Genetic Architecture of Resistance to Stripe Rust in a Global Winter Wheat Germplasm Collection

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ABSTRACT Virulence shifts in populations of Puccinia striiformis f. sp. tritici (Pst), the causal pathogen of wheat stripe rust, are a major challenge to resistance breeding. The majority of known resistance genes are already ineffective against current races of Pst, necessitating the identification and introgression of new sources of resistance. Germplasm core collections that reflect the range of genetic and phenotypic diversity of crop species are ideal platforms for examining the genetic architecture of complex traits such as resistance to stripe rust. We report the results of genetic characterization and genome-wide association analysis (GWAS) for resistance to stripe rust in a core subset of 1175 accessions in the National Small Grains Collection (NSGC) winter wheat germplasm collection, based on genotyping with the wheat 9K single nucleotide polymorphism (SNP) iSelect assay and phenotyping of seedling and adult plants under natural disease epidemics in four environments. High correlations among the field data translated into high heritability values within and across locations. Population structure was evident when accessions were grouped by stripe rust reaction. GWAS identified 127 resistance loci that were effective across at least two environments, including 20 with significant genome-wide adjusted P-values. Based on relative map positions of previously reported genes and QTL, five of the QTL with significant genome-wide adjusted P-values in this study represent potentially new loci. This study provides an overview of the diversity of Pst resistance in the NSGC winter wheat germplasm core collection, which can be exploited for diversification of stripe rust resistance in breeding programs.

KEYWORDS hexaploid wheat disease resistance yellow rust association mapping QTL-tag SNP genetics of immunity

GENETICS OF IMMUNITY

Wheat (Triticum aestivum L.) is the most widely cultivated cereal in the world with more than 215 million hectares planted annually (www.wheatinitiative.org). However, stripe rust (or yellow rust), caused by the basidiomycete fungus Puccinia striiformis Westend. f. sp. tritici Erikss (hereafter referred to as Pst), continues to pose a serious threat to global wheat production (Chen 2005, 2007; Hovmøller et al. 2008; Markell and Milus 2008; Milus et al. 2009; Wellings 2011). Controlling Pst through the use of foliar fungicides is less appealing in most cropping systems as it is costly, time sensitive, and complete control is impractical under severe epidemics. Therefore, improving the resistance of wheat varieties to cope with changing races of Pst is the preferred strategy for achieving global wheat demand. Resistance to Pst can be categorized as either race-specific (also referred to as all-stage resistance), which is controlled by genes with major effects, or race-nonspecific which is often expressed at the adult stages of growth (Lagudah 2011). Although cultivars with effective race-specific resistance genes are more attractive to farmers, growing cultivars with a single resistance gene often comes with a greater risk of emergence of a virulent race within a short period of time due to high selection pressure on the pathogen (Stubbs 1985; Kolmer et al. 2009; Jin et al. 2010; Wellings 2011; Hubbard et al. 2015). In 2000, the states of Arkansas and California suffered significant yield losses worth millions of dollars as new aggressive strains of Pst
emerged with virulence to the race-specific resistance gene Yr9 (Chen et al. 2002). These new strains of Pst are capable of attacking wheat growing in environments with warm temperatures that were previously considered unfavorable for development of the disease (Hovmoller et al. 2008; Måus et al. 2009). These challenges have triggered greater emphasis on the deployment of cultivars with combinations of race-nonspecific and race-specific resistance genes as a long term management strategy for Pst (Singh et al. 2000, 2005, 2011; Chen 2013; Chen et al. 2013; Hulbert and Pumphrey 2014). The slow-rusting resistance genes, Yr18, Yr29, and Yr66, and the high-temperature adult-plant (HTAP) resistance genes such as Yr36, Yr39, and Yr52 are examples of race-nonspecific resistance genes that have been shown to provide durable resistance for over 50 years (Johnson 1984; Qayoum and Line 1985; Chen et al. 2013; Singh et al. 2014). Slow-rusting resistance is characterized by the combined effect of an increased latent period and reduced uredinial size, infection frequency, and spore production (Parlevliet 1975; Ohm and Shaner 1976; Wilcoxen 1981), and HTAP resistance is characterized by increased effectiveness with increase in temperatures and growth stage (Line and Chen 1995; Chen 2013).

There are 67 officially named stripe rust resistance genes (Yr1–Yr67), and 42 with temporary designations according to the 2013 Catalogue of Gene Symbols for Wheat (McIntosh et al. 2013) and the 2013–2014 Supplement (http://wheat.pw.usda.gov/GG2/Triticum/wgc/2013/2013-Supplement.pdf). However, most of these cataloged race-specific genes are already ineffective against the group of post-2000 races of Pst (Chen et al. 2002). Frequent shifts in Pst populations dictate the strategy for deployment of the currently available resistance genes, and search for new sources of durable resistance. To date, over 200 loci are associated with resistance to stripe rust in wheat (Maccari et al. 2015). Most of the previous work involved in identifying these loci relied on classical linkage-mapping methods that are costly, characterized by poor resolution in QTL detection, and limit the number of alleles that can be studied simultaneously at any given locus (Flint-Garcia et al. 2003; Parisseaux and Bernardo 2004). Recent advances in genomic tools, including genome sequencing, high-density single nucleotide polymorphism (SNP) and genotyping by sequencing (GBS) markers, and statistical methods, have enabled the development of new approaches for mapping complex traits. Genome-wide association studies (GWAS) have emerged as the alternative approach, which maximize recent advances by exploiting cumulative recombination events that occur in the population and taking into account numerous alleles present in the population to identify significant marker-trait associations.

In wheat, GWAS has been successfully applied in mapping studies of several traits, including resistance to diseases (Gurung et al. 2011, 2014; Adhikari et al. 2011; Kollers et al. 2013; Baigain et al. 2015; Maccari et al. 2015; Gao et al. 2016). Although GWAS using polyploid wheat is often characterized by poor mapping resolution, because wheat is a self-pollinating crop with a relatively short evolutionary history (Dubcovsky and Dvorak 2007), the high levels of linkage disequilibrium (LD) in wheat significantly reduces the number of markers required for finding marker-trait associations (MTAs) (Chao et al. 2010). While overcoming the constraints inherent to linkage mapping, GWAS introduces several other drawbacks. Population stratification, if not accounted for in GWAS, often leads to spurious associations (Flint-Garcia et al. 2003; Yu et al. 2006; Kang et al. 2008; Stich et al. 2008). Another major drawback is the limited power of GWAS to detect rare variants with individual large effects, or multi-allelic variants with minor effects (Brachi et al. 2011; Zhang et al. 2012). Thus, in order to discover robust genotype-trait associations, it is necessary to use a large population size that represents the genetic variation of a crop species, dense sets of molecular markers tagging most of the nonrepetitive genome, and mixed linear models to account for spurious associations as a result of population structure.

Germplasm core collections that reflect the genetic and phenotypic diversity of crop species are ideal platforms for association studies. The United States Department of Agriculture Agricultural Research Service National Small Grains Collection (USDA-ARS NSGC) Research Unit in Aberdeen, Idaho (http://www.ars.usda.gov/Main/docs.htm?docid=21891) maintains a collection of germplasm representing the global diversity of cereals, including wheat and its various wild relatives. The wheat germplasm core collection, constructed based on phenotypic information and passport data such as geographical origin and breeding pedigrees, is comprised of over 5000 spring and winter wheat accessions, which include cultivars, cultivated materials, breeding lines, genetic stocks, and landraces from 116 countries. Evaluation and use of the wheat accessions from the NSGC germplasm core collection for identifying genotype-trait associations through GWAS is already showing promising results. For example, the recent use of a core subset of the spring wheat germplasm collection in a GWAS revealed 97 resistance loci that are effective against the current populations of Pst in the western United States, and provided a detailed insight into the distribution of resistance genes and frequencies of the resistance alleles in spring wheat subpopulations around the world (Maccari et al. 2015).

The NSGC wheat germplasm core collections are currently being evaluated extensively for resistance to a variety of diseases, abiotic stresses, and agronomic and morphological traits. Here, we report a GWAS for resistance to Pst in a core subset of the NSGC winter wheat germplasm collection, which was conducted in order to identify Pst resistance loci that may be useful and accessible to wheat breeding programs throughout the world.

**MATERIALS AND METHODS**

**Association mapping panel**

A core subset of the NSGC hexaploid winter wheat (*T. aestivum* ssp. *aestivum*) germplasm collection comprised of 1414 diverse accessions was evaluated for reactions to *Pst*. Out of the 1414 accessions, only 1221 were genotyped using the Infinium wheat SNP 9K iSelect assay. After preprocessing for missing data cutoff ≤ 10%, 1175 accessions, comprised of 174 breeding lines, 7 genetic stocks, 362 cultivars, 119 cultivated lines, and 513 landraces, were retained for the GWAS. The 1175 accessions originated from 67 countries in six continents, with Asia, Europe, North America, and South America contributing 34, 54, 6, and 3% of the accessions, respectively. Only four accessions are from both Africa and Oceania. The names of the accessions, their improvement status, and countries of origin are listed in Supplemental Material, File S1. The country-specific distribution of the accessions based on their improvement status is illustrated in Figure S1.

**Field-based evaluation of reaction to stripe rust**

The accessions were evaluated in two-way crossed nonreplicated field trials as single rows (0.75 m long and 0.25 m apart) under natural disease epidemics during the 2011–2012 and 2012–2013 crop seasons at Pullman (46.744653° N, 117.11554° W) in eastern Washington State, and during the 2011–2012 and 2013–2014 crop seasons at Mount Vernon (48.439138° N, 122.387459 W) in western Washington State. Both locations are characterized by recurrence of stripe rust epidemics as a result of oversummering and overwintering of *Pst* (Sharma-Poudyal et al. 2014) that supply sufficient inoculum. The two sites are 500 km apart and differ in weather patterns and populations of *Pst*. The predominant local *Pst* races commonly found in the two locations during the 2011–2014 crop seasons and their virulence status are described in
Table S1. The susceptible winter wheat ‘PS 279’ was planted in spreader rows bordering the nurseries to ensure sufficient inoculum for uniform disease infection. PS 279 was also used as the susceptible check every 20 rows. Reactions to stripe rust were recorded at flowering through grain filling stages of growth. Infection type (IT) was based on a 0–9 scale (Line and Qayoum 1992), with 0 = no visible sporulation; 2 = necrotic and/or chlorotic stripes with no sporulation; 3 = necrotic and/or chlorotic stripes with a trace of sporulation; 4, 5, and 6 = necrotic and/or chlorotic stripes with light, intermediate, and moderate sporulation, respectively; and 7, 8, and 9 = abundant sporulation with necrotic and/or chlorotic stripes, chlorosis behind the sporulation area, and no chlorosis or necrosis, respectively. On the basis of the 0–9 IT scale, accessions were grouped as highly resistant to moderately resistant (IT 0–3), intermediate (IT 4–6), and moderately susceptible to highly susceptible (IT 7–9). Disease severity (SEV) was estimated visually as percentage of infected leaf area when rust severities on flag leaves of the susceptible check reached 80–100%. Early onset of the disease epidemics in the 2011–2012 crop seasons in Mount Vernon allowed us to record IT at the tillering (seedling) stage of growth.

SNP genotyping

The method of DNA extraction was described in Maccaferri et al. (2015). Briefly, tissue samples were collected in 96-well coster boxes from 10-day-old seedlings, the same source of seeds used in the present study. Genomic DNA was extracted from tissues using the cetyltrimethyl ammonium bromide (CTAB) protocol (Stewart and Via 1993) at the USDA-ARS NSGC Research Unit. Chloroform:isoamyl alcohol (24:1) extraction was performed once. DNA in the aqueous phase was precipitated by adding ice-cold isopropanol, followed by washing of the DNA pellet with ice-cold 70% ethanol, and resuspension in 200 µl of TE buffer (pH 8.0).

The accessions were genotyped for 9K gene-based SNPs using the custom Infinium iSelect bead chip assay developed by the International Wheat Consortium (Cavanagh et al. 2013) on an iScan instrument following the manufacturer’s instructions, and the raw image data files were imported into the computer program GenomeStudio ver. 2011.11 (Illumina Inc., San Diego, CA) for SNP genotype calling. Before exporting genotype data, all calls were manually inspected to ensure call accuracy. The SNPs were ordered based on the published scaled map positions of the hexaploid wheat 9K consensus map (Cavanagh et al. 2013), with the chromosome arms of 4A, 5A, and 5B reoriented as described in Maccaferri et al. (2015).

Quality control

For quality control, datasets were filtered on the basis of 10% threshold for missing data. To account for inflated false MTAs, SNPs with minor allele frequency (MAF) < 0.05 were excluded. These data preprocessing steps yielded 5347 polymorphic SNPs, with 94% of the retained SNPs positioned on the hexaploid wheat consensus map.

For internal control, the SNP-derived KASP (Kompetitive Allele Specific PCR) assay wMAS000003 (http://www.cerealsdb.uk.net/cerealgeneomics/CerealsDB/indexNEW.php) for the slow-rusting resistance gene Yr18/Lr34 was incorporated into the GWAS. Marker data for the slow-rusting resistance gene Yr46/Lr67 (Forrest et al. 2014) was not used as an internal control because the frequency of the resistance allele was below the 5% MAF threshold. Marker screening for alleles associated with resistance gene Yr15 and the HTAP Yr36 resistance gene indicated that they were not present in any of the 1175 accessions based on the best available DNA markers developed by Distelfeld et al. (2006) and Ramirez-Gonzalez et al. (2015). Data for the control markers are summarized in File S2.

Linkage disequilibrium

Information on patterns of linkage disequilibrium (LD) is required for the design of GWAS, determining the interval defining a QTL, and tracking of alleles through marker-assisted selection (MAS) (Ardlie et al. 2002; Kim et al. 2007; Matther et al. 2007; Khatkar et al. 2008; Huang et al. 2010). We estimated the correlation coefficient ($r^2$) values for all pairwise comparison between intrachromosomal SNPs positioned on the 9K consensus map of hexaploid wheat (Cavanagh et al. 2013), and fitted a first-order natural smoothing spline regression model to visualize the LD as a function of genetic distance between markers on the same chromosome using the software JMP Genomics ver. 6.0 (SAS Institute Inc., Cary, NC). A critical value of LD $r^2$, as evidence of linkage, was derived from distribution of the unlinked $r^2$ following the approach of Breseghello and Sorrells (2006). The parametric 95th percentile of the unlinked $r^2$ distribution was taken as a population-specific critical value of $r^2$, beyond which the LD was likely to be caused by genetic linkage. The intersection of the spline curve fit to the $r^2$ of the syntenic SNP loci with this baseline was considered as the estimate of the extent of LD in the genome.

Population structure and genetic diversity

All 5347 SNPs were used for Fast Ward identity-by-descent clustering analysis algorithm implemented in JMP Genomics ver. 6.0. A subset of 1202 tag SNPs were used to assess population genetic structure using the model-based Bayesian clustering algorithm implemented in the computer program STRUCTURE ver. 2.3.3 (Pritchard et al. 2000) under the admixture model, assuming correlation allele frequencies and no informative priors, with five independent runs for each hypothetical number of subpopulations ($K$) ranging from 1–10. For each run, burn-in length of 50,000 in cycles followed by 100,000 Monte Carlo Markov Chain (MCMC) iterations was used. All other parameters were set at default. Based on the $\Delta K$ statistic approach (Evanno et al. 2005) implemented in STRUCTURE HARVESTER (Earl and vonHoldt 2012), the minimum and optimum numbers of $K$ were two and three, respectively. Population structure ($Q$) matrix values from different replicates of STRUCTURE runs were analyzed by CLUMPP ver. 1.1.2 (Jakobsson and Rosenberg 2007). Individual accessions were assigned to specific population groups based on highest membership coefficient values, and population assignment was plotted using DISTRUCT ver. 1.1 (Rosenberg 2004).

To further explore the population structure, we calculated fixation index ($F_{ST}$), a genetic distance measure for interpopulation differentiation compared to intrapopulation variation. All 5347 SNPs were used in the calculation with settings of $P$-value threshold $= 0.05$ and 10,000 permutations for testing the significance of the pairwise $F_{ST}$ values. We also assessed genetic diversity of the accessions in each subpopulation using PowerMarker ver. 3.25 (Liu and Muse 2005).

Statistical analyses

All statistical analyses were performed with SAS ver. 9.3 (SAS Institute Inc., Cary, NC). To keep with standard statistical nomenclature, years from field studies were represented by environments. The residual plots revealed a non-normal distribution of the data, and thus the data were transformed based on Box-Cox power transformation, to achieve near-normality. Analyses of variance of the transformed data were performed using the SAS Mixed Procedure (Proc Mixed). Estimates of variance components a non-normal distribution of the data, and thus the data were transformed based on Box-Cox power transformation, to achieve near-normality. Analyses of variance of the transformed data were performed using the SAS Mixed Procedure (Proc Mixed). Estimates of variance components were calculated by the restricted maximum likelihood method (Corbeil and Searle 1976), assuming full random model. Broad-sense heritability ($H^2$) was calculated by ratio of estimate of genetic variance to population mean variance. A model describing the data was:
GWAS was intended to identify loci that were either completely or partially masked by large-effect, or seedling resistance loci. Con-1104 accessions with IT-BLUE-ALL ranging from 3 to 9. Two of the environments, with 6 and 7 accessions, were reported.

Variances of the population mean is calculated as:

\[ \sigma^2_{ij} = \sigma^2_G + \frac{\sigma^2_E}{i} + \frac{\sigma^2_{GE}}{i} + \frac{\sigma^2}{i} \]

where \( i \) = number of environments, \( j \) = population size, \( l_i \) = environment variance \((\sigma^2_E)\), \( f_j \) = genotypic variance \((\sigma^2_G)\), \( l_f_j \) = genotype \( \times \) environment variance \((\sigma^2_{GE})\), and \( e_i \) = residual variance \((\sigma^2_e)\). Best linear unbiased estimates (BLUEs) were calculated across environments assuming fixed effects for genotypes. Transformed data from each environment, and BLUEs of data for each location (Mount Vernon = BLUE-MTV and Pullman = BLUE-PLM) and across environments (BLUE-ALL), were used for GWAS.

Genome-wide association analyses
Both the model-based Bayesian and distance-based hierarchical clustering algorithms revealed a strong population structure in the panel, and thus genome-wide association mapping tests for resistance to Pst were performed using a mixed linear model with both kinship \((K)\) and population structure \((Q)\) matrices as covariates. To eliminate linear dependence between columns of the Q matrix, the last column was removed prior to GWAS. All analyses were conducted using the R package GAPIT (Lipka et al. 2012). Loci that are significant in at least two of the environments, with \( P \leq 0.01 \) in at least one of the environments, were reported.

Our GWAS approach consisted of two steps: the first GWAS used all the 1175 accessions, and the second GWAS was performed on only 1104 accessions with IT-BLUE-ALL ranging from 3–9. The second GWAS was intended to identify loci that were either completely or partially masked by large-effect, or seedling resistance loci. Confidence intervals for the QTL identified in this study were established based on a \( \pm 2.3 \) cM genetic distance corresponding to the standard critical LD \( r^2 \) threshold of 0.3. Since the Benjamini-Hochberg FDR-controlling procedure provides less stringent control of Type I errors compared to familywise error rate (FWER) controlling procedures such as Bonferroni correction, we retested the individual effects of the loci with genome-wide significant associations using an analysis of variance (ANOVA) model with \( Q \) as a covariate. A previously developed integrated map (Maccaferri et al. 2015) was used to determine the relationships of the loci identified in this study with previously reported \( Yr \) genes and QTL. Names assigned to the QTL start with the prefix “\( Q \)" for QTL, followed by “\( Yr \)” for yellow rust, “\( wsu \)” for Washington State University, chromosome name, and number in cases where more than one QTL were reported per chromosome.

Characterization of loci with significant MTAs
To saturate QTL confidence intervals of significant MTAs in this study, we projected the 90K consensus map of hexaploid wheat onto the 9K consensus map using BioMercator ver. 4.2 (Sosnowski et al. 2012). Full nucleotide sequences of contigs, from which the anchor and projected SNPs within the QTL confidence intervals were derived, were retrieved from the Unité de Recherche Génomique Info (URGI) (http://wheat-urgi.versailles.inra.fr/) database. The contig sequences were used for a BLASTx search against nonredundant protein database for Brachypodium (Brachypodium distachyon) and rice (Oryza sativa L.) in Phytozone ver. 10.3 (http://phytozone.jgi.doe.gov/pz/portal.html#Search?show=LAST). Functional annotations of Brachypodium and rice proteins corresponding to the best hits were used as reference to infer the putative biological functions of the loci identified in the present study (data summarized in File S4).

Data availability
Both the genotypic and phenotypic information for the winter wheat diversity panel presented in this study are available at the T3 Wheat database (https://triticeaetoolbox.org/wheat/). The seeds are available to scientists worldwide through the National Small Grains Collection.
which is a component of the National Plant Germplasm System of the United States Department of Agriculture – Agricultural Research Service (http://www.ars.usda.gov/Main/site_main.htm?modecode=20-50-05-00).

RESULTS

Population structure and genetic diversity

The population genetic structure obtained from the Bayesian model-based and the Fast Ward distance-based hierarchical clustering methods revealed similar genetic variation in the accessions (Figure 2). The population of 1175 accessions was structured into two main subpopulation groups, henceforth referred to as groups 1 and 2, respectively. Group 1 further subdivides into 1A and 1B. The geographical distribution of the Bayesian model-based subpopulation groups is shown in Figure 1. European countries contributed 69% and 82% of the accessions in 1A and 1B, respectively. Cultivars and breeding lines from the US accounted for 88% of the group 1A accessions collected from the Americas, while breeding lines and landraces from Chile comprised 61% of the group 1B accessions originating from the Americas. Landraces comprised 89% of accessions in group 2, with Iran contributing 83% of the landraces.

Considerable variation in the level of admixture was evident across a large number of the accessions. Admixture among the cultivars, cultivated materials, and breeding lines in this study is consistent with our previous findings from a study on a core subset of the USDA spring wheat germplasm collection (Maccari et al. 2015). Nearly half of the landraces from Asia and Europe are classified as founders of landraces with membership probability ≥ 0.90. The majority of founders of the Asian and European landraces are from Iran, and Eastern and Southern European countries, respectively. Since information derived from population structure studies on germplasm core collections is not necessarily reflective of the range of the genetic diversity in specific countries or regions, a great deal of caution should always be exercised when interpreting the information.

Phenotypic distribution

The IT and SEV scores for the population maxima and minima for all environments are listed in Table 1. Wide variation occurred among the accessions in all environments, with Pst reactions ranging from very resistant to very susceptible. On the basis of the 0–9 scale for infection type, 35.5% of the accessions were very resistant to moderately resistant, 40.3% displayed intermediate resistance, and 24.2% were moderately susceptible to very susceptible. The susceptible check PS 279 displayed very high ITs (8–9) and SEVs (80–100%) in all environments. The population showed a non-normal distribution in the field for IT-BLUE-ALL and SEV-BLUE-ALL, with frequency distributions skewed toward resistant accessions. Based on tests of power transformations, the IT and SEV data were transformed by the square root and arcsine of data to achieve near-normality.

The zero estimates of genotype × environment interactions and the high Pearson correlation coefficient (r) values for IT and SEV suggest stability of genotypic performance across environments. The high correlation between IT and SEV in the two locations (Table 2 and Table S2) is also a strong indication of the reliability and consistency of the data collected from the nonreplicated field trials. Taken together, these observations translated into high heritability estimates for both IT and SEV in each of the locations and across all environments.

Table 1 Estimates of variance components of IT and SEV of stripe rust for the global winter wheat germplasm collection

| Environment | MTV | PLM | Across |
|-------------|-----|-----|--------|
| Minimum     | 0   | 0   | 2      |
| Maximum     | 9   | 100 | 9      |
| \( \sigma^2_{\text{IT}} \) | 0.002400*** | 0.001022*** | 0.001873*** |
| \( \sigma^2_{\text{SEV}} \) | 0.000341ns | 0.000096ns | 0.000064ns |
| \( \sigma^2_{\text{GE}} \) | 0.000000 | 0.000000 | 0.000000 |
| \( \sigma^2_{\text{G}} \) | 0.000919*** | 0.002285*** | 0.000594*** |
| \( H^2 \) | 0.79 | 0.90 | 0.85 |

Variance components were estimated from the random model using the restricted maximum likelihood method. \( \sigma^2_{\text{IT}} \), \( \sigma^2_{\text{SEV}} \), \( \sigma^2_{\text{GE}} \), and \( \sigma^2_{\text{G}} \) represent estimates of genetic, environment, genetic × environments and residual variance components. *** \( P < 0.0001 \). MTV, Mount Vernon; PLM, Pullman; IT, infection type; SEV, disease severity; \( H^2 \), broad-sense heritability estimate; ns, not significant.

Influence of population structure on reaction to Pst

Population structure clearly influenced reactions to Pst (Figure 2D and Figure 5, A and B). Based on the heat map for IT and SEV reactions of the individual accessions (Figure 2D), we found that highly susceptible accessions had very high cluster membership in subpopulation group 2. Similarly, most of the accessions clustering in subpopulation group 2 were characterized with high IT and SEV values (Figure 5, A and B). However, for subpopulation groups 1A and 1B the opposite was true, with 1B accessions displaying a relatively high level of resistance compared to 1A accessions. Population structure (Q) accounted for significant proportions of the observed variance for IT-BLUE-ALL and SEV-BLUE-ALL across environments (Table S7).

Similar global trends in reaction to Pst were also observed among accessions based on improvement status and geographic origin. As expected, cultivars had relatively higher numbers of favorable alleles, and displayed increased resistance to Pst (Figure S2, A and B). We also observed a high number of favorable alleles and high levels of resistance in the accessions from Europe and South America (Figure S2, D and E). These observations are a clear demonstration of the success of modern wheat breeding practices that include resistance gene pyramiding. The relatively lower number of favorable alleles and intermediate resistance displayed by cultivars and breeding lines from the US are likely a reflection that the majority of wheat acres in the US were not historically subjected to stripe rust epidemics (Sharma-Poudyal et al. 2014). Not surprisingly, landraces had the lowest number of favorable alleles, and generally high levels of susceptibility.

Linkage disequilibrium

The analysis of LD with mapped SNP markers was based on pairwise squared-allele frequency correlation (r²) for all intrachromosomal SNP
loci. SNPs mapped to the 9K wheat consensus map (Cavanagh et al. 2013) were grouped into 28 linkage groups, covering a total map length of 1871.19 cM, with average marker density of three markers per 1.0 cM. However, marker coverage for the D genome chromosomes was poor, with an average marker density of 2.6 cM per marker.

Across the 1175 accessions, the genome-wide LD generally declined with genetic distance (Figure 3, A and B). The median $r^2$ value for pairs of completely linked markers based on the 9K wheat consensus map was 0.77, with interquartile values ranging from 0.22–0.99. At the average distance ranging from 0.1–1.0 cM, LD decayed to $r^2$ 0.24. The median $r^2$ decreased to 0.08 for pairs of markers within 1.1–5.0 cM genetic distance. Based on the 95th percentile of the distribution of unlinked pairwise $r^2$ estimates across the genome, values of $r^2 > 0.16$ (corresponding to genetic distance < 5.1 cM) were probably due to genetic linkage. QTL confidence intervals were established based on a ± intermarker genetic distance of 2.3 cM corresponding to the standard critical LD $r^2$ of 0.3.

### Genome-wide association analyses

We compared the performance of two GWAS approaches: a general linear model (GLM) with population structure adjustment (Q matrix) vs. a mixed linear model (MLM) incorporating $K$ and $Q$ matrices. Distribution of observed $-\log_{10} P$-values from the GLM with Q approach separated early from the expected distribution under a model of no association, with significant inflation of nominal $P$-values leading to a high level of false positives. On the other hand, the MLM using $K$ and $Q$ as covariates eliminated the moderately significant $P$-values that showed departure from the expected distribution under GLM with Q, and thus was selected for performing GWAS in this study. In a diverse population such as the one in this study, phenological traits are also expected to influence reactions of the plants to Pst. To address this, we performed GWAS on heading date and plant height data recorded in 2012 and 2013 crop seasons at the Pullman location. All SNP markers in LD with Pst resistance QTL in this study that were also consistently associated with heading date and/or plant height in the two environments ($P < 0.05$) were excluded from the final list of QTL-tag and -associated SNPs.

Using the 1175 accessions, we identified 100 QTL based on significant MTAs to Pst in at least two of the four environments (Table S3). To identify QTL that were either completely or partially masked by major effect QTL, we performed a second GWAS using only 1104 accessions with IT-BLUE-ALL ranging from 3–9, which also detected 100 QTL with significant MTAs to Pst (Table S4). Excluding the common QTL between the two GWAS, the total number of QTL identified in this study is 127, with 24 showing significant associations at a genome-wide adjusted significance level of $P < 0.1$ based on the False Discovery Rate (FDR) correction method (Benjamini and Hochberg 1993). However, 20 of the QTL were selected for further characterization after the effects of the significant loci (adjusted $P < 0.1$) were retested using an ANOVA model that takes population structure into account. The 20 loci (QTL-tag SNPs) are located on chromosomes 1A (IWA5505 and IWA5215), 1B (IWA6963, IWA915), and IWA62, 2A (IWA2526 and IWA5824), 3A (IWA3401), 4A (IWA3981, IWA3774, IWA6697, IWA4651, and IWA3422), 4D (IWA3581), 5A (IWA5002), 5B (IWA5166), 6A (IWA8595), and 6B (IWA4169, IWA7527, and IWA349). Effects of the 20 loci and their pairwise interactions are summarized in Table 3 and Table S7.

The genetic variance explained by the individual QTL ranged from 1.02–6.60% for IT, and 1.09–5.84% for SEV (Table S3). The cumulative genetic variances explained by the 20 QTL were 28 and 26% for IT and SEV, respectively (Table S7). When significant pairwise interactions were factored into the model, the variations increased to 37% and 33% for IT and SEV, respectively. However, the significant effects of IWA62, IWA2526, IWA5824, IWA3401, IWA6697, IWA4651, and IWA3422, and IWA5166 were reversed when the pairwise interactions were included in the ANOVA model. In a similar study, Maccarferri et al. (2015) found that the reduction or elimination of significant effects of loci involved in pairwise interaction could be attributed to either negative effects of other interacting loci or the presence of the susceptibility alleles.

Marker loci IWA5505, IWA62, IWA3401, IWA3774, and IWA5002 also showed genome-wide significant associations with IT at the tillering (seedling) stage of growth at the Mount Vernon field site in 2012. Analysis of the field data collected at the seedling stage also revealed additional loci on 5B (IWA7815), 6D (IWA93), and 7A (IWA7592) that were not detected at the adult stages of growth. This can likely be explained by an increase in population of Pst race(s) with virulence on these genes.

Incorporation of the marker data for Yr18/1r34 into the GWAS as an internal control allowed us to detect significant association of Yr18 with IT and SEV only in Mount Vernon ($P < 0.05$). In our previous study on the spring wheat core collection, Yr18 was also associated with reactions to Pst in Mount Vernon but not in Pullman. We could not find linkage between KaspLr34 and any of the SNP markers on chromosome 7D (File S2) due to inadequate marker coverage of the D genome. Accessions carrying the Yr18 marker displayed a frequency distribution skewed toward susceptibility with 5.3, 43.4, and 51.3% of the accessions classified as resistant-moderately resistant, intermediate, and moderately susceptible-very susceptible (File S2). This is not surprising, since Yr18 is not expected to confer adequate resistance by itself and the effect of this gene has noted environmental sensitivity (Singh and Gupta 1992; Singh et al. 2000).

### Influence of number of favorable alleles on reaction to Pst

The total number of individual QTL (represented by QTL-tag SNPs) with significant associations with reactions to Pst in this present study is

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**Table 2** Pearson correlation coefficients among IT and SEV of stripe rust data for four environments

| Trait | MTV-IT-12 | MTV-IT-14 | PLM-IT-12 | PLM-IT-13 | MTV-SEV-12 | MTV-SEV-14 | PLM-SEV-12 | PLM-SEV-13 |
|-------|----------|----------|----------|----------|------------|------------|------------|------------|
| MTV-IT-12 | 1.00     | 0.75     | 0.81     | 0.69     | 0.86       | 0.74       | 0.74       | 0.67       |
| MTV-IT-14 | 1.00     | 0.77     | 0.76     | 0.82     | 0.94       | 0.79       | 0.79       | 0.75       |
| PLM-IT-12 | 1.00     | 0.79     | 0.82     | 0.75     | 0.88       | 0.76       | 0.76       | 0.76       |
| PLM-IT-13 | 1.00     | 0.76     | 0.74     | 0.82     | 0.91       |            |            |            |
| MTV-SEV-12 | 1.00     | 0.84     | 0.85     | 0.77     |            |            |            |            |
| MTV-SEV-14 | 1.00     | 0.79     | 0.76     | 0.82     | 0.91       |            |            |            |
| PLM-SEV-12 | 1.00     | 0.85     | 0.85     | 0.77     |            |            |            |            |
| PLM-SEV-13 | 1.00     | 0.85     | 0.85     | 0.77     |            |            |            |            |

All correlation coefficients are highly significant ($P < 0.0001$). MTV, Mount Vernon; IT, infection type; PLM, Pullman; SEV, disease severity.

$^a$ Trait: two locations (MTV, Mount Vernon, WA; PLM, Pullman, WA) and 3 yr (2012, 2013, and 2014).
higher frequencies of favorable alleles for Pst resistance were present at low frequencies. Resistance alleles were almost fixed for IWA5963, IWA3981, IWA6697, and IWA5381 in all the subpopulation groups. Similar distributions were also evident when accessions were grouped by improvement status and geographical regions of origin.

**Comparison of Pst resistance QTL and Yr genes**

Positions of QTL identified in this study relative to that of previously known Yr genes and QTL are visualized in Figure 6. Only 19 of the QTL in this study are positioned on the standardized chromosomes because IWA62 is not mapped. However, IWA62 loci are present in scaffolds from chromosomes 1A, 1B, and 1D based on TGACv1.3 genome assembly of T. aestivum cultivar ‘Chinese Spring’ generated by The Genome Analysis Center (TGAC), Norwich, UK. Loci identified in this study are highlighted in green, loci from our previous study on a core subset of the USDA spring wheat germplasm collection (Maccaveri et al. 2015) are highlighted in yellow, and common or overlapping loci between the two germplasm collections are highlighted in blue. Detailed description of Figure 6 is summarized in File S3.

Positions of SNPs tagging IWA3505 (QYr.wsu-1A.1), IWA824 (QYr.wsu-2A.2), IWA3981 (QYr.wsu-4A.1), IWA3774 (QYr.wsu-4A.2), and IWA349 (QYr.wsu-6B.3) do not correspond to that of any previously identified genes or QTL for resistance to Pst, and thus represent potentially novel loci. We previously reported IWA7257 as a potential new resistance locus (Maccaveri et al. 2015). However, recent mapping of IWA7257 (unpublished data) repositioned the locus in a region overlapping with previously reported APR QTL QYr.wgp-6B.1 in the cultivar ‘Stephens’ (Santra et al. 2008), QYr.sun-6B in the Australian cultivar ‘Jana’ (Bariana et al. 2010), and QYr.caas-6BS in the Chinese landrace ‘Pingyuan 50’ (Lan et al. 2010). The association of IWA7257 with resistance to Pst was also reported in a GWAS study on winter wheat from...
Figure 3 Genome-wide average linkage disequilibrium (LD) decay over genetic distance. (A) Plot of pair-wise SNP LD $r^2$ values as a function of intermarker map cm distance based on a reference consensus map (Cavanagh et al. 2013). The curve represents the model fit to LD decay. The red dashed line represents the standard critical $r^2$ = value (0.3) used for establishing QTL confidence intervals in this study; the blue solid line represents the population-specific critical $r^2$ value (0.16) beyond which LD is likely due to linkage; and the black dashed line represents the mean. (B) Boxplot showing LD $r^2$ values for categories of SNP pairwise map distances (A–I): 0, 0.1–1, 1.1–5, 5.1–10, 10.1–20, 20.1–30, 30.1–40, 40.1–50, and > 50 cm. SNP, single nucleotide polymorphism.

the Pacific Northwest region of the United States (Naruoka et al. 2015). Intervals where the remaining 13 QTL reside overlap with positions of either previously reported Yr genes or QTL for Pst resistance.

**Group 1:** IWA3215 (QYr.wsu-1A.2) maps within the confidence interval of QYr.tam-1AL identified in the hard red winter wheat ‘TAM 112’ (Bassett et al. 2014). However, the relationship between IWA3215 and the QTL reported in TAM 112 could not be established because a locus with marker-wise significant associations in this study, IWA4277, also overlaps with QYr.tam-1AL. IWA5963 (QYr.wsu-1B.1) maps to the gene-rich region of 1BS that includes YrAlp, Yr15, and YrH52 (Lin and Chen 2007, 2008), and the position of IWA5915 (QYr.wsu-1B.2) overlaps with the race-specific gene YrExp1 (Lin and Chen 2008). Based on the available KASP markers R5 and R8 for Yr15 (Ramirez-Gonzalez et al. 2015), the gene is not present in the 1175 winter wheat accessions. Also, YrH52 is not expected to be present in the T. aestivum accessions in this study as the gene was recently identified in T. dicoccoides (Feng et al. 1999). The post-2000 races of Pst in the Pacific Northwest region have high virulence on both YrAlp and YrExp1, which are derived from the spring wheat cultivars ‘Alpowa’ and ‘Express,’ respectively. The position of IWA5915 also overlaps with IWA3017, which showed significant marker-wise association in our previous study on the spring wheat germplasm collection (Maccarelli et al. 2015).

Several stripe rust resistance QTL have been mapped to the same 1BS region as IWA5963 (Quan et al. 2013; Case et al. 2014; Naruoka et al. 2015). The distance between IWA5963 and the closest QTL QYr.wgp-1B.1 (Naruoka et al. 2015) is 4.71 cm, suggesting that they may represent the same QTL. IWA5915 maps 28.33 cm proximal to the slow-rusting resistance gene Yr29/Lr46, and very close to the HTAP QTL QYrexp.wgp-1BL identified in Express (Lin and Chen 2009).

**Group 2:** IWA2526 (QYr.wsu-2A.1) should be different from Yr17 (Helguera et al. 2003) because Yr17 is no longer effective against the Pst populations in the Pacific Northwest (Wan et al. 2016). The position of IWA2526 also overlaps with the confidence intervals of QTL identified in the soft winter wheats ‘Pioneer 26R61’ (Hao et al. 2011) and ‘VA00W-38’ (Christopher et al. 2013). IWA2526 lies 36.20 cm proximal to QYr.ucw-2A.2, which we recently identified in a similar study (Maccarelli et al. 2015). The relationship between IWA2526 and the QTL reported in Pioneer 26R61 and VA00W-38 could not be established because a locus with marker-wise significant associations in this study, IWA5087, also overlaps with these QTL.

**Group 3:** IWA4301 (QYr.wsu-3A) maps in the same proximal region where Hou et al. (2015) identified a APR QTL, QYrdr.wgp-3AL, in the French winter wheat ‘Druchamp.’

**Group 4:** IWA6697 (QYr.wsu-4A.3) maps to a similar position as the previously reported IWA1034 (QYr.ucw-4A) (Maccarelli et al. 2015), and thus are likely representing the same locus. Positions of the seedling resistance gene Yr60 (Herrera-Foessel et al. 2015) and a QTL in the cultivar ‘Kariega’ (Prins et al. 2011) also overlap with the interval of IWA6697. Loci IWA4651 (QYr.wsu-4A.4) and IWA3422 (QYr.wsu-4A.4) reside in the 4AL/7BS translocation region where several QTL have been identified (Prins et al. 2011; Chen et al. 2012a; Vázquez et al. 2015). Although IWA5381 (QYr.wsu-4D) on 4DS overlaps with QYr.cim-4DS in the CIMMYT spring wheat cultivar ‘Pastor’ (Rosewarne et al. 2012), we could not establish a definitive relationship between the two QTL due to overlap of a linked marker-wise significant locus IWA2122 with QYr.cim-4DS.

**Group 5:** IWA5002 (QYr.wsu-5A) overlaps with positions of the APR QTL identified on the 5AL arm of the Chinese landrace Pingyuan 50 (Lan et al. 2010), hard red spring wheat PI 610750 (Lowe et al. 2011), and the spring wheat line ‘SH3A/CBRD’ (Ren et al. 2012). IWA5166 (QYr.wsu-5B) is located within the C.I. of QYr.cim-5BL identified on 5BL of the CIMMYT spring wheat ‘Chapio’ (Yang et al. 2013). IWA5166 is 58.87 cm proximal to the genome-wide significant seedling locus in this study, IWA7815, which corresponds to the race-specific gene YrExp2 (Lin and Chen 2008).

**Group 6:** IWA5959 (QYr.wsu-6A) maps to the proximal region of 6AL, where QTL for APR have also been reported in the cultivars Stephens (Vázquez et al. 2012) and ‘Avocet’ (William et al. 2006). However, the position of IWA7894, which showed marker-wise significant associations with Pst in this study (Table S3), also overlaps with QYrpLorr-6AL.
Table 3 QTL significantly associated with reactions to *Puccinia striiformis* f. sp. *tritici* in at least three environments (genome-wide adjusted \( P < 0.1 \))

| SNP   | Chr | Posb | Allelec | Seeding | Flowering to Grain Filling |
|-------|-----|------|---------|---------|---------------------------|
|       |     |      |         | IT      |   SEV         |
|       |     |      |         | MTV     |   MTV         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | MTV     | 2013         |
|       |     |      |         | PLM     | 2013         |
|       |     |      |         | BLUE    | 2013         |
|       |     |      |         | ALL     | 2013         |
|       |     |      |         | MTV     |   SEV         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | MTV     |   SEV         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | MTV     |   SEV         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | BLUE    | 2013         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | BLUE    | 2013         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2014    | 2014         |
|       |     |      |         | 2012    | 2012         |
|       |     |      |         | 2013    | 2013         |
|       |     |      |         | ALL     | 2013         |

Marker-wise significance: * \( P < 0.05 \); ** \( P < 0.01 \); *** \( P < 0.001 \). Genome-wide significance: **** Benjamini-Hochberg FDR adjusted \( P < 0.10 \). SNP, single nucleotide polymorphisms; IWA, loci in linkage disequilibrium and with significant association with reaction to *Pst*; Chr, chromosome; Pos, position; IT, infection type; SEV, disease severity; MTV, Mount Vernon; PLM, Pullman; BLUE, best linear unbiased estimates; ALL, across environments; NA, not applicable; ns, not significant.

in Stephens (Vazquez et al. 2012), and thus the relationship between *IWA8595* and *QYrl.orr-6AL* could not be established. The same region of *IWA8595* has also been associated with resistance to powdery mildew in the cultivar ‘Saar’ (Lillemo et al. 2008). In a RIL population derived from the rust-resistant winter wheat cultivar ‘Einstein’ and susceptible winter wheat ‘Tubbs,’ Vazquez et al. (2015) identified *QYrl.orr-6AL* in a region 7.67 cM distal to *IWA8585*, suggesting the presence of more than one QTL in the proximal region of 6AL. *IWA4169* (*QY.wrsw-6B.2*) resides in the same proximal region of 6BL where a QTL for APR has been identified in the CIMMYT spring wheat ‘Pavon 76’ (William et al. 2006) and the French winter wheat cultivar ‘Renan’ (Dedryver et al. 2009). The position of *IWA4169* also overlaps with *IWA2090*, which showed marker-wise significant association with *Pst* resistance in the spring wheat germplasm collection (Maccarelli et al. 2015).

**Saturation of QTL regions, and comparative analyses of the genomes of wheat, Brachypodium, and rice**

Enrichment of the regions of the QTL with known chromosomal locations by projecting the 90K consensus map SNPs (Wang et al. 2014) onto the 9K SNP consensus map (Cavanagh et al. 2013) increased the total number of SNPs to 979, with an average of 52 SNPs per QTL C.I. (Table S9). However, nucleotide sequences of contigs from which the SNPs were derived were available for only 490 SNPs in the wheat database of the Unité de Recherche Génomique Info, Institut National de la Recherche Agronomique in France. BLASTx searches against *Brachypodium* and rice databases assigned functional annotations to 398 of the loci, and 63 of the annotated protein domains correspond to either LRR-containing proteins or NB-ARC domains. Thirty-eight percent of these resistance-related protein domains are within the C.I. of *IWA4651* and *IWA3422*, which reside in the 4AL/7BS translocation region. Details of the 9K anchor SNPs and 90K projected SNPs, and information on transcripts corresponding to top hits with proteins in the *Brachypodium* and rice databases, are given in File S4.

**DISCUSSION**

**Population genetic structure and genetic diversity**

Knowledge about genetic diversity and population structure is essential for the utilization of core collections as genetic resources for germplasm development and the diversification of the gene pool in stripe rust resistance breeding programs. In this study, the distance-based hierarchical Fast Ward clustering approach revealed two distinct major genetic groups in the population that could be explained by geography (Figure 2, A and B). The first major group mostly consisted of accessions collected from Europe and the Americas (henceforth referred to as the Western Subpopulation), and the second major group comprised primarily accessions from Asia (henceforth referred to as the Asian Subpopulation). In a study on the USDA spring wheat core collection, Maccarelli et al. (2015) reported the presence of two major clusters (Asian and
Western) based on distance-based hierarchical clustering. Wang et al. (2013) also reported similar findings based on an analysis of genetic diversity in the wheat D genome. In the present study, 98% of accessions from the Americas clustered in the Western Subpopulation. Only 3% of accessions originating from Europe were found in the Asian Subpopulation. In contrast, a higher number of accessions originating from Asia (28%) were found in the Western Subpopulation. The observed pattern of distribution of the clustering-inferred groups is indicative of the path of distribution of *T. aestivum* following its domestication in the Fertile Crescent around 10,000 years ago (Dubcovsky and Dvorak 2007). As domesticated wheat spread from the Fertile Crescent to northwestern Europe, the crop was subjected to intensive natural selection, and enrichment of favorable alleles for key traits essential for successful adaptation in the new ecological environments.

The model-based Bayesian clustering algorithm implemented in the software STRUCTURE also revealed the presence of two distinct genetic groups that were similar to the distance-based clustering, with the ad hoc measure $\Delta K$ peaking at $K = 2$. However, we considered $K = 3$ as the optimum number representing genetic subgroups in the winter wheat germplasm collection as the population structures divided into well-defined subgroups, based on geographical regions of origin and improvement status (Figure 2, C, E, and F). Both subgroups 1A and 1B, which correspond to the Western Subpopulation, were dominated by accessions from Europe. The North American accessions mostly clustered in subgroup 1A, while a majority of the South American accessions were found in 1B. Accessions from the United States represent 96% of the North American accessions in 1A, and breeding lines from Chile account for 90% of the South American accessions in 1B. Subgroup 2, which corresponds to the distance-based Asian Subpopulation, is comprised mostly of landraces from Iran.

To explore the genetic diversity and relationships among and within the STRUCURE clustering-inferred groups, various diversity statistics were assessed. $F_{ST}$ estimates revealed high levels of genetic differentiation between the Western Subpopulations and the Asian Subpopulation (Table S8). In contrast, minor genetic differences were observed between the 1A and 1B Western Subpopulations. The similar polymorphism information content (PIC) values of the Western Subpopulations (Table S5 and Table S6) illustrate the likely presence of a large number of accessions in both groups, with identical alleles in most of the genomic regions with low gene diversity as a result of continuous allele reshuffling and selections through modern breeding practices. This explanation is further supported by the observed considerable variation in the level of admixture in the Western Subpopulations (Figure 2C). The relatively low PIC value in the Asian Subpopulation is not surprising as it is comprised of a large number of genetically similar landraces (Figure 2B) as a result of shared ancestry, which is reflected in the high average membership coefficient value of 0.88 for accessions in the model-based subgroup 2 (Figure 2D). Compared to studies using multi-allelic markers (Roussel et al. 2004; Zhang et al. 2010; Chen et al. 2012b), the PIC values in this study are generally low; this is because, for biallelic markers such as SNPs, the PIC is not expected to exceed the 0.5 value for equal allele frequencies.

The variance component among the subpopulations accounted for 21.78% of the genetic variation (Table S8). Although the $F_{ST}$-based differences between groups were highly significant ($P < 0.0001$), the within-subgroup component of the variance was predominant, accounting for 78.22% of the total variation. The low within-Asian Subpopulation variance component further confirms findings from the $F_{ST}$ analysis that, among the defined subpopulations, the Asian subpopulation was the most distinct.

**Genome-wide association analyses**

In this study, we used a mixed linear model with $K$ and $Q$ as covariates to account for the strong influence of population structure on $Pst$ reaction (Figure 5, A and B). The two-step GWAS approach employed in this study identified a total of 127 loci on all chromosomes (except 7D) with significant associations with reactions to $Pst$ in at least two environments. A more stringent approach that included FDR multiple testing correction, followed by retesting of the effects of QTL with genome-wide significant associations (adjusted $P < 0.1$) using ANOVA with population structure ($Q$) as a covariate, yielded a total of 20 loci (Table 3). Among the 20 loci, nine and 11 were detected in the GWAS using all accessions and accessions with IT-BLUE-ALL $\geq$ 3, respectively.

To test whether the stringency of the analyses in this study created false-negatives or whether known resistance loci could not be detected due to poor marker coverage in some regions of the genome, we included marker data for the slow-rusting resistance gene complex *Yr18/Lr34/Pm38* in the GWAS. Detection of *Yr18* KASP marker-$Pst$ significant associations at the Mount Vernon location (File S2) is consistent with the findings of Maccaferri et al. (2015), who also reported marker-wise significant associations of *Yr18* with $Pst$ reactions at Mount Vernon and Davis locations. The *Yr18* KASP marker is not in LD with any of the chromosome 7D SNP markers, suggesting that the number of resistance loci in the germplasm in this study could increase with higher density of SNP markers on the array from the wheat D genome chromosomes with low LD.

The individual effects of the QTL on the observed variations ranged from 0.86–6.6%, consistent with the complex nature of the wheat-$Pst$-pathosystem, which involves numerous types of resistance genes with varying levels of effects on the pathogen. Given the heterogeneity of the germplasm collection and the likely effects of developmental genes on the observed variations in reactions to $Pst$, well-designed seedling tests are required to determine the identity of the QTL overlapping with positions of known seedling genes. The cumulative effect of the QTL with genome-wide significant associations in this study explained up to...
### Table 4: Frequencies of stripe rust resistance alleles

| Chr | Pos  | SNP IWA | Alleles | Panel | Subpopulation | Breeding | Genetic | Cultivar | Cultivated | Landrace | Origin | Mean reaction to stripe rust |
|-----|------|---------|---------|-------|---------------|----------|---------|----------|------------|----------|--------|-----------------------------|
|     |      |         |         |       |               |          |         |          |            |          |        | Infection type (0–9 scale) |
|     |      |         |         |       |               |          |         |          |            |          |        | Disease severity (0–100%) |

- **Chr:** chromosome; **Pos:** position; **SNP:** single nucleotide polymorphisms; **IWA:** loci in linkage disequilibrium and with significant association with reaction to *Pst*; **ACIMPT:** population structure subgroups in improvement status of accessions.
- **Alleles** underscored.
- **Panel** indicates association with reaction to *Pst*.
- **Origin** indicates geographical origin.
- **Mean reaction to stripe rust** indicates the average reaction of the accessions to stripe rust.

**Notes:**
- **Scaled position from hexaploid wheat consensus map (Cavanagh et al. 2013).**
- **SNP indexes from Illumina iSelect 9K wheat assay (Cavanagh et al. 2013).** SNPs in **bold** were identified in the analyses using the 1104 accessions with IT scores ≥ 3, and **bold and underlined** SNPs were detected in both the first and second approaches for GWAS. Loci in LD and with significant association with reaction to *Pst* (IWA): 5505/475/4934, 3215/none, 5963/none, 5915/5749, 62/none, 2526/none, 5824/5495, 3401/2263/2264/2265/2266, 3981/none, 3774/none, 4169/none, 5002/none, 516/none, 8595/none, 4651/none, 3422/none, 5381/none, 5002/none, 516/none, 8595/none, 4169/none, 7257/none, 349/none.

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in the model also resulted in Type I error, ANOVA model, suggesting that the FDR multiple testing corrections and SEV-BLUE-ALL increased from 28% to 58% and 26% to 63%, MTAs. However, the proportion of the variations for IT-BLUE-ALL their detection is more prone to Type II error, Similar observations were also reported by Maccaferri increasing with increase in the number of favorable alleles (Figure 4). The number of QTL significantly associated with Pst resistance (adjusted \( P < 0.1 \)) in this study is twice that in our previous study (Maccaferri et al. 2015). This could be explained by the use of a larger sample size, and the less stringent FDR multiple corrections testing to minimize the chance of any false MTAs. The post-2000 Pst populations in Pullman and Mount Vernon have high virulence on most of the cataloged and temporary designated Yr genes (Chen 2005, 2007; Wan and Chen 2014; Wan et al. 2016), and thus the QTL identified in this study should be useful to breeding programs that deploy combinations of both race-specific and race-nonspecific genes.

### Influence of population structure on Pst reaction

The Asian Subpopulation was generally more susceptible to Pst compared to the Western Subpopulations. This was not unexpected, as three-quarters of the Western Subpopulation accessions represent cultivars, cultivated lines, breeding lines, and genetic stocks that have been subjected to genome manipulation through modern breeding practices. However, differences in reaction to Pst have also been observed between the Western Subpopulations, with 1B exhibiting a relatively higher level of resistance. These differences could be attributed to enrichment of favorable Pst resistance alleles in subpopulation 1B at the six key loci IWA3215, IWA5915, IWA2526, IWA5824, IWA4651, and IWA5166 (Table 4). The effect of these six key loci is even more pronounced when the accessions are grouped by geographic region of origin, with very high frequencies found in accessions from Western and Northern Europe, and South America. In contrast, very low frequencies of favorable alleles at the six loci were observed in the Asian Subpopulation, which is comprised of 89% landraces. This information can allow breeders in regions of Eastern and Southern Europe, and North America, where frequencies of subpopulation 1A are high, to develop strategies to utilize subpopulation 1B accessions as genetic resources for diversification of the stripe rust resistance gene pool in their programs.

### Extent of LD and its implications for comparative genomics

We observed a slower rate of LD decay in the winter wheat population compared to that reported by Maccaferri et al. (2015). This is because the winter wheat germplasm collection in this study is less diverse when compared to the spring wheat core collection, which is comprised of seven model-based subpopulation groups (Maccaferri et al. 2015). The high level of diversity in the spring wheat germplasm collection is not unexpected, since the crop is cultivated over a wider geographical area with different climatic conditions. Because of the relatively slower rate of LD decay, the genetic distance of \( \pm 2.3 \text{ cM} \) used for establishing confidence intervals for QTL in this study is also relatively higher compared to the \( \pm 1.6 \text{ cM} \) used in the spring wheat core collection (Maccaferri et al. 2015). To enrich the regions of the QTL with additional SNP markers, we projected the 90K Illumina data (Wang et al. 2014) onto the 9K reference map (Cavanagh et al. 2013). This led to a fourfold increase in the number of SNP markers per QTL C.I., enabling collinearity analyses between the genome of wheat and the genomes of Brachypodium and rice (File S4). In addition to CloudBlast, the Brachypodium and rice protein databases in Phytozone were also used for similarity searches using sequences of wheat contigs from which the SNPs were derived. Putative functions of candidate genes in wheat were inferred based on the functional annotations of proteins of the two model species. Among the wheat SNPs with corresponding annotated Brachypodium or rice proteins, 16% were classified as R genes, with high concentration in the 4AL/7BS translocation region where IWA4651, and IWA3422 (Table S9) reside. Interestingly, comparative analyses between the 4AL/7BS translocation and the genomes of Brachypodium and rice also revealed a large number of NB-LRR and LRR-receptor-like kinases in the collinear regions of the model species. The 4AL/7BS translocation region harboring IWA4651 and IWA3422 showed collinearity with 9.0–47.0
Mb region in chromosome 4 of *Brachypodium*, and a 2.0–2.7 Mb region in chromosome 11 of rice. The presence of a high number of R genes in collinear regions of the model species suggests that these resistance loci represent a conserved cluster of genes that evolved during ancient times, predating the event of this translocation in wheat. In a study of a segregating population derived from the durum wheat cultivars ‘Cresco’ and ‘Pedroso’, Marone et al. (2013) identified a QTL for resistance to powdery mildew in the same region of IWA4651, suggesting a potential pleiotropic effect of this locus. Associations of the 4AL/7BS translocation with other adaptive traits have also been reported (Nezhad et al. 2012; Zanke et al. 2014), further confirming the selective advantage provided by this translocation during the course of evolution and the domestication of wheat crop. With the currently available dense wheat SNP and GBS markers, our understanding of the importance of this translocation will improve as more results from GWAS on other traits become publicly available.

Enrichment of the QTL confidence intervals could facilitate identification of haplotypes associated with resistance alleles, development of molecular markers, and construction of high-density genetic maps as a first step toward cloning of the defense-related candidate genes identified in this study. It is noteworthy that we could not find sequences coding for disease resistance-related proteins within the C.I. of some of the QTL in this study. This may be due, in part, to ascertainment bias where our QTL-tag SNPs are relatively far from the target loci, or the functional annotations of some of the genes are resistance factors that are not known.

**Relative position-based comparison of Pst resistance QTL and Yr genes**

Relationships of the 20 loci with genome-wide significant associations (adjusted \( P < 0.1 \)) in this study to previously mapped QTL and Yr genes are summarized in Figure 6 and File S3. Although the focus in this study is on the 20 loci with genome-wide significant associations, the remaining 107 loci can be used as references for future QTL validation work. The reported positions of the QTL in this study may not be precise in some cases due to lack of sufficient common markers among the maps used for developing the integrated map. An integrated map with more accurate positions would be possible as more genetic maps constructed from SSR, DArT, SNP, and GBS markers become available.

The confidence intervals of QTL tagged by SNPs IWA5505, IWA5824, IWA3981, IWA3774, and IWA349 do not overlap with positions of any of the previously reported QTL or Yr genes, and thus are likely novel resistance loci. IWA5824 and the QTL QYr.ucw-2A.3 (IWA424) reported by Maccaferri et al. (2015) may represent two closely linked genes near the centromeric region of 2AS, since the distance between them is more than 4.6 cM. The locus IWA3774 is
present in very low frequencies in all the population structure subgroups (Table 4), providing an opportunity for exploiting this QTL for diversification of germplasm in stripe rust resistance breeding programs. *IWA62* is not in LD with any of the SNP loci on the Group 1 chromosomes, and thus we cannot estimate its map positions on 1A, 1B, and 1D. We could not establish definitive relationships between previously mapped QTL overlapping with the positions of QTL tagged by IWA3215, IWA2526, IWA5381, and IWA8595, because their confidence intervals also overlap with positions of other QTL with marker-wise significant associations in this study. Based on the integrated map developed by Maccaferri et al. (2015), the QTL tagged by IWA5963, IWA5915, IWA3401, IWA6697, IWA3422, IWA5002, IWA5166, IWA8595, IWA7257, IWA4169, and IWA6561 are likely corresponding to alleles of previously reported QTL (File S3).

In our previous study on a global collection of spring wheat accessions, we reported association of *IWA7257* with *Pst* resistance (Maccaferri et al. 2015). In that same study, we also identified a QTL *QYr.ucw-4A* tagged by IWA1034 in a similar position as *IWA6697*, suggesting these are likely alleles of the same locus. The confidence intervals of *IWA5915* and *IWA4169* also overlap with the respective positions of *IWA3017* and *IWA2090*, which were significant at marker-wise *P*-value in the spring wheat germplasm collection. Similarly, the QTL *QYr.ucw-1B* and *QYr.ucw-2A.3* with genome-wide association in our previous study also overlap with the respective positions of *IWA1791* and *IWA690*, which showed marker-wise significant associations in this study. The observed overlaps between marker-wise and experiment-wise trait associations is a clear indication that the use of multiple correction procedures to account for false positive associations in the two studies also increased the likelihood of false negatives, supporting our decision to also report the marker-wise trait associations (Table S3 and Table S4). The testing strategy we employed in both studies is further validated by the identification of 26 similar genomic regions with marker-wise trait associations on chromosomes 1A, 2A, 2B, 3A, 3B, 4A, 4B, 5B, 6A, 6B, and 7B in both the winter and spring wheat germplasm collections.

**Conclusion**

The overview of the allele composition of the USDA winter wheat germplasm core collection anchored to SNP markers presented in this study can be exploited by breeders for targeted pyramiding of the alleles. Furthermore, the data in this study can be utilized for designing GWAS and genomic selection cross-validation experiments to predict performance of genotypes under high *Pst* pressure, while simultaneously analyzing the effect of all the 127 QTL on genomic selection. The identification of resistance genes as major contributing factors to the genetic background without other QTL.

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