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Linkage between RFLP molecular markers and the dwarfing genes *Rht-B1* and *Rht-D1* in wheat

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Identifying genotypes carrying the dwarfing genes Rht-B1 and Rht-D1 would be of great interest for wheat (*Triticum aestivum* L. em Thell) breeding. Two RFLP loci were found to be linked to those two genes, Xpsr 144-4B with Rht-B1 on chromosome 4BS and Xglk578-4D with Rht-D1 on chromosome 4DS, by genotyping two F₂ populations, Renan (Rht-B1b) × Camp-Rémy and Rendez-Vous (Rht-D1b) × Roazon. Utilisation of these markers in breeding schemes is discussed.

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Reducing plant height in wheat is of great economical importance because it makes it possible to simultaneously increase lodging resistance, and thus stability of yield, and harvest index. Because many genetic factors which influence development, morphology or vigour will have effects on final plant height, this character involves most of the 21 chromosomes in wheat (SNAPE et al. 1977). Several genes have already been identified and "short" alleles have been designated Rht (for Reduced height) relative to their "tall" alleles rht. The Norin 10 dwarfing alleles Rht1 and Rht2 have been used extensively in the breeding schemes during the last 40 years due to the lack of consequences on yielding ability. In CIMMYT, BOR-LAUG (1968) made the first successful crosses between Norin 10-Brevor 14 and tall "daylength insensitive" wheats in 1955. The varieties Pitic 62 and Penjamo 62 were released to Mexican farmers in 1962. Since the registration in 1964 of the semi-dwarf variety 'Gaines' (VOGEL 1964), the proportion of varieties carrying dwarfing alleles is in constant increase and, at the present time, reaches more than 80% of the world registrations.

Dwarf wheat plants are classified into two groups following their reaction to the application of exogenous gibberellin (GA; REID 1986). The GA insensitive character of Norin 10 was first mentioned by ALLAN et al. (1959), and genes influencing this trait were named *Gai* (for GA insensitivity; MCINTOSH 1979). Genetic linkage analysis between *Rht1* and *Gai1* on one hand, and *Rht2* and *Gai2* on the other, made by GALE and MARSHALL (1975) and GALE and GREGORY (1977) showed absence of recombination. It was considered that the *Rht* and *Gai* genes were identical and that they conferred, by pleiotropic effects, both reduced height and gibberellin insensitivity (WORLAND et al. 1987). *Rht1* and *Rht2* act by reducing internode length without reducing spike morphology (ALLAN et al. 1968). Both are semi-dominant (GALE and YOUSSEFIAN 1985) and have similar effects on plant height (reduction of about 20 cm) even though *Rht2* seems to have a stronger effect than *Rht1* (ALLAN 1970; BÖRNER et al. 1993). Numerous alleles inducing different levels of dwarfism have already been identified for either *Rht1* by GALE et al. (1975; *Rht3*), WORLAND and PETROVIC (1988; *Rht1S*), WORLAND and SAYERS (1995; *Rht1(B.dw)*), or *Rht2* by BÖRNER and METTIN (1988; *Rht10*).

Combination of Rht1 and Rht2 in the same genotype generally produces a strong effect on plant height which, most of the time, induces a decrease of the agronomical value of the variety (ALLAN 1983). In addition, crosses between lines carrying different dwarfing genes generate a wide segregation for height which leads to the rejection of the majority of the plants only because they are too tall or too short. It would then be advisable to cross lines with the same dwarfing gene, especially for recurrent breeding. A large variation in height reduces the efficiency of breeding because of strong competition effects between tall and short plants resulting in difficulties in evaluating populations segregating for height. Unfortunately, the test using GA is unable to distinguish the lines carrying Rht1 from those with Rht2. Test crosses with lines from a known genotype (Rht1 or *Rht2*) must be done to identify the gene(s) of new lines.

Since 1989, dense RFLP linkage maps of the hexaploid wheat *Triticum aestivum* L. em. Thell. have been developed from interspecific crosses for the ho-

moeologous groups 1 (VAN DEYNZE et al. 1995), 2 (DEVOS et al. 1993; NELSON et al. 1995a), 3 (DEVOS et al. 1992; NELSON et al. 1995b), 4 and 5 (XIE et al. 1993; DEVOS et al. 1995; NELSON et al. 1995c), 6 (JIA et al. 1996; MARINO et al. 1996) and 7 (CHAO et al. 1989: NELSON et al. 1995c) and from an intervarietal cross (CADALEN et al. 1997). These maps would be useful to identify markers linked to Rht1 and to Rht2 for an accurate selection of the plants having these dwarf alleles. Previous studies indicate that these two genes are carried on the short arms of the homoeologous group 4. Using telocentric mapping, it was shown that Rht1 is located at 13 map units from the centromere on chromosome 4BS while Rht2 falls 15 map units from the centromere on the chromosome 4DS (MCVITTIE et al. 1978).

The aim of this study was to look for markers linked to these two genes, using segregation in two F_2 populations. We have thus focused our study on chromosomes 4B and 4D. In the present study, we will use the conventional and newly proposed nomenclature of the GA insensitive *Rht* alleles in wheat (BÖRNER et al. 1996) i.e., *Rht-Bla* and *Rht-Dla* (formerly *rht1* and *rht2*) and *Rht-Blb* and *Rht-Dlb* (formerly *Rht1* and *Rht2*).

MATERIAL AND METHODS

Plant material

Two F_2 populations were elaborated at the INRA Plant Breeding Station at Rennes. The first one, of 116 plants, was from the cross between the cultivars Renan (*Rht-B1b*) and Camp-Rémy, and height was ranging between 58 and 105 cm. The second one, of 115 individuals, was produced from the cross between the cultivars Rendez-Vous (*Rht-D1b*) and Roazon, and height was ranging between 61 and 127 cm.

Plant height evaluation

Plant height of F_2 plants and of the four parents (Renan, Camp-Rémy, Rendez-Vous, Roazon) was scored under field conditions at INRA Station at Rennes in 1995. The genotype at the loci XRht-B1 and XRht-D1 was determined in the greenhouse in 1996 on F_3 seedlings, using a method derived from that of GALE and GREGORY (1977). Two replications of 20 seeds were sown in wet sand in seed trays and placed into controlled environments to ensure even germination and to avoid etiolation. The seeds were then irrigated once a day with a solution containing 5 mg/l of GA₃. Measurements were taken 12–15 days after sowing, and plantlets were divided into two groups. Those homozygous for Rht-B1b or Rht-D1b ("short" alleles) present short sheaths and first leaves,

while those heterozygous (Rht-Bla/Rht-Blb or Rht-Dla/Rht-Dlb) or homozygous for Rht-Bla or Rht-Dla ("tall" alleles) have long sheaths and first leaves twice as long as short plants.

Molecular markers and RFLP analysis

The probes used in this study are given in Table 1. The techniques for DNA extraction, digestion, electrophoresis, blotting, and hybridization were described by CADALEN et al. (1997). The protocol using non-radioactive probes was detailed in LU et al. (1994) and SOURDILLE et al. (1996).

One microsatellite (GWM165; PLASCHKE et al. 1996) was also mapped on these populations. PCR reactions were carried out in a final volume of 50 µl in a Perkin-Elmer 9600 thermocycler. The reaction buffer contained 100 ng of template DNA, 0.2 mM of each deoxynucleotide, 1.5 mM MgCl₂, 500 nM of each primer, and 1 unit of Taq-DNA-polymerase (Boehringer). Thirty-five cycles with 1 min at 96°C, 1 min at 60°C (in some cases 57°C), and 2 min at 72°C were realised, followed by a final elongation step of 10 min at 72°C. Polyacrylamide (6%) denaturing gels (0.4 mm thick) were prepared with 1X TBE buffer (90 mM Tris, 90 mM boric acid, 2 mM EDTA, pH 8.2) and cast between glass plates, one previously treated with Sigmacote (Sigma) and the other with Silane A-174 (Sigma). From each sample, 1.6 µl were loaded together with 0.8 µl of loading buffer (10 mM NaOH, 0.05% bromophenol blue, 0.05 % xylen cyanol, in 95 % deionized formamide) after a 5 min denaturation in boiling water. Samples were then run for 2 h at 2000 V, 36 mA and 61 W. DNA was revealed by a silver staining method derived from the protocol of BASSAM et al. (1991). The gel was first fixed for 20 min in 10% acetic acid, rinsed three times with water and stained in a solution containing 0.1% silver nitrate and 0.05%formaldehyde for 30 min. After a quick rinse (not exceeding 10 s) microsatellites were revealed by adding the developer (3 % sodium carbonate, 0.05 %

Table 1. DNA probes and microsatellites used in this study

Probes	References
GLK335	LIU and TSUNEWAKI (1991)
GLK556	LIU and TSUNEWAKI (1991)
GLK578	LIU and TSUNEWAKI (1991)
GWM165	PLASCHKE et al. (1996)
PSR144	GALE et al. (1995)
FBA41	QUETIER (personal gift)
FBA177	QUETIER (personal gift)
FBA211	QUETIER (personal gift)
FBB58	QUETIER (personal gift)



Fig. 1. Maps of wheat chromosomes 4B and 4D obtained from the F_2 populations Renan × Camp-Rémy (a) and Rendez-Vous × Roazon (b). Markers are in LOD 3.0 order. Distances on the left of the chromosome are in cM according to KOSAMBI (1944) and numbers in brackets indicate recombination values. Approximate position of the centromere (c) is indicated with an arrow.

formaldehyde, 2 mg/l sodium thiosulfate). Image development was conducted at a maximum temperature of 12°C during 3 to 10 min. The reaction was stopped by addition of acetic acid (10%), and the gel was rinsed in water and dried at room temperature.

Mapping and linkage analysis

The maps obtained from the F_2 populations were constructed using Mapmaker/exp version 3.06 (LAN-DER et al. 1987). The linkage groups were established by calculating recombination frequencies with the following conditions: thresholds for LOD = 3 and $\theta = 0.35$. Association between molecular markers and plant height (QTLs) was evaluated by the "marker regression" method (KEARSEY and HYNE 1994) computed with Splus software (BECKER et al. 1992).

RESULTS

The averages of plant height of the cultivars Renan (Re), Rendez-Vous (RV), Camp-Rémy (CR) and Roazon (Rz) scored at the Plant Breeding Station at Rennes were 91, 80, 93, and 92 cm respectively. The parents of the F_2 populations (Re × CR and RV × Rz) were not highly contrasted in height.

Mapping of Rht-B1 and Rht-D1

The GA₃ test allowed identification of the genotype for each F_2 plant in both populations (Re × CR and $RV \times Rz$). The two genes have been considered as genetic markers and their localisation on the maps is given in Fig. 1a and 1b. Assignation of the linkage groups to the chromosomes 4B and 4D was realised using a microsatellite (GWM165) which gave homoeologous loci on each of the three chromosomes of the group 4 (PLASCHKE et al. 1996). The loci on chromosomes 4B and 4D were both polymorphic for both F_2 populations. The Rht-B1 gene was located between Xpsr144-4B and Xglk556-4B. Nevertheless, the linkage was not very strong since the gene was 13 cM distal to the former and 43 cM distal to the latter (Fig. 1a). On the other hand, the gene Rht-D1 was very close to Xglk578-4D (3 cM), but we did not find a more distal marker beyond this gene. The level of polymorphism detected between the two parents of this cross was low (RV and Rz: 18% of polymorphic probes). This, combined with the poor polymorphism generally detected on the D genome (CHAO et al. 1989; CADALEN et al. 1997) can explain our result.

The marker regression method conducted on the data from the $\text{Re} \times \text{CR}$ F₂ population detected the



Fig. 2 a and b. Detection of loci associated with plant height on the F_2 population Renan × Camp-Rémy using the regression method (KEARSEY and HYNE 1994). Only partial maps (around 90 cM long) are given. Positions of the anchor markers are given in cM along the abscissa according to KOSAMBI (1944). The long arm of each chromosome is toward the right of the graphics and the short arm toward the left. The origin (0) indicates the first marker of the short arm. The graphs represent the *F* tests for each marker (vertical bars). The curve indicates the changes in residual mean square for various putative QTL positions as defined by KEARSEY and HYNE (1994). (a) all markers, (b) when omitting XRht-B1 from the analysis.

same QTL (Fig. 2) with or without including the *Xrht-B1* locus in the analysis. This QTL was located at the same position in both cases. This locus explained only 9% of the variability for plant height on this population and the locus Xglk556-4B was not found to be significantly associated with this trait (*F* probability 0.15). The positive allele (increasing plant height) came from CR, which was consistent with the fact that Re carried the *Rht-B1b* allele. The additive value was 5.4 cm while the dominance effect was only 0.4 cm. This result was not expected since *Rht-B1* was described as semi-dominant (GALE and YOUSSE-FIAN 1985).

DISCUSSION

Molecular markers linked to dwarfing genes would allow more rational breeding schemes concerning plant height. In this study, we have found two loci (Xpsr144-4B and Xglk578-4D) associated with two such genes (Rht-B1 and Rht-D1, respectively). In the case of Rht-B1, information from the two loci flanking the gene (Xpsr144-4B and Xglk556-4B) should be necessary for integration of these molecular markers in a breeding scheme. The gene Rht-D1 was not framed between two markers. It is located on the D genome (chromosome 4D), which was often mentioned to be less polymorphic than the A and B genomes (CHAO et al. 1989; KAM-MORGAN et al. 1989; LIU and TSUNEWAKI 1991; CADALEN et al. 1997). This may explain the fact that we found only few markers on this chromosome.

It was unexpected to find so weak additive and dominance values (5.4 cm and 0.4 cm, respectively) on the Re \times CR F₂ population. Perhaps, they are due to epistatic effects between *Rht-B1* and other regions on the genome, which reduce the effect of *Rht-B1*.

Two loci found to be associated with these genes were revealed by a unique microsatellite (GWM165). Microsatellites have several advantages versus RFLP markers: (1) the level of polymorphism detected is much higher (e.g., PIC values for the three loci for this microsatellite on the A, B, and D genomes, respectively: 0.42, 0.81, 0.42) than the level of polymorphism of RFLP (mean of PIC value 0.3); (2) each locus is genome-specific and, in our case, previous chromosome assignment done on aneuploid lines (PLASCHKE et al. 1996) allowed each linkage group to be attributed to a chromosome.

Our results should be validated on a core collection of genotypes with the probes PSR144 and GLK578. We will probably be confronted with the lack of

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crosatellites could be a way to solve this problem.

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