# ACCEPTED ARTICLE Genome-wide association mapping revealed SNP alleles associated with resistance to cereal cyst nematode (*Heterodera filipjevi*) in wheat

Z. Majd Taheri<sup>1, 4\*,</sup> Z. Tanha Maafi<sup>1</sup>, K. Nazari<sup>2</sup>, K. Zaynali Nezhad<sup>3</sup>, F. Rakhshandehroo<sup>4</sup>, and A. A. Dababat<sup>5</sup>

7 1. Iranian Research Institute of Plant Protection, Agricultural Research, Education and Extension Organization
 8 (AREEO), Tehran, Islamic Republic of Iran.

- 9 2. Regional Cereal Rust Research Center, Aegean Agricultural Research Institute, P.K. 9, Menemen, Izmir,
   10 Turkey.
- 3. Plant Breeding and Biotechnology Department, Gorgan University of Agricultural Sciences and Natural Resources, Gorgan, Islamic Republic of Iran.
- 4. Department of Plant Protection, College of Agricultural Sciences and Food Industries, Science and Research
   Branch, Islamic Azad University, Tehran, Islamic Republic of Iran.

15 5. International Maize and Wheat Improvement Center, Emek, Ankara, Turkey.

16 \*Corresponding author; e-mail: <u>Majdtaheri@vahoo.com</u>

#### 18 Abstract

4

5

6

17

Resistance traits are economically important in crops in terms of accessibility to promising 19 resistant germplasms. This study was conducted to evaluate SNP marker-trait association for 20 cereal cyst nematode (CCN), Heterodera filipjevi in a large number of natural bread wheat 21 populations. Phenotypic data analysed using GLM (Generalized Linear Model) indicated 22 significant differences among the landrace accessions for resistance to H. filipjevi. The 23 24 genotyping was performed by 152K SNP chip on 188 accessions. After filtering, 10,471 polymorphic SNPs were employed for Genome Wide Association Study (GWAS). Population 25 26 structure among the wheat genotypes were investigated using 840 well distinct SNP markers. Two sub-populations were revealed by structure software, and eleven markers were found to 27 be significantly (p-value < 0.001) associated with resistance to H. filipjevi on chromosomes 28 2A, 3B, 4A, 4B, 5A, 5B, 5D, and 6B. The linkage disequilibrium analysis for all significantly 29 associated SNPs showed that markers on chromosomes 4A and 4B were in high intra-30 chromosomal linkage disequilibrium, and consequently, eight markers were recommended as 31 strongly associated with resistance to H. filipjevi. The present study demonstrated valuable 32 sources of resistance in the studied wheat genotypes against a widespread and important species 33 of CCNs. The associated markers could be used in molecular breeding programs of bread 34 wheat. 35

36 Keywords: Association mapping, Heterodera filipjevi, GWAS, SNP, Wheat.

## INTRODUCTION

Cereal Cyst Nematodes, CCNs (*Heterodera* spp.) are one of the most important causal agents of yield losses on wheat annually, hence its global importance is known in most wheat-growing

37

38

39

areas (Smiley et al., 2017; Toumi et al., 2018). The genus Heterodera is divided into nine 41 groups based on morphological and molecular characteristics (Handoo and Subbotin, 2018), in 42 which *H. filipjevi* is one of the most important species belonging to Avenae group. Host plants 43 of H. filipjevi include wheat, rye, barley, corn, and many grasses (Smiley et al., 2017). Yield 44 losses caused by H. filipjevi in three winter wheat cultivars in Iran were estimated to be 20.4 45 to 24.8% (Karimipour Fard et al., 2018). Wheat is one of the world's most commonly used 46 cereal grains growing all over the world and feeding more than 40% of the world population. 47 Amongst the different types of wheat grain, bread wheat (Triticum aestivum L., AABBDD) is 48 49 the most economically important crop and the world's most widely cultivated cereal. It is originated from hybridization between Triticum urartu (AA) and Aegilops speltoides-related 50 species (BB), forming Triticum turgidum ssp. dicoccoides, and again hybridized between 51 Triticum turgidum ssp. durum (AABB) and Aegilops tauschii (DD), forming the modern 52 hexaploid bread wheat (AABBDD). 53

Resistant cultivars are often regarded as one of the most effective tools for controlling CCNs. 54 Many sources have been reported and reviewed for conferring resistance measures. Important 55 sources of resistance genes were revealed in landrace varieties by identifying many resistance 56 Cre genes. In recent years, different types of molecular markers have been applied in plants 57 58 such as Restriction fragment length polymorphisms (RFLPs), microsatellites or simple sequence repeats (SSRs), expressed sequence tags (ESTs), cleaved amplified polymorphic 59 60 sequence (CAPS), randomly amplified polymorphic DNA (RAPD), amplified fragment length polymorphisms (AFLPs), inter simple sequence repeat (ISSR), Diversity arrays technology 61 62 (DArT) and single nucleotide polymorphism (SNP) (Dhingani et al., 2015). In genetic studies, single nucleotide polymorphisms (SNPs) are one of the most effective tools. SNPs are more 63 64 powerful in estimating population structure which are abundant in the genome.

In recent years, research on wheat genome recorded 90K SNP by the new Infinium to 500K 65 and 4 M in Illumina shortgun WGS array (Avni et al., 2014; Wang et al., 2017; Lai et al., 66 2015). Association mapping (AM), is known extremely for the identity of markers associated 67 traits based on linkage disequilibrium (LD) in plants. AM has been applied to discovery of 68 quantitative trait loci (QTL) on chromosomes in range of crop species. To date, QTL regions 69 70 on different chromosomes were detected in association with particular traits using AM in wheat 71 such as pre-harvest sprouting resistance, low  $\alpha$ -amylase and seed color (Rabieyan *et al.*, 2022) 72 and grain-associated traits (Wang et al., 2017), resistance to CCNs (Heterodera spp.), resistance to root lesion nematode (Pratylenchus spp.) and resistance to crown rot (Dababat et 73 al., 2016; Erginbas-Orakci et al., 2018; Kumar et al, 2021; Sohail et al., 2022). 74

75 Several QTLs have been suggested to affect on resistance to *H. filipjevi*. The first survey of

76 QTLs conferring resistance to H. filipjevi in wheat reported eleven QTLs on chromosomes

1AL, 2AS, 2BL, 2D, 3AL, 3BL, 4AS, 4AL, 5BL, 6B, 6D and 7BL (Pariyar *et al.*, 2016;
Dababat *et al.*, 2021).

The aim of the present study was to *a*: find marker-trait associations within 188 wheat genotypes collected from West Asia-North Africa, WANA, *b*: identify SNPs associated with resistance to *H. filipjevi* in wheat, *c*: combine analyses of phenotypic data and association mapping.

83

# 84 MATERIAL AND METHODS

## 85 Plant Materials and Inoculum Preparation

86 A total of 223 wheat accessions originating mostly from West Asia and North Africa (WANA countries) with three wheat cultivars as susceptible and resistant controls were used to evaluate 87 their resistance to H. filipjevi. It is worth to note that 188 accessions out of 223 accessions used 88 89 for phenotyping indicated sufficient DNA quality for SNP calls. The wheat accessions were 90 provided by the International Center for Agricultural Research in the Dry Areas (ICARDA), and were originated from Afghanistan (7), China (1), Iran (164), Iraq (3), Morocco (1), Pakistan 91 (7) and Syria (5) countries. The pedigree of the 188 wheat genotypes used in this study is given 92 in supplementary Table.1. For the preparation of inoculum, the collecting of nematodes, 93 extracting, identifying, incubation of the cysts and obtaining infective juveniles were conducted 94 95 as described by Majd Taheri et al., (2019).

96

## 97 Phenotyping Assessment

The phenotypic evaluation was performed in a growth chamber at the Iranian Research 98 99 Institute of Plant Protection (IRIPP). Wheat seeds of each accession, were sterilized, germinated and planted in a plastic tube filled with a mixture of sand, field soil, and organic 100 101 matter (70:29:1, v:v:v) arranged in a completely randomized design with five replications. The wheat cultivars Bezostaya and Sonmez were chosen as the susceptible and resistant checks, 102 103 respectively. Each plant was inoculated with 1 mL of inoculum containing 500 fresh second stage juveniles in a water suspension. After nine weeks, the level of resistance was counted and 104 105 categorized into four groups based on the number of white females and cysts, Resistant (R)  $\leq$ 3; Moderately resistant (MR) = 3-7; Susceptible (S) = 7-20; Highly Susceptible (HS)  $\ge 20$ 106 according to Sharma et al., (2013). Normality of data and Homogeneity of variances were 107 examined with Shapiro-Wilk test and Levine's test, respectively. All phenotypic data were 108

analysed using Generalized Linear Model (GLM) using statistical software SAS v9.4 and mean
separation was conducted using Duncan's Multiple Range Test.

111

## 112 Genotyping and Data Preprocessing

Genomic DNA was extracted from fresh leaves using a modified CTAB 113 (cetyltrimethylammonium bromide) method as described by Saghai-Maroof et al. (1984). 114 Samples were genotyped by genotyping-by-sequencing (GBS) and Diversity Arrays 115 Technology (DArT) (Sansaloni et al., 2011) using 152K SNP panel at the Genetic Analysis 116 Service for Agriculture (SAGA) at the International Maize and Wheat Improvement Center 117 (CIMMYT), Mexico. The quality of genotypic data were curated by removing SNPs with 118 minor allele frequency (MAF) less than 0.05 and missing data more than 20% from the 119 120 subsequent analysis (Bhatta et al. 2018), and the heterozygous data were considered as missing data (Mourad et al. 2018; Pariyar et al. 2016), which left a set of 10,471 polymorphic SNP 121 markers with known chromosomal position (based on Chinese spring map of IWGSC RefSeq 122 v1.0 assembly (Appels *et al.*, 2018)) for further analysis. 123

124

## 125 Analysis of Population Structure

The 840 SNP markers were selected based on physical position on chromosomes (A, B and D) from the total 10,471 markers with known chromosomal positions. Population structure analysis was performed using a Bayesian model in software STRUCTURE v2.3.4 (Pritchard *et al.*, 2000), where number of populations (K) were assumed from 1 to 10 using 100,000 burn iterations followed by 100,000 Markov-Chain Monte Carlo (MCMC) iterations. Process was repeated 5 times for each K. Output was visualized using STRUCTURE harvester and the optimal K value was identified based on the LnP(D) and Evanno's  $\Delta$ K (Evanno *et al.*, 2005).

133 134

# Linkage Disequilibrium Association mapping

Linkage disequilibrium and Genome-Wide Association Study, GWAS were implemented 135 using 10,471 SNPs with known chromosomal positions. Chinese Spring genome map IWGSC 136 RefSeq v1.0 assembly was used as the reference genome (Appels et al., 2018). A mean pairwise 137 r for the 21 chromosomes was determined. The LD heat maps plot for significantly associated 138 SNPs was constructed by using Haploview software 4.2 (Broad Institute, Cambridge, MA). 139 140 GWAS was conducted using the General linear model (GLM) and Mixed linear model (MLM) (Q+K) in TASSEL v. 5.2.51 (Bradbury et al., 2007). The Q matrix was adapted from the K=2 141 for association mapping for controlling spurious results due to population stratification as a 142 major issue in GWAS. TASSEL software was employed to estimate kinship matrix and the 143

association analyses were carried out to generate Manhattan and quantile-quantile plots (Q–Q
plot). A threshold P-value of 0.001 (–log10P= 3) was applied to declare significant SNPs for
marker-trait association results. The phenotypic variation (R<sup>2</sup>) was estimated for significant
markers. To reduce the false discovery rate, FDR was implemented at 0.001 level in SAS v 9.4
(SAS Institute Inc., Cary, NC, United States).

149

## 150 **RESULTS AND DISCUSSION**

Wild relatives of wheat are important sources of disease resistance. In recent years, different 151 types of molecular markers have been applied to study the genetic traits in many crops i.e., 152 barley (Hordeum vulgare L.), maize (Zea mays L.), potato (Solanum tuberosum L.), rice (Oryza 153 sativa L.), soybean (Glycine max (L.) Merr.), sorghum (Sorghum bicolor L.) Moench), tomato 154 155 (Lycopersicon esculentum Mill.) and wheat (Triticum aestivum L.). SNP chips were mostly applied in GWAS which makes it easier to identify QTLs associated with certain traits. Our 156 raw data and variances were normal and homogeneous, respectively. The analyses of 157 phenotypic data revealed significant differences among the accessions for resistance to H. 158 159 filipjevi (Table 1). The 35 % of wheat accessions showed resistant (R) reaction to H. filipjevi, 44% of the accessions were moderately resistant (MR) and 21% were susceptible (S) (Figure 160 1). Most of the Iranian genotypes indicated moderately resistant (45%) trait (Majd Taheri et 161 al., 2019). Of the 10,471 SNPs found to be highly associated with resistance to H. filipjevi, 162 4,096 (39%), 4,739 (45%), and 1,636 (16%) SNPs were recorded on the AA, BB, and DD 163 genomes, respectively (Figure 2). AA and BB genomes have a higher distribution of SNPs than 164 the DD genome, this finding is in agreement with similar studies (Wen et al., 2017; Gahlaut et 165 al., 2019; Rabieyan et al., 2022; Tehseen et al., 2022). The minimum number of SNPs were 166 associated with resistance to H. filipjevi from chromosome 4D (147 SNPs) and most numbers 167 of SNPs were from 2B (887 SNPs). Population structure analysis implemented using 840 168 markers, indicated two possible subpopulations, based on the clear peak at k=2 (Figure 3). The 169 first and second group consisted of 62% and 38% of the wheat accessions, respectively. We 170 found significant differences among the genotypes for resistance to H. filipjevi. The genetic 171 diversity of wheat genotypes from our previous experiment revealed the suitability of this 172 group of wheat genotypes for association mapping studies (Majd Taheri et al., 2019). 173

Using 10,471 SNPs, linkage disequilibrium (LD) was determined by calculating squared correlation coefficient ( $r^2$ ) for the 21 chromosomes. We applied a mixed linear model (MLM) and General linear model (GLM) in GWAS analysis. QQ-plots and Manhattan plots of the GWAS results of both GLM and MLM analysis were compared for resistance trait which are

- shown in Figure 4. Based on the obtained QQ-plot from GLM and MLM models, the Q-Q plot
- 179 of GLM shows deviations from the slope line, demonstrating the loci which deviate from the
- 180 null hypotheses and indicating significant positive marker-trait association which makes GLM
- 181 **as a better approach.** Manhattan plots represent the profile of the P-value of SNPs in Figure 5.
- 182 A total of 11 SNPs significantly associated with resistance to *H. filipjevi* trait and crossed the 183 false detection rate (FDR) at p < 0.001 were identified. The phenotypic variation ( $r^2$ ) explained
- 184 by the individual SNPs ranged from 7 to 13% (Table 2).
- So far, some significant Marker-trait associations (MTAs) were identified on wheat 185 186 chromosomes to agronomic characteristics and diseases. This collection of wheat genotypes has not been utilized for resistance studies to cereal cyst nematode so far, however, GWAS of 187 diverse panels against *H. filipjevi* was done by Pariyar et al., (2016) and Dababat et al., (2021). 188 In the present study, 11 markers were significantly (p-value < 0.001) associating with resistance 189 to *H. filipjevi* which were detected on chromosomes No 2A, 3B, 4A, 4B, 5A. 5B, 5D and 6B. 190 The linkage disequilibrium (LD) analysis for all significantly associated SNPs showed that 3 191 markers on 4A and 2 markers on 4B Chromosomes were in high intra-chromosomal LD, hence 192 the 11 SNPs could be reduced to 8. It is noteworthy that the D genome carries only one of all 193 identified MTAs in this study, likely implies the low level of diversity in the D genome 194 195 originated from the late hybridization of Aegilops tauschii during the evolution of common wheat (Gahlaut et al., 2019). A previous GWAS have demonstrated 11 QTLs on chromosomes 196 197 1AL, 2AS, 2BL, 3AL, 3BL, 4AS, 4AL, 5BL and 7BL (Pariyar et al., 2016). Another study identified QTLs on chromosomes 1A, 2A, 2B, 2D, 3A, 6B, and 6D were detected using a mixed 198 linear model (MLM) (Dababat et al., 2021). Fourteen genes for resistance to CCN have been 199 identified which include the following: Cre1, Cre2, Cre3, Cre4, Cre5, Cre6, Cre7, Cre8, Cre9, 200 201 CreR, CreV, CreX, CreY and CreZ (Ali et al., 2019; Kishii, 2019; Dababat et al., 2021). CCN 202 resistance genes Cre1, Cre2, Cre3, Cre5 (syn. CreX), Cre6, Cre8 and CreR identified in wheat 203 and its relatives on chromosome 2B, 2A, 2D, 2A, 5A, 6B and 6D, respectively (Slootmaker et al., 1974; Asiedu et al., 1990; Eastwood et al., 1991; Delibes et al., 1993; Jahier et al., 1996; 204 Paull et al., 1998; Ogbonnaya et al., 2001). Our results demonstrated three QTLs (on 2A, 5A 205 and 6B) found on chromosomes with identified resistance genes. Surprisingly Cre8 gene as a 206 207 resistance gene to CCN, H. avenae was mapped by Williams et al., (2003) on chromosome 6B, 208 moreover the effective role of Cre8 in conferring of resistance to CCN, H. fili pjevi in wheat 209 was emphasized by Imren *et al.*, (2013). Our finding suggests that the marker identified in this study may be present in the genomic region of the Cre8 gene, however further evidences are 210 needed to confirm the exact loci. Similar to the present study some QTLs that confer resistance 211

against other cereal nematodes were recently reported on wheat, *i.e.* in *H. avenae* on
chromosomes 5A, 5B, 5D and 6B, in root lesion nematodes, *P. neglectus* on chromosomes 3B,
4A and 6B, *P. thornei* on chromosomes 2A, 3B and 5B (Dababat *et al.*, 2016).

Pleiotropic effect resistance genes to multiple races (HG types) of soybean cyst nematode 215 (SCN), H. glycines and Root knot Nematode (Meloidogyne incognita) was revealed in soybean 216 line 438489B which carrying multi-nematode resistance gene package (Vuong et al., 2011). 217 Recently two QT controlling reniform nematode, RN (Rotylenchulus reniformis) resistance 218 were identified in the SCN resistance gene GmSNAP18 at the rhg1 locus and its paralog 219 GmSNAP11 in soybean line 438489B (Usovsky et al. 2021). Hence, there is a possibility of 220 similarities between identified QTLs which necessitates further experiments to determine 221 common QTL between two or more pathogens. Importantly, it is obvious that a QTL with 222 capability of inducing resistance to plant against several traits, is a valuable resource in 223 breeding programs. In conclusion, in this study, 188 wheat accessions were applied to perform 224 association mapping, and 10,471 SNPs used for GWAS after filtering according to the MAF, 225 missing data, and heterozygous data. We estimated the phenotypic and genotypic parameters 226 for resistance trait and eleven significantly associated SNP markers were detected by GLM. 227 Based on the results, the use of populations from different genetic backgrounds provide further 228 229 progress in identifying valid QTLs. The findings of present study demonstrated valuable sources of resistance in the studied wheat genotypes to a widespread and important species of 230 231 CCN in some areas of the crescent fertile region for inclusion in future breeding programs by 232 new resistance gene resources.

233

238

# 234 ACKNOWLEDGEMENTS

We are particularly grateful to the Iranian Research Institute of Plant Protection (IRIPP),
International Center for Agricultural Research in the Dry Areas (ICARDA) and International
Maize and Wheat Improvement Center (CIMMYT) for valuable supports.

239 **REFERENCES** 

Ali MA, Shahzadi M, Zahoor A, Dababat AA, Toktay H, Bakhsh A, Nawaz MA and Li H.
2019. Resistance to cereal cyst nematodes in wheat and barley: an emphasis on classical and
modern approaches. *Int. J. Mol. Sci.*, 20(2): 432.

Appels, R., Eversole, K., Stein, N., Feuillet, C., Keller, B., Rogers, J., ... and Ronen, G. 2018.
Shifting the limits in wheat research and breeding using a fully annotated reference genome. *Science.*, 361(6403).

- Asiedu R, Fisher JM and Driscoll CJ. 1990. Resistance to *Heterodera avenae* in the rye
  genome of triticale. *Theor. Appl. Genet.*, **79:** 331-336.
- Avni R, Nave M, Eilam T, Sela H, Alekperov C, Peleg Z, Dvorak J, Korol A and Distelfeld
  A. 2014. Ultra-dense genetic map of durum wheat×wild emmer wheat developed using the
  90K iSelect SNP genotyping assay. *Mol. Breed.*, 34: 1549-1562.
- Bhatta M, Morgounov A, Belamkar V, Poland J, Baenziger PS. 2018. Unlocking the novel
  genetic diversity and population structure of synthetic Hexaploid wheat. *BMC Genom.*,19(1):591.
- Bradbury, P.J., Zhang, Z., Kroon, D.E., Casstevens, T.M., Ramdoss, Y., Buckler, E.S. 2007.
  TASSEL: Software for association mapping of complex traits in diverse samples. *Bioinformatics.*, 23: 2633–2635.
- 257 Dababat AA, Ferney G, Erginbas-Orakci G, Dreisigacker S, Imren M, Toktay H, Elekcioglu
- H, Mekete T, Nicol J, Ansari O and Ogbonnaya F. 2016. Association analysis of resistance to
- 259 cereal cyst nematodes (Heterodera avenae) & root lesion nematodes (Pratylenchus neglectus
- & *P.* thornei) in CIMMYT advanced spring wheat lines for semi-arid conditions. *Breed. Sci.*,
  66: 692-702.
- 262 Dababat A, Rehman-Arif MA, Toktay H, Atiya O, Shokat S, E-Orakci G, Imren M, Singh S.
- 263 2021. A GWAS to identify the cereal cyst nematode (*Heterodera filipjevi*) resistance loci in
  264 diverse wheat prebreeding lines. *J. Appl. Genet.*, 62: 93-98.
- 265 Delibes A, Romero D, Aguaded S, Duce A, Mena M, Lopez-Brana I, Andres MF, Martin-
- Sanchez JA and Garcia-Olmedo F. 1993. Resistance to cereal cyst nematode (*Heterodera avenae* Woll.) transferred from the wild grass *Aegilops ventricosa* to hexaploid wheat by a stepping-stone procedure. *Theor. Appl. Genet.*, 87: 402-408.
- Dhingani RM, Umrania VV, Tomar RS, Parakhia MV and Golakiya B. 2015. Introduction to
  QTL mapping in plants. *Ann. Plant Sci*, 4: 1072-1079.
- Eastwood R, Lagudah E, Appels R, Hannah M, Kollmorgen J. 1991. *Triticum tauschii*: a
  novel source of resistance to cereal cyst nematode (*Heterodera avenae*). *Aust. J. Agric. Res.*,
  42: 69–77.
- Erginbas-Orakci G, Sehgal D, Sohail Q, Ogbonnaya F, Dreisigacker S, Pariyar SR and
  Dababat AA. 2018. Identification of novel quantitative trait loci linked to crown rot resistance
  in spring wheat. *Int. J. Mol. Sci.*, **19:** 2666.
- Evanno G, Regnaut S and Goudet J. 2005. Detecting the number of clusters of individuals
  using the software STRUCTURE: a simulation study. *Mol. Ecol.*, 14: 2611-2620.

- Gahlaut V, Jaiswal V, Singh S, Balyan HS and Gupta PK. 2019. Multi-locus genome wide
  association mapping for yield and its contributing traits in hexaploid wheat under different
  water regimes. *Sci. Rep.*, **9**: 19486.
- Handoo ZA and Subbotin SA. 2018. Taxonomy, identification and principal species. In Perry
- 283 RN, Moens M and Jones JT (eds) Cyst nematodes. CAB International, Wallingford., p. 365284 397.
- Imren M, Toktay H, Bozbuğa R, Dababat A, Elekcioğlu H. 2013. Pathotype determination of
- 286 the cereal cyst nematode, Heterodera avenae (Wollenweber, 1924) in the Eastern
- 287 Mediterranean Region in Turkey. *Turk Entomol Derg.*,**1:** 13-19.
- Jahier J, Tanguy AM, Abelard P and Rivoal R. 1996. Utilisation of deletions to localise a
- gene for resistance to the cereal cyst nematode, Heterodera avenae, on an Aegilops ventricosa
- 290 chromosome. *Plant Breed.*, **115:** 282-284.
- 291 Karimipour Fard H, Pourjam E, Tanha Maafi Z, Safaie N. 2018. Assessment of yield loss of
- wheat cultivars caused by *Heterodera filipjevi* under field conditions. *J. Phytopathol.*, 166:
  293 299-304.
- Kishii, M. 2019. An update of recent use of *Aegilops* species in wheat breeding. *Front. Plant Sci.*, **10**: 585.
- 296 Kumar D, Sharma S, Sharma R, Pundir S, Singh VK, Chaturvedi D, Singh B, Kumar S and
- 297 Sharma S. 2021. Genome-wide association study in hexaploid wheat identifies novel genomic
- regions associated with resistance to root lesion nematode (*Pratylenchus thornei*). *Sci. Rep.*,
  11: 3572.
- Lai K, Lorenc MT, Lee HC, Berkman PJ, Bayer PE, Visendi P, Ruperao P, Fitzgerald TL, Zander M, Chan CH, Manoli S, Stiller J, Batley J and Edwards D. 2015. Identification and characterization of more than 4 million intervarietal SNPs across the group 7 chromosomes of
- 303 bread wheat. *Plant Biotechnol. J.*, **13**(1):97–104.
- Majd Taheri Z, Maafi ZT, Nazari K, Nezhad KZ, Rakhshandehroo F and Dababat AA. 2019.
  Combined study on genetic diversity of wheat genotypes using SNP marker and phenotypic
  reaction to *Heterodera filipjevi. Genet. Resour. Crop Evol.*, 66: 1791-1811.
- Mourad AMI, Sallam A, Belamkar V, Wegulo S, Bowden R, Jin Y, Mahdy E, Bakheit B, ElWafaa AA, Poland J, Baenziger PS. 2018. Genome-wide association study for identification
  and validation of novel SNP markers for Sr6 stem rust resistance gene in bread wheat. Front *Plant Sci.*, 9:380.

- Ogbonnaya FC, Seah S, Delibes A, Jahier J, Lopez-Brana I, Eastwood RF and Lagudah ES.
- 312 2001. Molecular-genetic characterisation of a new nematode resistance gene in wheat. *Theor.*
- 313 Appl. Genet., **102:** 623-629.
- Pariyar SR, Dababat AA, Sannemann W, Erginbas-Orakci G, Elashry A, Siddique S,
- 316 identifies resistance to the cereal cyst nematode *Heterodera filipjevi*. *Phytopathol.*, **106**: 1128-

Morgounov A, Leon J and Grundler FM. 2016. Genome wide association study in wheat

317 1138.

- Paull JG, Chalmers KJ, Karakousis A, Kretschmer JM, Manning S and Langridge P. 1998.
- Genetic diversity in Australian wheat varieties and breeding material based on RFLP data. *Theor. Appl. Genet.*, 97: 435-446.
- Pritchard JK, Stephens M and Donnelly P. 2000. Inference of population structure using
  multilocus genotype data. *Genetics.*, 155: 945-959.
- Rabieyan E, Bihamta MR, Esmaeilzadeh Moghaddam M, Mohammadi V and Alipour H.
- 324 2022. Genome-wide association mapping and genomic prediction for pre-harvest sprouting
- resistance, low α-amylase and seed color in Iranian bread wheat. *BMC Plant Biol.*, **22:** 300.
- Saghai-Maroof MA, Soliman K, Jorgensen RA and Allard RW. 1984. Ribosomal DNA
  spacer-length polymorphisms in barley: Mendelian inheritance, chromosomal location, &
  population dynamics. *Proc. Natl. Acad. Sci.*, 81: 8014-8018
- Sansaloni C, Petroli C, Jaccoud D, Carling J, Detering F, Grattapaglia D and Kilian A. 2011.
- 330 Diversity Arrays Technology (DArT) and next-generation sequencing combined: genome-
- wide, high throughput, highly informative genotyping for molecular breeding of Eucalyptus. *BMC Proc.*, 5: 54.
- 333 Sharma P, Saini M, Gupta OP, Gupta N, Singh AK, Selvakumar R, Tiwari V and Sharma I.
- 2013. Tracking of cereal cyst nematode resistance genes in wheat using diagnostic markers. *J. wheat res.*, **5:** 35-40.
- Slootmaker LAJ, Lange W, Jochemsen G and Schepers J. 1974. Monosomic analysis in bread
  wheat of resistance to cereal root eelworm. *Euphytica.*, 23: 497-503.
- Smiley RW, Dababat AA, Iqbal S, Jones MGK, Tanha Maafi Z, Peng D, Subbotin SA and
  Waeyenberge L. 2017. Cereal cyst nematodes: a complex and destructive group of *Heterodera*species. *Plant Dis.*, **101:** 1692-1720.
- 341 Sohail Q, Erginbas-Orakci G, Ozdemir F, Jighly A, Dreisigacker S, Bektas H, Birisik N,
- 342 Ozkan H and Dababat AA. 2022. Genome-Wide Association Study of Root-Lesion Nematodes
- 343 *Pratylenchus* Species and Crown Rot *Fusarium culmorum* in Bread Wheat. *Life.*, **12(3):** 372.

- 344 Tehseen MM, Tonk FA, Tosun M, Istipliler D, Amri A, Sansaloni CP, Kurtulus E, Mubarik
- 345 MS and Nazari K. 2022. Exploring the Genetic Diversity and Population Structure of Wheat
- Landrace Population Conserved at ICARDA Genebank. *Front. Genet.*, **13**: 900572.
- Toumi F, Waeyenberge L, Viaene N, Dababat AA, Nicol J, Ogbonnaya F and Moens M.
- 348 2018. Cereal cyst nematodes: Importance, distribution, identification, quantification, and
- 349 control. Eur. J. Plant Pathol., 150:1-20.
- 350 Vuong, T. D., Sleper, D. A., Shannon, J. G., Wu, X., and Nguyen, H. T. 2011. Confirmation
- 351 of quantitative trait loci for resistance to multiple-HG types of soybean cyst nematode
- 352 (*Heterodera glycines* Ichinohe). *Euphytica.*, 181, 101.
- Usovsky M, Lakhssassi N, Patil GB, Vuong TD, Piya S, Hewezi T, Robbins RT, Stupar RM,
- 354 Meksem K and Nguyen HT. 2021. Dissecting nematode resistance regions in soybean revealed
- 355 pleiotropic effect of soybean cyst and reniform nematode resistance genes. Plant Genome.,
- 356 **14(2): e20083.**
- 357 Wang SX, Zhu YL, Zhang DX, Shao H, Liu P, Hu JB, Zhang H, Zhang HP, Chang CH, Lu
- J, Xia XH, Sun GL and Ma CX. 2017. Genome wide association study for grain yield and
  related traits in elite wheat varieties and advanced lines using SNP markers. *Plos One.*, **12** (**11**):
  e0188662.
- Wen W, He Z, Gao F, Liu J, Jin H, Zhai S, Qu Y and Xia X. 2017. A high-density consensus
  map of common wheat integrating four mapping populations scanned by the 90K SNP Array. *Front. Plant Sci.*, 8: 1389.
- Williams KJ, Lewis JG, Bogacki P, Pallotta MA, Willsmore KL, Kuchel H and Wallwork H
  .2003. Mapping of a QTL contributing to cereal cyst nematode tolerance & resistance in wheat. *Aust. J. Agric. Res.*, 54: 731-737.
- 367 368

369

370 371 372



Fig 1. Phenotypic responses of wheat accessions to *Heterodera filipjevi* based on the number
of white females and cysts (Resistant≤ 3; Moderately resistant= 3-7; Susceptible= 7-20).

387

Table 1. Analysis of variance of the reaction of wheat genotypes to *Heterodera filipjevi* using
 Generalized Linear Model (GLM).

Source	Degrees of freedom	Mean of square	F value	<b>Pr</b> > <b>F</b>
Genotype	225	1.15	3.60	< 0.0001
Error	904	0.32	-	-
CV <sup>a</sup>	-	25.09	-	-
<sup>a</sup> Coefficient of va	riation.			

390 391

Table 2. Single nucleotide polymorphisms (SNPs) significantly associated with resistance to
 *Heterodera filipjevi*.

No.	SNP Marker	CHR	POS (bp)	FDR	P value	Allele	Allelic effect	$R^{2}$ (%)	cM
1	3034005	2A	7919418	0.00060	0.00007	T/C	-0.49	10	8
2	1106119	3B	183514671	0.00070	0.00034	T/G	0.89	13	-
3	2262587	4A	374449717	0.00060	0.00017	C/T	0.42	10	-
4	1220611	4A	430870112	0.00070	0.00046	T/C	-0.41	8	-
5	2266236	4A	433757511	0.00070	0.00037	G/A	-0.40	8	27
6	1128101	4B	605941582	0.00100	0.00081	G/A	-0.49	8	-
7	1244896	4B	608261318	0.00060	0.00023	C/G	0.52	9	45
8	1098989	5A	480788416	0.00100	0.00097	G/C	0.57	7	51
9	1209179	5B	506087964	0.00060	0.00011	A/G	-0.91	9	48
10	2260283	5D	380840282	0.00100	0.00096	G/A	0.52	7	-
11	1091272	6B	79045627	0.00070	0.00051	A/G	-0.42	8	23
394	CHR: Chromosor	ne; POS:	Position; FDR:	False Disco	very Rate;	R <sup>2</sup> : Effect	t due to genetic	variation:	<mark>Cm:</mark>
395	Centimorgan.								



#### 

Fig 2. Genome origin (A, B and D) of tested wheat SNPs of tested wheat genotypes. Yellow and blue columns represent highest and lowest numbers of SNPs, respectively.





Fig 3. A: Graph of delta K values showing highest probability at number of groups (K=2) and B: Estimated population structure of 188 wheat genotypes on k=2. 



Fig 4. QQ (Quantile-Quantile) plots, Red line represents the observed P values using the GLM (Q) model and green line represents the observed P values using the MLM (Q + K) model.



#### 424

433

434

Fig 5. Manhattan plots of *P* values showing genomic region of wheat genotypes associated with *Heterodera filipjevi* resistance. The X-axis represents the position of markers over the wheat chromosomes and Y-axis represents -log10 (P-values) of the marker-trait association. Each Point in the plot represents a SNP marker. The red line represents the threshold for genome-wide significance. Markers with -log10 (P-values) above the threshold are candidates.

# 431 با Heterodera filipjevi با مقاومت گندم به نماتد سیستی غلات SNP مرتبط با مقاومت گندم به نماتد سیستی غلات 432 استفاده از نقشه یابی ارتباطی گسترده ژنوم

## چکیدہ

وجود صفت مقاومت در گیاهان از لحاظ دسترسی به ژرم پلاسمهای مقاوم امید بخش حائز اهمیت اقتصادی است. این 435 مطالعه به منظور بررسی ارتباط نشانگر - صفت در تعداد زیادی از جمعیتهای گندم نان نسبت به نماتد سیستی غلات 436 Heterodera filipjevi انجام شد. نتایج حاصل از تجزیه و تحلیل آماری داده ای فنوتییی با استفاده از مدل خطی تعمیم 437 يافته (GLM) نشان داد، ژنوتيپها از لحاظ واکنش مقاومت به نماتد از اختلاف معنىدارى برخوردار هستند. ارزيابى 438 ژنوتیپی با استفاده از یک تر اشه SNP 152 H صورت گرفت. پس از اعمال کنترل کیفیت روی مجموعه داده−ها، تعداد 439 10471 نشانگر SNP برای نقشهیابی ارتباطی گسترده ژنوم (GWAM) استفاده شد. آنالیز ساختار جمعیت با استفاده از 440 840 نشانگر SNP، جمعیت مورد مطالعه را به دو زیر جمعیت طبقه بندی نمود. بازده نشانگر متعلق به هشت جایگاه 441 ژنی به طور معنی داری (p-value <) polue () در ارتباط با صفت مقاومت به نماتد روی کروموزوم-های A2، 3B، 442 A، 4B، 4A، 5B، 5B، 5B، 5B، 5B، 4E شناسایی شدند. از میان 11 نشانگر شناسایی شده، سه نشانگر روی کروموزوم A A، 443 دو نشانگر روی کروموزوم B 4 از میزان عدم تعادل پیوستگی بالایی برخوردار بودند. لذا تعداد 11 نشانگر شناسایی 444 شده، به هشت نشانگر کاهش یافت مطالعه حاضر منابع ارزشمندی از مقاومت به نماتد سیستی غلات را در ژنوتیپهای 445 گندم نشان داد. نشانگرهای مرتبط را میتوان در برنامههای اصلاح مولکولی گندم نان استفاده کرد. 446