Fusarium head blight (FHB), primarily caused by *Fusarium graminearum* and *F. culmorum* is one of the major diseases of wheat worldwide, resulting in significant loss in terms of both grain yield and quality (Bai & Shaner 1994; Parry et al. 1995; Waldron et al. 1999). The disease is favoured by warm humid conditions during anthesis and the early stages of kernel development (Gilbert & Tekauz 2000). FHB can cause bleached spikes, spikelet sterility, poor seed filling, low-test weight and tombstone seeds, and in epidemic years the yield losses can range from 10 to 40% (Wang 1996). Furthermore, the grain is often contaminated with mycotoxins such as deoxynivalenol (DON) which is a significant health risk for humans and animals (Gilbert & Tekauz 2000). FHB also affects the milling and baking quality of wheat (Bechtel et al. 1985).

FHB resistance is a quantitatively inherited trait controlled, in many spring wheat varieties by a few genes of major effect plus several genes of more minor effect (Liu et al. 2005). Although fungicides and good agronomic practices can limit damage, the use of genetic resistance is considered to be a key component in the management of FHB (Bai & Shaner 1994). The accepted model of resistance to FHB in wheat is resistance to initial infection (type I) and resistance to spread within the spikelet (type II) (Schroeder & Christensen 1963), although additional components of FHB resistance have been proposed (Bai & Shaner 1994; Yang 1994; Mcmullen et al. 1997). To date, immunity to FHB has not been identified and there is still a relative paucity of potent resistance available to plant breeders. Resistance from the Chinese spring
wheat Sumai 3 (Mesterhazy 1983; Chen et al. 1997; Buerstmayr et al. 2002) and its derivatives has been deployed in breeding programmes for several years, and, as a result, this source tends to predominate worldwide (Del Blanco et al. 2003). Use of a limited number of host resistance gene sources imposes a strong selective pressure for virulent pathogen strains (Gervais et al. 2003) and consequently there is a need to identify additional sources of FHB resistance.

Phenotyping for FHB resistance is costly and time-consuming and the expression of FHB resistance is greatly influenced by environmental factors such as temperature, humidity and crop stage at the time of inoculation (Bai & Shaner 1994; Parry et al. 1995; Miedaner et al. 2001; Klahr et al. 2007). In addition, previous authors have reported that morphological characters including plant height (PH), awns and spike compactness (Steiner et al. 2004) are linked to, or have a pleiotropic effect on FHB resistance in wheat. Such an association with physical characteristics further complicates efforts to understand the physiological basis of FHB resistance.

The Brazilian spring wheat Frontana has the pedigree Fronteira/Mentana and is known to have moderate type I and type II resistance (Singh et al. 1995; Buerstmayr et al. 1996; Van Ginkel et al. 1996; Steiner et al. 2004). In addition to this, in vitro experiments suggest that Frontana may be able to degrade and tolerate higher levels of DON (Miller & Arnison 1986; Wang & Miller 1988). Previous mapping studies indicate that resistance in Frontana is conditioned by several QTL of minor effect (Steiner et al. 2004; Mardi et al. 2006). RL4137 is a Canadian spring wheat derived from Frontana and was found to be highly resistant to FHB in preliminary trials at the John Innes Centre. A mapping study was undertaken to identify and characterise FHB resistance in recombinant inbred lines (RILs) derived from a cross between RL4137 and the moderately resistant variety Timgalen and to study the relationship between FHB resistance traits (disease symptoms and weight of infected spikelets) and some developmental and morphological traits (plant height and measurements of awns).

MATERIALS AND METHODS

Mapping population, markers and genotyping

Ninety F₅ recombinant inbred lines (RILs) from a cross between the FHB resistant variety RL4137 (Frontana/3/McMurachy/Exchange//2*Redman/4/Thatcher*6/Kenya Farmer) and the moderately resistant variety Timgalen (Aguillera/Kenya//Marroqui/3/Supremo/4/Gabo/5/Winglen), were examined in this study along with the parental lines. The population was originally developed at the John Innes Centre and used to study pre-harvest sprouting resistance (Bassoi & Flintham 2005).

Phenotyping

The RILs and the parents, RL4137 and Timgalen, were phenotyped in two polytunnel and field experiments. The polytunnel experiments were conducted at the John Innes Centre (JIC), Norwich, UK in 1999 (J1999) and 2000 (J2000). The field trials were conducted in 2006 at the National Institute of Agricultural Botany (NIAB), Cambridge (N2006) and Central Science Laboratory (CSL), York, UK (C2006). All the lines were tested in J2000, C2006 and N2006 experiments but, due to a shortage of seed, only half of the population was tested in J1999. In both field and polytunnel experiments, the lines were grouped according to flowering time (early, medium and late) and used in the study. Twelve seeds from each recombinant inbred line and parents RL4137 and Timgalen were germinated in Petri-dishes for 48 h at 4°C in total darkness in 10 ml of distilled water. After germination, they were planted in pots in special John Innes cereal mix (3 barrows of loam, 1 barrow of grit, 1 bale of medium peat, 5 ½ lb. Oscomote, 5 lb. Chalk and 1 bushel of tap water.) adjusted to pH 8.0, with one seed per pot and grown in a polytunnel (National Polytunnels, Preston, UK) allowing protection from the rain, but not from the wind, atmospheric humidity or relative sun warmth. Pots containing individual plants were arranged in a randomised complete block design with three plants per block, prior to inoculation. The field experiments at NIAB (N2006) and CSL (C2006) were conducted in small hand sown plots-each plot comprised of 1 row of approximately 40 cm using a randomized complete block design with two replicates.

The inoculum preparation, plant husbandry, trial set-up and disease assessments were carried out as described in Gosman et al (2005). Plants were spray inoculated at mid-anthesis (growth stage 65, Zadoks et al. (1974) with a conidial suspension (1 × 10⁵ conidia ml⁻¹) of a DON-producing isolate.
of *F. culmorum*. Inoculum was amended with 0.05% Tween 20 prior to use. In the field, inoculation was carried out in the evening with a knapsack sprayer and plants in the polytunnel experiments were inoculated with a hand-held sprayer. In the polytunnel, inoculated spikes were covered with cellophane crossing bags for 72 h post inoculation to maintain high humidity.

Disease was assessed as the percentage (0 to 100%) of visually infected spikelets (Gosman *et al.* 2005). In the polytunnel trials, disease was assessed four times at 7, 14, 21 and 28 days post inoculation (dpi). The lines in the N2006 trial were assessed at 20, 28 and 34 dpi. The area under the disease progress curve (AUDPC) was calculated (Buerstmayr *et al.* 2000) and used in subsequent analysis. The disease levels were low at C2006 trial where a single assessment was made. The weight of infected spikelets (WIS) was determined at harvest after counting the number of spikelets per head and weighing heads to provide an additional FHB resistance trait. In addition to this, the effect of DON toxin on the growth and germination of seeds (Petritox test) was measured as described in Gosman *et al.* (2005) and the area under germination curve (DON-AUG) was measured. Due to the high cost associated with the test, only half of the population and the parents RL4137 and Timgalen were examined for DON response. The DON-AUG data was only used in QTL detection and no further statistical analysis was performed on this data. The overall AUDPC was calculated from J1999, J2000 and N2006 and was treated as another experiment (pooled AUDPC) for QTL analysis (the C2006 data were omitted as only a single score was taken).

Morphological traits were assessed in the J2000 experiment. Awns were scored on a 1 to 5 scale (1 = no awns, 5 = long awns). The plant height (PH) to the top of the spike in cm was recorded during mid-anthesis.

**Statistical analysis**

All the statistical analyses were performed using GenStat for Windows 9th edition (copy right Lawes Agricultural Trust, Rothamsted Experimental Station, UK). Analysis of variance (ANOVA) was carried out using the generalised linear model (GLM) of regression analysis. The experimental repeatability ($h^2$) was estimated from the ANOVA using the formula:

$$h^2 = \sigma_e^2/\left[\sigma_G^2 + (\sigma_e^2/r)\right]$$

where:

- $\sigma_G^2$ – genetic variance
- $\sigma_e^2$ – environmental variance
- $r$ – number of replicates per genotype (Nyquist 1991).

**Markers, map construction and QTL analysis**

A genetic linkage map was constructed using simple sequence repeat (SSR) markers, amplified fragment length polymorphism (AFLP) and diversity arrays technology (DArT™) markers, a sequence independent, relatively cheap and high through put dominant marker system (Jaccoud *et al.* 2001, Triticarte Private Limited, Australia, http://www.triticarte.com.au/content/wheat_diversity_analysis.html) which has been used in crop plants. Simple sequence repeat (SSR) loci of known map location (primer prefix XpSp and Xgwm from Bryan *et al.* (1997) and Roden *et al.* (1998), respectively) were used. The PCR master mix comprised 20 µl containing 60 ng of DNA, 0.2 µM each of forward and reverse primers, 0.13 mM of dNTPs, 2 U of *Taq* DNA polymerase (Roche). The PCR profile was as follows: 94°C for 2 min followed by 30 cycles of 94°C for 1 min, 51°C for 1 min (ramp 0.5°C/s) and 72°C for 2 min (ramp 0.5°C/s) with a final extension of 72°C for 4 min. AFLPs were amplified as described by Vos *et al.* (1995) using 81 combinations of MseI and PstI primers, and the amplified products were visualised by silver staining. The linkage map was constructed using JoinMap (version 3.0) (Van Ooijen & Voorips 2001) and the map distances were calculated using the Kosambi mapping function (Kosambi 1944). The map comprised a total of 341 loci of which, 90 were AFLPs, 15 SSRs and 236 DArT markers. The SSRs and DArT markers of known map location permitted the assignment of linkage groups (LGs) to chromosomes according to the previously published wheat genetic maps (Somers *et al.* 2004; Semagn *et al.* 2007; http://www.triticarte.com.au/pdf/TriticartewhtmapalignV1-2.xls). QTL detection and mapping was carried out using MapQTL® 4.0 (Van Ooijen 2004). For each trait, one-way ANOVA (Kruskall-Wallis test) was performed to detect the association between markers and traits individually. In a second step, interval mapping (IM) was performed to identify the major QTL. Automatic cofactor selection was used to fit the multiple QTL model (MQM) (backward elimination ($P > 0.02$)). For each trait, a permutation test
(1000 iterations) was performed to identify the LOD threshold corresponding to a genome-wide false discovery rate of 5% ($P < 0.05$). Based on the permutation tests, a threshold LOD value of 3 was used to declare the presence of QTL. The QTL that explained more than 10% of the variance ($R^2$) in at least one environment/experiment were arbitrarily classified as major QTL and those explaining less than 10% as minor QTL. Linkage Maps were drawn using MapChart (Voorrips 2002).

**RESULTS**

**Phenotyping of RILs**

The parents, RL4137 and Timgalen, and the progeny were phenotyped for FHB resistance in the polytunnel (J1999 and J2000) and field experiments (N2006 and C2006). The frequency distribution of AUDPC was continuous for all the experiments and distribution was slightly skewed towards the resistant parent RL4137. The frequency distribution of pooled AUDPC is shown in Figure 1. Summary statistics of disease and other FHB related traits recorded in different experiments are shown in Table 1.

The correlations between N2006 and polytunnel experiments for AUDPC were highly significant ($P < 0.001$) and the correlation coefficient ranged from 0.5 (between J2000 and N2006) to 0.7 (J1999 and J2000) (Table 2). The correlation between C2006 and N2006 was also highly significant ($P = 0.005$). WIS had a significant negative relationship with AUDPC except for C2006 ($P = 0.082$) and the correlation coefficient ranged from −0.29 to −0.80 over experiments. Correlation coefficients between PH and AUDPC were in all experiments negative but this was only significant for J2000, N2006 and for the pooled data and the correlation coefficient ranged from −0.12 (C2006) to −0.47 (N2006). Significantly positive was the relation between PH and WIS. The trait “awns” showed non-significant relationship with AUDPC, but significantly negative association with PH ($r = −0.46$). For AUDPC, the genotypic variance was highly significant in all the experiments ($P < 0.001$) (Table 3). The experimental

![Figure 1. Frequency distribution of pooled AUDPC of Fusarium head blight in RL4137 X Timgalen RIL population](image)

Table 1. Summary statistics for Fusarium head blight (FHB) disease assessed in J1999, J2000 and N2006 and other FHB related traits recorded in J2000 trial

<table>
<thead>
<tr>
<th>Experiment and trait recorded</th>
<th>RL4137</th>
<th>Timgalen</th>
<th>Mid-parent value</th>
<th>Population mean</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>J1999-AUDPC</td>
<td>545.64</td>
<td>1161.08</td>
<td>853.36</td>
<td>835.9</td>
<td>50–1796.7</td>
</tr>
<tr>
<td>J2000-AUDPC</td>
<td>357.38</td>
<td>983.85</td>
<td>670.62</td>
<td>477.4</td>
<td>0–1540</td>
</tr>
<tr>
<td>N2006-AUDPC</td>
<td>75.31</td>
<td>435.88</td>
<td>255.59</td>
<td>276.7</td>
<td>12–835</td>
</tr>
<tr>
<td>C2006 (%)</td>
<td>0.25</td>
<td>1.5</td>
<td>0.75</td>
<td>1</td>
<td>0–7.75</td>
</tr>
<tr>
<td>J2000 – spikelet weight (g)</td>
<td>0.08</td>
<td>0.04</td>
<td>0.06</td>
<td>0.59</td>
<td>0.04–1.49</td>
</tr>
<tr>
<td>J2000 – awns</td>
<td>1</td>
<td>5</td>
<td>2.5</td>
<td>4.336</td>
<td>1–5</td>
</tr>
<tr>
<td>J2000 – plant height (cm)</td>
<td>162.3</td>
<td>97.7</td>
<td>130.0</td>
<td>121.1</td>
<td>73–170</td>
</tr>
</tbody>
</table>


Awns were measured on 1 to 5 scale.
The linkage map

The genetic map comprised of a total of 341 loci mapped onto 44 LGs and the map included 15 SSRs, 90 AFLPs and 236 DArT loci covering a genetic distance of 973 cM. LGs were assigned to chromosomes either based on the consensus wheat DArT maps from the Triticarte website and/or using the marker information from previously published maps. Genetic distance and marker coverage varied between genomes. There was a good coverage of the B genome with 198 markers spanning 571 cM. The A genome map contained 99 markers with the map length of about 282 cM however, the D genome map was less well covered and contained only 40 markers spanning 119 cM. There were two anonymous small LGs with two markers each and no LG corresponding to 5A.

FHB QTL

QTL analysis was performed on AUDPC values for individual experiments and for data pooled across experiments. The pooled average AUDPC from the three experiments was treated as another environment. The analysis identified a total of seven putative QTL for AUDPC on chromosomes 1B, 2B, 3A, 6A, 6B, 7A and 7D (Table 4 and Figure 2). Of these, five QTL were detected above the LOD score 3. Alleles from RL4137 contributed the positive effect for all the QTL except that on 6B. The QTL *Qfhs.jic-2b* (LOD = 2.1 to 5.2; \( R^2 = 8.4 \) to 21.5) contributed by RL4137 and *Qfhs.jic-6b* (LOD = 2.4 to 3.8; \( R^2 = 9.2 \) to 14) contributed by the alleles from Timgalen were the major QTL detected in more than one environment indicating that they are relatively stable. QTL for WIS, the yield associated FHB trait, were detected on chromosomes 2B and 6A with RL4137 contributing the positive effect alleles (greater WIS) in each case (Table 4). In the N2006 field trial a major QTL on 3A was detected, contributed by RL4137. QTL analysis for DON tolerance identified two minor QTL (contributed by alleles from RL4137) one
QTL for morphological traits

QTL for PH (positive effect of “greater height” allele from RL4137) and “awns” (positive effect of “long awns” allele from Timgalen) were coincident on 2B and associated with the AFLP marker S13/M23G (Tables 4 and 5; Figure 2).

Co-incidence of trait QTL

The QTL for PH (LOD = 15.9; $R^2 = 47.9$), WIS (LOD = 3.5; $R^2 = 33.9$) and “awns” (LOD = 3.4; $R^2 = 14.6$) co-localized with major FHB resistance QTL Qfhs.jic-2b (LOD = 2.1 to 5.2; $R^2 = 8.4$ to 24.2) (Table 4 and Figure 2). RL4137 contributed the positive effect alleles for the FHB (greater resistance), PH (greater height) and WIS (greater weight) while Timgalen contributed the positive effect alleles for “awns”.

DISCUSSION

To date, almost 50 QTL studies have been published on FHB resistance in hexaploid wheat. Many of these are either on Sumai 3 or other varieties of Chinese origin with a limited number of studies on known resistance sources from elsewhere. The red-grained, awnless, Canadian FHB resistant spring wheat RL4137, showing “extremely” great plant height, was derived from the Brazilian cultivar Frontana. Timgalen is a white-grained Australian spring wheat that has moderate resistance to FHB. It carries a large introgressed segment on 2B from Triticum timopheevi (AAGG) (Devos et al. 1993). Classical genetic studies suggest that a
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</tr>
</thead>
<tbody>
<tr>
<td>Qfhs.jic-1b</td>
<td>wPt-6425</td>
<td>51.3</td>
<td>RL4137</td>
<td>2.6</td>
<td>9.1</td>
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<td>2.1</td>
<td>9.2</td>
<td>2.4</td>
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<td>Qfhs.jic-2b</td>
<td>wPt-5292</td>
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<td>RL4137</td>
<td>4.8</td>
<td>21.5</td>
<td>4.4</td>
<td>2.1</td>
<td>8.4</td>
<td>2.4</td>
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<tr>
<td>Qfhs.jic-2b</td>
<td>S24/M16i</td>
<td>6.2</td>
<td>RL4137</td>
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<td>4.4</td>
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<td>8.4</td>
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<td>RL4137</td>
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</tr>
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<td>2.1</td>
<td>9.1</td>
<td>2.4</td>
<td>2.1</td>
<td>9.2</td>
<td>2.4</td>
</tr>
<tr>
<td>Qfhs.jic-6a</td>
<td>S14/M20M</td>
<td>5.3</td>
<td>RL4137</td>
<td>7.4</td>
<td>14.3</td>
<td>7.4</td>
<td>14.3</td>
<td>7.4</td>
<td>14.3</td>
</tr>
<tr>
<td>WIS</td>
<td>S13/M23G</td>
<td>5.5</td>
<td>RL4137</td>
<td>3.5</td>
<td>33.9</td>
<td>3.5</td>
<td>33.9</td>
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<td>33.9</td>
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<tr>
<td>Qfhs.jic-2b</td>
<td>wPt-9132</td>
<td>12.3</td>
<td>RL4137</td>
<td>2.1</td>
<td>23.3</td>
<td>2.1</td>
<td>23.3</td>
<td>2.1</td>
<td>23.3</td>
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<tr>
<td>DON-AUG</td>
<td>S24/M16i</td>
<td>6.2</td>
<td>RL4137</td>
<td>2.3</td>
<td>20.3</td>
<td>2.3</td>
<td>20.3</td>
<td>2.3</td>
<td>20.3</td>
</tr>
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</table>

- J = JIC, N = NIAB, r² = Percentage phenotypic variance explained, a = LOD below the permutation test threshold for significance, AUDPC = area under disease progress curve, WIS = weight of infected spikelets, DON = deoxynivalenol, AUG = area under growth curve.
minimum of two or three additive genes control FHB resistance in Frontana (Singh et al. 1995; Vanginkel et al. 1996). In the current study, RILs from a cross between RL4137 and Timgalen were screened for FHB resistance in two polytunnel and two field experiments using a DON-producing isolate of *F. culmorum*. Generally, distribution of AUDPC was continuous and was marginally skewed towards RL4137. Positive transgression of FHB resistance was observed indicating that both parents contributed unique alleles of positive effect. In the present study, the RL4137 FHB
QTL of greatest effect was on 2B (Qfhs.jic-2b), explaining up to 24% of the phenotypic variance. The shift in the QTL peak for C2006 could due to the introgression of the large chromosomal segment on 2B from *T. timopheevi* into RL4137 (Devos *et al*. 1993) and this made it difficult to reliably position FHB QTL on 2B. RL4137 also contributed a putative FHB QTL of minor effect on 1B (Qfhs.jic-1b) and less environmentally stable, but more potent QTL on 3A, 6A and 7A. The moderately resistant variety Timgalen contributed a QTL of major effect on 6B (Qfhs.jic-6b) and, together, these accounted for a total of about 40% the total phenotypic variance. It is not uncommon for a moderately resistant or susceptible parent to contribute alleles for resistance. For example, Waldron *et al*. (1999) reported a QTL for FHB resistance from Stoa, a moderately susceptible parent and Shen *et al*. (2003), found that Alondra, a FHB susceptible parent contributed an allele for resistance. In addition to visual disease, a second FHB-associated trait was assessed. Two QTL (2B and 6A) contributed by alleles from RL4137 explained most of the variation for WIS (weight of infected spikelets) (> 74%) in the population. The QTL on 2B co-localized with that for the major FHB QTL in this population.

Despite Frontana being acknowledged as a useful source of FHB resistance (Van Ginkel *et al*. 1996), only two QTL studies have been reported to date (Steiner *et al*. 2004; Mardi *et al*. 2006). Steiner *et al*. (2004) reported stable QTL for field resistance on 3A and 5A and less stable QTL on 2B and 6B in a Frontana/Remus cross. For type II resistance, these researchers found only a single QTL of small effect on 2B. In addition, a QTL of small effect on 1BL was contributed by Remus, the susceptible parent. Subsequently, Mardi *et al*. (2006) confirmed the QTL on 3A and detected an additional QTL on 7AS. They also detected a QTL on 1BL originating from Seri82, the susceptible parent of their mapping population. The results of the present QTL analysis partially support the findings of these earlier studies, but also highlight some inconsistencies that may reflect differences in the genotype of the susceptible parent and/or the methodologies employed. The 3A QTL for field resistance reported in both previous studies was detected in the N2006 field trial of the present study but, significantly, was not observed in the C2006 field trial where disease pressure was very low or the polytunnel trials where disease pressure was very high. This highlights the importance of appropriate testing for the detection of some FHB QTL that are, perhaps, only effective under particular conditions. We observed FHB QTL on 2B and 7A, in common with the reports of Steiner *et al*. (2004) and Mardi *et al*. (2006), respectively. Both previous studies found Frontana to carry alleles on 1B that were deleterious for FHB resistance compared to the other parent. In contrast, alleles on 1B from Frontana were marginally more beneficial than those of Timgalen. This probably reflects differences in the genotypes of the susceptible parent in each study.

In addition to physiological, or active, components of FHB resistance, morphological traits including PH, and the presence and size of awns have previously been reported to be associated with FHB resistance, either through linkage or pleiotropy. For this reason PH and awnedness were also measured to assess their relationship to FHB resistance in the present materials. Studying the relationship between FHB resistance and other important agronomic traits is crucial for the development of resistant cultivars. Breeding programmes can be seriously hampered if FHB resistance is linked to undesirable traits. For example, tall stature has widely been assumed to be an FHB escape mechanism reducing spike infec-

<table>
<thead>
<tr>
<th>Trait</th>
<th>Closest marker</th>
<th>Chr.</th>
<th>Position</th>
<th>Origin</th>
<th>LOD</th>
<th>r²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plant height (cm)</td>
<td>S13/M23G</td>
<td>2B</td>
<td>5.5</td>
<td>RL4137</td>
<td>15.9</td>
<td>47.9</td>
</tr>
<tr>
<td>Plant height (cm)</td>
<td>wPt-7524</td>
<td>4A</td>
<td>3.4</td>
<td>Timgalen</td>
<td>2.9</td>
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<td>Plant height (cm)</td>
<td>wPt-9454</td>
<td>5B</td>
<td>57.8</td>
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<td>5.9</td>
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<td>Awns</td>
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<td>2B</td>
<td>5.5</td>
<td>Timgalen</td>
<td>3.4</td>
<td>14.6</td>
</tr>
</tbody>
</table>

r² = percentage phenotypic variance explained, Chr. = chromosome, a = LOD below the permutation test threshold for significance.
tion through reduced spore transfer by rain splash and by providing a less favourable micro-climate for spore germination on the spike (Miedaner 1997; Hilton et al. 1999; Klahr et al. 2007). However, tall cultivars are prone to lodging and have a low harvest index. In the current study, plant height (PH) had a significant negative correlation with AUDPC. QTL mapping detected a major PH QTL, which co-localized with a major FHB resistant QTL on 2B with alleles for a positive effect both contributed by RL4137. QTL for these two traits were also detected on 2B by Steiner et al. (2004). Some studies have revealed a more complex relationship between FHB resistance and PH (Schmolke et al. 2005; Draeger et al. 2007; Srinivasachary et al. 2008a,b). These studies suggested that co-incidence of QTL for FHB resistance and PH has a genetic basis, e.g. linkage or pleiotropy, rather than being the result of escape. The observation that the FHB QTL on 1B and 6B were not associated with PH QTL and that the minor PH QTL on 4A and 5B, did not co-localise with any other FHB resistance QTL. The FHB QTL Qfhs.jic-2b remained highly significant even after co-variance analysis was carried out on the AUDPC data adjusted in relation to PH. The existence of FHB resistance that is independent of PH is of importance since it allows for the selection of resistant cultivars of any height.

The precise role of awns in FHB resistance is not clear and the literature is often contradictory. Tamburic-Ilnicic et al. (2007) found that awned genotypes had a lower FHB index than awnless genotypes while Ban and Suenaga (2000) reported that fully awned genotypes were more resistant than tip-awned genotypes. Snijders (1990) suggested that awns could be used as a marker to select FHB resistant lines among progenies from crosses in which the resistant parent carries awns. However, Mesterhazy (1995) found that the presence of awns was associated with higher levels of disease under field conditions but, interestingly, not under conditions of artificial inoculation. In the current study, a major QTL for awns was also found to co-localise in repulsion phase (associated with the long awn allele from Timgalen) with the major FHB resistance QTL on 2B contributed by RL4137. The present result, combined with that of other studies suggests that QTL for both FHB resistance/susceptibility and awns/no awns are located on 2B and that they are sufficiently closely linked to provide a means to enhance FHB resistance in breeding programmes by selecting for either presence or absence of awns as appropriate depending upon the association in the original FHB resistance donor.

The trichothecene mycotoxin deoxynivalenol (DON) has been shown to be required by F. graminearum to facilitate spread, from the point of infection, into adjacent spikelets (Desjardins et al. 1996; Bai et al. 2002). Furthermore, Lemmens et al. (2005) found that a major QTL for DON tolerance co-localised with that for type II resistance on chromosome 3B supporting the view that type II resistance is associated with resistance to DON. Most of the FHB resistance in Frontana appears to be of type I (Steiner et al. 2004). However, early studies reported that Frontana possesses resistance to DON (Miller & Arnison 1986). Germination in the presence of DON toxin, known as in vitro DON tolerance, has previously been reported to be associated with FHB resistance in wheat (Lemmens et al. 1994; Gosman et al. 2005). Interestingly, in the current study, QTL analysis for DON tolerance identified two minor QTL one each on 2B and 7A both of which co-localized with the FHB resistance QTL contributed by RL4137. These findings correlate with those from previous studies that reported moderate correlation between FHB symptom development, DON accumulation in kernels and in vitro DON tolerance (Lemmens et al. 1994, 1997). This probably reflects a relatively minor role for type II resistance in RL4137 as was reported by Steiner et al. (2004) in Frontana. Interestingly, Steiner et al. (2004) also reported that the most consistent QTL for Type II resistance in Frontana was on 2B.

The current study identified two major consistent QTL one each on 2B and 6A, contributed by alleles from RL4137 and Timgalen, respectively. The QTL for the FHB related traits co-localized with those for PH and awns on 2B. Co-localization of QTL may result from linkage or pleiotropy. However, at the level of resolution afforded by the present mapping population, linkage or pleiotropy could not be distinguished. The development of iso-genic lines carrying individual FHB resistance loci in a common genetic background may provide the opportunity to distinguish linkage from pleiotropy in the relationship between FHB resistance and morphological traits. Unfortunately, the 2B chromosome of Timgalen carries a large DNA segment from T. timopheevi (Devos et al. 1993),
which reduces recombination making it impractical to attempt to undertake precise mapping of the RL4137 2B QTL in a Timgalen background. Additional crosses to an alternative FHB susceptible parent will be required to differentiate between linkage and pleiotropy for these traits.

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