AB-QTL analysis in winter wheat: II. Genetic analysis of seedling and field resistance against leaf rust in a wheat advanced backcross population

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Abstract The present study aimed to localize exotic quantitative trait locus (QTL) alleles for the improvement of leaf rust (P. triticina) resistance in an advanced backcross (AB) population, B22, which is derived from a cross between the winter wheat cultivar Batis (Triticum aestivum) and the synthetic wheat accession Syn022L. The latter was developed from hybridization of T. turgidum ssp. dicoccoides and T. tauschii. Altogether, 250 BC2F3 lines of B22 were assessed for seedling resistance against the leaf rust isolate 77WxR under controlled conditions. In addition, field resistance against leaf rust was evaluated by assessing symptom severity under natural infestation across multiple environments. Simultaneously, population B22 was genotyped with a total of 97 SSR markers, distributed over the wheat A, B and D genomes. The phenotype and genotype data were subjected to QTL analysis by applying a 3-factorial mixed model analysis of variance including the marker genotype as a fixed effect and the environments, the lines and the marker by environment interactions as random effects. The QTL analysis revealed six putative QTLs for seedling resistance and seven for field resistance. For seedling resistance, the effects of exotic QTL alleles improved resistance at all detected loci. The maximum decrease of disease symptoms (−46.3%) was associated with marker locus Xbarc149 on chromosome 1D. For field resistance, two loci had stable main effects across environments and five loci exhibited marker by environment interaction effects. The strongest effects were detected at marker locus Xbarc149 on chromosome 1D, at which the exotic allele decreased seedling symptoms by 46.3% and field symptoms by 43.6%, respectively. Some of the detected QTLs co-localized with known resistance genes, while others appear to be as novel resistance loci. Our findings indicate, that the exotic wheat accession Syn022L may be useful for the improvement of leaf rust resistance in cultivated wheat.

Introduction

Wheat leaf rust, caused by the obligate parasitic fungus Puccinia triticina Eriks (syn. P. recondita Roberg ex Desmaz. f. sp. tritici), is considered to be the most widely distributed wheat disease in the world (Wiese 1987; Gupta et al. 2006). The pathogen interferes with wheat plants at all developmental stages and serious yield losses of up to 62.7% have been reported in susceptible cultivars (Singh et al. 1998). Fungicides and different cultural practices can be used to reduce yield losses. However, the development
of resistant cultivars is the most efficient, sustainable and economic method to prevent leaf rust epidemics.

Research on leaf rust resistance in wheat has mainly concentrated on race-specific resistance that is conferred by genes with major effects and recognized by characteristic low-infection types. A battery of resistance genes (R-genes), some of which are derived from rye and from wild relatives of wheat, is known and has been catalogued by McIntosh et al. (1995, 2003). Of these R-genes, \( Lr21 \), \( Lr10 \) and \( Lr1 \) have been cloned (Huang et al. 2003a; Feuillet et al. 2003; Cloutier et al. 2007). Resistance contributed by R-genes typically remains effective for only a few years because of its dependency on the avirulence of one or a few genotypes of the relevant pathogen. Therefore, it has been suggested that the available resistance genes in hexaploid cultivated wheat have been nearly exhausted (Bai and Knott 1994). On the other hand, adult plant resistance has the potential to provide durable disease control. This form of resistance is controlled either by genes with major effects in terms of hypersensitive reaction like \( Lr12 \) or by slow rusting or partial resistances genes like \( Lr34 \). The parameters that cause slow rusting of a cultivar are longer latent period, low receptivity, as well as small uredial size and reduced quantity of spore production (Drijver and Pretorius 1989). To date, a number of adult plant resistance genes have been reported, of which \( Lr34 \) and \( Lr46 \) are the most important slow rusting genes (German and Kolmer 1992; Singh et al. 1998; Spielmeyer et al. 2005). \( Lr34 \) has been used in different breeding programs in the world because of its significance in conferring durable resistance in combinations with other genes (Dyck 1991; Singh 1992; Rosewarne et al. 2006). Currently, there is an urgent need to understand the genetic basis and quantification of such adult plant resistances as well as a quest for additional genes that could contribute to durable resistance.

Resistance genes introduced from wild species into elite cultivars have usually been genetically mapped in balanced populations of progeny derived from an early generation (e.g., \( F_2 \) plants or families or \( F_1 \)-derived doubled haploids). Such populations contain an equal proportion of exotic and elite genotypes, and deleterious effects of exotic alleles may mask the desired target gene effect. The advanced backcross quantitative trait locus (AB-QTL) strategy, developed by Tanksley and Nelson (1996), has been proposed for introgression of exotic QTL alleles from a donor accession into elite germplasm. The method proved effective in detecting additive, dominant, partially dominant and overdominant QTLs. Repeated backcrossing with the elite parent decreases the number and size of the exotic introgressions, which in turn reduces the burden of linkage drag from deleterious exotic QTL alleles. Near isogenic lines can be developed readily and used for QTL verification and QTL cloning (Frary et al. 2000). AB-QTL analyses have been successfully executed to locate favorable exotic QTL alleles that could improve agronomic traits in wheat (Huang et al. 2003b, 2004; Narasimhamoorthy et al. 2006; Liu et al. 2006; Kunert et al. 2007). Leonova et al. (2007) applied the AB-QTL strategy to improve leaf rust resistance in wheat. Using AB progeny derived from six exoticintrogressions originating from \( T. timopheevii \) and \( T. tauschii \), they detected exotic QTL alleles conferring resistance in seedlings and/or in field plots on chromosomes 2B, 2D and 6B. The strongest effect, both for seedlings and in the field was localized to a QTL on chromosome 2B that explained more than 49% of the phenotypic variation.

In our study, we performed an AB-QTL analysis in order to identify QTLs associated with seedling as well as field resistance against \( P. triticina \) in the \( BC_2F_3 \) population B22, which is derived from a cross between the elite winter wheat cultivar Batis and the exotic wheat accession Syn022L.

**Materials and methods**

**Development of the mapping population B22**

The winter wheat cultivar Batis was crossed with the synthetic wheat accession Syn022L and two backcrosses were made to Batis, using the \( F_1 \) and \( BC_1F_1 \) plants as the maternal parents. As described by Kunert et al. (2007), the resulting \( BC_2F_1 \) plants were self-pollinated, and single seed descent was used in order to obtain 250 \( BC_2F_3 \) plants. Progeny of each \( BC_2F_3 \) plant were propagated in bulk until \( BC_2F_3 \) to increase the number of seeds for multi-environment testing. The recurrent parent Batis is an elite cultivar from Fr. Strube Saatzucht KG originating from the cross Bert \( \times [(\text{Maris Huntsman} \times \text{Glaucus}) \times \text{Urban}] \). The primary synthetic wheat parent Syn022L was kindly provided by Professor W. Lange from the Centre for Plant Breeding and Reproduction Research (CPR-DLO), Agriculture University Wageningen, The Netherlands. Syn022L was developed by hybridization of \( T. turgidum \) ssp. \( dicoccoides \) (4n, accession G168-1-2-4BM) and \( T. tauschii \) (2n, accession Cam L, see Lange and Jochemsen 1992a, b) and serves as the donor of wild alleles for the wheat A, B and D genomes.

**Evaluation of seedling resistance against 77WxR**

The \( BC_2F_{3,5} \) lines of population B22 were sown in 96-cell trays in two randomized complete blocks with individual lines replicated three times within each block. Both parents were included and replicated five times per block as controls. Initially, the susceptible cultivar Borenos was...
inoculated with *P. triticina* pathotype 77WxR with the help of a mechanical duster. Later, the fresh spores from the pre-infected seedlings of Borenos were trickled onto two-week-old seedlings of population B22. The infected plants were kept for 24 h in the dark at 20°C and 80% relative humidity to allow germination and penetration of spores, and then transferred to a growth chamber with 16 h light per day at 25°C and 80% relative humidity. After 10 days, the seedlings were screened for leaf rust symptoms using a scale from 0 to 4. The scale values are interpreted as follows: 0: no uredia or hypersensitive flecks without uredia, 1: small uredia surrounded by necrosis, 2: small uredia surrounded by chlorosis, 3: uredia of moderate size that might associate with chlorosis, 4: large uredia without chlorosis (modified after Long and Kolmer 1989). The virulence and avirulence pattern of the isolate is listed in Table 2. Inoculum, space and technical assistance were kindly provided by the Federal Centre for Breeding Research on Cultivated Plants (BAZ) Quedlinburg, Germany.

**Evaluation of field resistance**

The evaluation of field resistance was carried out under natural occurrence of leaf rust in seasons 2004 (04) and 2005 (05) at five different environments in Germany. The AB lines were sown in a randomized complete block design at the experimental stations Dikopshof, University of Bonn (D04 and D05), Rosenthal, Limagrain-Nickerson GmbH (R05) and Söllingen, Fr. Strube Saatzucht KG (S04 and S05). The parent Batis was included in the experiment with eight replications per environment. The synthetic parent Syn022L was not sown in the field due to lack of sufficient seed for multi-environment trials, but it had previously shown a resistant response against the natural occurrence of leaf rust in the field (data not shown). Each AB line and the control were sown with 310–360 kernels/occurrence of leaf rust in the field (data not shown). Each parent Syn022L was not sown in the field due to lack of sufficient seed for multi-environment trials, but it had previously shown a resistant response against the natural occurrence of leaf rust in seasons 2004 (04) and 2005 (05) at five different environments in Germany. The synthetic parent Batis was included in the experiment with eight replications per environment.

The genetic constitution of the original BC2F3 plants was analyzed with simple sequence repeat (SSR) markers. Almost 200 published SSRs from different sources (BARC, Song et al. 2005; CFA and CFD, Guyomarc’h et al. 2002; http://www.wheat.pw.usda.gov, GDM, Pestsova et al. 2000; GWM, Röder et al. 1998; WMC, Gupta et al. 2002) were tested and their chromosomal locations were assigned according to the consensus genetic map of Somers et al. (2004). Polymerase chain reaction (PCR) was used to amplify the SSR markers with the tailed primer method, where the forward primer was extended at the 5′-end by the M13 universal forward sequence. The PCR polymerization was performed in 20 µl final volume reactions containing 5 µl of the template DNA (ca. 20 ng/µl), 0.5 µl of *Taq* polymerase (5 units/µl), 0.75 µl of dNTP (2 mM), 0.5 µl of 25 mM MgCl2, 2 µl of 10× PCR buffer (500 mM KCl, 100 mM Tris–HCl, pH 9.0, 1.0% Triton X-100), 0.2 µl of each forward and reverse primer (10 µM) and 0.5 µl (1 µM) of the M13 universal forward primer. The latter primer was labeled at the 5′-end with either IRD700 or IRD800 for visualization. The amplified DNA fragments were separated electrophoretically using the Li-Cor DNA sequencer 4200, Li-Cor, Bad Homburg.

**QTL analysis**

Following von Korff et al. (2005), a marker analysis was executed in the general linear model (GLM) procedure of SAS version 9.1 (SAS Institute 2004) by using a 3-factorial analysis of variance (ANOVA) with a mixed hierarchical model in order to detect QTLs associated with seedling and field resistance. The two models applied are:

\[
y_{ijklm} = \mu + L_j(M_i) + B_k + M_i \cdot B_k + e_{ijkl} \tag{1}
\]

for seedling resistance and

\[
y_{ijklm} = \mu + M_i + L_j(M_i) + E_k + M_i \cdot E_k + e_{ijkl} \tag{2}
\]

for field resistance, where *µ* is the general mean, *M* is the fixed effect of the *i*-th marker genotype, *L* is the random effect of the *j*-th BC2F3 line nested in the *i*-th marker genotype, *E* is the random effect of the *k*-th environment, *M* × *E* is the random interaction effect of the *i*-th marker and *k*-th environment and *e* is the error of *Y*. In model [1], the random factor block (*B*) was used instead of *E* in order to test the reproducibility of a marker effect between blocks.

For field resistance, marker main effects (*M*) and marker by environment interactions (*M* × *E*) with *P* < 0.01 were accepted as putative QTLs. In contrast, for seedling resistance only marker main effects with *P* < 0.01 were accepted as putative QTLs, because marker by block interactions were considered as an indication of a lack of reproducibility. Markers detecting similar effects were considered to represent a single QTL if they were within 20 cM of each other (Pillen et al. 2003).

The proportion of genetic variance (*R*²) explained by marker main effects or marker by environment interaction effects was calculated as described by von Korff et al. (2005). The relative performance of the homozygous exotic genotype RP[exot] as a measure of trait improvement due
to the substitution of the elite alleles (Batis) by the exotic alleles (Syn022L) was calculated according to the formula:

$$RP_{\text{exot}} = \frac{[aa] - [AA]}{AA} \times 100.$$  

For each marker locus, [aa] and [AA] represent the least square means of disease scores of the homozygous exotic and the homozygous elite genotype, respectively, calculated across all environments and blocks, respectively.

**Results**

**Seedling resistance**

The seedling infection scores of population B22 and both parents, averaged across blocks, are displayed in Fig. 1. The donor parent Syn022L exhibited a completely resistant response in the form of a hypersensitive reaction, whereas the recurrent parent Batis displayed a highly susceptible phenotype against isolate 77WxR. In population B22, the disease scores ranged from 0 (highly resistant) to 3.5 (highly susceptible) with a mean value of 2.5. As expected, the standard deviation was higher in population B22 than in the two controls, Batis and Syn022L.

**Field resistance**

Leaf rust symptoms in the field were scored in five environments (Table 1). The disease severity varied considerably across the environments and the mean disease scores of population B22 ranged from 1.7 at S04 to 4.4 at D05. An almost similar trend between the environments was observed for the recurrent parent Batis. Significantly different responses between population B22 and Batis were recorded at S04 and S05. In both cases, B22 showed higher mean disease scores than Batis. The standard deviation in all environments was higher in population B22 than in Batis.

**QTL detection**

The QTLs for seedling and field resistance that were identified by applying a 3-factorial mixed model ANOVA are summarized in Table 3 and their chromosomal location is drawn in Fig. 2.

**Leaf rust seedling resistance**

Six QTLs were detected for seedling resistance against the leaf rust isolate 77WxR. At each of these, the exotic allele improved the resistance and its effect was stable across both blocks. The QTLs were located on the A genome (chromosome 6A), the B genome (chromosomes 1B and 4B) and the D genome (chromosomes 1D, 4D and 6D). The strongest effect of an exotic allele was detected at locus Xbarc149 on chromosome 1D where the disease score decreased by 46.3%. Here, 49.0% of the genetic variation was explained by the marker main effect. At QTLs QLrs.B22-6A and QLrs.B22-4B, the exotic alleles reduced disease severity by 28.9 and 27.3%, respectively. On chromosome 1B, favorable exotic alleles were detected at marker loci Xbarc8 and Xgwm11. The latter was associated with a 24.7% reduction in disease severity. Due to their close linkage, they were interpreted as a single QTL, QLrs.B22-1B. An almost 21% reduction in disease severity was associated with an exotic allele at QLrs.B22-6D. A minor QTL effect was detected on chromosome 4D, where the exotic allele accounted for 15.0% decrease of leaf rust symptoms.

**Leaf rust resistance in the field**

Seven QTLs were detected for field resistance to leaf rust under natural infection across five environments. These QTLs were identified on the A, B and D genomes and appeared as marker main effects at two loci and as marker by environment \((M \times E)\) interaction effects at five loci. As for seedling resistance, the strongest QTL effect was detected at locus Xbarc149 (QLr.B22-1D). Here, the exotic allele was associated with a reduction of leaf rust symptoms by 43.6%, explaining 14.6% of the genetic variation for field resistance. At this QTL, and at QLr.B22-7D, at which the exotic allele reduced symptoms by 20.9%, the effect of the exotic allele was favorable in all environments but fluctuated in magnitude across the environments (Fig. 3), resulting in significant \(M \times E\) interaction effects. At three QTLs, QLr.B22-1A, QLr.B22-2D and QLr.B22-6D, the effects of the exotic alleles were favorable in some environments but unfavorable in other environments.
At two other QTLs, the exotic alleles had consistently unfavorable effects: QLr.B22-1B and QLr.B22-3D, at which the exotic alleles increased disease severity by 27.2 and 18.1%, respectively.

**Table 1** Least square means (Lsmeans) and standard deviations of field resistance scores against leaf rust for population B22 and the recurrent parent Batis across five environments

| Environment | Batis  | Population B22 |
|-------------|-------|----------------|
|             | Lsmeansa | SDb | Lsmeansa | SDb | Diff | Minimum | Maximum |
| D04         | 1.9c   | 0.4  | 1.9c     | 1.1  | ns   | 1       | 9       |
| D05         | 4.9a   | 1.4  | 4.4a     | 2.2  | ns   | 1       | 9       |
| R05         | 2.5b   | 0.5  | 3.1b     | 1.3  | ns   | 1       | 9       |
| S04         | 1.2d   | 0.5  | 1.7c     | 1.0  | *    | 1       | 8       |
| S05         | 2.9b   | 0.4  | 4.3a     | 1.7  | *    | 1       | 9       |
| Average     | 2.4    | 0.6  | 2.9      | 1.5  | ns   |         |         |

* ns not significant

* Calculated by a Tukey–Kramer test with $P < 0.05$

a Significant differences between environments are indicated by different letters at $P < 0.05$ after Student–Newman–Keuls (SNK) grouping implemented in the general linear model (GLM) procedure of SAS (SAS Institute 2004)

b Standard deviation

c Significant differences between Batis and B22

(Fig. 3). At two other QTLs, the exotic alleles had consistently unfavorable effects: QLr.B22-1B and QLr.B22-3D, at which the exotic alleles increased disease severity by 27.2 and 18.1%, respectively.

**Discussion**

An AB-QTL strategy, employing the AB generation BC$_2$F$_3$, was applied in order to identify exotic QTL alleles from the synthetic wheat accession Syn022L that could contribute to resistance against leaf rust.

Assignment of QTLs for seedling and field resistance and their comparison to known resistance genes and QTLs

In population B22, QTLs for resistance against leaf rust were identified in 11 chromosome regions on wheat genomes A, B and D (see Table 3). In six of these regions, the exotic alleles consistently showed superiority over the elite genotype, indicating their potential use for wheat leaf rust resistance breeding.

The strongest effect for both leaf rust seedling and field resistance was present on chromosome 1D at marker locus Xbarc149 (QLrs.B22-1D and QLr.B22-1D). Here the exotic allele was associated with 46.3 and 43.6% reduction in leaf rust symptoms, respectively. A number of leaf rust resistance genes had already been mapped to chromosome 1D. McIntosh (1998) reported genes which are effective against leaf rust on the short arms of wheat group 1 chromosomes. In addition, Cox et al. (1994) described three leaf rust resistance genes, Lr41, Lr42 and Lr43, on chromosome 1D, which were transferred from *T. tauschii* to common wheat. The position of Lr21, a member of the NBS-LRR resistance gene family (Huang et al. 2003a), on chromosome arm 1DS (Jones et al. 1990) agrees with the chromosomal location of the 1D QTL in population B22. For Lr21, the USDA Cereal Disease Laboratory (http://www.ars.usda.gov/Main/docs.htm?docid=10342) recorded a low infection score (0–2) at seedling stage and immunity at adult plant stage. When we inoculated the Thatcher nearly isogenic line containing Lr21 with our leaf rust isolate 77WxR, the reaction was scored as virulent (see Table 2). The fact that our QTL on chromosome 1D is active both in seedling and in field experiments can be interpreted in two ways. Either a new allele of Lr21 which has different specificities than that present in Thatcher-Lr21 might be active in population B22 or a new resistance gene, independent of Lr21, is located on chromosome arm 1DS. In future, the genotyping of the Lr21 sequence in population B22 will be used to validate if this gene could explain the QTL effect in B22. Locus Xbarc149 is the only case in population B22 where the exotic allele consistently contributed to resistance both at the seedling stage and in the field experiment. Leonova et al. (2007) also reported a leaf rust resistance gene with effects at the seedling stage and in a field experiment. That gene was transferred from *T. timopheevii* to wheat chromosome 2B.

Some other QTLs located in population B22 potentially correspond to previously mapped resistance genes or QTLs (see Table 3). For seedling resistance, QLrs.B22-1B might correspond to Lr26 which was transferred from the Imperial rye (Secale cereale) chromosome 1R to wheat chromosome 1B (Singh et al. 1990) and QLrs.B22-6D may correspond to a QTL which was located on chromosome 6D by Nelson et al. (1997). QLrs.B22-4B is on the same chromosome as Lr31 (Singh et al. 1999) but Lr31 is
functional only in conjunction with \( Lr27 \) on chromosome 3B. The fact that no QTL in B22 has been identified on chromosome 3B might indicate that QLrs.B22-4B does not correspond to \( Lr31 \) and thus could be a unique resistance source. For field resistance, QTLs mapped in B22 may correspond to \( Lr34 \) on 7DS (Spielmeyer et al. 2005) and a QTL mapped on 6D by Nelson et al. (1997). Singh et al. (1993) reported that the \( Lr34 \) gene confers a quantitative resistance to leaf rust and Schnurbusch et al. (2004) found a similar QTL, which confers a durable slow rusting phenotype in a population derived from the cross Arina × Forno. Syn022L may contribute a further resistance allele at the \( Lr34 \) locus.

In population B22, two main QTLs were mapped at which the exotic alleles on chromosomes 1B and 3D increased leaf rust symptoms in the field. It can, thus, be concluded that the recipient parent of B22, the wheat cultivar Batis, possesses at least two resistance genes. Winzeler et al. (2000) reported that the adult plant resistance gene \( Lr13 \) and so far unidentified resistance sources are present in Batis. However, the effect of \( Lr13 \), mapped on chromosome arm 2BS, could not be verified.

![Chromosomal map of population B22 showing 13 putative QTLs for leaf rust seedling and field resistance. The putative QTLs are presented at the right to the associated marker where the highest \( F \)-value has been calculated. The QTLs are designated according to Table 3.](image-url)
in B22, either because the \textit{P. triticina} races that differentiate \textit{Lr13} were not present at the environments tested or because the Batis gene at \textit{Lr13} is matched by a similar resistance gene at this locus present in the exotic donor Syn022L.

Marker by environment interaction effects

Of the seven putative QTLs for field resistance, five (71\%) showed an $M \times E$ interaction effect in this study (see Table 3 and Fig. 3). This is in contrast to the findings of Keller et al. (1999), who identified three of 18 QTLs showing QTL $\times$ environment interactions while studying powdery mildew resistance across five environments. An obvious reason for the higher number of $M \times E$ interactions in our study may be the fact that the selected environments were located far from each other representing different climatic conditions, altitudes and types of soil in contrary to the study of Keller et al. (1999) who investigated powdery mildew in five environments located close to each other near Zürich, at the same altitude of 450–550 m above the sea level on loamy soils. Moreover, they

### Table 2

| Lr-gene | Infection type$^a$ | +/−  | Lr-gene | Infection type$^a$ | +/−  | Lr-gene | Infection type$^a$ | +/−  |
|---------|--------------------|------|---------|--------------------|------|---------|--------------------|------|
| 1       | 3                  | +    | 15      | 3                  | +    | 30      | 2                  | +    |
| 2a      | 3                  | +    | 16      | 3                  | +    | 32      | 3                  | +    |
| 2b      | 3                  | +    | 17a     | 3                  | +    | 33      | 3                  | +    |
| 2c      | 3                  | +    | 18      | 3                  | +    | 35      | 3                  | +    |
| 3       | 0                  | −    | 19      | 0                  | −    | 37      | 3                  | +    |
| 3bg     | 0                  | −    | 20      | 3                  | +    | 38      | 0                  | −    |
| 3ka     | 0                  | −    | 21      | 2                  | +    | 44      | 2                  | +    |
| 9       | 0                  | −    | 23      | 3                  | +    | 52      | 3                  | +    |
| 10      | 2                  | +    | 24      | 0                  | −    | B       | 3                  | +    |
| 11      | 3                  | +    | 25      | 0                  | −    | Controls; |                    |      |
| 13      | 3                  | +    | 26      | 0                  | −    | Thatcher | 3                  | +    |
| 14a     | 2                  | +    | 28      | 3                  | +    | Batis    | 3                  | +    |
| 14b     | 3                  | +    | 29      | 0                  | −    | Syn022L  | 0                  | −    |

Data source: Dr. V. Lind, Federal Center for Breeding Research on Cultivated Plants (BAZ), Quedlinburg, Germany

$^a$ Seedling infection was conducted and scored as explained in “Materials and methods”

$^b$ Virulence (+)/avirulence (−) pattern of isolate 77WxR
reported a medium disease pressure in their studies in both natural and artificial inoculations and they used growth regulators to further decrease the sampling and testing errors. In this study, the field experiments were grown under natural infestation and without the use of growth regulators. 

M9E interactions could be higher in our study because of the diversity of pathogens present in the environments investigated. Furthermore, disease symptoms depend primarily on the environmental condition, i.e., temperature, humidity, light, etc., which presumably were not constant across all environments tested. However, the differentiation of QTLs into stable and environment-dependent QTLs has important implications for marker-assisted selection. In future, the stable QTLs could be used for marker-assisted selection across all environments while the environment-dependent QTLs may be restricted to specific environments.

**Outlook**

The present study was designed to identify exotic QTL alleles which affect leaf rust seedling and field resistance in the BC2F3 population B22. Altogether, QTLs were detected in 11 chromosome regions. At most loci, the exotic alleles improved resistance. QTLs, at marker Xbarc149 on chromosome 1D, QLrs.B22-1D and QLr.B22-1D, explained high proportions of the genetic variance for seedling resistance (49.0%) and field resistance (14.6%). In future, BC2F3 lines of population B22 which possess exotic resistance QTL alleles can be utilized to generate QTL-
bearing nearly-isogenic lines (NILs) via backcrossing with Batis and marker-assisted selection. These NILs can be used to verify the QTL effects in new field experiments. If verified, the NILs can be directly used both for development of improved varieties and for map-based cloning of the QTL-underlying resistance genes.

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