014) and anchored to the Chinese
290 Spring genome assembly v1.0 (Appels *et al.* 2018). SNP markers with minor allele frequency <
291 0.01, missing data > 85%, or heterozygosity > 15% were removed from the analysis. After
292 filtering, we retained 29,480 SNPs markers that were used to investigate the population structure

293 through principal component analysis (PCA), genome-wide association analysis (GWAS), and
294 GS. In addition, GBS data was used to run a bioinformatics pipeline to predict the presence or
295 absence of the translocated segment on chromomere 7DL carrying the *Bdv2* gene for each entry
296 (Table S1). The prediction was done based on a modified alien predict pipeline (Gao *et al.*
297 2021). Briefly, alien or wheat specific tags were counted in the 7DL region and tabulated using
298 a training set of cultivars or lines that are known to be *Bdv2* positive and negative. A simple
299 classification was done based on alien to wheat tag counts ratios.

300

301 **Genome-Wide Association Analysis**

302 The GWAS analysis was performed with a mixed linear model implemented in the ‘GAPIT’ R
303 package (Lipka *et al.* 2012) that includes principal components to account for population
304 structure as fixed effects and the individuals to explain familial relatedness as random effects,

305

$$306 \quad y = X\beta + Zu_i + e \quad \text{[Eq. 5]}$$

307

308 where y is the vector of phenotypic BLUPs, X and Z are the incidence matrix of β and u_i ,
309 respectively, with u_i assumed $\sim N(0, 2K_i\sigma_i^2)$ where K is the individual kinship matrix, and e is
310 the vector of random residual effects with $\sim N(0, I\sigma_e^2)$, where I is the identity matrix and σ_e^2 is
311 the unknown residual variance. The false discovery rate correction with an experimental
312 significance level value of 0.01 was used to assess marker-trait associations. Manhattan plots
313 were generated with ‘CMplot’ package in R software (Yin 2020). PCA using GBS-SNPs was
314 performed in R language. Eigenvalues and eigenvectors were computed with ‘e’ function using
315 ‘A.mat’ function and the ‘mean’ imputation method of ‘rrBLUP’ package (Endelman, 2011). To
316 declare a quantitative trait locus (QTL) we considered only the regions having several SNP
317 markers in linkage disequilibrium, clearly showing a peak. We did not consider regions with a
318 single SNP above the significant threshold as a QTL.

319

320 **Genomic Selection**

321 Using data from the five seasons, GS models using the genomic best linear unbiased predictor
322 (G-BLUP) were developed to assess predictive ability. A five-fold cross-validation method was
323 used to assess model accuracy where the data set was split into five sets based on season, with

324 four seasons forming the training set and the fifth season serving as prediction set. This process
325 was repeated until all seasons were predicted. Along with predicting all other seasons from each
326 season, a model was evaluated with a leave-two-out cross-validation strategy. This strategy was
327 used to get a better mix of years with and without disease incidence, where the training
328 population consisted of three seasons, and the remaining two seasons were predicted from the
329 combined training population. The GS model was fitted with the training population using
330 ‘rrBLUP’ *kin.blup* function (Endelman 2011), the GS model equation was,

$$331$$
$$332 \quad y = Wg + \varepsilon \quad \text{[Eq. 6]}$$
$$333$$

334 where y is a vector of phenotypic BLUPs, W is the design matrix of g , g is the vector of
335 genotypic values $\sim N(0, K\sigma_g^2)$ and ε is the vector of residual errors (Endelman 2011).

336 Predictive ability was assessed using Pearson’s correlation (r) between the predicted value (G-
337 BLUP) and the BLUP for the respective phenotype. In addition, for both GS strategies we also
338 tested the effect of adding the genotype of the *Bdv2* loci as a fixed effect cofactor, using the
339 model,

$$340$$
$$341 \quad y = \mu + X\beta + Wg + \varepsilon \quad \text{[Eq. 7]}$$
$$342$$

343 which combines parameters described in equation 6 and X is the matrix ($n \times 1$) of individual
344 observation for presence or absence of *Bdv2* and β is the fixed effect for the *Bdv2* measurements.

345

346 **Results**

347

348 **Phenotypic Data**

349 We analyzed five years of BYD field-screening nurseries (seasons 2015-16 to 2019-20)
350 characterizing a total of 381 wheat lines. The disease pressure and the expression of BYD
351 associated symptoms varied each season, however, we were able to observe a significant effect
352 of the insecticide treatment in all seasons (Figure 1). Across all seasons, BYD symptoms were
353 lower on the insecticide treated plots and both PHT_M and GY increased compared to the non-
354 treated control. Season 2016-17 had the most conducive conditions for BYD screening, resulting

355 in high average severity and a larger difference between mean values for the treated vs untreated
356 plots for all the collected traits (Figure 1). There was general consistency in order across all
357 seasons with the susceptible check ‘Art’ ranked among the highest in BYD severity (Figure S2).

358
359 Phenotypic correlations between the traits showed a negative correlation between BYD and GY
360 for all the seasons and a negative or no correlation between BYD and $PTHT_M$ (Figure S3). The
361 same correlation trends were observed under insecticide treated and untreated plots. Broad-sense
362 heritability was moderate to high for all the traits, ranging between 0.21 and 0.79 for the
363 insecticide treated plots and between 0.41 and 0.84 for the untreated plots. Across all traits, the
364 untreated insecticide replications showed higher H^2 values, with season 2016 – 17 showing the
365 highest values (Figure 2).

366
367 For the HTP data collected (Table 2), we obtained three different parameters (θ_1 , θ_2 , and θ_3) for
368 both $PTHT_D$ and NDVI after fitting a logistic regression model using the data collected during
369 the experiments (2015-16 season data was not included due to lack of data quality) (Figure S1).

370 Correlations between these parameters and the phenotypic traits collected manually were
371 different for all the traits (Figure S3). For the insecticide untreated plots, BYD resulted in a
372 negative correlation with θ_{2NDVI} and a positive correlation with θ_{3NDVI} , in most of the field
373 seasons. We did not find a clear correlation pattern between BYD and $PTHT_D$. For $PTHT_M$ we
374 detected a positive correlation with θ_{1PTHT_D} across all seasons, and for GY we observed a
375 positive correlation with θ_{1NDVI} and θ_{2NDVI} , and a negative correlation with θ_{3NDVI} (Figure S3).

376

377 **Prediction of *Bdv2* Resistance Gene**

378 We used GBS data to genotype the *Bdv2* resistance gene located on a translocation segment from
379 intermediate wheatgrass on chromosome 7DL of bread wheat. In total, 33 of the 346 wheat lines
380 carried the *Th. intermedium* chromosomal translocation with *Bdv2* (Table S1). Interestingly, 28
381 of these *Bdv2* lines belonged to the same breeding cycle, entering the advanced yield nursery
382 stage of the KSU breeding program in the 2017 – 18 season. Furthermore, only 7 pedigrees are
383 represented within the 28 *Bdv2* entries, meaning that these lines are highly related. The
384 remaining 5 *Bdv2* lines were distributed in 2015 – 16 (n=3), 2018 – 19 (n=1), and 2019 – 20
385 (n=1), and none of the lines from the season 2016 – 17 had the presence of *Bdv2* (Table S1).

386

387 **Population Structure**

388 We studied the population structure of 346 wheat lines using 29,480 GBS-derived SNP markers.
389 The PCA did not reveal a strong pattern of population structure (Figure 3). Moreover, the
390 variation explained by the first two principal components (4 and 3%, respectively) also supports
391 the hypothesis of minimal population structure within a single breeding program. We observed
392 that most of the wheat cultivars released by KSU breeding program were located outside the
393 cluster grouping all the breeding lines (Figure 3A). Lines with the presence of *Bdv2* clustered
394 together (Figure 3B), likely due to a related pedigree to the original source, and we did not
395 identify any evident pattern for BYD severity associated with the population structure (Figure
396 3C).

397

398 **Genome-Wide Association Analysis**

399 To investigate the genetic architecture of BYD we performed GWAS analyses for all collected
400 traits using the BLUP values for 346 lines and 29,480 SNP markers. The first two principal
401 components from PCA and the kinship matrix were included in the mixed model to account for
402 population structure and genetic relatedness. We found significant marker-trait associations for
403 BYD severity on chromosomes 5AS, 7AL, and 7DL (Figure 4A). The highest peak was
404 observed on the proximal end of chromosome 7DL, located at 571 Mbp – 637 Mbp. To test the
405 hypothesis that this association was explained by the resistance gene *Bdv2* (located on
406 chromosome 7DL), we investigated the haplotypes defined by the 16 SNP markers associated
407 with BYD severity and were able to identify two haplotypes that exactly matched the presence or
408 absence of *Bdv2* (Fig 4A). This same region was mapped using BYD severity and the presence
409 or absence of *Bdv2* as a fixed covariate (Figure 4B). This analysis (Figure 4B) also detected a
410 peak on chromosome 7AL. Lastly, we explored the effect of *Bdv2* on both BYD BLUEs and
411 BLUPs, and we observed that the presence of *Bdv2* had a positive effect in reducing the disease
412 severity by approximately 10% (Figure 5A). The significant peak on chromosome 5AS, located
413 at 46 Mbp – 103 Mbp, was explained by 10 SNP markers, comprising two main haplotypes, one
414 of them associated with reduced BYD severity (Fig 5B). When we combined the different 5AS
415 haplotypes with *Bdv2*, we observed that the presence of *Bdv2* had a positive effect, reducing the
416 levels of BYD when combined with both 5AS haplotypes (Figure 5C), and suggesting an

417 additive effect. Compared to the associations found for *Bdv2* (Fig 4B), we did not find any
418 strong evidence of marker trait associations for the other evaluated traits (Figure S4).

419

420 **Genomic Selection**

421 To evaluate the potential of GS to predict BYD disease severity, we fit several GS models to the
422 phenotypic BLUPs of BYD, PTH_M, and reduction in GY. Across all traits, to determine
423 predictive ability we used a five-fold cross validation where prediction ability ranged from -0.08-
424 0.26. There was relatively good predictive ability for BYD severity ranging between 0.06 –
425 0.26, in comparison with PTHT_M and reduction in GY resulting in a lower range from 0.02 – 0.17
426 and -0.08 – 0.2, respectively (Figure 6). Evaluating the conformation of the training population,
427 we observed that when including 2016-17 season, prediction abilities were the highest for BYD
428 but the lowest for the other two traits, implying that season 2016 – 17 was either a good season
429 to train the prediction models or a difficult season to predict based on available data.

430

431 To further investigate the power of GS, we developed models using a leave-two-out strategy,
432 where two seasons were excluded from the training population and used as the testing
433 population. We fitted GS models for all possible two-season combinations. This strategy
434 resulted in slightly smaller training populations which decreased overall predictive ability
435 (Figure 6). This result was evident for BYD predictions where excluding two seasons had a larger
436 negative impact.

437

438 Lastly, we evaluated the effect of adding information about the genotype of the *Bdv2* resistance
439 gene as a phenotypic fixed covariate into the GS models. There were differences in the effect of
440 *Bdv2* on the predictive ability across BYD severity, PTHT_M, and GY, showing a large effect for
441 predicting BYD but almost no effect for PTHT_M and reduction in GY (Figure 6). The improved
442 predictive ability for BYD was clearly reflected with the decrease of prediction ability obtained
443 when season 2017 – 18 was excluded from the training population since most of the lines with
444 the presence of *Bdv2* were evaluated in that season.

445

446 **Discussion**

447

448 **Phenotypic Data**

449 The success of breeding for BYD resistance is highly impacted by the ability to precisely
450 characterize breeding material and disease symptoms. Even though BYD is spread worldwide,
451 its incidence in a given year depends on several factors such as aphid pressure, planting date, and
452 environmental conditions (e.g., temperature, rainfall, frost, etc.). In this study, we evaluated
453 winter wheat advanced breeding lines during five seasons implementing a rigorous field-testing
454 approach, that ultimately enabled us to consistently have plots contrasting with BYD infection
455 and uninfected or low incident plots. Moreover, by using large yield-size plots we were able to
456 calculate the reduction in GY and use this parameter as an estimate of field resistance.

457

458 The expression of BYD symptoms, however, was highly inconsistent during the different
459 seasons. Seasons 2015-16 and 2016-17 showed the best expression of the disease symptoms,
460 supported by the wide range of BYD severity between treated and untreated replications (Figure
461 1). Interestingly, both these seasons were conducted in the same experimental field (Table 1),
462 suggesting that this location could favor the expression of BYD. Moreover, weather conditions
463 were variable for all the seasons, suggesting that these had a huge impact on the disease
464 occurrence. While temperature records were similar for all the seasons, precipitation records did
465 show some differences. Season 2017 – 18 was dryer than normal, with 34% less precipitation
466 than the 30 years historical average (1981 – 2010). On the other hand, season 2018-19 was
467 wetter than normal, with 58% more precipitation than the 30 years historical average (Table S2).

468

469 **High-Throughput Phenotyping**

470 Evaluating BYD resistance using visual phenotypic selection can be challenging due to the
471 complex nature of the disease and rater variability (Poland and Nelson 2011). The use of HTP
472 with UAS is gaining popularity within breeding programs because it further improves selection
473 based on classical phenotyping. Accurate phenotyping is crucial for understanding the genetic
474 basis of quantitative and complex traits like BYD. In this study, we used HTP to complement
475 the visual BYD scoring. This tool improved our capacity for rapid, non-destructive, and non-
476 biased evaluation of large field-scale numbers of entries for BYD resistance. We were able to

477 determine strong correlation patterns between visual BYD severity and HTP derived parameters
478 (Figure S3). However, none of the traits collected with UAS had a common genetic base with
479 BYD severity (Figure 4 and Figure S4). Disease scoring using HTP is scaling fast among
480 breeding programs; however, how to effectively use this data remains challenging. Some studies
481 have shown that data collected with sensor-based tools can be substituted to improve classical
482 disease visual evaluation (Sankaran *et al.* 2010; Kumar *et al.* 2016; Zheng *et al.* 2018); however,
483 to the best of our knowledge this study is the first attempt to characterize BYD in wheat using
484 HTP.

485

486 **Genome-Wide Association Analysis**

487 Using GWAS we detected QTLs on chromosomes 5AS, 7AL, and 7DL for BYD severity BLUPs
488 values. Using GBS tags that mapped to known alien fragments, we confirmed *Bdv2* resistance
489 gene was located at 7DL, and confirmed that the 7DL QTL was explained by the presence of the
490 *Bdv2* resistance gene. Even though only 33 wheat lines were positive for the presence of *Bdv2*,
491 we still had enough power to detect its effect, suggesting that *Bdv2* has a strong effect on BYD
492 under Kansas field conditions (Figure 5). The associations on chromosome 7AL, observed for
493 both BYD severity and *Bdv2*, suggest that the SNP markers on the 7AL peak may be miss-
494 anchored markers that should have mapped to 7DL. The relatively high heritability values
495 obtained for the untreated replications (Figure 2) allowed us to detect a minor QTL on 5AS.
496 Marza *et al.* (2005) reported a QTL at 38cM on the short arm of chromosome 5A associated with
497 yellowing symptoms caused by BYD, and it is possible that this is the same region yet more data
498 is needed to confirm if these QTLs are the same. The only other study reporting GWAS for BYD
499 in wheat was able to identify several markers associated with BYD resistance on chromosomes
500 2A, 2B, 6A, and 7A (Choudhury *et al.* 2019b). However, most of the association were explained
501 by individual SNP markers, and to date do not have any definitive biological link. GWAS
502 results for the other traits used in this study did not discover genomic regions associated with the
503 traits (Figure S4). Taken together, these results suggest that BYD resistance is not controlled by
504 any large effect loci that could easily be incorporated into the breeding program, thus GS could
505 be an efficient way to enhance BYD resistance.

506

507 **Genomic Selection**

508 We evaluated several different GS models to identify the best approach for predicting BYD
509 (Figure 6). Overall, we observed some trends including i) incorporating years with consistent
510 BYD disease data in the training population increased the model predictive ability, ii) predicting
511 years with high disease pressure is difficult, iii) using major effect QTL, such as *Bdv2*, had
512 increased prediction performance, suggesting that it is responsible for much of the predictive
513 power. These results suggest that GS based on G-BLUP with *Bdv2* as fixed effects would lead to
514 the greatest genetic gain for BYD breeding. Using selected major QTL as a fixed effect to
515 improve GS models was suggested in a simulation study (Bernardo 2014) and demonstrated with
516 empirical studies (Rutkoski *et al.* 2014). Nonetheless, using *Bdv2* as a fixed effect in our GS
517 strategies did not consistently improve the predictive ability for PTH_M or reduction in GY (Rice
518 and Lipka 2019). However, there was not a consistent distribution of *Bdv2* allele across the
519 cohorts. BYD predictions were low compared to other disease (reviewed by Poland and
520 Rutkoski 2016). However, since this is the first report of GS for BYD resistance in wheat, we do
521 not have similar results to make better comparisons. BYD has traditionally been reported to
522 have low H^2 (Tola and Kronstad 1984; Choudhury *et al.*, 2019b) and in this study, even with
523 well managed plots that often had H^2 approaching 0.8, we still had difficulty reproducing
524 these results year to year as evidence of the challenge of studying this pathosystem. Moreover,
525 the correlation between HTP parameters and BYD phenotypes was interesting, but not sufficient
526 to be useful in combination with GS in the germplasm tested.

527

528 **Conclusions**

529 We were able to show that *Bdv2* has a major effect controlling BYD resistance in the KSU
530 breeding germplasm. Apart from the known *Bdv2* and a potentially novel 5AS region, we did
531 not find evidence of other regions controlling BYD resistance supporting the hypothesis of
532 limited resistance available in the current wheat gene pool and the highly polygenic nature of the
533 trait. Moreover, our study was the first attempt to characterize and improve BYD field-
534 phenotyping using HTP and apply GS to predict the disease. HTP traits showed strong
535 correlation patterns with BYD severity, however, none of these parameters shared a common
536 genetic architecture with BYD severity. The GS predictive ability results that we found in this
537 study open the door for further improvement and testing GS implementation for breeding for

538 BYD resistance. Continuing the improvement of BYD characterization and the search of new
539 sources of resistance using species related to wheat, will be crucial to broadening the resistant
540 genes available to introgress into wheat germplasm.

541

542

543 **Data Availability Statement**

544 Supplemental material, including raw and analyzed phenotypic data, genotypic data,
545 supplementary tables and figures, and basic plot scripts are available at Dryad
546 doi:10.5061/dryad.ncjsxkswd (temporary link: [https://datadryad.org/stash/share/xkKdr62QYB-](https://datadryad.org/stash/share/xkKdr62QYB-YA93mkzq18_4yFUgwdnvv0uZUyuEHpAI)
547 [YA93mkzq18_4yFUgwdnvv0uZUyuEHpAI](https://datadryad.org/stash/share/xkKdr62QYB-YA93mkzq18_4yFUgwdnvv0uZUyuEHpAI)) and GitHub

548 https://github.com/umngao/wsm1_bdv2

549 **Supplementary data description**

550 **Table S1** – List of wheat entries phenotypically evaluated in the study. The table includes the
551 type of entry (cultivar or breeding line), the season that the entry was evaluated, the result for the
552 prediction of the presence/absence of the segment carrying the resistance gene *Bdv2*, and the best
553 linear unbiased predictors (BLUPs) for all the phenotypic traits collected.

554 **Table S2** – Precipitation (inches) during the five field seasons in Riley County, KS, where
555 Rocky Ford and Ashland Bottoms experimental units are located. Normal temperature is defined
556 as a 30–year average from 1981 – 2010. Data was obtained from Kansas State University
557 (<http://climate.k-state.edu/precip/county/>)

558 **Figure S1** – Growth trajectories and adjustment of the non-linear regression model of wheat
559 lines for A-B) normalized difference vegetation index (NDVI) and C-D) digital plant height
560 (meters). The data used correspond to season 2016 – 17 phenotypic data. Calendar days is the
561 number of days starting at January 1, 2017.

562 **Figure S2** – Boxplots showing the phenotypic response of the wheat checks ‘Art’ (susceptible)
563 and ‘Everest’ (tolerant) for A) barley yellow dwarf (BYD) disease severity (%), B) manual plant
564 height (PTHT_M) (m) and C) grain yield (GY) (tons/ha). Adjusted phenotypic values are shown
565 for both insecticide treatment replications (treated and untreated).

566 **Figure S3** – Scatterplots showing distribution and Pearson’s correlation values for the
567 phenotypic traits studied during all the field seasons under two insecticide treatments (treated

568 and untreated). A-B) season 2016 – 17, C-D) season 2017 – 18, E-F) season 2018 – 19, and G-H)
569 season 2019 – 20.

570 **Figure S4** – Manhattan plots showing genome-wide association analysis (GWAS) results for the
571 phenotypic traits collected during the study.

572

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579

580

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Table 1 – Field experimental details for the five wheat nurseries

Season	2015 – 2016	2016 – 2017	2017 – 2018	2018 – 2019	2019 – 2020
Location	Rocky Ford farm		Ashland Bottoms farm		
	39°13'45.60" N, 96°34'41.21" W		39°07'53.76" N, 96°37'05.20" W		
Planting Date	Sep. 17, 2015	Sep. 12, 2016	Sep. 19, 2017	Sep. 17, 2018	Sep. 17, 2019
Number of Entries	68	52	81	81	107
Number of Plots	504	360	400	392	684
Field Design	split-plot with insecticide treatment as major factor effect and wheat genotype as secondary factor				
Replications	3	3	2	2	2
Plot Size	6 rows plots - 1.5 m × 2.4 m				
BYD Evaluation	April 28, 2016	May 12, 2017	May 19, 2018	May 13, 2019	May 19, 2020
Harvesting Date	June 20, 2016	June 19, 2017	June 23, 2018	June 28, 2019	June 25, 2020

Table 2 – Dates of high-throughput phenotypic data collection and details of image acquisition in the five wheat nurseries screened for BYD, Kansas, USA (2015-2020).

Season	2015 – 2016	2016 – 2017	2017 – 2018	2018 – 2019	2019 – 2020	
UAS Platform	PheMU		DJI Matrice 100			
Imaging Sensor	multiple digital single-lens reflex (DSLR) cameras		MicaSense RedEdge-M			
Flight/Pass speed	0.3–0.5 m/s		2 m/s			
Flight Dates				2019-04-01		
				2019-04-09		
		2017-03-28		2019-04-19		
		2017-04-13	2018-03-30	2019-04-26	2020-03-20	
		2017-05-01	2018-04-04	2019-05-02	2020-04-11	
		2016-03-31	2017-05-09	2018-04-12	2019-05-10	2020-04-23
		2016-04-07	2017-05-21	2018-04-19	2019-05-15	2020-05-03
		2016-04-14	2017-05-23	2018-04-23	2019-05-23	2020-05-19
		2016-05-06	2017-05-30	2018-05-16	2019-05-31	2020-06-05
			2017-06-05	2018-06-13	2019-06-05	2020-06-11
			2017-06-13		2019-06-12	
					2019-06-17	
	Flight/Pass altitude	0.5 m above the canopy		20 m AGL		
	In-Air Flight Duration	NA		~ 11–14 min		

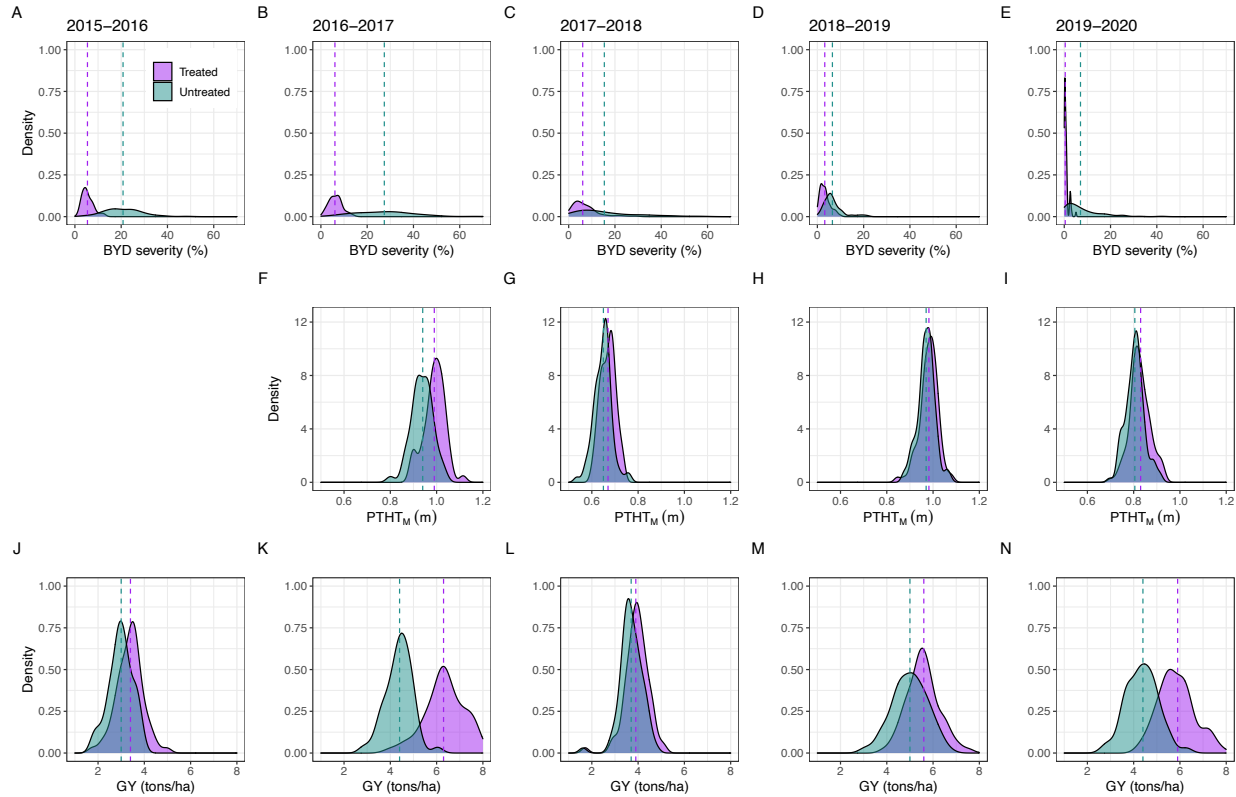


Figure 1 Adjusted phenotypic values for the traits collected manually for five different field seasons (2015-2016 to 2019-2020). A-E) barley yellow dwarf (BYD) severity (%) characterized as the typical visual symptoms of yellowing/purpling on leaves using a 0 – 100% visual scale, F-I) manual plant height/stunting (PTHT_M) (meters), note that the trait was not recorded for the 2015 – 2016 season, and J-N) grain yield (GY) (tons/ha). Insecticide-treated and untreated replications are represented by purple and green, respectively. The dashed line represents the mean value for the trait in each treatment

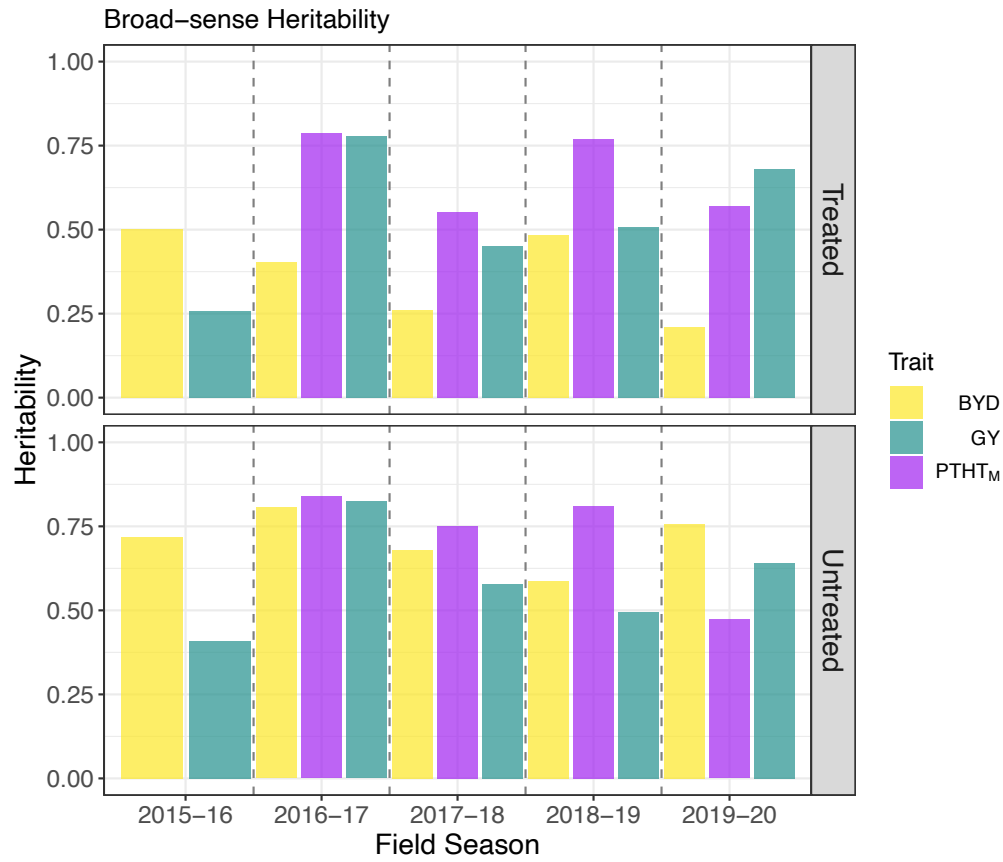


Figure 2 Broad-sense heritability of wheat phenotypic traits collected manually, including visual barley yellow dwarf (BYD) score, plant height (PTHT_M) and grain yield (GY) during five different field seasons under two insecticide treatments.

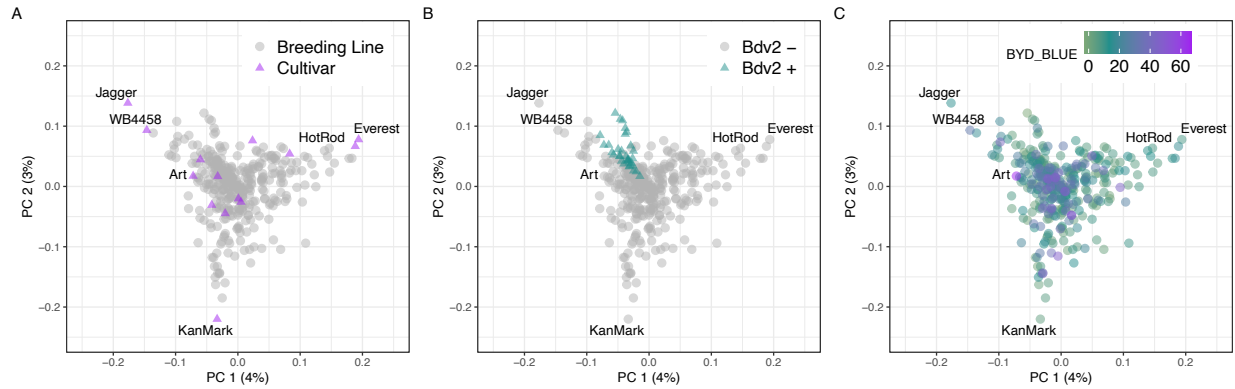


Figure 3 Scatterplot of the first two principal component axis, made from principal component analysis on the marker matrix, $n = 357$ wheat lines, markers = 29,480. Each data point represents an individual wheat line that is color-coded by A) breeding status, B) prediction of *Bdv2* presence/absence, and C) adjusted mean for BYD severity (BYD BLUE) scored visually. Total variance explained by each principal component (PC) is listed on the axis.

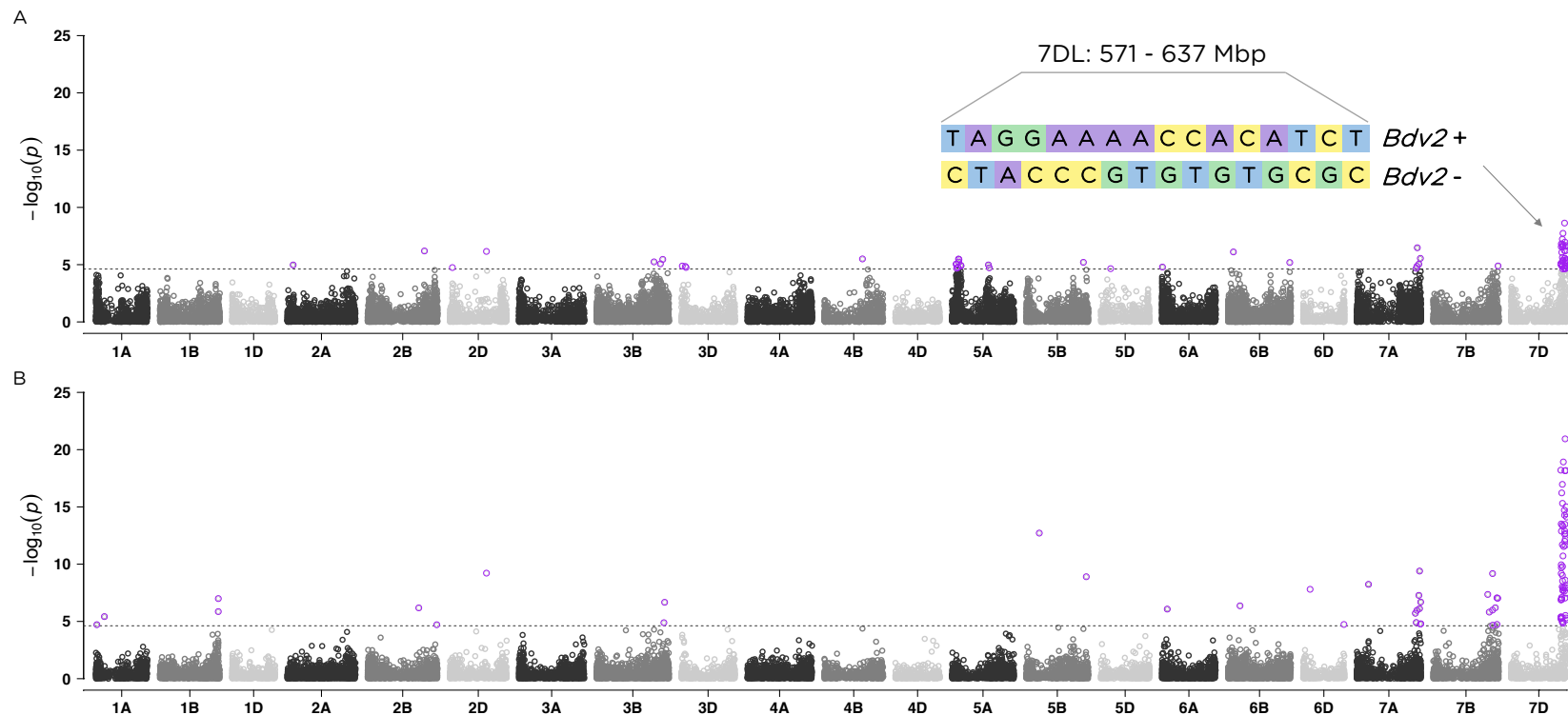


Figure 4 Manhattan plots showing the marker-trait associations using 346 wheat accessions and 29,480 SNP markers obtained with genotyping-by-sequencing (GBS) for A) BYD severity and B) presence/absence of *Bdv2* resistance gene. The 21 labeled wheat chromosomes with physical positions are on the x-axis and y-axis is the $-\log_{10}$ of the p-value for each SNP marker. Horizontal dashed lines represent the false discovery rate threshold at 0.01 level and data points highlighted in purple and above the threshold represent SNPs significantly associated with the trait. In panel a, the length of the region and the haplotypes defined by the significant SNP markers is displayed.

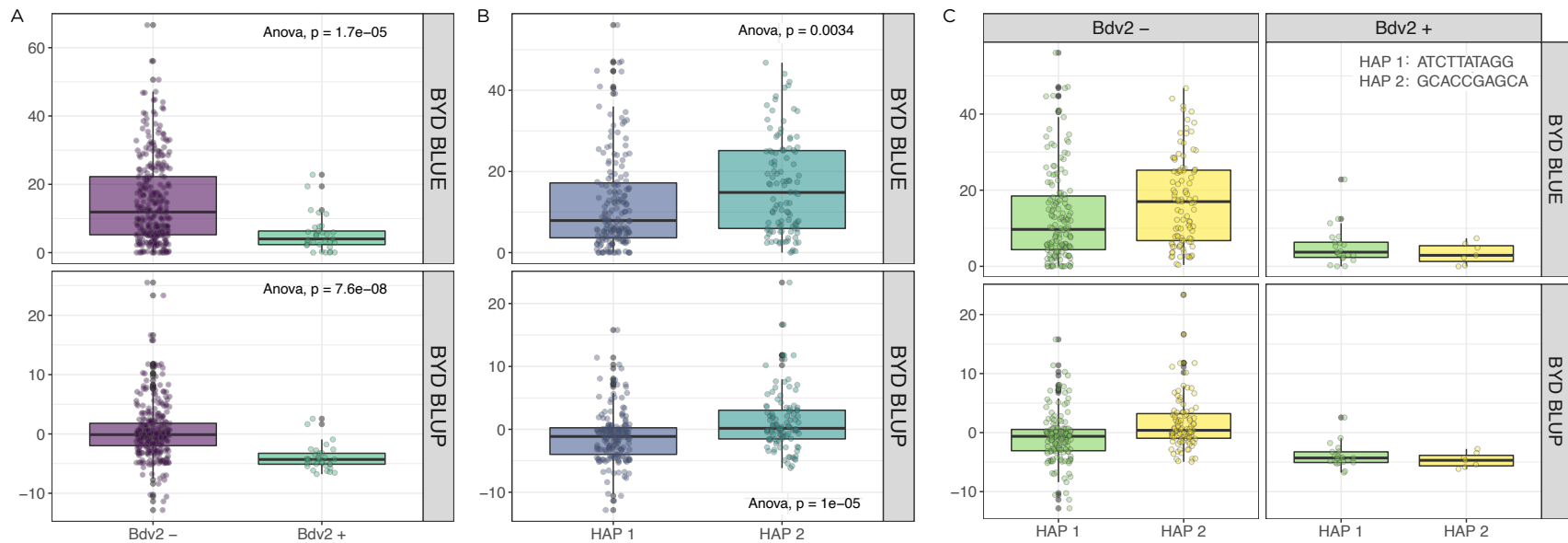


Figure 5 Measurement of barley yellow dwarf disease severity in wheat based on certain haplotype effects were panel A) represents the translocation segment carrying the resistance gene Bdv2, B) displays the two haplotypes for the significant region on chromosome 5AS, and C) shows the combination of Bdv2 resistance gene and 5A haplotype. Boxplots showing the significant reduction of BYD disease severity by averaging the phenotypic best linear unbiased estimated (BLUE) or best linear unbiased predicted (BLUP) values for the lines.

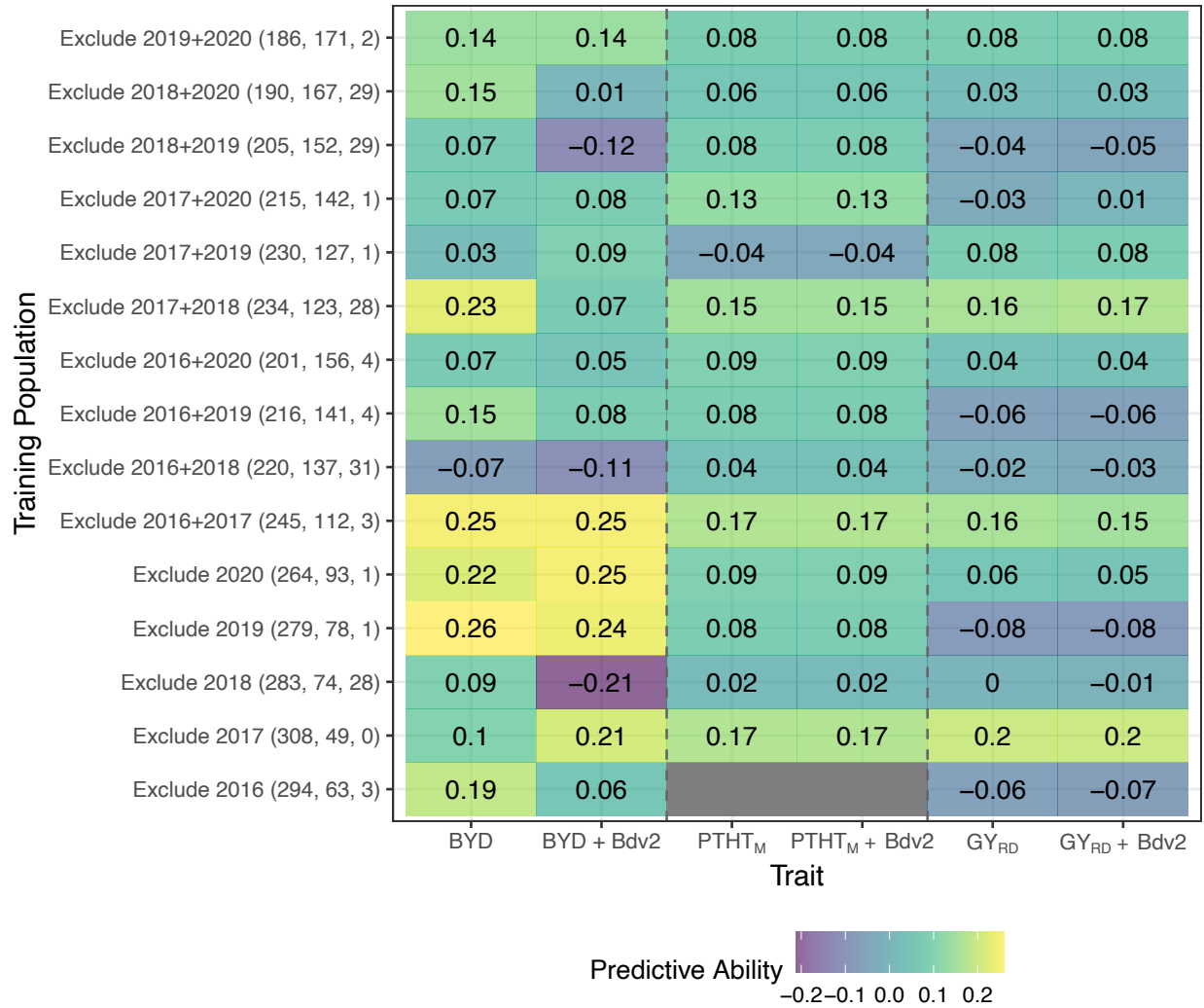


Figure 6 Genomic selection model predictive ability where each column represents one trait, and each row shows the conformation of the training population including size of training and testing population and number of lines with presence of *Bdv2* resistance gene. The value in each cell represents the predictive ability which is the correlation between the GS predicted value (GBLUP) and the phenotypic best linear unbiased predictor (BLUP).