Mining the Vavilov wheat diversity panel for new sources of adult plant resistance to stripe rust

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Received: 20 October 2021 / Accepted: 12 January 2022 / Published online: 3 February 2022 © The Author(s) 2022

Abstract

Key message Multi-year evaluation of the Vavilov wheat diversity panel identified new sources of adult plant resistance to stripe rust. Genome-wide association studies revealed the key genomic regions influencing resistance, including seven novel loci.

Abstract Wheat stripe rust (YR) caused by Puccinia striiformis f. sp. tritici (Pst) poses a significant threat to global food security. Resistance genes commonly found in many wheat varieties have been rendered ineffective due to the rapid evolution of the pathogen. To identify novel sources of adult plant resistance (APR), 292 accessions from the N.I. Vavilov Institute of Plant Genetic Resources, Saint Petersburg, Russia, were screened for known APR genes (i.e. Yr18, Yr29, Yr46, Yr33, Yr39 and Yr59) using linked polymerase chain reaction (PCR) molecular markers. Accessions were evaluated against Pst (pathotype 134 E16 A + Yr17 + Yr27) at seedling and adult plant stages across multiple years (2014, 2015 and 2016) in Australia. Phenotypic analyses identified 132 lines that potentially carry novel sources of APR to YR. Genome-wide association studies (GWAS) identified 68 significant marker–trait associations (P < 0.001) for YR resistance, representing 47 independent quantitative trait loci (QTL) regions. Fourteen genomic regions overlapped with previously reported Yr genes, including Yr29, Yr56, Yr5, Yr43, Yr57, Yr30, Yr46, Yr47, Yr35, Yr36, Yrxy1, Yr59, Yr52 and YrYL. In total, seven QTL (positioned on chromosomes 1D, 2A, 3A, 3D, 5D, 7B and 7D) did not collocate with previously reported genes or QTL, indicating the presence of promising novel resistance factors. Overall, the Vavilov diversity panel provides a rich source of new alleles which could be used to broaden the genetic bases of YR resistance in modern wheat varieties.

Introduction

Wheat is one of the most widely cultivated cereal crops worldwide, with an annual production of approximately 761 million tonnes (Crop Prospects and Food situation 2020). Global wheat yields are threatened by climate change (Asseng et al. 2015; Barlow et al. 2015) and rapidly evolving pathogens causing serious outbreaks of diseases, including rusts (Chaves et al. 2013). Among the rusts, stripe rust (YR) caused by Puccinia striiformis f.sp. tritici (Pst) is an economically important disease that has caused several major epidemics worldwide, resulting in significant production losses (Sanin and Nazarova 2010; Hovmoller et al. 2011; Ellis et al. 2014b; Xia et al. 2016a; Rahmatov 2016; Ali et al. 2017). Historically, YR was prevalent in cooler climates; however, the majority of wheat-growing areas in the world have now become prone to YR (Ali et al. 2014; Hubbard et al. 2015). Currently, 88% of global wheat production is
under threat to YR, which accounts for annual losses of more than one billion US dollars (Beddow et al. 2015). Deployment of YR-resistant varieties is the preferred method of rust disease management because it is cost-effective and reduces the reliance on fungicides (Chen 2005a).

Genetic resistance to YR is broadly categorized into two major classes: All stage resistance (ASR) (e.g. R-genes) and adult plant resistance (APR-genes). ASR is often underpinned by a single gene with a large effect that provides effective resistance at all stages of plant growth. The R-gene interacts with the pathogen in a gene-for-gene relationship (Flor 1971) and therefore is commonly referred to as “race-specific” resistance (Ellis et al. 2014). When deployed in a variety grown on a large scale, strong selection pressure is exerted on the pathogen population to select for mutations that overcome the resistance mechanism. Therefore, this type of resistance is often rendered ineffective within just 3–5 years. In contrast, APR is typically controlled by multiple genes, each with minor or partial effect, and is usually best expressed at adult growth stages. Most of the well-characterized APR genes are race nonspecific and usually confer a “slow rusting” phenotype, which is known to be more durable (Lagudah 2011a; Ellis et al. 2014a; Mundt 2014). Importantly, APR genes can contribute to high levels of resistance through additive or epistatic effects (Sorensen et al. 2014). Some of the genes, such as pleiotropic APRs and high temperature adult plant (HTAP) resistance genes, are highly valuable in breeding programmes. For instance, APR genes Yr18, Yr29, Yr30 and Yr46 confer pleiotropic resistance to YR, leaf rust, stem rust and powdery mildew of wheat (Lan et al. 2015), while Yr18, Yr29, Yr36, Yr39 and Yr52 exhibit HTAP resistance (Chen 2013b).

To date, 83 Yr resistance genes (Yr1–83) have been officially designated, along with 47 genes that have been temporarily named (McIntosh et al. 2019). According to the available information, most are classified as R-genes, whereas only 18 are classified as APR genes (Wu et al. 2016). Notably, only three APR genes Yr18/Lr34 (Krattinger et al. 2009), Yr36 (Fu et al. 2009), and Yr46 (Moore et al. 2015a) have been cloned to date (Liu et al. 2015). The majority of the R-genes, which have been deployed in various wheat varieties, are no longer effective due to the emergence of virulent pathotypes (Hovmoller et al. 2011). For instance, a large number of predominant races evolved from the year 2000 onwards and displayed added virulence to numerous resistance genes such as Yr2, Yr6, Yr7, Yr8, Yr9, Yr10, Yr17, Yr27, Yr43 and Yr44 (Wan and Chen 2014). Virulence was also reported for some of the APR genes (Sorensen et al. 2014). Therefore, an additional level of durable genetic resistance could be achieved by pyramiding both seedling and APR genes in future varieties (Mundt 2014). Hence, the discovery of new sources of genetic resistance is a priority for wheat research, and the successful integration of new technologies in crop improvement programmes is important to achieve long-term rust control.

Plant genetic resources that are stored in gene banks worldwide are a valuable source of genetic diversity for biotic and abiotic stresses (Rao 2004b). Among these, wheat landraces and wild relatives are valuable sources of novel alleles for YR resistance (Sthapit et al. 2014, 2017; Manickavelu et al. 2016; Pasam et al. 2017). In previous studies, three important Yr genes (Yr47, Yr51 and Yr57) have been successfully characterized from wheat landraces in the Watkins collection (Bansal et al. 2011; Randhawa et al. 2015). Another historically important germplasm collection is preserved at the N.I. Vavilov Institute of plant genetic resources (VIR) in Saint Petersburg, Russia, which holds ~38,430 wheat accessions. The collection comprises 76% bread wheat, 16% durum wheat and 7.9% wild and primitive wheats from diverse geographical origins (i.e. Africa, east and west Asia, the USA, Canada, Central and South America, Europe) and of diverse biological status, including wild forms, local cultivars, breeding lines, mutants and artificial allopolyploids (Mitrofanova 2012). While the VIR collection is yet to be explored for YR resistance, previous studies have highlighted the genetic variation for a number of biotic stresses (Mitrofanova 2012; Sadovaya et al. 2015; Riaz et al. 2016a, 2018).

To unravel the genetic architecture of rust resistance, biparental linkage mapping studies are traditionally performed (Yang et al. 2017). However, high costs associated with population development, poor mapping resolution due to low recombination and the constraint of low allelic diversity are some of the limitations associated with the linkage mapping approach (Flint-Garcia, 2013). Alternatively, a GWAS (Genome-wide association study) approach can be applied to a collection of accessions or a natural population. It offers broader allele coverage and higher mapping resolution due to historical recombination events among the panel of lines (Brachi et al., 2011a). This helps to localise the association signals to smaller regions within the chromosome and supports more efficient identification of candidate gene(s). However, applying GWAS to germplasm collections can be challenging because of population structure, which can result in spurious correlations between markers and traits (Gupta 2014; Kulwal et al. 2019; Yang et al. 2017). Another drawback is the low detection power of rare alleles with larger effects or multi-allelic variants with minor effects. Thus, an ideal GWAS analysis requires a large population size, high marker density and a mixed linear model to detect true genotype–phenotype associations (Bulli et al. 2016a). GWAS was initially successfully implemented for rust resistance in hexaploid wheat by Crossa et al. (2007b). Since then, it has been widely used in several studies to detect genomic regions associated with YR resistance in spring, winter, synthetic wheat germplasm, and landraces (Zegeye et al. 2014;
Jighly et al. 2015; Maccaferri et al. 2015b; Naruoka et al. 2015; Bulli et al. 2016a; Godoy et al. 2017; Liu et al. 2017). In this study, we evaluated 292 hexaploid bread wheat accessions from the VIR against Pst to identify novel sources of YR resistance. We applied GWAS to identify key genomic regions that could support the development of new cultivars incorporating durable resistance to YR.

Materials and methods

Plant material

This study examined the Vavilov wheat diversity panel for resistance to YR. The composition of the panel was previously described by Riaz et al. (2017). Briefly, it includes 292 bread wheat accessions, including 136 landraces, 32 cultivars, 10 breeding lines and 115 lines with unknown cultivation classification, which were collected over a 70-year period (Online Resource 1). A total of 206 lines have origin information, whereas the origin is unknown for the remaining 89 lines. Notably, the diversity panel is both morphologically and genetically more diverse compared to Australian and CIMMYT elite wheat materials (Riaz et al. 2017).

Pathogen

Pathotype 134 E16 A+ was used to screen the diversity panel because it was the most virulent and widespread pathotype in Australia during the experimental period. The pathotype 134 E16 A+ was first detected in Western Australia in 2002 where it initially displayed virulence for Yr6, Yr7, Yr8, Yr9, and YrA (Wellings et al. 2003) and gradually acquired virulence for Yr10 and Yr17 (Wellings 2007). Virulence for Yr27 was first reported in 2008, and combined virulence for Yr17 and Yr27 in pathotype E16A+Yr17+Yr27+ was detected in 2010 (Randhawa et al. 2015). PCR marker screening for known APR genes

A subset of 283 wheat accessions were screened for known APR genes. The accessions were screened by a PCR reaction, using markers corresponding to Yr18, Yr29, Yr46, Yr33, Yr39 and Yr59. Marker screening for Yr18, Yr29 and Yr46 (Riaz et al. 2016b) was performed with gene-specific cleaved amplified polymorphic sequence (CAPS) markers cssfr5 (Lagudah et al. 2009), csLV46 (unpublished) and a gene-specific single-nucleotide polymorphic marker SNP1-TM4 (Moore et al. 2015b), respectively. The genes Yr33 and Yr59 were evaluated, each with two closely linked markers to detect their presence or absence. SSR markers gwm111 and gwm437 were used to identify Yr33 (Zahravi et al. 2003), whereas the SSR marker barc32 and resistance gene analog polymorphism (RGAP) marker wmc557 were used to identify Yr59 (Zhou et al. 2014). The Yr39 gene was evaluated using the RGAP marker wgp45 (Lin and Chen 2007). PCR was carried out in a 20 µl reaction containing up to 100 ng of genomic DNA, 1X GoTaq Flexi buffer, 1.5 mM MgCl2, 200 µM dNTP, 200 nM of both forward and reverse primers and 1U of Taq polymerase (M829B, Promega, the USA). A touch down PCR cycling condition was used as follows: denaturation at 94 ºC for 30 s; annealing at 62 ºC for 30 s, decreasing by 0.5 ºC per cycle; extension at 72 ºC for 1 min followed by repeating these steps for 10 cycles; after enrichment, the programme continued for 29 cycles as follows: 94 ºC for 30 s, 58 ºC for 30 s and 72 ºC for 40 s. Amplified products were evaluated using agarose gel electrophoresis.

Phenotyping for seedling resistance

All accessions were evaluated for Pst resistance at the seedling stage in a glasshouse (GH) at the University of Queensland, St Lucia, Queensland, Australia. The experiment was conducted using a complete randomised design with three replicates. Three to four seeds per accession were sown at four different positions around the circumference of 140 mm ANOVApots® filled with potting media. Seedlings were raised under glasshouse conditions maintained at day/night temperature of 22/17 ºC using a natural 12 h diurnal photoperiod. Eleven-day-old seedlings (i.e. two-leaf stage) were inoculated with Pst pathotype 134E16 A+Yr17+Yr27 according to the air-brush protocol reported by Hickey et al. (2012). Seedlings were evaluated after 14 days post-inoculation based on 0–4 Stakman et al. (1962) scale which contains both numbers (e.g. 0, 1…4) and symbols (e.g. +). The numbers represent the disease score: highly resistant (HR, 0–1), moderately resistant (MR, 2), moderately susceptible (MS, 3), and susceptible (S, 4).

Phenotyping for adult plant resistance

A total of 292 accessions of the Vavilov wheat diversity panel were evaluated across three years (2014, 2015 and 2016) at The University of Queensland Gatton Research Station, Queensland, Australia. Wheat lines were sown in the field as un-replicated hill plots. The hill plots were 0.5 m apart along the direction of sowing and in two rows positioned between two spreader rows of the very susceptible genotype, Morocco (Online Resource 2a). To establish YR in the nursery, wheat seedlings that were raised in the greenhouse, infected with Pst, and were transplanted among the spreader rows approximately one month after sowing the nursery. The reaction to YR was evaluated using a modified Cobb’s scale (Peterson et al., 1948) and grouped on a 0–9 scale in 2014, a 1–9 scale (Bariàna et al., 2007) was used.
in 2015 and 2016, the lower score values on both scales indicated increased resistance (Online Resource 2b). Disease assessments were performed at different time-points from heading (Zadoks 50) to grain filling stage (Zadoks 85). Specifically, data were collected at three time-points in 2014 and two time-points in 2015 and 2016. The seven phenotypic datasets were referred to as: Field_2014_1, Field_2014_2, Field_2014_3, Field_2015_1, Field_2015_2, Field_2016_1, and Field_2016_2.

**Genotyping**

DNA of each accession was extracted according to the recommended DAIS protocol (www.diversityarrays.com) and genotyped using the Genotyping-by-sequencing (GBS) Diversity Array Technology (DArTseq) platform (Li et al., 2015). A total of 56,306 raw silicoDArT markers were returned, of which 34,311 were polymorphic in the panel. Only markers positioned on the current DArTseq consensus map (i.e. 14,228 polymorphic markers) were selected and filtered based on 10% and 15% threshold for missing data for markers and accessions, respectively. Markers with a minor allele frequency (MAF) <0.05 were also excluded. Additionally, three markers specific for known APR genes (Yr18, Yr29 and Yr46) were included. In total, 13,934 high-quality markers and 292 accessions were retained and used for GWAS. These markers were ordered based on the genetic map positions in a high-resolution DArTseq consensus map (version 4.0), developed by Diversity Array Technology Inc., Canberra, ACT, Australia.

**Data analysis**

Prior to analysis, seedling resistance scores (Stakman 0–4) were converted to the 0–9 scale where, 0 = immune and 9 = very susceptible, using a conversion table (Ziems et al., 2014). The IT was converted as follows: 0::n:, 1 − 1, 1 + 2 − 2, 2 + 2 +, 3 − 3, 3 +, 3 + + and 4 were coded as 0, 0.5, 1, 2.5, 3, 3.5, 4, 5, 6, 6.5, 7, 8, 8.5 and 9, respectively. The accessions were grouped based on the phenotypic data (IT – Infection Type) of the seedlings; highly resistant (IT = 0–3), intermediate (IT = 4–6) and highly susceptible (IT = 7–9). Accessions representing landraces, cultivars, breeding lines and the unclassified group displaying susceptible reactions at the seedling stage (IT = 7–9) were selected (132 lines) for further analysis to identify lines carrying potentially novel APR genes.

Data analyses were performed using R (R Core Team 2019). To investigate groups of accessions within the Vavilov wheat diversity panel, hierarchical cluster analysis was performed using passport data (including biological status and origin) and the YR disease response data. Squared Euclidean distances were used to calculate the dissimilarity matrix, and the tree was created according to Ward agglomerative clustering criterion. To compare the results across the seedling and adult plant experiments, correlation analyses were performed using mean disease severity (IT) scores. Principal component analysis (PCA) was performed using all datasets, and results were displayed as a biplot to investigate the correlation of disease responses across experiments.

**Identification of novel sources of APR**

To identify accessions that most likely carried novel APR loci, disease response and PCR marker screening results were considered. First, the accessions which displayed high levels of susceptibility at the seedling stage (disease score ≥7) were selected assuming they lacked effective seedling resistance genes. Next, the accessions that carried one or more previously characterized APR gene(s) (Yr18, Yr29, Yr46, Yr33, Yr39 and Yr59) were excluded. Finally, of the remaining accessions, those that displayed high levels of resistance at the adult plant stage were selected (n = 48).

**Genome-wide association studies**

The genetic diversity and population structure of the Vavilov wheat diversity panel were previously described by Riaz et al. (2016a, b). GWAS was performed using a compressed mixed linear model (MLM) (Yu et al. 2006) implemented in the R package known as genome association and prediction integrated tool (GAPIT) (Lipka et al. 2012). To minimise spurious associations due to population structure and relatedness, the Q matrix was used as a fixed effect, and the K matrix was used to fit a random genetic effect (Gupta et al. 2014). To decrease the Type II error rate, a relaxed significance threshold for marker–trait associations (MTA) of –log10(P) > 3.0 was used, as previously employed in other GWAS studies examining the Vavilov panel (Riaz et al. 2018; Dinglasan et al. 2019). Markers detected in different assays, but in the same chromosomal position within a 1 cM window were considered as the same QTL. To confirm the independence of closely positioned QTL, the local LD (linkage disequilibrium) value for associated markers was calculated in R using the ‘genetics’ package (Warnes et al. 2012). For each significantly associated marker, the resistance allele was assigned based on the direction of the allele effect on the resistance score value (Riaz et al. 2018). For example, if the “1” marker allele had a negative effect (e.g. -1.2), it was considered the resistance allele and the “0” marker allele was considered the susceptibility allele.
Comparison of QTL with previously reported Yr genes (QTL) and identification of putative candidate genes

To compare the position of the 47 QTL detected in this study with previously reported Yr genes/QTL, we projected the genetic positions on to the integrated consensus map developed by Maccaferri et al. (2015b) using the MapChart software version 3.2 (Voorrips, 2002) following the co-location strategy described in Ziem et al. (2014). Briefly, all the positions of QTL detected in this study (DArT wheat consensus map version 4) were positioned on the Maccaferri map using two approaches: 1) directly if the marker was present in both maps, or 2) projection using flanking markers in common across the maps. Once all QTL were positioned on the Maccaferri consensus map, the genomic regions were compared with previously reported Yr genes/QTL. The integrated consensus map includes markers from the 9 K (Cavanagh et al., 2013) and 90 K (Wang et al., 2014b) consensus maps, the tetraploid consensus map (Maccaferri et al., 2015a), the Synthetic × Opata DH GBS map (Saintenac et al. 2013), the Diversity Array Technology (DArT) integrated map (http://www.diversityarrays.com/search/node/wheat%20DArT%20map), SSR consensus map (Somers et al. 2004) and the Synthetic × Opata ITMI BARC SSR map (Song et al. 2005). In addition, recently reported genes were also projected onto the integrated map. All QTL regions were compared with the genes/QTL reported by Chen (2017). QTL were considered novel when they were positioned more than 10 cM away from a previously reported gene or QTL. The sequences for flanking markers associated with QTL identified in this study were BLAST searched against the Ensemble Plant genome-centric portal to find the physical position based on the wheat genome assembly IWGSC Ref-Sequence v1.1 (IWGSC 2018). For five DArTSeq markers (1,255,550, 981,525, 1,236,960, 3,026,338, 1,159,261), the 10+ genome assembly browsers at GrainGenes (http://www.graingenenes.org) were used to retrieve the physical position.

Assigning and stacking of alleles associated with novel APR QTL

Seven newly discovered APR QTL were further analysed to determine their effect on disease response. To remove the masking effect of major resistance genes, only accesses displaying susceptibility at the seedling stage (i.e. IT scores > 7) were selected for these analyses. The difference between the mean disease responses for groups of accessions carrying different combinations of resistance alleles was tested for significance based on Tukey’s test for multiple comparisons with a family-wise error rate of 5%. To investigate the trend between disease response and the accumulation of resistance alleles for the seven APR QTL, a relative disease index (YRi) was calculated across field trials according to:

\[ YRi = \frac{\sum_{k} individualDis.Score[k]}{meanDis.Score[k]} \]

where YRi is the disease index of a line, in experiment k in relation to the population mean over n field experiments. A disease index below 0 reflects a high level of resistance to YR across the trials, whereas values above 0 imply susceptibility.

Results

Analysis of stripe rust response at seedling and adult plant stages

In total, 292 accesses were tested for their IT at the seedling stage and the results revealed that 39% of the accesses showed highly resistant reactions (IT = 0–3), 16% moderately resistant reactions (IT = 4–6) and 45% were susceptible (IT = 7–9). Most accesses displaying susceptibility were of unknown origin (n = 50; Online Resource 1), followed by accesses from Russia (n = 27), India (n = 17) and Pakistan (n = 11). In total, 132 lines displayed susceptible reactions to YR at the seedling stage, and therefore, these accesses were considered potential candidates for carrying uncharacterised APR genes. Under field conditions, the accesses displayed a wide range of responses across the three years (Fig. 1). In 2014 and 2015, more than 80% of the tested lines (n = 284) displayed resistant to moderately resistant reactions, whereas in 2016, only 58% of the tested lines (n = 247) showed resistant reactions of which 43% displayed intermediate resistant phenotypes (Fig. 1).

Weak positive correlations were observed between seedling and most field-based assessments; for example, Field_2014_3 (r = 0.33, P < 0.001) and Field_2015_2 (r = 0.30, P < 0.001). The seedling disease response showed a moderate correlation with Field_2016_2 (r = 0.53, P < 0.001). Conversely, stronger correlations were observed between the field assessments; for example, Field_2015_1 and Field_2016_2 (r = 0.81, P < 0.001). In Online Resource 3, the accesses displayed in black are seedling-susceptible and lack known genes, thus representing accesses that could carry uncharacterised APR.

Cluster analysis using phenotype and passport data

Ward’s hierarchical agglomerative cluster analysis was performed for 238 wheat accessions using trait data, including biological status (Unclassified, Classified – Breeding line, Cultivar, Landrace), origin of collection
and YR disease response in the glasshouse and field experiments. The analysis revealed three major clusters divided into eight subgroups (Fig. 2 and Online Resource 1). Cluster I comprised accessions belonging only to the unclassified group (i.e. unknown geographical information and unknown biological status). All the accessions in this cluster were grouped under the APR and moderate resistance (MR) category (Subgroups 2, 3 and 5). The two other major clusters (Cluster II and III) comprised accessions belonging to a combination of classified and unclassified groups. Cluster II comprised accessions carrying APR (Subgroups 6 and 8) and accessions that were generally susceptible (Subgroup 4). Accessions in Subgroup 6 were more resistant and stable across the years compared to accessions belonging to Subgroup 8. Cluster III comprised accessions carrying mainly APR (Subgroup 7) and ASR (Subgroup 1).

Identification of novel sources of APR

Based on the results from YR assessments and PCR marker screening, 48 lines were deemed to carry potentially novel sources of APR to YR (Online resource 3). This included lines classified as unknown origin (n = 18), landraces (n = 14), cultivars (n = 10) and breeding lines (n = 6). The landraces were mainly from India (n = 3), Spain (n = 2), Russia (n = 2), and Kazakhstan (n = 2), although landraces from Pakistan, Turkey, Azerbaijan, Tajikistan, and Armenia were also included in the set. Most cultivars (n = 8) and breeding lines (n = 5) that carried potentially novel sources of APR were from Russia. The 10 most promising lines were selected for crossing to elite cultivars. These showed a high degree of susceptibility at the seedling stage and intermediate levels of resistance at the adult plant stage (Fig. 3). The majority of these lines were from the unclassified group (n = 6), followed by a landrace from Pakistan (WLA–043).
A total of 68 significant marker–trait associations were detected \((-\log_{10}(p\text{ value}) > 3)\) which represented 47 different QTL regions (Table 1, Online Resource 4). Fifty-seven markers were associated with resistance at the adult plant stage, and many \((n = 26)\) were detected only under the environmental conditions experienced in 2014. Nine markers were associated with resistance at the seedling stage, while only one marker was detected at both seedling and adult plant stage. Based on the chromosomal position and LD between adjacent markers, the QTL represented 41 APR loci and 6 seedling resistance loci. Comparison of these QTL regions with the previously reported \(Yr\) genes and QTL identified seven genomic regions as potentially novel QTL associated with \(YR\) resistance at the adult plant stage (Online Resource 5, 6). Notably, 14 QTL overlapped with the positions of previously reported \(Yr\) genes: \(Yr29\), \(Yr5\), \(Yr43\), \(Yr57\), \(Yr30\), \(Yr51\), \(Yr48\), \(Yr47\), \(Yr35\), \(Yr36\), \(Yrxy1\), \(Yr59\), \(Yr52\) and \(YrYL\) on chromosome 1B, 2B, 3B, 4A, 5B, 6B, 7A, 7B and 7D, suggesting they could be linked to, or represent, those known resistance factors (Online resource 6).

Six QTL regions defined by 10 markers (2A, 2B, 2D, 3A, 3B and 5A) were associated with seedling resistance. Of these QTL, only four were detected at the seedling stage, while the remaining two QTL \((qNV.Yr-2B.2\) and \(qNV.Yr-3B\)) were detected at both seedling and adult plant growth stages. The QTL \(qNV.Yr-2B.2\) included seven markers within the 88.5–89.9 cM region, which were associated with resistance across test environments (i.e. seedling, field trials in 2015 and 2016). It also co-located with the seedling resistance gene \(Yr43\). The remaining three QTL were detected only in one adult plant environment (besides the seedling stage). The QTL with the strongest association was \(qNV\).

**Marker–trait associations**

Fig. 2 Ward’s hierarchical agglomerative clustering of 238 Vavilov wheat lines generated from biological status (Unclassified, Classified–Breeding line, Cultivar, Landrace), origin of collection and stripe rust disease response in the glasshouse and field experiments. Dendrogram height (0–35) is represented by the horizontal line. Eight subgroups were delineated (height 10) which represent level of stripe rust resistance.

Fig. 3 Seedling and adult plant responses to stripe rust for the 10 lines most promising lines carrying potentially novel sources of adult plant resistance (APR). Orange bars represent the disease response at the seedling stage, whereas yellow, green and brown represent disease responses at the adult plant stage in 2014, 2015 and 2016, respectively. Avocet and the APR near-isogenic lines (Avocet + \(Yr18\) and Avocet + \(Yr29\)) were evaluated at the adult stage in 2014 only.
Table 1  Summary of the stripe rust resistance QTL identified at both the seedling and adult plant stage in the Vavilov wheat diversity panel

| QTL name      | Peak Marker a | Chr | Position (cM)b | Position (Mb)c | Phenotypic dataset(s)d | − log₁₀(P) | R. allele e | Effect on trait | Gene/QTL | References                          |
|---------------|---------------|-----|----------------|----------------|------------------------|------------|-------------|----------------|----------|-------------------------------------|
| qNV.Yr-1B-1   | 1,219,437     | 1B  | 241.91         | 660.6          | Field_2_2016           | 3.14       | 1           | − 0.2          | Previously reported | Maccaferri et al. 2015a, b |
| qNV.Yr-1B-2   | 1,119,286     | 1B  | 269.26         | 677.4          | Field_2_2016           | 3.34       | 0           | − 0.2          | Lan et al. 2014, Maccaferri et al. 2015a, b |
| qNV.Yr-1D-1   | 1,862,252     | 1D  | 29.72          | 21             | Field_3_2014           | 3.17       | 1           | − 0.8          | New                  |                      |
| qNV.Yr-1D-2   | 1,229,892     | 1D  | 47.7           | 35.7           | Field_2_2016           | 3.23       | 1           | − 0.22         | Previously reported | Maccaferri et al. 2015a, b |
| qNV.Yr-2A.1   | 2,295,954     | 2A  | 9.93           | 12.2           | Field_1_2015           | 3.2        | 1           | 0.05           | Previously reported | Maccaferri et al., 2015a, b |
| qNV.Yr-2A.2   | 1,255,550     | 2A  | 24.56          | 33.6           | Seedling               | 3.5        | 1           | 0.05           |                      | Bansal et al. 2014, Basnet et al. 2014a, b |
| qNV.Yr-2A.3   | 1,093,102     | 2A  | 46.78          | 49.5           | Field_2_2015           | 3.78       | 1           | − 0.2          | Previously reported | Naraoka et al. 2015 |
| qNV.Yr-2A.4   | 1,122,504     | 2A  | 60.52          | 87             | Field_1_2015           | 3.59       | 1           | 0.08           | Previously reported | Bulli et al. 2016 |
| qNV.Yr-2A.5   | 1,862,406     | 2A  | 68.57          | 256.2          | Field_2_2015           | 3.45       | 1           | 0.11           | New                  |                      |
| qNV.Yr-2A.6   | 1,183,380     | 2A  | 124.25         | 761.1          | Field_2_2015           | 3.6        | 0           | 0.23           | Previously reported | Dedryver et al. 2009 |
| qNV.Yr-2B.1   | 2,289,349     | 2B  | 78.31          | 583.7          | Field_2_2016           | 3.45       | 1           | 0.29           | Yr5                  | Godoy et al. 2017 |
| qNV.Yr-2B.2   | 1,235,357     | 2B  | 88.54          | 763            | Field_2_2016           | 3.19       | 0           | 0.37           | Yr43                 | Maccaferri et al. 2015a, b, Xu et al. 2013 |
| qNV.Yr-2B.3   | 1,236,960     | 2B  | 107.02         | 783.9          | Field_2_2015           | 4.33       | 0           | − 0.33         | Previously reported | Crossa et al. 2007, Buerstmayr et al. 2014 |
| qNV.Yr-2D.1   | 2,246,033     | 2D  | 70.85          | 790.6          | Field_2_2014           | 4.11       | 1           | − 0.57         |                      |                      |
| qNV.Yr-2D.2   | 2,244,346     | 2D  | 70.85          | 790.6          | Field_2_2014           | 3.01       | 0           | − 0.12         |                      |                      |
| qNV.Yr-3A.1   | 1,123,104     | 3A  | 44.75          | 170.1          | Seedling               | 3.22       | 1           | 0.33           | Previously reported | Zegeye et al. 2014 |
| qNV.Yr-3A.2   | 2,253,468     | 3A  | 49.23          | 574.3          | Field_1_2015           | 3.41       | 0           | 0.06           | Previously reported | Pasam et al. 2017 |
| qNV.Yr-3A.3   | 2,295,884     | 3A  | 53.26          | 600.3          | Field_3_2014           | 3.09       | 1           | 1.5            | Previously reported | Hou et al. 2015 |
| qNV.Yr-3A.4   | 1,116,501     | 3A  | 109.54         | 100.4          | Field_2_2015           | 3.23       | 0           | − 0.11         | Previously reported | Rosewarne et al. 2013 |
| QTL name    | Peak Marker | Chr | Position (cM) | Position (Mb) | Phenotypic dataset(s) | $-\log_{10}(P)$ | R. allele | Effect on trait | Gene/QTL | References                  |
|-------------|-------------|-----|---------------|---------------|-----------------------|----------------|-----------|----------------|-----------|-----------------------------|
| qNV.Yr-3A.5 | 1,237,203   | 3A  | 142.13        | 738.9         | Field_1_2014          | 4.38           | 0         | 0.6            | New       | Lan et al. 2014             |
| qNV.Yr-3B   | 2,307,351   | 3B  | 4.97          | 5.3           | Field_1_2016 Field_2_2016 | 4.28           | 0         | $-0.23$        | Yr57      | Basnet et al. 2014a         |
|             | 1,239,212   |     | 5.25          | 5.3           | Field_1_2016 Field_2_2016 | 3.83           | 0         | $-0.02$        | Yr30      |                             |
| qNV.Yr-3D.1 | 992,483     | 3D  | 59.49         | 374           | Field_1_2014          | 3.88           | 1         | $-0.41$        | New       |                             |
| qNV.Yr-3D.2 | 1,228,106   | 3D  | 123.84        | 582.5         | Field_2_2014          | 3.45           | 0         | 3.73           | Previously reported | Basnet et al. 2014a |
| qNV.Yr-4A.1 | 1,120,960   | 4A  | 96.08         | 668.4         | Field_2_2014 Field_2_2014 | 3.76           | 1         | 1.02           | Previously reported | Bulli et al. 2016, Zegeye et al. 2014 |
| qNV.Yr-4A.2 | 1,131,408   | 4A  | 117.26        | 252.7         | Field_2-2016 Field_2_2016 | 3.79           | 0         | 0.42           | Previously reported | Vazquez et al. 2012 |
| qNV.Yr-4B.1 | 3,021,792   | 4D  | 43.44         | 158.4         | Field_2_2016 Field_2_2014 | 3.74           | 0         | 0.3            | Yr46      | Herrera-Foessel et al. 2011 |
| qNV.Yr-4A.2 |             |     |               |               | Field_1_2014          | 3.4            |           | $-0.07$        |           |                             |
| qNV.Yr-5A   | 1,117,919   | 4D  | 54.74         | 439.3         | Field_1_2014          | 3.08           | 1         | 1.19           | Previously reported | Ren et al. 2012a, b |
| qNV.Yr-5B.1 | 1,127,003   | 5A  | 58.39         | 524.8         | Seedling              | 4.8            | 0         | $-0.03$        | Previously reported | Manickavelu et al. 2016, Lan et al. 2014 |
| qNV.Yr-5B.2 | 1,088,238   | 5B  | 12.41         | 15.3          | Field_2_2015 Field_2_2016 | 3.26           | 0         | NA             | Yr47      | Bansal et al. 2013         |
| qNV.Yr-5B.3 | 1,125,706   | 5B  | 22.69         | 42.3          | Field_2_2016 Field_2_2015 | 3.07           | 1         | $-0.01$        | Previously reported | Hao et al. 2011 |
|             | 1,104,944   | 5B  | 27.83         | 54.2          | Field_2_2015 Field_2_2015 | 3.07           | 0         | 0.04           | Previously reported | Yang et al. 2013, Lu et al. 2014 |
| qNV.Yr-5B.4 | 1,128,792   | 5B  | 29.32         | 53            | Field_2_2015 Field_2_2014 | 3.03           | 0         | 0.01           | Previously reported | Lu et al. 2009, Vazquez et al. 2015 |
| qNV.Yr-6A   | 991,465     | 5D  | 58.61         | 424           | Field_1_2014 Field_3_2014 | 3.62           | 0         | 0.14           | New       |                             |
| qNV.Yr-6B.1 | 3,022,417   | 6A  | 100.76        | 616.3         | Field_2_2016 Field_2_2016 | 3.61           | 0         | $-0.01$        | Previously reported | Vazquez et al. 2012, Bulli et al. 2016 |
| qNV.Yr-6B.1 | 1,302,727   | 6B  | 2.41          | 9.8           | Field_2_2014 Field_2_2014 | 3.64           | 0         | $-1.78$        | Yr35      | Prins et al. 2011, Bansal et al. 2013 |
|             | 1,097,180   |     | 2.41          | 0.3           | Field_1_2015 Field_1_2015 | 3.19           | 0         | $-0.2$         | Yr35      | Bansal et al. 2013         |
| qNV.Yr-6B.2 | 1,121,581   |     | 4.36          | 18            | Field_1_2014 Field_1_2014 | 3.89           | 1         | 1.06           | Previously reported | Prins et al. 2011, Bansal et al. 2013 |
| qNV.Yr-6B.2 | 2,290,856   | 6B  | 23.19         | 117.1         | Field_1_2015 Field_1_2015 | 3.63           | 1         | 0.12           | Yr36      | Pasam et al. 2017, Santra et al. 2008 |
| qNV.Yr-6B.4 | 1,058,394   | 6B  | 35.34         | 576.5         | Field_2_2016 Field_3_2014 | 3.55           | 0         | $-0.09$        | Previously reported | Klarquist et al. 2016 |
| qNV.Yr-6B.5 | 2,293,763   | 6B  | 79.73         | 710.6         | Field_3_2014 Field_1_2015 | 3.07           | 1         | $-3.44$        | Previously reported | Basnet et al. 2014a, Rosewarne et al. 2013 |
|             | 1,109,194   |     | 82.95         | 715.8         | Field_1_2015 Field_1_2015 | 3.19           | 0         | $-0.37$        |           |                             |
Table 1 (continued)

| QTL name  | Peak Marker | Chr | Position (cM) | Position (Mb) | Phenotypic dataset(s) | $-\log_{10}(P)$ | R. allele | Effect on trait | Gene/QTL | References                                      |
|-----------|-------------|-----|---------------|---------------|------------------------|-----------------|-----------|-----------------|-----------|------------------------------------------------|
| qNV.Yr-7A.1 | 1,244,188 | 7A  | 96.07         | 648.3         | Field_2_2016           | 3.28            | 0         | 0.15            | Yrxy1     | Zhou et al. 2011, Singh et al., 2014a, b, Zegeye et al. 2014 |
| qNV.Yr-7A.2 | 1,090,077  | 7A  | 149.56        | 712.3         | Field_2_2015           | 3.34            | 0         | 0.15            | Yrxy1     | Rosewarne et al. 2013 Vazquez et al. 2012 |
| qNV.Yr-7B.1 | 1,126,816  | 7B  | 20.97         | 170.5         | Field_2_2014           | 3.26            | 1         | 2.31            | New       | Zhou et al. 2014 |
| qNV.Yr-7B.2 | 1,059,624  | 7B  | 38.66         | 138.7         | Field_2_2014           | 3.23            | 1         | 0.12            | Previous reported Suenaga et al. 2003 |
| qNV.Yr-7B.3 | 1,259,637  | 7B  | 38.65         | 129.5         | Field_1_2014           | 3.17            | 0         | 0.01            | Previous reported Pasam et al. 2017 |
| qNV.Yr-7B.4 | 1,066,421  | 7B  | 118.47        | 718.1         | Field_1_2014           | 3.36            | 1         | 0.14            | Yr52      | Ren et al. 2012a, b, Zhou et al. 2014 |
| qNV.Yr-7D.1 | 1,105,401  | 7D  | 56.58         | 65.9          | Field_2_2015           | 3.3             | 1         | 0.27            | YrYL      | Wu et al. 2016 |
| qNV.Yr-7D.2 | 2,303,363  | 7D  | 103.6         | 103.6         | Field_1_2016           | 3.03            | 1         | 0.04            | New       | Zhou et al. 2014 |

*Most strongly associated marker for the QTL

*Chromosome position (cM) of the QTL on the DArTseq wheat consensus map (v 4.0)

*Chromosome position (Mb) of the QTL on the physical map (Ref-Sequence v1.1 (IWGSC 2018), 10+ genome assembly browsers—GrainGenes)

*The dataset for which the QTL was detected

*Allele associated with resistance
Yr-3B \((-\log_{10} P \geq 4)\) with larger marker effects on disease response \((-0.23)\). This QTL co-located with partial resistance (and pleiotropic) gene \(Yr30\) and the seedling resistance gene \(Yr57\), which provides resistance against the Australian pathotype \(134\text{E}16\ A+ Yr17 + Yr27\). However, due to the narrow genomic interval between these genes on chromosome 3B, we were unable to determine whether \(Yr30\), \(Yr57\) or both were present in this QTL.

A total of 41 QTL (57 markers) were considered “APR QTL” and were significantly associated with IT in one or multiple environments (Table 1). A total of 26, 23 and 16 markers were detected in 2014, 2015 and 2016, respectively, and clustered into different QTL regions. Comparison of the detected loci with the previously reported \(Yr\) genes and QTL (using the integrated map from Maccafferri et al. 2015b) revealed that 41 genomic regions likely represented previously reported genes and/or QTL (Table 1, Online Resource 5, 6 and 7). Interestingly, genomic regions harbouring the HTAP resistance genes \(Yr36\), \(Yr59\) and \(Yr52\) were detected in the 2014 and 2015 field environments, whereas APR genes \(Yr29\) and \(Yr30\) were detected in 2016. Notably, high average maximum temperatures \((28.5 \degree C - 31.4 \degree C)\) were experienced during the YR assessment periods in 2014 and 2015 compared to 2016.

Seven QTL (chromosomes 1D, 2A, 3A, 3D, 5D, 7B and 7D) were detected at the adult plant stage and appeared to be novel (Table 1). The QTL \(qNV.Yr-7D.2\) was detected under environmental conditions in 2016, whereas all other QTL in the D genome were detected in 2014. The QTL \(qNV.Yr-2A.5\) and \(qNV.Yr-5D\) were significant in multiple field environments, while \(qNV.Yr-1D-1\), \(qNV.Yr-3A.5\), \(qNV.Yr-3D.1\), \(qNV.Yr-7B.1\), and \(QYr.uq-7D.2\) were detected only in a single field environment.

### Allele stacking effects and geographical distribution of the novel APR QTL

The seven novel APR QTL were investigated for their potential additive effect to reduce disease severity. A total of 132 accessions that were seedling susceptible were selected for the analysis. To further verify allelic effects in the subset, the mean resistance level of lines carrying alleles associated with resistance was compared with those carrying alleles for susceptibility at each locus. Significant allelic effects were verified through t-tests for \(qNV.Yr-2A.5\), \(qNV.Yr-3A.5\), \(qNV.Yr-3D.1\), and \(qNV.Yr-7D.2\), whereas no significant effects were detected for \(qNV.Yr-1D-1\), \(qNV.Yr-5D\) and \(qNV.Yr-7B.1\) (Fig. 4). To investigate the accumulation effect of the seven novel APR loci for YR resistance, lines were grouped based on the number of QTL present in each line. A clear decreasing trend in the field relative disease index was observed as the number of resistance alleles increased from one to three (Fig. 5a). None of the Vavilov lines combined more than three resistance alleles. Notably, most of the lines carrying three novel resistance loci \((n = 23)\) showed a disease index lower than zero. The novel alleles occurred singly or in combination in accessions from India, Kazakhstan, Ukraine, and Russia (Fig. 5b). Further, countries from central Asia to west Asia such as Uzbekistan, Tajikistan, Armenia, and Georgia also carried combinations of 2–3 novel alleles.
Discussion

Vavilov wheat accessions offer diverse sources of stripe rust resistance

In this study, the Vavilov wheat accessions displayed a wide range of YR responses in both seedling and adult plant stages across multiple environments. At the seedling stage, about 55% of the accessions displayed high levels of resistance (39%, IT = 0–3) to intermediate levels of resistance (16%, IT = 4–6). Based on field assessments in 2014, 83% of the lines also showed resistance at the adult plant stage, indicating the presence of strong seedling resistance or combination of effective seedling and APR to YR. Alternatively, these resistant accessions could carry multiple APR genes, as the presence of just 2 or 3 minor genes can confer “near-immune” levels of resistance (Bariana et al. 2010; Singh et al. 2014b). Some genes are known to have a synergistic effect on disease resistance, such as Lr34/Yr18 (German and Kolmer 1992), which directly contributes resistance to leaf rust, YR and powdery mildew (Rinaldo et al. 2017), while also contributing resistance to stem rust due to additive and epistatic interactions (Hiebert et al. 2016). These types of genes are more valuable when breeding for rust resistance because they are less likely to be selected and deployed alone, and some genes can enhance resistance levels to more than one economically important disease.

A total of 45% of the accessions (n = 132) displayed susceptibility at the seedling stage, but resistance at adult growth stages. Considering the diversity of the germplasm, it suggests a high frequency of multiple APR genes. Despite using the same pathotype each year, the correlation between disease response varied across the three years of field evaluations. This highlights the role of genotype × environment interactions influencing the expression of resistance. The low temperature and high rainfall in 2016 compared to 2014 and 2015 likely favoured disease development and could be the reason for increased susceptibility in 2016. Although, this does not rule
out other factors related to seasonal variation such as differences in physiological development across seasons (Wang et al. 2014) and abiotic stress (Guo et al. 2008). Such effects are well described for the Lr34/Yr18 gene, where studies have found that gene expression is highly influenced by the genetic background, environmental factors and level of infection (Risk et al. 2012).

Screening for known APR genes (Yr18, Yr29, Yr46, Yr33, Yr39, and Yr59) using linked molecular markers revealed some interesting relationships when accession passport information was explored. For instance, the presence of one or more gene combinations of Yr59 and Yr46 in almost all accessions from India and Pakistan suggests that these APR genes are common in germplasm originating from these regions. A total of the 88 (65%) lines that displayed strong resistant responses in both seedling and adult plant stages were classified as landraces. Previous studies have also found that wheat landraces offer a rich source of disease resistance loci (Tala et al. 2011; Pasam et al. 2017).

By combining the field screening data with results from marker screening for known APR genes, we identified accessions that carried potentially novel resistance alleles. Interestingly, a number of lines originating from European and Asian countries such as Russia, Kazakhstan, Armenia, Ukraine, Tajikistan, Spain, Uzbekistan, Turkey, Mongolia, and China showed an absence of known APR genes and clustered in the APR category, which suggests they likely carry novel resistance alleles. The top 10 lines deemed to carry potentially novel sources of APR were from Russia, Pakistan, Chile, and the group of unknown origin. The Russian accessions WLA–302 (1978) and WLA–087 (1990) displayed intermediate resistance across the three years of evaluation. The landrace WLA–043 from Pakistan collected in 1936 was found to be a good source of APR to YR across the years. Pakistan is a diversity hotspot for wheat due to its wide agroecological zones, where many landraces that are genetically diverse with rare alleles have been found, particularly in the Himalayan region (Hirano et al. 2008). Evaluation of 5,700 accessions from USDA—ARS National Small Grain Collection (NSGS) showed that Chile is another important geographic centre of rust resistance with the highest number of accessions (28%) for stem rust resistance determined at the adult plant stage (Bonman et al. 2007). In this study, the cultivar WLA–300 collected from Chile in 1963 was identified as one of the best accessions for APR to YR. The top 10 accessions identified in this study should be prioritised for bi-parental mapping studies to characterise the underlying resistance loci.

GWAS reveals 47 genomic regions associated with resistance

This study identified 68 DArTseq markers, representing 47 QTL regions, that were significantly ($P < 0.001$) associated with seedling and adult plant resistance to YR. Notably, a larger number of marker–trait associations ($n = 26$) were detected under field conditions in 2014, which represented 21 QTL regions. In contrast, the markers detected in 2016 ($n = 15$) represented 14 different genomic regions and 13 of them were unique to the field environment in 2016. Therefore, the field conditions experienced in 2016 resulted in the detection of many loci that were either not expressed or weakly expressed in the 2014 and 2015 field environments because GWAS was unable to detect them. This highlights the importance of testing material for rust resistance across multiple environments.

Among the 6 QTL detected at the seedling stage, a major haplotype block comprising seven markers was identified on chromosome 2B (qNo.Yr-2B.2). These markers were significantly associated with resistance at seedling, adult plant or both stages, suggesting the possible presence of more than one locus within this region. Previously, four different genes (Yr5, Yr44, Yr53 and Yr43) have all been mapped in this region within 35.1 cM (Xu et al. 2013). Based on the positions of key markers, Yr43 is most likely the underlying gene for qNo.Yr-2B.2. It should be noted that virulence for Yr43 and Yr44 occurs in many Pst races that emerged from 2000 onwards in the USA (Chen 2005; Wan and Chen 2014), while Yr5 and Yr53 have remained effective against all races. Therefore, it is important to study this QTL in detail to understand the specific gene/genes which confer resistance to the Australian pathotype 134E16 A + Yr17 + Yr27. Such information would be very useful for breeding programmes.

Three well-characterised APR multi-resistance loci were detected in this study, including Lr46/Yr29/Sr58/Pm39/Ln2 on chromosome 1BS, Sr2/Yr30/Lr27/Pm on 3BL and Lr67/Yr46/Pm46/Sr55/Ltn3 on 4DS. These are the most commonly introduced and selected genes in global wheat breeding programmes. Interestingly, effects associated with these genes were detected in the 2016 field environment that experienced relatively lower temperatures and higher rainfall, in comparison to the 2014 and 2015 seasons. Apart from these genes, only a few QTL were detected in 2016 compared to the other years. The QTL (QYrst.orr-1B and QYr.ucw-1D) reported by Maccarelli et al. (2015a) and QTL (QYrst.orr-4A and QYrst.orr-6A) reported by Vazquez et al. (2012) were also detected in the present study in 2016. The resistance conferred by these QTL may be further studied in detail by analysing two seedling-susceptible landraces from Armenia (WLA–249) and Russia (WLA–315). Further, the HTAP resistance genes such as Yr36 (Uauy et al. 2005), Yr52 (Ren et al. 2012a, b) and Yr59 (Zhou et al. 2014) coincided with QTL detected in 2014 and 2015 field environments which were much warmer than 2016, which further suggests these regions could be underpinned by HTAP resistance loci.
Other important QTL detected in this study were qNV.Yr-3B and qNV.Yr-7B.4 (Table 1). The first QTL, qNV.Yr-3B, was detected at the seedling stage; however, this QTL was also detected at the adult stage under field conditions in 2016. qNV.Yr-3B co-located with seedling resistance gene Yr57 and adult plant pleiotropic gene Yr30. Yr57 is a broadly effective against Australian pre- and post-2000 Pst pathotypes 104 E137A +, 108E141A +, 110 E143A +, 134 E16A +, 134 E16A + Yr17 +, 134 E16A + Yr17 + Yr27 + and 150 E16A + (Randawa 2015), whereas Yr30 is closely linked to Sr2 and confers partial resistance to YR. These two genes are closely linked (<5 cM apart), which could be exploited for gene pyramiding (Randawa 2015). The second QTL, qNV.Yr-7B.4, were represented by DArT marker 1,066,421 in the field in 2014. Notably, this region on chromosome 7B harbours two HTAP genes, Yr52 (Ren et al. 2012a, b) and Yr59 (Zhou et al. 2014), and two temporally designated genes YrC591 (Li et al. 2008) and YrZH84 (Li et al. 2006). Yr52 confers HTAP resistance, which is highly consistent and partial in nature, making it an attractive target for breeding programmes targeting durable rust resistance (Ren et al. 2012a, b). Yr59 is another important HTAP resistance gene, which is linked to YrZH84 (Zhou et al. 2014) and is effective against all known North American Pst races; thus, it is considered highly valuable for resistance breeding.

Novel QTL for adult plant resistance

Seven new QTL for YR resistance were identified on chromosomes 1D, 2A, 3A, 3D, 5D, 7B and 7D. These genomic regions were associated with different levels of resistance across the three years of field assessments. qNV.Yr-2A.5 (chromosome 2A) and qNV.Yr-5D (chromosome 5D) were associated with resistance in 2014 and 2015, while other QTL were detected only in a single environment. Therefore, these QTL may be environment-specific or convey smaller effects. Notably, the majority of novel QTL detected in this study were located in the D-genome. Aegilops tauschii, the D-genome progenitor of hexaploid wheat, is considered a valuable source of YR resistance (Liu et al. 2013). However, until now, only a few resistance loci have been identified in the D-genome. For instance, Yr28 (Singh et al. 2000), slow rusting APR gene Yr46 (Herrera foessel. 2011) and YrAs2388R (Zhang et al. 2019) have been mapped on chromosome 4D. Therefore, the novel QTL detected in the current study provides valuable sources for APR. Accession numbers WLA-106 (Ukraine), WLA-124 (Russia) for qNV.Yr-1D-1, WLA-43 (Pakistan) for qNV.Yr-3D.1 and qNV.Yr-5D, and WLA-114 (Azerbaijan) for QYr.uq-7D.2 represent valuable donor lines that can be used for pre-breeding or future research seeking to fine-map or clone the underlying gene(s).

Novel APR QTL confer additive effects to reduce disease

We demonstrated that pyramiding of novel alleles associated with APR can significantly reduce YR disease. As a group, APR lines that carry a combination of 2–3 resistance alleles show stronger levels of resistance compared to lines carrying just one resistance allele. However, allele effects are variable depending on the QTL, the QTL combination, genetic background and environmental conditions. For example, WLA-043 from Pakistan carries three QTL (3A, 3D and 5D) and showed stable resistance across years in comparison with WLA-279 from Kazakhstan which also carries three QTL (1D, 2A and 3A). It has been well described that some resistance genes (such as Yr18, Yr30 and Yr39) can increase the effectiveness of other R genes or weak APR genes (Chen et al., 2013, Ellis et al., 2014). While Yr18 and Yr49 confer additive effects for YR resistance, increased stem rust severity was detected in Chinese Spring due to the inactivation of Lr34 gene in the presence of Sr2 (Ellis et al. 2014). Therefore, it is important to understand gene interactions to incorporate effective disease resistance into wheat varieties.

Conclusions

This is the first study to document YR resistance in diverse wheat accessions sourced from the VIR. GWAS identified 47 genomic regions including 68 markers that were significantly associated with resistance. Over 90% of the QTL detected in this study aligned with previously reported genes and QTL, which highlights the power of association mapping and accuracy of resistance loci that have been identified in the present study. The molecular markers linked to QTL identified in this study will be valuable for further validation of resistance loci and potentially future marker-assisted selection breeding approaches. Further, insight into the distribution of resistance alleles and their frequencies in relation to geographical origin could aid selection of specific sources of resistance in breeding programmes. The significant effect of pyramiding novel APR alleles identified in this study underscores the potential for breeding wheat varieties with improved YR resistance.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00122-022-04037-8.

Acknowledgements DJ was supported by an Australian Postgraduate Award. LTH was supported by a Discovery early career researcher award (DECRAs) Fellowship from the Australian research council (ARC) and an Early Career Researcher grant from UQ. We give thanks to the past and present members of the N.I. Vavilov Institute of Plant Genetic Resources in St Petersburg, Russia, for preserving and sharing the historic wheat germplasm. We are grateful to the technical staff at UQ Central Glasshouse Facilities at St. Lucia and the UQ
Contribution: DJ, AR and LTH conceived and designed the study; DJ, AR, NA and WLN collected the data; DJ wrote the manuscript; DJ, EGD, KVF, SA, LZ, OA, EA, SKP and IG contributed to data analyses and interpretation of results; GP guided the design and establishment of rust screening nurseries in Queensland, Australia; all authors read and revised the manuscript.

Funding: Open Access funding enabled and organized by CAUL and its Member Institutions.

Data availability: The datasets generated during and/or analysed during the current study (i.e., phenotypes, genotypes, and mapping data) are available from the corresponding author on reasonable request.

Conflict of interest: The authors declare that they have no conflict of interest.

Ethical approval: This article does not contain any studies with human participants or animals performed by any of the authors.

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