Transfer of new leaf rust resistance genes from diploid *T. monococcum* and *T. boeoticum* to *T. aestivum*

Kaur S¹, Chhuneja P¹, Dhaliwal HS², and Singh K¹
¹Dept. of Plant Breeding, Genetics and Biotechnology, Punjab Agricultural University, Ludhiana 141004, India
²Dept. of Biotechnology, Indian Institute of Technology, Roorkee, India

ABSTRACT

The diploid ‘A’ genome progenitor gene pool of wheat, comprising three closely related species *T. monococcum* ssp *monococcum* (*T. monococcum*), *T. monococcoccum* ssp *aegilopoides* (*T. boeoticum*) and *T. urartu*, harbours useful genes for many economically important traits, including resistance to leaf rust. *T. monococcum* acc. paul4078 and *T. boeoticum* acc. pau 5088 are immune to all the leaf rust races at adult plant stage and so is the RIL population generated from the cross these two accessions. At seedling stage these lines were resistant to leaf rust races 77-2 and 77-5 but population segregated for resistance against race 104-2. *T. monococcum* and two RILs designated as RIL130 and RIL101 were crossed to susceptible *T. durum* parent, N59. Rust resistance was suppresses in the F₁. Further crossing and backcrossing with hexaploid wheat cultivars WL 711 and PBW 343 led to identification of resistant progenies. One BC₂F₂ population of the cross N59/RIL130//3*PBW343 when tested at seedling stage showed single gene segregation (489R: 142S χ² = 2.0). Similarly another BC₂F₂ population of the cross N59/T. *monococcum*/3*WL711 also showed single gene segregation (153R:66S, χ² = 3.0) at seedling stage. Both the populations are cytologically stable with 42 chromosomes indicating a stable leaf rust resistance gene transfer from *T. monococcum* to hexaploid wheat. Bulk segregant analysis of the BC₂F₂ population of the cross N59/*T. monococcum*/3*WL711 with 38 polymorphic SSRs indicated that the seedling resistance gene of *T. monococcum* is located on chromosome 5A. As none of the designated *Lr* genes map on 5A, the gene in question may be a novel one.

INTRODUCTION

Among the three rust diseases of wheat, leaf rust caused by *Puccinia recondita* Rob ex Desm. f. sp. *tritici* is the most common, affecting the wheat production worldwide. An effective, economical and environmentally safe method to control epidemics of leaf rust is the cultivation of resistant cultivars. Until now more than 50 leaf rust resistance genes (*Lr1-Lr58*) have been identified (McIntosh et al. 2005; Kuraparth et al. 2007, Chhuneja et al 2007) and most of these, especially the ones identified from the cultivated germplasm, are ineffective against the recently evolved pathotypes. This necessitates search for new sources of resistance. The wild relatives of wheat are rich sources of resistance genes particularly for biotic stresses. The diploid ‘A’ genome progenitor gene pool of wheat, comprising three closely related species *T. monococcum* ssp *monococcum* (*T. monococcum*), *T. monococcoccum* ssp *aegilopoides* (*T. boeoticum*) and *T. urartu*, harbour useful genes for many economically important traits (Feldman and Sears 1981; Dhaliwal et al. 1993; Hussien et al. 1997, Qiu et al. 2005) including resistance to leaf rust. Transfer of these genes into to hexaploid wheat generally require the use of *T. durum* as bridging species. However, presence of suppressor loci on the B genome of *T. durum* present a major hurdle in transferring useful variability from diploid to hexaploid wheats (Chhuneja et al 2008). Identification of DNA markers linked to the desirable genes at the diploid level could facilitate their transfer to hexaploid wheat (Yao et al. 2007). A spring type *T. monococcum* acc. paul14087 and *T. boeoticum* acc. pau 5088 has maintained high levels of resistance to a number of wheat diseases including leaf rust in Punjab (India) over years (Dhaliwal et al. 2003). This report describes the transfer of leaf rust resistance of *T. monococcum* and a RIL (RIL130 - derived from cross of *T. monococcum/T. boeoticum*) to bread wheat and its tagging with microsatellite markers using bulk segregant analysis.

MATERIAL AND METHODS

The plant material consisted of *T. monococcum* acc. 14087 (*Tm14087*), *T. boeoticum* acc. 5088 (*TB5088*) and a set of 121 recombinant inbred lines (RILs) derived from their intercross. Mapping of leaf rust resistance was not possible because the parents and the population showed immunity to leaf rust under field conditions at adult plant stage. Hence efforts were made to transfer these genes to hexaploid wheat first. For this *Tm14087* and RIL130 were crossed to a susceptible *T. durum* CV N59. The F₁ plants were crossed to WL 711 and a widely grown bread wheat cultivar PBW 343. The three-way F₁ thus generated was backcrossed to respective recurrent parents WL711 and PBW 343 to obtain BC₁F₁ generation. The BC₁F₁ plants resistant to leaf rust at seedling stage were identified, selfed and chromosomally stable BC₂F₂ plants identified. The BC₂F₂ and BC₂F₃ populations of N59/RIL130//2*PBW343 and BC₁F₂ and BC₂F₃ progenies of N59/Tm14087//3*WL711 were screened against leaf rust at seedling and against a mixture of pathotypes at adult plant stage. Chi-square (χ²) test was applied to fit appropriate genetic ratio in F₂ and F₃ generations.
For mapping the leaf rust resistance genes transferred from *T. monococcum*, DNA was extracted from individual BC$_2$F$_2$ plants of the cross N59/RIL130//2*PBW343 and BC$_2$F$_2$ plants of N59/Tm14087//3*WL711.

For screening against leaf rust at seedling stage, race 77-5 (with avirulence/virulence formula P$_Lr$9, Lr18, Lr19, Lr24, Lr25, Lr28, Lr29, Lr32, Lr41, Lr45/pLr1, Lr2, Lr3, Lr10, Lr11, Lr12, Lr13, Lr14, Lr15, Lr16, Lr17, Lr18, Lr20, Lr21, Lr22, Lr23, Lr26, Lr27+Lr31, Lr33, Lr34, Lr36, Lr37, Lr42, Lr43, Lr44, Lr46, Lr48, Lr49) was used. For seedling tests, 10-15 seedlings of each parental line, and the segregating generations were sown in plastic trays containing mixture of farmyard manure and sandy loam in equal proportion. In each tray, six rows of experimental material and seventh row of susceptible cultivar Agra Local were sown. First leaf of seven-day-old seedling(s) of each plant was inoculated individually with uridiniospores - talc mixture keeping inoculum density of 6-8 uridiniospores over a microscopic field area of 2.92mm$^2$. The inoculated seedlings were incubated in a dark chamber maintained at 20°C±1°C at 100 percent relative humidity for 16 hours. After incubation, the trays were shifted to glass houses maintained at 20°C±2°C. Fourteen days after inoculation, the infection types on seedlings using the scale proposed by Stakman et al. (1962). Disease severity for all parents and populations was assessed by growing adult plants of these in open experimental field during third week of November, which is the normal sowing time for wheat crop in Northern India. This material was repeatedly spray inoculated every morning with mixture of uridiniospores of leaf rust races (including race 77-5), suspended in water (1 gram inoculum per 10 litres of water, using one drop of Tween-20 as dispersant). The inoculations started during first week of January and continued until rust started appearing on susceptible lines early February. Spreader rows were planted all around the population to ensure an effective disease spread.

At seedling stage plants with the disease reaction 0-2 were considered as resistant and the ones with 3 and above as susceptible. At adult plant stage observations on leaf rust severity were recorded as percentage of leaf area covered with rust according to a modification of the Cobb scale as described by Peterson et al. (1948). The plants with disease severity 0 to 10MR were classified as resistant and the ones with 10MS or more were classified as susceptible.

**RESULT AND DISCUSSION**

**TRANSFER OF LEAF RUST RESISTANT GENES**

*Tm14087* and RIL130 showed infection type of 0, and 1, respectively at the seedling stage and at adult plant stage in the field these were immune. The recipient hexaploid wheat cultivars WL711 and PBW343 were susceptible, giving an infection type of 33+ and 3 at seedling stage and leaf rust severity of 80S and 40S, respectively at the adult plant stage. *T. durum* cv N59 was also susceptible both at the seedling as well as the adult plant stages.

While transferring leaf rust resistance from *T. monococcum*, one major problem encountered was suppression of the resistance in F$_1$ of *T. durum* cv N59/ *T. monococcum* and *T. durum* cv N59/ *T. monococcum/ *T. aestivum* cv PBW343 (WL711). It was later confirmed that the loci suppressing resistance in F$_1$ generations are present on B genome chromosomes (Chhuneja et al. 2008). Leaf rust resistant plants, however, were recovered in the BC$_1$F$_2$ generation of the crosses, *T. durum* cv N59/*T. monococcum/ *T. aestivum* cv PBW343 and *T. durum* cv N59/ RIL130// WL711. The resistant plants were further, either selfed or backcrossed to the recurrent bread wheat parent and BC$_1$F$_2$, BC$_1$F$_3$, BC$_2$F$_2$, and BC$_2$F$_3$ progenies generated. Segregation pattern of these progenies is presented in Table 1. At seedling stage BC$_2$F$_2$ progeny of the cross N59/RIL130//2*PBW343 segregated in 3:1 ratio ($\chi^2 = 2.0$ and the BC$_1$F$_2$ progeny showed 1:2:1 segregation ($\chi^2 = 4.63$). This confirms the transfer of a single dominant seedling resistance gene from RIL130 into hexaploid wheat cultivar PBW 343. Similarly, BC$_2$F$_2$ progeny of the cross N59/Tm14087//3*WL711 showed 3:1 segregation ($\chi^2 = 3.0$) and the BC$_1$F$_2$ progeny showed 1:2:1 segregation ($\chi^2 = 0.99$). This again confirms transfer of a single dominant gene from *T. monococcum* to hexaploid wheat. Allelic test for these genes is in progress to ascertain whether the gene transferred from RIL130 is same as that from *T. monococcum* or from *T. boeoticum*. Also the populations are being analysed for SSR markers for mapping of these genes. Bulk segregant analysis of BC$_2$F$_2$ progeny of the cross N59/Tm14087//3*WL711 indicates that the gene for leaf rust resistance from *T. monococcum* may be located on chromosome 5A.

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<th>Table 1. Segregation pattern in various generations of the crosses N59/RIL130//2<em>PBW343 and N59/Tm14087//3</em>WL711</th>
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<td><strong>Progeny</strong></td>
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Res = resistant/ homozygous resistant, Sus = susceptible/ homozygous susceptible, Seg = segregating
REFERENCES


