Molecular markers for leaf rust resistance gene Lr42 in wheat

Zhengli Liu, Robert L. Bowden and Guihua Bai

How to cite this manuscript

If you make reference to this version of the manuscript, use the following information:


Published Version Information


Copyright: © Crop Science Society of America


Publisher’s Link: https://www.crops.org/publications/cs/articles/53/4/1566
Molecular Markers for Leaf Rust Resistance Gene \textit{Lr42} in Wheat

Zhengli Liu, Robert L. Bowden and Guihua Bai,

Z. Liu, Institute of Millet Crops, Hebei Academy of Agricultural & Forestry Sciences, Shijiazhuang, Hebei, 050031 China; Z. Liu and G. Bai, Dep. of Agronomy, Kansas State University; R. L. Bowden and G. Bai, USDA-ARS, Hard Winter Wheat Genetics Research Unit, Manhattan, KS 66506

Received ------. *Corresponding author (guihua.bai@ars.usda.gov).

Abbreviations: Infection type, IT; short arm of chromosome 1D, 1DS; simple sequence repeats, SSR; quantitative trait locus, QTL; marker-assisted selection, MAS.
Abstract

Wheat leaf rust, caused by *Puccinia triticina* Eriks., is an important wheat foliar disease worldwide. Growing cultivars incorporating genetic resistance is one of the most effective approaches for disease control. Leaf rust resistance gene *Lr42* was identified from a wheat relative, *Aegilops tauschii* Coss, and has been transferred into hard winter wheat. A previous study identified two markers closely linked to the gene on the short arm of chromosome 1D (1DS) using a near isogenic population, but flanking markers for *Lr42* were not found. In this study, a new mapping population was developed from a cross between ‘KS93U50’ (a *Lr42* carrier) and a susceptible parent, ‘Morocco’. An F2 population was analyzed with all simple sequence repeat (SSR) markers available from chromosome 1D and F2 plants and F3 families were evaluated for seedling resistance to isolate PNMR, a rust isolate avirulent to *Lr42*. The F2 and F3 rust data showed that *Lr42* was recessive. Seven markers formed the linkage group on 1DS. The *Lr42* region was flanked by two simple sequence repeat (SSR) markers; *Xwmc432* and *Xgdm33* at 17 cM apart. The results confirmed that *Lr42* is positioned on the distal end of chromosome 1DS. The flanking markers for *Lr42* should be useful for map-based cloning and marker-assisted pyramiding of *Lr42* with other leaf rust resistance genes.
Introduction

Leaf rust of wheat (*Triticum aestivum* L.), caused by *Puccinia triticina* Eriks., is one of the most important wheat foliar diseases worldwide (Kolmer, 1996). Severe leaf rust epidemics can cause yield losses up to 40% (Knott, 1989). Growing genetically resistant cultivars is one of the most effective strategies to minimize yield losses due to the disease. To date, at least 71 genes for resistance to leaf rust have been catalogued in wheat (Singh et al., 2012). Of these, about half were introgressed from related species. Many of them were from *Ae. tauschii* Coss, the D genome donor of common wheat, including *Lr21* (1DS), *Lr22a* (2DS), *Lr32* (3DS), *Lr39* (2DS), and *Lr42* (1DS) (Rowland and Kerber, 1974, Kerber, 1987, Gill et al., 1991, Cox et al., 1994).

Most *Lr* genes confer race-specific seedling resistance and are vulnerable to defeat by new virulent races. Greater durability of resistance could be achieved through combinations of race-specific genes or by using race-nonspecific resistance genes, such as *Lr34* and *Lr46* (Kolmer et al, 2008a & b). However, such genes provide low levels of resistance when deployed alone or singly (Zhang et al., 2008). A third option is to combine both race-specific and race-nonspecific resistance.

Combining race-specific and race-nonspecific resistance genes in a single cultivar could significantly improve both durability and the level of resistance (Kolmer et al., 2008a, Zhang et al., 2008). For instance, in spring wheat, the combination of *Lr16, Lr23* (race-specific resistance genes) and *Lr34* (a race-nonspecific resistance gene) has provided effective and durable resistance for many years (Kolmer et al., 2008a & b). *Genes Lr34* and *Lr13* were demonstrated to enhance the level of resistance synergistically when in combination with other leaf rust resistance genes (Germán and
Kolmer, 1992; Kolmer, 1992). Two recent papers also demonstrated the ability of quantitative race-nonspecific resistance to extend the usefulness of qualitative race-specific resistance genes (Brun et al, 2010; Pailloix et al, 2009).

Stacking of several leaf rust resistance genes using phenotypic selection is difficult. A variety of rust races are often not available that could distinguish the presence or absence of different genes. If some genes in the stack confer high resistance, they can mask the phenotypic expression of other genes. Molecular markers can make it possible to stack several genes simultaneously in a single genetic background, therefore, identifying closely linked markers for each resistance gene is essential for successful use of these genes in breeding.

*Lr42* is a gene conferring rust resistance at both seedling and adult plant stages and remains effective against all leaf rust races reported to date (Sun et al., 2010). Lines containing *Lr42* have been used as a parent in some breeding programs with success (Bacon et al., 2006; Singh et al., 2007). Previous work located *Lr42* on 1DS (Cox et al., 1994) and found that *Lr42* also played a significant role in increasing wheat yield and kernel size, apparently due to the resistance (Martin et al., 2003). Using molecular mapping, Czembor et al, (2008) located *Lr42* to a different chromosome (3D) using DArT markers and reported that the gene was dominant. Sun et al. (2010) further mapped *Lr42* to the distal end of 1DS using SSR markers and the same near-isogenic population developed by Martin et al. (2003), and identified two closely linked markers. Thus additional mapping work using different populations is necessary to validate the chromosome location of *Lr42*. The current study examined a population derived from the cross ‘KS93U50’ x ‘Morocco’ to: 1) confirm the gene location in a new population, 2)
study the inheritance pattern of the gene, 3) identify flanking markers linked to the gene for marker-assisted selection (MAS).

Materials and Methods

Plant materials and rust inoculation

The mapping population was developed by crossing ‘KS93U50’ and ‘Morocco’.

‘KS93U50’ is a hard red winter wheat accession containing Lr42 selected from ‘KS91WGRC11’ (PI 566668, pedigree ‘Century’* 3/T. tauschii TA2450) and provided by the Wheat Genetic and Genomic Resources Center, Kansas State University, Manhattan KS. Morocco is an old soft white spring cultivar originally collected from North Africa and frequently used as a susceptible check for leaf, stem, and stripe rust.

A total of 371 F2 plants derived from the cross ‘KS93U50’ x ‘Morocco’ was inoculated with rust race PNMR. PNMR is avirulent to ‘KS93U50’, but virulent to several susceptible cultivars tested (Sun et al., 2010). All F2 plants and both parents were planted in 72-cell plastic flats and inoculated with PNMR at the two-leaf stage. Before inoculation, rust spores from a liquid N2 tank were heat shocked in water bath at 40°C for 5 min to break dormancy. The spores were suspended in Soltrol 170 light oil (Chevron Phillips Chemical Company, Woodlands, TX) and misted uniformly over the seedling leaves using a pressure sprayer. After inoculation, the plants were incubated in a moist chamber at 20 °C for 24 h before being moved to a greenhouse bench for rust establishment. The plants were grown at 20 °C supplemented with 10 h daylight. The rust infection type (IT) on a 1 to 4 scale as described previously (Stakman et al., 1962; Roelfs and Martens, 1988), was recorded at 10 d after inoculation and rechecked 2 d
later for confirmation. After rust data were collected, all F$_2$ plants were transplanted to 13 cm by 13 cm plastic pots for seed increase. Seeds from 361 F$_2$ plants (10 F$_2$ plants did not set any seed) were harvested individually and planted in 72-cell plastic growth trays for rust evaluation. Six seeds per F$_{2:3}$ family were planted and the experiments had two replications. Six seeds per parent also were planted in each tray as control. Rust inoculation and disease scoring for the parents and F$_{2:3}$ families were the same as for the F$_2$ population. Based on parent reactions to PNMR, plants with IT$\leq$2 were classified as resistant and IT$\geq$2$^+$ as susceptible for segregation analysis. The segregation ratio from F$_{2:3}$ families was used to determine the genotypes of F$_2$ plants and the combined data from both generations were used to determine inheritance pattern.

**Marker genotyping and data analysis**

After rust evaluation, newly developed disease-free wheat leaf tissue was collected in 1.1 mL 96-deepwell plates for DNA isolation. Tissue samples were immediately dried in a freeze-dryer (ThermoSavant, Holbrook, NY) for 48 h and ground to fine powder using a Mixer Mill (MM300, Rotsch, Germany) by shaking the tubes at 25 times per s for 5 min with a 3.2-mm stainless steel bead in each well. Genomic DNA was extracted using a modified cetyltrimethyl ammonium bromide method (Sun et al., 2009). Because $Lr42$ was previously located on chromosome 1D (Cox et al., 1994), 27 SSR primers on chromosome 1D (Somers et al., 2004; Song et al., 2005, Roder et al., 1998) were selected to screen the parents for polymorphism. PCR amplifications were performed in a Tetrad Peltier DNA Engine (Bio-Rad Lab, Hercules, CA). A 12 µL PCR mixture containing 1.2 µL of 10X PCR buffer (Bioline, Taunton, MA), 2.5 mM of MgCl$_2$, 200
µM of each dNTP, 50 nM of forward M13-tailed primer, 250 nM of reverse primer and 200 nM of M13 fluorescent-dye-labeled primer, 0.6 U of Taq DNA polymerase, and about 50 ng template DNA. A touchdown PCR program was used for PCR amplification.

In brief, the reaction was incubated at 95 ℃ for 5 min then continued for 5 cycles of 1 min at 96 ℃, 5 min at 68 ℃ with a decrease of 2 ℃ in each subsequent cycle, and 1 min at 72 ℃. For another 5 cycles, the annealing temperature started at 58 ℃ for 2 min with a decrease of 2 ℃ for each subsequent cycle. Reactions then went through an additional 25 cycles of 1 min at 96 ℃, 1 min at 50 ℃, and 1 min at 72 ℃ with a final extension at 72 ℃ for 5 min. PCR products were analyzed on an ABI PRISM 3730xl DNA Analyzer (Applied Biosystems, Foster City, CA). Data were analyzed using GeneMarker software ver. 1.6 (SoftGenetics LLC. State College PA), and manually checked twice to remove ambiguous data.

A linkage map was constructed using Kosambi (1944) function and MultiPoint package (available at http://www.MultiQTL.com). A logarithm of odds (LOD) threshold of 3 was used to identify linkage groups.

Results

Ten days after inoculation with PNMR, the susceptible parent ‘Morocco’ showed severe rust symptoms with IT ≥ 2+, while most of plants of the resistant parent, ‘KS93U50’, only showed tiny pustules surrounded by necrosis and with IT ≤ 2. In F₂ and F₃ populations, IT ranged from 1 to 4 for different genotypes. Because most of the plants of the susceptible parent had IT ≥ 3 with only a few plants that had IT = 2+, plants with IT ≥ 2+ were classified as susceptible genotypes and plants with IT ≤ 2
were classified as resistant genotypes. Of 371 F₂ plants evaluated 282 were susceptible and 89 were resistant, fitting a 3:1 ratio ($x^2=0.202, P=0.653$), indicating that $Lr42$ behaved as recessive in the ‘KS93U50’ x ‘Morocco’ population. The F₂:₃ family segregation fit a 1 resistant: 2 segregating: 1 susceptible ratio ($x^2=7.693, p=0.0214$), with slight distortion toward susceptible families (Table 1). In each segregating family, most of the plants were susceptible, which confirmed that $Lr42$ is recessive.

Of the 27 SSR markers available on chromosome 1D, 11 proved polymorphic between two parents. Eight markers were mapped to one linkage group and three others were unlinked. Two of the eight markers, Xbarc229 and Xwmc336, showed significant segregation distortion and were removed from the linkage group. The map with six markers covered 68.5 cM on chromosome 1D (Fig. 1). $Lr42$ was mapped between Xwmc432 and Xgdm33 at 4 cM from Xwmc432. A slight segregation distortion was observed for the markers in the region of $Lr42$ (Table 2).

Of the primers for these markers, Gdm33 amplified a fragment of 177 bp in ‘KS93U50’ and a fragment of 145 bp in ‘Morocco’. Primer Wmc432 amplified two fragments in each of the two parents, 204 and 212 bp from ‘KS93U50’, and 203 and 218 from ‘Morocco’, but only the larger fragments were segregating in the population and mapped in the $Lr42$ region.

**Discussion**

In this study, most F₂ plants were susceptible and segregation indicated a single recessive gene for resistance. Segregation in the F₃ population was consistent with the premise that resistance to rust isolate PNMR due to $Lr42$ in KS93U50 was recessive.
This result disagrees with previous reports that \textit{Lr42} was dominant (Czembor et al., 2008) or partially dominant (Cox et al., 1994). The discrepancy could result from different isolates and genetic materials used in different studies. Kolmer and Dyck (1994) demonstrated that expression of resistance genes could range from complete dominance to complete recessiveness and the expression of resistance and avirulence genes in wheat leaf rust system is highly dependent on the genotypes of the host lines and pathogen isolates used. In this study, ‘KS93U50’ was used as the source of \textit{Lr42} that was a selection from ‘KS91WGRC11’, while ‘KS91WGRC11’ was used in the other studies (Czembor et al. 2008, Cox et al. 1994). Pathogen inocula used in these studies might also be different. In this study, several local isolates were evaluated using a set of parents and near-isogenic lines and we found that PNMR could clearly differentiate \textit{Lr42} from various susceptible genotypes (Sun et al., 2010). Thus it was the isolate of choice used in this study. However, for direct resistance evaluation in segregating breeding materials, this isolate may not be ideal because phenotypic differences are not sufficient to distinguish heterozygous from homozygous susceptible plants.

The closest marker to \textit{Lr42} is \textit{Xwmc432}, which confirmed the previous report (Sun et al., 2010). In the previous study, a backcross derived homozygous near-isogenic population was used and disease scores were treated as binary data. In this study, \textit{F}_2 was used for mapping. The genetic distance between \textit{Lr42} and \textit{Xwmc432} is about 4 cM longer in this study than in the previous study. In this study, \textit{F}_3 rust data agreed with \textit{F}_2 in general, but heterozygous status of some \textit{F}_2 plants could not be detected due to limited number of \textit{F}_3 plants per family were evaluated for rust resistance, which might contribute to expended genetic distance between \textit{Lr42} and \textit{Xwmc432}. In addition, we
identified one new marker, \textit{Xgdm33}, distal to \textit{Lr42}. \textit{Xwmc432} and \textit{Xgdm33} defined \textit{Lr42} in a 17.8 cM region. Thus they can be used as flanking markers for further fine mapping in the region to facilitate map-based cloning of the gene.

Two markers developed in this study are good candidate markers for MAS of \textit{Lr42}. Primer \textit{Wmc432} amplified two fragments in both parents, but only the 212 bp fragment from ‘\textit{KS93U50}’ and the 218 bp fragment from ‘\textit{Morocco}’ associated with \textit{Lr42}, as in a previous study (Sun et al., 2010). This marker should be the first choice for MAS. Because allele size differences between the two parents were relatively large for \textit{Xwmc432} and \textit{Xgwm33}, they can be analyzed in an agarose gel in breeding programs. A high concentration agarose gel is recommended when using \textit{Xwmc432}.

In this study, four markers closely linked to \textit{Lr42} were positioned on chromosome 1DS based on the previously reported linkage maps (Somers et al., 2004, Song et al., 2005), which disagrees with Czembor et al. (2008). However, our results are in agreement with the previous report by Sun et al. (2010) despite different types of populations (near-isogenic verse F2 populations) and susceptible parents used in the two studies. The discrepancy among different studies could be due to difference in resistant parents used in different studies as discussed previously. Thus, if the markers developed in this study are to be used for screening \textit{Lr42} gene, it is advised that ‘\textit{KS93U50}’ be used as the source of resistance.

\textbf{Acknowledgements}

This project was partially supported by the National Research Initiative Competitive Grants CAP project 2011-68002-30029 from the USDA National Institute of Food and
Agriculture and USDA-ARS project 5430-21000-006-00D. The authors would like to thank Dr. Paul St. Amand, USDA Central Small Grain Genotyping Center, and Dr. Chengsong Zhu, Department of Agronomy, Kansas State University, Manhattan KS, for technical assistance. Mention of trade names or commercial products in this article is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the U.S. Department of Agriculture. USDA is an equal opportunity provider and employer. This is contribution No. 12-414-J from the Kansas Agricultural Experiment Station, Manhattan, KS.

References


Mackay, L. McIntyre, P. Sharp. 739-740 (824pp).


resistance and awns to agronomic and grain quality performance in winter wheat.

Crop Sci. 43:1712-1717.


races of *Puccinia graminis var. tritici*. US Department of Agricultural Publications E617. USDA, Washington, D.C.


Table 1. Segregation of *Lr42* in F$_2$ and F$_{2:3}$ populations derived from a cross of ‘KS93U50’ x ‘Morocco’ when inoculated with *Puccinia triticina* Eriks. isolate PNMR at seedling stage

<table>
<thead>
<tr>
<th>Generation</th>
<th>No. of Lines</th>
<th>Observed ratio†</th>
<th>Expected ratio†</th>
<th>$X^2$</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>F2</td>
<td>371</td>
<td>89:282</td>
<td>1:3</td>
<td>0.202</td>
<td>0.653</td>
</tr>
<tr>
<td>F3</td>
<td>361</td>
<td>86:167:113</td>
<td>1:2:1</td>
<td>7.693</td>
<td>0.021</td>
</tr>
</tbody>
</table>

† F2 ratio is resistant:susceptible; F3 ratio is resistant:segregating:susceptible
Table 2 Segregation of marker alleles located on chromosome 1D.

<table>
<thead>
<tr>
<th>Marker</th>
<th>No. of Lines</th>
<th>Observed ratio †</th>
<th>Expected ratio †</th>
<th>$X^2$</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Xgdm33$</td>
<td>359</td>
<td>76:174:109</td>
<td>1:2:1</td>
<td>6.404</td>
<td>0.041</td>
</tr>
<tr>
<td>$Xwmc432$</td>
<td>360</td>
<td>83:167:110</td>
<td>1:2:1</td>
<td>5.928</td>
<td>0.052</td>
</tr>
<tr>
<td>$Xcfd0015$</td>
<td>364</td>
<td>84:169:111</td>
<td>1:2:1</td>
<td>5.863</td>
<td>0.053</td>
</tr>
<tr>
<td>$Xgwm337$</td>
<td>355</td>
<td>80:188:87</td>
<td>1:2:1</td>
<td>1.518</td>
<td>0.468</td>
</tr>
<tr>
<td>$Xgdm60$</td>
<td>359</td>
<td>89:183:87</td>
<td>1:2:1</td>
<td>0.159</td>
<td>0.924</td>
</tr>
<tr>
<td>$Xgwm458$</td>
<td>357</td>
<td>93:173:91</td>
<td>1:2:1</td>
<td>0.361</td>
<td>0.835</td>
</tr>
</tbody>
</table>

† Ratio is resistant:segregating:susceptible
Figure 1. A genetic linkage map developed from the F₂ population of ‘KS93U50’ x ‘Morocco’ showing Lr42 on the short arm of the chromosome 1D of wheat.