Interactions Between \textit{Lr67} or \textit{Lr34} and Other Leaf Rust Resistance Genes in Wheat (\textit{Triticum aestivum})

Brent D. McCallum* and Colin W. Hiebert

Agriculture and Agri-Food Canada, Morden Research and Development Centre, Morden, MB, Canada

The wheat multi-pest resistance genes \textit{Lr67} and \textit{Lr34} are similar in that they both condition resistance to many diseases, in a non-race-specific manner, and code for cellular transporters. \textit{Lr34} plays a critical role in breeding wheat for disease resistance in large part because it interacts with other resistance genes to result in effective and durable resistance. To determine if \textit{Lr67} interacts with other resistance genes in a similar manner as \textit{Lr34} six different doubled haploid populations were developed which segregated for either \textit{Lr67} or \textit{Lr34} along with a second resistance gene, either \textit{Lr13}, \textit{Lr16}, or \textit{Lr32}. The presence or absence of each of these genes in the progeny lines was determined by molecular marker analysis. These six populations were tested for leaf rust field resistance in the same environments to compare the effects of \textit{Lr34} and \textit{Lr67} alone, and in combination with \textit{Lr13}, \textit{Lr16} or \textit{Lr32}. \textit{Lr67} and \textit{Lr34} significantly reduced the levels of rust severity, \textit{Lr34} showed a significant interaction with \textit{Lr13} but \textit{Lr67} did not. Both genes interacted with \textit{Lr16}, and \textit{Lr67} had a significant interaction with \textit{Lr32}. This analysis demonstrates the similar effect of \textit{Lr67}, as seen with \textit{Lr34}, on the interaction with other resistance genes to give a better level of resistance than with single resistance genes. While \textit{Lr67} is not widely deployed in agriculture, it could play an important role in disease resistance in future wheat cultivars.

Keywords: resistance, pyramid, gene, combinations, interaction, durable

INTRODUCTION

Wheat leaf rust is a very common and destructive disease of wheat internationally (Huerta-Espino et al., 2011) and in Canada (McCallum et al., 2016). Genetic resistance has proven effective in controlling this disease, however the \textit{Puccinia triticina} Eriks. pathogen population has evolved virulence for most of the race-specific resistance genes widely deployed in wheat cultivars (McCallum et al., 2016). The race non-specific resistance gene \textit{Lr34} has been used widely over many years in wheat cultivars and has remained effective. Canadian wheat cultivars commonly carry \textit{Lr34} (McCallum et al., 2011) and it is frequently present in other wheat cultivars throughout the world. It also confers resistance to other diseases including stripe rust (\textit{Yr18}, Singh, 1992), stem rust (\textit{Sr57}, Dyck et al., 1985; Hiebert et al., 2010), powdery mildew (\textit{Pm38}, Spielmeyer et al., 2005), and virus diseases (\textit{Bdv1}, Singh, 1993).

One important feature of \textit{Lr34} is that it interacts with other leaf rust resistance genes to give better levels of resistance. Ezzahiri and Roelfs (1989) determined that the durable and
effective adult plant resistance in Era wheat was controlled by the interaction of Lr13 and Lr34. These authors crossed plants of Era with the susceptible cultivar Baart, and they tested 473 and 367 F₁ derived lines in the F₂ generation in Minnesota USA and Morocco, respectively. They found that Lr34 significantly enhanced the level of resistance conditioned by Lr13 but the effect of Lr34 on its own was not detected.

German and Kolmer (1992) crossed the Thatcher-Lr34 near-isogenic line (NIL) with other Thatcher NILs containing different resistance genes. From F₂ families ten to 16 plants with the lowest infection type when inoculated with P. triticina race 1 were selected and grown to maturity. Seed from these selected F₂ plants was then grown in a rust nursery and the five or six most resistant F₂ lines, homozygous for Lr34 were selected and harvested. Two homozygous F₂ lines per cross were tested as seedlings in the greenhouse and three to four plants of one F₂ line were tested as adults in greenhouse tests. Selected lines with Lr34 and a second leaf rust resistance gene were also field tested over two years. These authors found that Lr34 enhanced resistance, both at the seedling and adult plant stages, in combination with many effective resistance genes, and this also resulted in lower adult plant infection types and leaf rust severity levels in the field. However, combinations involving Lr34 with less effective or ineffective genes had the same level of resistance as Thatcher-Lr34.

Kloppers and Pretorius (1997) investigated two gene combinations of Lr13, Lr34, and Lr37. For glasshouse studies, they used a single F₂ line from crosses between the pairs of Thatcher isolines that contained each of these resistance genes. These same F₂ lines and six to eight sister lines from the same crosses were compared in field trials. They found that the two gene lines generally had better resistance as measured by latent period, field resistance, and the microscopic development of fungal structures. Interestingly, they also noted significant variation between sister lines for the Lr34 + Lr13 gene combination in which partial resistance was found in field trials. No variation was found among the sister lines from the gene combinations involving Lr37 since the level of resistance was nearly complete.

The wheat leaf rust resistance gene Lr67 is also race non-specific, is only effective at the adult plant stage, and confers multi-pest resistance to stripe rust (Hiebert et al., 2010; Herrera-Foessel et al., 2011) along with stem rust and powdery mildew like Lr34 (Herrera-Foessel et al., 2014). Both Lr34 and Lr67 have been cloned and code for different types of cellular transporters (Krattinger et al., 2009; Moore et al., 2015). Mutations in either gene resulted in mutants that were susceptible to leaf, stem, and stripe rust (Spielmeyer et al., 2013). With combinations of most leaf rust resistance genes, the severity of disease observed is similar to the most effective of the genes involved. However, Lr34 interacts with other leaf rust resistance genes, and combinations of genes involving Lr34 are more resistant than any of the genes involved.

Some of the most common leaf rust resistance genes in Canadian wheat are Lr2a, Lr10, Lr13, Lr14a, Lr16, Lr21, and Lr34 (McCallum et al., 2016). In this study, we choose to determine the interactions between both Lr67 and Lr34 with each of the genes; Lr13, Lr16, and Lr32. They represent a range of effectiveness from mostly ineffective (Lr13) to highly effective (Lr32). Both Lr13 and Lr16 are in many Canadian wheat cultivars, such as Carberry (Bokore et al., 2022), because popular wheat cultivars grown in the recent past like AC Barrie and AC Domain have either or both genes and donated these genes to the current generation of wheat cultivars. Lr13 is relatively ineffective against the Canadian population of P. triticina, as nearly all isolates are virulent to Lr13 (McCallum et al., 2021). However, it may still have an effect on reducing leaf rust severity in combination with other genes, such as Lr34 and Lr67. Complete virulence to Lr16 is rare in Canada (McCallum et al., 2021) but nearly all isolates have an intermediate level of virulence and combinations of Lr16 with genes, such as Lr34 and Lr46, are fairly effective at reducing leaf rust severity (Bokore et al., 2022). In contrast, Lr32 is a very effective leaf rust resistance gene in Canada with no virulence detected to date (McCallum et al., 2021); however, it has not yet been deployed in any Canadian wheat cultivars. The Thatcher near-isogenic lines containing Lr13 (RL4031), Lr16 (RL6005), Lr32 (RL6086), Lr34 (RL6058), and Lr13 + Lr34 (RL6114) had annual rust severity averages in inoculated nurseries in Manitoba Canada over the years 2003–2021 of 76.6, 65.6, 30.8, 23.2, and 10.3%, respectively, compared with Thatcher at 81.9% (B. McCallum unpublished).

Given the many similarities between Lr67 and Lr34, the objective of this study was to determine if Lr67 also interacts with other resistance genes. To test this we developed six doubled haploid populations from the crosses with either single gene lines with Lr34 or Lr67 and each of the near-isogenic lines with either Lr13, Lr16, or Lr32. Each progeny line was genotyped with molecular markers to determine if the line had the resistant or susceptible allele of each gene involved in the population, except for Lr13 which was determined by rust testing at the adult plant stage. Progeny from these crosses were field tested over four years to determine the resistance level of lines in each phenotypic class; susceptible, those having the resistant allele for either Lr34 or Lr67 alone, those only having the resistant allele of the second resistance gene (Lr13, Lr16 or Lr32), and those with both genes.

MATERIALS AND METHODS

Populations

Doubled haploid (DH) populations were developed from the crosses between Thatcher near-isogenic lines with Lr34 (RL6058), Lr67 (RL6077), Lr13 (RL4031), Lr16 (RL6005), and a Katepwa backcross line with Lr32 (BW196R). The progeny populations consisted of 78 lines (Thatcher-Lr13/Thatcher-Lr34), 74 lines (Thatcher-Lr13/Thatcher-Lr67), 58 lines (Thatcher-Lr16/Thatcher-Lr34), 85 lines (Thatcher-Lr16/Thatcher-Lr67), 114 lines (Thatcher-Lr34/BW196R), and 113 lines (Thatcher-Lr67/BW196R). DH populations were generated using the maize pollination described by Thomas et al. (1997) except a single dicamba (100 ppm) treatment was used by placing a large drop with a
syringe between the primary and secondary florets (all other rows were removed from each spikelet prior to emasculation) the day after pollination.

**Marker Analysis**

To classify the progeny for the presence or absence of the genes targeted in each population, DNA markers were used to classify Lr34, Lr32, Lr16, and Lr32. The Lr34 locus was genotyped using a PCR marker, calND11, that targets an indel in the Lr34 gene sequence (Dakouri et al., 2010). Both Lr67 and Lr16 were classified based on closely linked SNP markers, csSNP856 (Forrest et al., 2014) and kwm742 (Kassa et al., 2017) respectively. SSR markers wmc43 and barc135 were used to detect the presence of Lr32 (Thomas et al., 2010). PCR products for calND11 and SSR markers for Lr32 were resolved using an ABI 3100 genetic analyzer (Applied Biosystems) as described by Somers et al. (2004). To genotype the SNP markers for Lr67 and Lr16, KASP assays were performed as described by Kassa et al. (2016).

Given that current markers for Lr13 are not tightly linked, Lr13 was classified based on indoor leaf rust assays. For both populations that segregated for Lr13, two plants per line were grown to the adult plant stage in the greenhouse then inoculated with the Lr13 avirulent P. triticina isolate 1–1 BBBD. While these populations also segregated for Lr34 or Lr67, the presence of Lr13 resulted in a clear and highly resistant reaction phenotype (‘1’-infection type as described by McCallum et al., 2021) that was not seen in those lines with Lr34 or Lr67 or susceptible lines which had more susceptible pustule types. Therefore all lines could therefore be scored as having either the resistant or the susceptible allele for Lr13.

**Leaf Rust Field Resistance**

These populations were grown in leaf rust inoculated, irrigated, field nurseries at Morden Manitoba during four years 2012–2015, with two replications per year, except in 2012 in which a single row was planted for each line. Progeny from the populations lines Thatcher-Lr13/Thatcher-Lr34 and Thatcher-Lr13/Thatcher-Lr67 were tested for an additional two field seasons in 2016 and 2017, with two replicates per season. Each line was seeded in approximately 1 m rows. Spreader rows of susceptible wheat were planted at regular intervals to help the epidemic develop and infect the test lines. Spreader rows were inoculated a few times each year with a mixture of urediniospores in Soltrol mineral oil. The inoculum was a mixture of P. triticina virulence phenotypes, representative of those found in western Canada the previous year (McCallum et al., 2018, 2019, 2020, 2021). The lines were assessed for the level of leaf rust infection on the flag leaves using a 0–100% modified Cobb scale (Peterson et al., 1948). They were also assessed for pustule type (R-MR-MS-S) but only the severity data were used for analysis since this was a better measure of the proportion of the flag leaves infected with leaf rust.

Leaf rust severity percentage ratings were converted to proportions for analysis, then back to percentages for presentation. Data from each population were analyzed separately with SAS 9.3 (SAS Institute Inc.) using PROC GLIMMIX (beta distribution) with the presence or absence of the genes, and their interaction, in each progeny line of the population as dependent variables and replication within each year as the random variable. Within each population, the groups of lines with all the possible gene combinations were compared pairwise to each other using LSMEANS.

**RESULTS**

The effects of both Lr13 and Lr34 were significant in the Lr13/Lr34 population, as was the interaction between Lr13 and Lr34 (Table 1). The lines that had both genes had the lowest level of leaf rust severity (17.7%), followed by lines with only Lr34 (23.8%), lines with only Lr13 (84.3%), and lines with neither gene (85.7%) (Table 1; Figure 1). When these four classes of lines were compared against each other, each class was significantly different from the others at the p < 0.01 level, except lines with only Lr13 which were not different from lines with neither gene (Table 2). In the Lr13/Lr67 population, the effect of Lr67 was significant, but not that of Lr13. In contrast to the Lr13/Lr34 population, there was no significant interaction between the two genes. In this population, lines with both genes had a similar level of leaf rust severity (36.2%) compared to lines with only Lr67 (37.0%) and lines that only had Lr13 were similar (80.5%) with lines that had neither gene (79.0%) (Tables 1, 2; Figure 2).

Both Lr16 and Lr34 were significant in reducing leaf rust severity in the Lr16/Lr34 population, and the interaction between the genes was also significant (p < 0.01). Lines with both genes had the lowest leaf rust severity (24.6%), followed by lines with only Lr34 (34.7%), lines with only Lr16 however had a similar level of leaf rust severity (82.2%) as the lines with neither gene (80.3%) (Tables 1, 2; Figure 3). Similarly, in the Lr16/Lr67 population, both Lr16 and Lr67 were significant in reducing leaf rust severity, and their interaction was significant (p < 0.01). Again lines with both genes had the lowest level of severity (38.0%), followed by lines with only Lr67 (50.1%) and lines with only Lr16 were similar (82.0%) to lines with neither gene (82.3%) (Tables 1, 2; Figure 4).

In the Lr32/Lr34 population, both genes significantly reduced leaf rust severity (p<0.01); however, their interaction was not significant. Lines with both genes were very resistant (3.1%), followed by lines with only Lr34 (16.9%) or only Lr32 (38.1%), compared to lines with neither gene (82.6%) (Table 1; Figure 5). However, each class of lines was significantly different from the other classes (Table 2). Similarly for the Lr32/Lr67 population both genes significantly reduced leaf rust severity (p<0.01), their interaction was however significant (p<0.01). Again lines with both genes had the lowest leaf rust severity (2.1%), followed by those with only Lr67 (23.6%), those with only Lr32 (29.7%), and those with neither gene (77.0%) (Table 1; Figure 6). Each class of line was also significantly different from all the other classes (Table 2).

**DISCUSSION**

This study compared how Lr34 and Lr67 interact in combination with other resistance genes in progeny populations. They were
each paired with Lr13, Lr16, or Lr32 in populations segregating for one of those genes and either Lr34 or Lr67. In the populations that segregated for Lr13, the effect of Lr13 was significant in the population with Lr34 but not with the population involving Lr67. There was a significant interaction between Lr13 and Lr34 whereas there was no significant interaction between Lr13 and Lr67. It appears that Lr34 and Lr67 differ in their interaction with Lr13. The lack of interaction with Lr67 may reflect the marginal resistance provided by Lr13 and that fact that most of the virulence phenotypes in Canada are virulent on Lr13. During the years of the field tests the frequency of virulence to Lr13 was close to 100% (McCallum et al., 2018, 2019, 2020, 2021). Both Lr34 and Lr67 were effective in reducing the leaf rust.
severity, but *Lr34* had a larger effect on reducing leaf rust compared to *Lr67* which may reflect a different in their interactive magnitudes. The *Lr13 + Lr34* gene combination appeared to be more effective than either gene alone in this study, and in previous studies (Ezzahiri and Roelfs, 1989; German and Kolmer, 1992; Kloppers and Pretorius, 1997).
The resistance gene *Lr16* was more effective than *Lr13* in reducing the severity of leaf rust. While the frequency of virulence to *Lr16* is very low (near 0%) (McCallum et al., 2021), most isolates have an intermediate response to *Lr16* and the Thatcher-*Lr16* wheat line is fairly susceptible in field trials as the long term leaf rust severity average of the Thatcher isoline with *Lr16* was 65.6% compared with Thatcher at 81.9% (B. McCallum unpublished). Overall *Lr16* had a significant effect on leaf rust in these populations and had significant interactions with both *Lr34* and *Lr67*.
The effect of \( Lr16 \) is mainly in its interaction with either \( Lr34 \) or \( Lr67 \), because the lines with both \( Lr16 \) and either \( Lr34 \) or \( Lr67 \) were significantly more resistant than lines with just \( Lr34 \) or \( Lr67 \), however, the lines with \( Lr16 \) alone were not significantly different from lines with neither gene in this study (Table 2). Both \( Lr34 \) and \( Lr67 \) interacted with \( Lr16 \) to produce an enhanced resistance, even though lines with \( Lr16 \) alone were not significantly different from susceptible lines. This may reflect on the ability of \( Lr16 \) to interact with other resistance genes, as it has been shown previously to do with genes like \( Lr13 \) (Samborski and Dyck, 1982) and both \( Lr34 \) and \( Lr46 \) in Carberry (Bokore et al., 2022).

This effect of enhancement was also seen with the populations involving \( Lr32 \). Alone \( Lr32 \) was very effective in reducing the severity of leaf rust, as were both \( Lr34 \) and \( Lr67 \). The lines with two gene combinations of \( Lr32 + Lr34 \) and \( Lr32 + Lr67 \) were even more resistant than any of these genes alone. Virulence has not been detected in Canada to \( Lr32 \) and the Thatcher-\( Lr32 \) line is moderately resistant in field trials. Both \( Lr34 \) and \( Lr67 \) had the ability to enhance the resistance of \( Lr32 \) when in combination with this resistance gene. However, the interaction was only significant between \( Lr32 \) and \( Lr67 \) (Table 1). The parental line containing \( Lr32 \) in these crosses also contained \( Lr13 \), which segregated in both populations, although its presence or absence was not determined it would have been distributed evenly between the phenotypic classes. The effects of \( Lr34 \) and \( Lr67 \) were stronger in these populations, than in the other populations analyzed, which could reflect the fact that \( Lr13 \) was segregating in these crosses and potentially interacting with the other leaf rust resistance genes.

Overall \( Lr67 \) behaved similarly to \( Lr34 \) in this study. Both genes consistently reduced the level of leaf rust in each population, \( Lr67 \) did not interact with \( Lr13 \), which was only effective on its own in the \( Lr34 \) population, however, both did have a significant interaction with \( Lr16 \). The effect of \( Lr16 \) was significant overall but it appeared to be effective only when in combination with either \( Lr34 \) or \( Lr67 \) to result in a significantly lower level of leaf rust than with either gene alone. Both genes also significantly reduced leaf rust when combined with the effective resistance gene \( Lr32 \).

Interactions between race-specific leaf rust resistance genes and \( Lr34 \) have been analyzed in previous studies. The studies by Kloppers and Pretorius (1997) and German and Kolmer (1992) both showed that lines carrying \( Lr34 \) plus a race-specific gene had lower disease severity than either gene singly. The data presented here show the same trend. The most direct comparison between these three studies is the interaction between \( Lr34 \) and \( Lr13 \) as this combination was present in all of the studies. German and Kolmer (1992) report what appears to be the strongest interaction between \( Lr13 \) and \( Lr34 \). However, there are some key differences between how these studies were conducted. German and Kolmer (1992) selected a single \( F_4 \) line that was homozygous for \( Lr34 \) and \( Lr13 \) and they reportedly selected the most resistant homozygous line for field testing. This was also done for the other gene combinations in their study. Kloppers and Pretorius (1997) analyzed six lines with \( Lr34 \) and \( Lr13 \) and they found varying responses between lines. At the time of their final field rating, the severities of the six lines ranged from 10 to 50%. The interaction would look different if they had only used the most resistant line.
Similarly, in our study, DH lines carrying \( Lr34 \) and \( Lr13 \) had mean severities ranging from 7 to 37%. There was at least some range of severity levels among the progeny lines for all the various gene combinations generated in this study. While Thatcher near-isogenic lines were used primarily as the parental lines in this study, intercrossing these lines resulted in significant variation between lines with the same major gene combinations, similar to the variation found by Kloppers and Pretorius (1997) with the \( Lr13 + Lr34 \) sister lines. Using multiple lines or populations gives a more accurate representation of the interactions between genes as all of the other factors will also be segregating and correlated errors are minimized. In the present study, we also compared DH lines carrying each gene singly for the same reason as when the NILs were developed, the best phenotypes were selected which may not best represent the resistance conferred by the \( Lr \) gene in question.

Both \( Lr34 \) and \( Lr67 \), along with a third multi-pest non-race-specific adult plant resistance gene \( Lr46 \), are important components of resistance in CIMMYT and Mexican wheat cultivars (Singh et al., 2008; Huerta-Espino et al., 2020). Deployed alone these adult plant multi-pest resistance genes did not confer adequate resistance, but combinations of 4–5 genes usually results in near immunity levels of resistance (Singh et al., 2011). Interestingly, \( Lr67 \) was deployed in many Mexican wheat cultivars developed in the 1950s, but not in later cultivars due to chance parental selection in which only \( Lr34 \) was used. Since the donor lines for \( Lr34 \) (RL6058) and \( Lr67 \) (RL6077) showed similar resistance phenotypes, both lines were initially thought to have \( Lr34 \) and were used interchangeably as a source for \( Lr34 \) in the 1950s by the CIMMYT wheat breeding program (Huerta-Espino et al., 2020).

Gene pyramids involving \( Lr34 \) are also common in Canadian wheat cultivars (McCallum et al., 2011; Toth et al., 2018). The highly resistant Canadian cultivar Pasqua contains five resistance genes including \( Lr34 \) (Dyck, 1993). \( Lr34 \) appears to be key to its high level of resistance, as progeny lines derived from Pasqua with the four other resistance genes were fairly susceptible (McCallum and Thomas, 2014). Similarly, the high level of durable resistance in the cultivar Carberry is conditioned by the combination of \( Lr2a, Lr16, Lr23, Lr13, Lr34, \) and \( Lr46 \) (Bokore et al., 2022), in which the key is the interaction of \( Lr34 \) and \( Lr46 \) with the other resistance genes. These multi-pest resistance genes along with others, such as \( Sr2 \), appear to function very well in combinations with other genes to condition effective and durable resistance, often by boosting the effect of other resistance genes (Ellis et al., 2014). Randhawa et al. (2018) found that \( Lr34/Yr18/Sr57 \) interacted with \( Lr68 \) to reduce leaf rust and \( Sr2/Yr30 \) to reduce rust severity to leaf, stem, and stripe rust in a segregating population.

It appears that \( Lr67 \) could play a similar and important role in leaf rust resistance, like \( Lr34 \) or \( Lr46 \), if it was combined with other resistance genes, such as \( Lr16 \) and \( Lr32 \), in which it could interact to result in lower levels of leaf rust severity and improved durability of resistance. However, \( Lr67 \) failed to show the same significant interaction with \( Lr13 \) that \( Lr34 \) demonstrated, and the effect of \( Lr34 \) alone was stronger than that of \( Lr67 \) in each of the pairs of populations. In contrast, the interaction between \( Lr32 \) and \( Lr67 \) was significant whereas that between \( Lr32 \) and \( Lr34 \) was not. While \( Lr67 \) was deployed in many CIMMYT wheat cultivars from the 1950s (Huerta-Espino et al., 2020), it is not deployed in Canadian wheat cultivars to date. If it was deployed in Canada and other countries it could improve the rust resistance and durability of future wheat cultivars.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article, further inquiries can be directed to the corresponding author.

**AUTHOR CONTRIBUTIONS**

BM and CH planned the experiments, analyzed the data, and co-wrote the manuscript. CH developed the populations analyzed in this study and conducted the molecular marker analysis of all progeny lines. BM conducted the leaf rust phenotyping of the populations. All authors contributed to the article and approved the submitted version.

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**REFERENCES**

Bokore, F. E., Knox, R. E., Hiebert, C. W., Cuthbert, R. D., DePauw, R. M., Meyer, B., et al. (2022). A combination of leaf rust resistance genes, including \( Lr34 \) and \( Lr46 \), is the key to the durable resistance of the Canadian wheat cultivar. *Carberry. Front. Plant Sci.* 12:775383. doi: 10.3389/fpls.2021.775383

Dukouri, A., McCallum, B. D., Walichnowski, A. Z., and Cloutier, S. (2010). Fine-mapping of the leaf rust \( Lr34 \) locus in *Triticum aestivum* (L.) and characterization of large germplasm collections support the ABC transporter as essential for gene function. *Theor. Appl. Genet.* 121, 373–384. doi: 10.1007/s00122-010-1316-7

Dyck, P. L. (1993). The inheritance of leaf rust resistance in the wheat cultivar Pasqua. *Can. J. Plant Sci.* 73, 903–906. doi: 10.4141/cjps93-118

Ellis, J. G., Lagudah, E. S., Spielmeyer, W., and Dodds, P. N. (2014). The past, present and future of breeding rust resistant wheat. *Fron Pl Sci.* 5:641. doi: 10.3389/fpls.2014.00641

Ezzahiri, B., and Roelfs, A. P. (1989). Inheritance and expression of adult plant resistance to leaf rust in era wheat. *Plant Dis.* 73, 549–551. doi: 10.1094/PD-73-0549

Forrest, K., Pujol, V., Bulli, P., Pumphrey, M., Wellings, C., Herrera-Foessel, S., et al. (2014). Development of a SNP marker assay for the \( Lr67 \) gene of wheat using a genotyping by sequencing approach. *Mol. Breeding* 34, 2109–2118. doi: 10.1007/s11032-014-0166-4
McCallum, B. D., Seto-Goh, P., Foster, A., Rosa, S., and Xue, A. (2014). Lr34 Is the Key to the Durable Leaf Rust Resistance in the Canadian Cultivar Pasqua. Borlaug Global Rust Initiative Technical Workshop. Obregon, Mexico.

McCallum, B. D., Reimer, E., McNabb, W., Foster, A., and Xue, A. (2011). Physiologic specialization of Puccinia triticina, the causal agent of wheat leaf rust, in Canada in 2015-2019. Can. J. Plant Pathol. 43, S333–S346. doi: 10.1007/s10323-015-0218-6

McCallum, B. D., Reimer, E., McNabb, W., Foster, A., and Xue, A. (2020). Physiologic specialization of Puccinia triticina, the causal agent of wheat leaf rust, in Canada in 2014. Can. J. Plant Pathol. 42, 520–526. doi: 10.1007/s10323-020-1273-0

McCallum, B. D., Seto-Goh, P., Foster, A., and Xue, A. (2018). Physiologic specialization of Puccinia triticina, the causal agent of wheat leaf rust, in Canada in 2012. Can. J. Plant Pathol. 40, 434–441. doi: 10.1007/s10323-018-0195-2

Moore, J. W., Herrera-Foessel, S., Lan, C., Schnippenkoetter, W., Ayliffe, M., Huerta-Espino, J., et al. (2015). A recently evolved hexose transporter variant confers resistance to multiple pathogens in wheat. Nat. Genet. 47, 1494–1498. doi: 10.1038/ng.3439

Peterson, R. E., Campbell, A. B., and Hannah, A. E. (1948). A diagrammatic scale for estimating rust intensity on leaves and stems of cereals. Can. J. Res. 26, 496–500. doi: 10.1139/cjr48-033

Randhawa, M. S., Lan, C., Basnet, B. R., Bhavani, S., Huerta-Espino, J., Forrest, K. L., et al. (2018). Interactions among genes Sr2/Yr30, Lr34/Yr18/ Sr57 and Lr68 confer enhanced adult plant resistance to rust diseases in common wheat (Triticum aestivum L.) line ‘Arul’. Aust. J. Crop. Sci. 12, 1023–1033. doi: 10.21475/aics.18.12.06.PNE105

Samborski, D. I., and Dyck, P. L. (1982). Enhancement of resistance to Puccinia recondita by interactions of resistance genes in wheat. Can. J. Plant Pathol. 4, 152–156. doi: 10.1002/0470752708.ch22

Singh, R. P. (1992). Genetic association of leaf rust resistance gene Lr34 with adult plant resistance to stripe rust in bread wheat. Phytopathology, 82, 835–838. doi: 10.1094/Phyto-82-835

Singh, R. P. (1993). Genetic association of gene Bdv1 for tolerance to barley yellow dwarf virus with genes Lr34 and Yr18 for adult plant resistance to rusts in bread wheat. Plant Dis. 77, 1103–1106. doi: 10.1094/PD-77-1103

Singh, R. P., Huerta-Espino, J., Bhavani, S., Herrera-Foessel, S. A., Singh, D., Singh, P. K., et al. (2011). Race non-specific resistance to rust diseases in CIMMYT spring wheats. Euphytica 179, 173–186. doi: 10.1007/s10681-010-9932-9

Singh, R. P., Huerta-Espino, J., and William, M. (2008). “Breeding for resistance to biotic stresses,” in Plant Breeding: The Arnel R. Hallauer International Symposium. eds. R. Kendall and M. L. Lamkey (Hoboken: Blackwell Publishing), 310–322. doi: 10.1002/9780470752708.ch22

Somers, D. J., Isaac, S., and Edwards, K. (2004). A high density microsatellite consensus map for bread wheat (Triticum aestivum L.). Theor. Appl. Genet. 109, 1105–1114. doi: 10.1007/s00122-004-1740-7

Spielmeyer, W., Mago, R., Wellings, C., and Ayliffe, M. (2013). Lr67 and Lr34 rust resistance genes have much in common - they confer broad spectrum resistance to multiple pathogens in wheat. BMC Plant Biol. 13:96.

Spielmeyer, W., McIntosh, R. A., Kolmer, J. A., and Lagudah, E. S. (2005). Powdery mildew resistance and Lr34/Yr18 genes for durable resistance to leaf and stripe rust co segregate at a locus on the short arm of chromosome 7D of wheat. Theor. Appl. Genet. 111, 731–735. doi: 10.1007/s00122-005-2058-9

Thomas, J., Chen, Q., and Howes, N. (1997). Chromosome doubling of haploids of resistant CIMMYT spring wheats. Euphytica 173, 281–291. doi: 10.1007/s10681-001-0013-5

Thomas, J., Nilimogoda, S., Hiebert, C., McCallum, B., Humphreys, G., and DePauw, R. (2010). Genetic markers and leaf rust resistance of the wheat gene Lr32. Crop Sci. 50, 2310–2317. doi: 10.2136/csc2010.02.0065

Toth, J., Pandurangan, S., Burt, A., Mitchell Fetch, J., and Kumar, S. (2018). Marker-assisted breeding of hexaploid spring wheat in the Canadian prairies. Can. J. Plant Sci. 99, 111–127. doi: 10.1139/cjps-2018-0183

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