Construction of a Practical SCAR Marker Linked to Clubroot Resistance in Chinese Cabbage, with Intensive Analysis of HC352b Genes

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A restriction fragment length polymorphism (RFLP) marker, HC352b, has identified as closely linked to the CRa locus responsible for clubroot resistance (CR) in a study dissecting CR loci in Chinese cabbage (Brassica rapa L. ssp pekinensis). Unfortunately, the RFLP pattern, including HC352b detected by the cDNA clone HC352, was complicated and confused in its interpretation because it represented multiple-copy loci. To provide a practical HC352b marker for CR breeding programs, a sequence characterized amplified region (SCAR) marker was constructed along the analysis of HC352 genes, followed by its evaluation for CR selection. Schematic characterization of signals was achieved by identification of HC352 homologous genes with their restriction sites, employing CRa-positive and -negative doubled haploid (DH) lines. Genomic sequence information from a set of HC352 homologous genes was analyzed to identify the CRa linked paralog HC352b, followed by the successful designation of a SCAR marker. This correctly predicted the CR phenotypes of all tested individuals of F2 and back-crossed progenies.

Key Words: clubroot resistance, HC352, paralog, sequence characterized amplified region (SCAR), CRa.

Introduction

Clubroot is one of the major diseases of Brassicaceae. It is caused by the parasitic infection of Plasmodiophora brassicae, a soilborne fungus, in the stem root of plants (Ludwig-Müller, 1999). Chinese cabbage (Brassica rapa L. ssp pekinensis) is highly susceptible to disease and frequently suffers destructive damage in infected fields. Several breeding programs have been successfully carried out to develop clubroot resistance (CR) cultivars of Chinese cabbage using European turnips (B. rapa ssp. rapifera) as sources of resistance (Hirai et al., 2004; Matsumoto et al., 1998). CR in turnips is considered to be qualitative and is grouped into several genotypical categories corresponding to the pathotype of P. brassicae (Matsumoto et al., 2005). It is important to monitor the inheritance of these resistant genes using molecular markers in the breeding of Chinese cabbage (Kuginuki et al., 1997; Matsumoto et al., 2005; Suwabe et al., 2003). CRa is the responsible locus for CR in a dominant manner. HC352b was reported to be the most useful DNA marker linked to CRa, together with E49380 (Matsumoto et al., 2005). The random amplified polymorphic DNA (RAPD) marker E49380 is located 4.6 cM from CRa with coupling linkage. HC352b was mapped at 2.9 cM from CRa on the opposite side of E49380 and appeared to have repulsion linkage. HC352b was established as an RFLP marker with a probe derived from cDNA clone HC352 (Matsumoto et al., 1998). Although specific HC352b signals detected by cDNA show tight linkage to clubroot resistance, many unlinked signals were detected simultaneously. The appearance patterns of non-linked signals were complicated and confused, obstructing the practical use of HC352b in breeding programs. To construct a sequence characterized amplified region (SCAR) marker on a HC352b locus with the characteristics of usefulness, clarity and reproducibility, we analyzed the genomic information of the HC352 loci intensively in two doubled haploid (DH) lines, T136-8 (CRa) and K10 (a susceptible allele of CRa).
Materials and Methods

Plant materials
As source material with CRa, a DH line, T136-8 (B. rapa L. ssp pekinensis), was used (Matsumoto et al., 2005). DH lines of B. rapa L. ssp pekinensis, K10 and Q5, were used as the parent as sources of a susceptible allele of CRa. K10 originated from the “CR Kanko” and does not have CRa. Q5 is a typical clubroot-susceptible line. The original resistance source of the parental line T136-8 is a member of the European Clubroot Differential (ECD) series, ECD 02 (B. campestris ssp. rapifera line AAbbCC).

Fourteen and 90 individuals of F2 progeny between T136-8 × K10 and T136-8 × Q5 were used for DNA analysis and CR tests. F1 between T136-8 and Q5 was crossed with Q5 to produce back-crossed (BC) populations. Each F2 and BC individual was selfed to produce the next generation for segregation analysis.

DNA extraction
DNA was extracted from the leaves of parental lines, F2 and BC1 plants, according to the cetyl trimethyl ammonium bromide (CTAB) method (Murray and Thompson, 1980).

RFLP analysis
RFLP analyses were performed as previously described using the cDNA clone HC352 (Matsumoto et al., 1998) and its subclones composed of EcoRI-digested fragments. Conditions of electrophoresis, membrane transfer, probe preparation, hybridization, washing and signal detection were described previously (Matsumoto et al., 1998).

Cloning and sequencing
Regular procedures were used for cloning and sequencing DNA fragments (Nakanishi et al., 2005). A cloning vector pXcm, the vector for the TA cloning of PCR fragments, was used with a Ligation Kit (Nippon Gene, Tokyo, Japan). Sequencing was performed according to the manufacturer’s instructions (ABI PRISM 310, Applied Biosystems, CA, USA). Analysis of sequence information was performed using the Genetyx software package (Software Development, Tokyo, Japan) and the BLAST search with DDBJ (http://blast.ddbj.nig.ac.jp/top-j.html, October 11, 2007). The sequence data produced in this report were submitted to DDBJ.

PCR
The standard protocol was used as follows: In a 10 μL reaction solution were mixed 3 ng of template DNA, 0.5 mM of forward and reverse primer, respectively, 0.2 mM dNTPs, 1 μL of 10 × reaction buffer, 1.0 mM MgCl2, and 0.5 unit of Taq DNA polymerase (Promega Corporation, Madison, USA). Amplification was performed in a thermal cycler programmed as follows: 94°C for 4 min, then 40 cycles at 94°C for 30 s, at an annealing temperature for 1 min and at 72°C for 1 min, followed by a final extension of 7 min at 72°C. The annealing temperature for F7-R11 was 38°C, and for F16-R13 it was 46°C. Sequences of F7 and R11 were 5'-TGCTACACCAAAAGATTGAGAT-3' and 5'-GGATCTCTTTCTGAATGTGATG-3'. F16 and R13 were designed as 5'-CTTTATAATGGCTACTATTTA-3' and 5'-TGCTCATGAGTGTATAACTA-3'.

CR tests
Inoculation tests for resistance to clubroot were as previously reported (Matsumoto et al., 1998). As an inoculation source, we used the M85 isolate of P. brassicaceae, which was identified as race 2 according to Williams’s classification (Williams, 1966). S elfed progeny of 14 and 90 F2 plants derived from T136-8 × K10 and T136-8 × Q5, and 21 BC1 plants produced from F1 × Q5 were used for CR tests. Twenty to 40 seeds of each F3 and the BC1S1 population were sown in the pathogen-inoculated soil. Disease infection was rated 30 days after sowing. Evaluation of CR was based on the previous report (Matsumoto et al., 2005). F2 and BC1 individuals that segregated no susceptible plants in their selfed progeny were scored as homozygotes (A) for CRa, while those that segregated both resistant and susceptible plants were assessed as heterozygotes (H). F3 and BC1 individuals whose selfed progeny were thoroughly galled were scored as homozygotes (B) for susceptibility.

Results
Genomic information from RFLP analysis
Hybridization patterns of HC352 on the genomic DNAs of T136-8, K10, and their F2 progeny were analyzed. T136-8 has CRa, only one locus with exclusive responsibility for CR in the cultivar. As shown in Figure 1A, HC352 detected several fragments that were both in the lanes of K10 and T136-8.

For detailed analysis, HC352 was divided into two subclones at the EcoRI site located in the cDNA (Fig. 1B), and hybridization patterns were also developed using these subclones (Fig. 2 and data not shown). Using subclone 2 as a probe, 4 kb and 5 kb signals are clearly segregated, suggesting physical maps of HC352 genes. A summary of signals detected in parental lines is interpreted as in the scheme shown in Figure 1B. It was necessary to assume that two or more loci were detected by HC352 in order to explain all of the hybridized signals. Here we judged one of these loci as HC352b that was found using other lines of Chinese cabbage (Matsumoto et al., 2005). The other locus should be HC352a.

Signals of 0.8 kb and 5 kb in length would reflect the existence of the HC352b-T allele. Unfortunately, however, if we use the full length of the HC352 cDNA clone, the signal at 0.8 kb is somewhat faint, and the
signal at 5 kb is overlapped by another signal derived from HC352a-T.

**Sequence information from cDNA clone HC352**

The cDNA clone HC352 and two subclones were successfully sequenced and submitted to DDBJ.
Database searching analysis showed high homology with genes encoding RNase L inhibitor proteins. Two genes, At4g19210 and At3g13640, of Arabidopsis thaliana have the highest homology, and sequence AY185357 was recorded from Chinese cabbage. Since AY185357 was 3’ partial EST, unfortunately it was not sufficient to be referred for the primer design. In the A. thaliana genome, two genes coding the RNase L inhibitor protein are located separately, i.e., in chromosomes IV and III. This coincides with our scheme of the HC352 regions.

**Sequence information from the HC352 regions of genomic DNA**

Primers F7 and R11 were designed on the positions of highly conserved sequences with two introns in the target sequence. This primer set brought a couple of fragments with PCR using genomic DNA (Fig. 3). The discrete fragments in Figure 3 were tentatively named T-L, T-S, K-L, and K-S, and were sequenced after cloning (T-L: AB302979; K-L: AB302980; T-S: AB302981; K-S: AB302982). All of these fragments had good homology to part of the cDNA clone HC352, with two extra sequences considered to be introns. T-L has higher homology to K-L (98.8%) than to T-S (84.7%). Namely T-L and K-L appear allelic, and T-S and K-S could be allelic. T-S and K-S were assumed to be CRa linked (see Discussion), and were used for the further design of primers.

**Construction of the HC352b-SCAR marker**

HC352b-SCAR was designed to be amplified by T-S specific primer set F16-R13, on the assumption that the diversity of T-S vs K-S is the substantial polymorphism of HC352b. PCR with F2 from T136-8 × K10 successfully amplified targeted fragments only from genomic DNAs of F2 individuals that have the HC352b signal in RFLP analysis (Figs. 2 and 4). Finally, linkage of the HC352b-SCAR marker to the CRa phenotype was demonstrated using the F2 population from T136-8 × Q5 and the BC population from F1 × Q5 (Table 1). The HC352b-SCAR positiveness and the CR against M85 isolate (CRa phenotype) were indicated to be tightly linked.

**Discussion**

In this report, we established the co-dominant SCAR marker HC352b-SCAR tightly linked to CRa, with intensive analysis of HC352-related genes. On the other hand, RAPD marker E49 was positioned at 4.6 cM from CRa with coupling linkage (Matsumoto et al., 2005). HC352b-SCAR would be positioned at 2.9 cM from CRa and 7.5 cM from E49, respectively, if it correctly detected “HC352b” as we expected. This result means that CRa was closely put between dominant and co-dominant markers, providing efficient selection of CR plants in CR breeding programs.

**Table 1.** Linkage between HC352b-SCAR and CRa phenotype.

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<th>Populations</th>
<th>HC352b-SCAR</th>
<th>CRa phenotype</th>
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F1 and F2 plants were derived from T136-8 × Q5. BC1 plants were produced from F1 × Q5. CRa phenotype was scored with CR tests using each F1 and BC1 populations. A: CRa homozygous; H: CRa heterozygous; B: Susceptible allele of CRa.

**Fig. 3.** PCR results using F7-R11 primers. Templates are T: genomic DNA of T136-8; K: genomic DNA of K10; C: HC352 cDNA clone. Fragments indicated as T-L, T-S, K-L, and K-S are recovered from agarose gel for further analysis (see text). M: 1 kb DNA Ladder.

**Fig. 4.** Detection of HC352b-SCAR. HC352b-SCAR was detected with PCR using F16-R13 primer set. Material plants are the same as in Figure 2.
The hybridization patterns of RFLP analysis coincided with reported data using segregated populations from crosses among several B. rapa cultivars and lines, including fodder turnips with a slight change in the size of fragments (Matsumoto et al., 2005). The 4 kb fragment in Figure 1 and 2 correspond to the 4.4 kb fragments detected in the susceptible lines as reported (Matsumoto et al., 1998, 2005). The scheme of restriction sites proposed in this report could explain those reported RFLP patterns, with the possibility that the scheme would also be applicable to other cultivars of Chinese cabbage and to turnips.

HC352 cDNA-specific primers were designed at several positions referring to sequences of the A. thaliana homologue. Amplified products using cDNA-specific primers are single fragments in every combination, with no polymorphism between T-1 and K-1, indicating that the cDNA and T-1/K-1 could be derived from allelic genes. Therefore, the cDNA clone HC352 and PCR fragments T-1 and K-1 were possibly derived from HC352a. We concluded that a simple effort to make direct conversion of the sequence information derived from the HC352 cDNA clone to the SCAR marker is not fruitful.

Analysis of the genomic sequence brought more insight into the constitution of HC352-related genes, supporting the scheme in Figure 1B. Here, the genomic region HC352b-T in Figure 1B may correspond to the T-5 fragment in Figure 3, and HC352b-K may correspond to K-5. HC352a-T to T-1 and HC352a-K to K-1 appear to be derived from allelic genes. K-5 is also a counterpart of T-5. In contrast, “HC352a” and “HC352b” genes are diverse genes that are related to each other as paralogs, having originated from the duplication of an ancestral gene. To confirm this possibility, the full sequences of HC352 genes are required.

Recently, large-scale duplications or multiplications in the genome of B. rapa were indicated in linkage map analysis (Kim et al., 2006). Syntenic analysis with A. thaliana also provides evidence of multiplication in the Chinese cabbage genome (Saito et al., 2006). This information makes us cautious regarding the cross-detection of paralogical genes in studies of marker-assisted breeding. To overcome this problem, re-sequencing of homologous genes might be a powerful tool, as in this work. The employment of divergence and polymorphism uncovered by intensive sequencing could become a stable and secure strategy.

**Acknowledgements**

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**Literature Cited**


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