Full Length Research Paper

Detection of homozygosity in near isogenic Lines of non-susceptible to Zhenjiang strain of densonucleosis virus in silkworm

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The near isogenic lines (NILs) of non-susceptible to DNV-Z were bred through backcrossing successively using L10 as donor parent and Js as recurrent parent. The homozygosity of the NILs was detected using the SSR markers from the SSR linkage map. The results showed that the ratio of the linkage groups from the recurrent parent in the NILs increased rapidly along with the increase of the backcrossing generations. After 6 generations of backcrossing, it was selfed and was fed with DNV-Z. The linkage groups of 9 of the 5 BC_6 pairs, whose 1/4 offspring were non-susceptible to DNV-Z, were all replaced by the recurrent parent except the linkage group that held nsd-Z, and only part of one linkage group from the other one individual had not been replaced. According to the markers linked to nsd-Z, Fl0316 had not been replaced in all of the 10 individuals, but crossover happened in 2 individuals between Fl0568 and nsd-Z.

Key words: Silkworm, Non-susceptible to DNV-Z, Near Isogenic line, SSR marker, Homozygosity and Linkage group.

INTRODUCTION

Near isogenetic lines (NILs) mean a group of lines that are genetically identical except at one or a few loci. A NIL is developed by several backcrosses (more than six generations) between the recurrent and donor (which take the aim marker) parent. In each generation, the individuals which are similar to the recurrent parent except the aim gene should be selected and are backcrossed to the recurrent parent. Self-crossing is taken out after backcrossing for more than 6 generations, and the aim gene is homozygous. When the NIL is completed, it is genetically identical to the recurrent parent except the region close to the aim gene locus on the chromosome. Molecular markers linked to the aim gene can be identified using this NIL, and the molecular linkage map can be parallelism to the traditional mutant linkage map. Young et al. (1988) Identified DNA markers linked to the Tm-2a gene in tomato using isogenic lines. Martin et al. (1991) used random markers and NILs to identify markers linked to a pseudomonas resistance gene in tomato. Abe et al. (1998, 2000) bred the NILs of nsd-1 and nsd-2, then identified and mapped several RAPD markers linked to the nsd-1 or nsd-2 allele successively. Li et al. (2001) constructed the near-isogenic lines of nsd-Z and had identified RAPD markers linked to nsd-Z. Li et al. (2003) had constructed NILs for all 28 linkage groups by successive backcrossing method.

Bombyx mori densonucleosis viruses (BmDNVs) belong to the genus Iteravirus of the family Paroviridae (Lü, 1998). BmDNVs, which cause serious economic damage to sericulture, were discovered in 1976. They comprise several strains, such as Ina virus, Saku virus, Yamanashi virus and Zhenjiang (China) virus (DNV-Z). As distinguished by Watanabe et al. (1986), Ina virus belongs to the DNV-1 type, and Saku virus and Yamanashi virus belong to the DNV-2 type. Iwashita et al. (1983) discussed the chemical characteristics of DNV-Z. Qian et al.
Figure 1. The flow chart of construction the NIL of nsd-Z.

1985) demonstrated that it is similar to Saku virus. BmDNVs multiply only in the nuclei of the columnar cells of the larval midgut epithelium (Guo et al., 1985; Seki and Iwashita, 1983). Certain silkworm strains are highly susceptible, while other strains are non-susceptible (completely resistant) to DNV-1, DNV-2 or DNV-Z even though they are fed a high concentration of virus inocula. It was reported that a recessive gene, nsd-1, and another dominant one, Nid-1, control the non-susceptibility to DNV-1 severally (Watanabe and Maeda, 1981; Eguchi et al., 1991). nsd-2, a recessive gene conferring resistance to DNV-2, was shown to be different from nsd-1 (Abe et al., 2000). Hu et al. (1984) demonstrated that a third recessive gene, nsd-Z, controls the non-susceptibility to DNV-Z. RAPD markers linked to nsd-Z had been screened (Li et al., 2001) and the molecular linkage map of nsd-Z by SSR markers had been constructed (Li et al., 2006). Two SSR markers, Fl0316 and Fl0568, were the nearest markers on its both sides; they were 4.4cM and 11.7cM from nsd-Z respectively.

In this research, NILs of nsd-Z was bred by backcrossing to recurrent parent successively. SSR markers were selected to detect the homozygosity in the NILs to inspect the replacement of the linkage groups of the donor parent.

**MATERIALS AND METHODS**

**Silkworm strains**

The donor parent, L10 strain (nsd-Z/nd-Z), non-susceptible to DNV-Z, and the recurrent parent, Js strain (+/nsd-Z), susceptible to DNV-Z, were maintained by single pair mating in SRI-CAAS.

**Virus inoculation and serological diagnosis**

Dried silkworm midgut that contained DNV-Z was ground in distilled water until it turned into a dense solution. The solution was filtered through gauze and centrifuged at 3500 rpm for 20 min. After adding an equal volume of 7% acetic acid to the supernatant, the solution was incubated at 25°C for 40 min, followed by adjustment to pH 7.0 and progressive dilution to 0.5% solutions of tissue. The bioassays were conducted at 25°C. DNV-Z was fed to newly hatched larvae for the first instar, followed by rearing on uncontaminated fresh mulberry leaves since the second instar. Normally, silkworms that are susceptible to DNV-Z develop very slowly and cannot survive through the third instar. All of the infected silkworms were checked for DNV by PCR methods (Hou et al., 2005).

**Construction of near allele lines**

To construct a near allele line homozygous for the nsd-Z in a Js background, for the crossing over experiments, a female of Js was crossed with a male of L10, then females of Js were mated to (Js×L10) F1 males or other backcross generations (6 in all) to eliminate the chromosomes of L10 except the region close to the nsd-Z locus. Because of its being recessive, the nsd-Z allele would be lost if the methods of selection were not efficient during the backcrossing. Therefore, in order to ensure that the nsd-Z allele would not be lost, we used a single male moth, which carried the nsd-Z allele, to mate successively with two females of different races from the BC1 generation. In other words, the male moth was first mated to a female moth of Js, and then successively mated to another female moth of L10. The eggs from the two females were reared separately, and the offspring from the backcross to L10 were fed with DNV-Z to test for the presence of nsd-Z. If all of those individuals died, all of its offspring (including those crossed to Js) were discarded. On the contrary, if 50% of the offspring obtained from the backcross to L10 survived, the offspring obtained from the same male moth crossed to Js were reared. Selfing was adopted in the B6 generation (Figure 1).

**DNA extraction**

DNA samples were extracted from the moth. The moth was ground with a mechanical homogenizer in a micro-centrifuge tube and suspended in DNA extraction buffer [50 mM Tris-HCl (pH 8.0), 100 mM NaCl, and 2 mM EDTA] that contained 150 µg/ml proteinase K. After digestion with proteinase K at 50°C for 8 - 10 h, phenol-chloroform extraction was carried out and the DNA was recovered.
Table 1. The achievement of the parents and backcrossing generations.

<table>
<thead>
<tr>
<th>Race/generation</th>
<th>Rearing season</th>
<th>Duration of 5th instar (D:H)</th>
<th>Duration of feeding period (D:H)</th>
<th>Cocoon weight (g)</th>
<th>Cocoon shell weight (g)</th>
<th>Cocoon shell ratio (%)</th>
<th>Ratio of cocoon mortality (%)</th>
<th>Larva-pupa survival rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>2003Au</td>
<td>6:18</td>
<td>21:06</td>
<td>1.283</td>
<td>0.227</td>
<td>17.69</td>
<td>4.32</td>
<td>91.33</td>
</tr>
<tr>
<td>BC1</td>
<td>2004Sp</td>
<td>7:00</td>
<td>22:12</td>
<td>1.591</td>
<td>0.362</td>
<td>22.75</td>
<td>3.87</td>
<td>96.00</td>
</tr>
<tr>
<td>BC2</td>
<td>2004Su</td>
<td>7:12</td>
<td>23:06</td>
<td>1.741</td>
<td>0.411</td>
<td>23.61</td>
<td>3.45</td>
<td>95.67</td>
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<tr>
<td>BC3</td>
<td>2004Au</td>
<td>8:00</td>
<td>23:18</td>
<td>1.882</td>
<td>0.452</td>
<td>24.02</td>
<td>2.23</td>
<td>96.86</td>
</tr>
<tr>
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<td>2004Au</td>
<td>7:12</td>
<td>23:00</td>
<td>1.688</td>
<td>0.401</td>
<td>23.76</td>
<td>4.67</td>
<td>91.23</td>
</tr>
<tr>
<td>BC5</td>
<td>2005Sp</td>
<td>8:12</td>
<td>25:00</td>
<td>1.822</td>
<td>0.441</td>
<td>24.20</td>
<td>4.32</td>
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<td>BC6</td>
<td>2005Su</td>
<td>7:18</td>
<td>24:00</td>
<td>1.791</td>
<td>0.428</td>
<td>23.90</td>
<td>8.55</td>
<td>82.89</td>
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<td>BC5F2</td>
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<td>8:00</td>
<td>24:18</td>
<td>1.876</td>
<td>0.463</td>
<td>24.68</td>
<td>6.75</td>
<td>89.73</td>
</tr>
<tr>
<td>BC5F3</td>
<td>2005LAu</td>
<td>8:12</td>
<td>25:00</td>
<td>1.968</td>
<td>0.491</td>
<td>24.95</td>
<td>2.67</td>
<td>96.59</td>
</tr>
<tr>
<td>Js</td>
<td>2005LAu</td>
<td>8:00</td>
<td>24:18</td>
<td>1.861</td>
<td>0.458</td>
<td>24.61</td>
<td>7.26</td>
<td>88.06</td>
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<tr>
<td>L10</td>
<td>2005Sp</td>
<td>4:00</td>
<td>17:08</td>
<td>0.630</td>
<td>0.070</td>
<td>11.11</td>
<td>2.90</td>
<td>97.00</td>
</tr>
</tbody>
</table>

*D (Day); H Hour; Sp (Spring); Su Summer; Au (Autumn); Lau Late autumn.

by isopropanol precipitation. The purified DNA was dissolved in 0.1 x TE buffer (pH 8.0). The DNA concentration was measured spectrophotometrically (BioPhotometer, Eppendorf) and the samples were diluted to a concentration of 10 ng/µl for use in the PCR analysis.

Methods for detection the homozygosity of the NIL

Two to four polymorphic SSR markers were selected from the constructed SSR linkage map (Miao et al., 2005). In each backcrossing generation, 3 male moths which had been demonstrated to harbor nsd-Z gene and had been selected to generate next generation were detected for their genotype of these SSR markers.

PCR and fragment analyses

The PCR reactions were performed using the Flexigene Cycler (Techne, England). The following PCR conditions were used for the micro-satellite loci: (1) 95ºC for 3 min, 63ºC for 40 s, and 72ºC for 1 min, followed by 14 cycles of 94ºC for 45 s, fourteen-step touchdown decreasing by 0.5ºC at each step to 56ºC (40 s), and 72ºC for 1 min; (2) the conditions for the last 24 cycles were 94ºC for 40 s, 56ºC for 40 s, and 72ºC for 1 min, followed by (3) a final elongation step of 10 min, with extension at 72ºC. The PCR was performed in a final volume of 15 µl that contained 10 mM Tris–HCl (pH 8.4), 50 mM KCl, 1.5 mM MgCl₂, 0.2 mM of each dNTP, 0.2 µM of each primer, 20 ng of genomic DNA, 0.5 U Taq polymerase, and distilled deionized water. The PCR products at about 0.5 ng (0.01 pM) per sample were analyzed in an ABI377 DNA sequencer (ABI PRISM) 2.5.

RESULTS

The characters of the NIL

All of the individuals in these NILs expressed the same characters as Js except the aim gene, nsd-Z. The egg color, larva color, larva body form, cocoon shape, and other characters such as economic characters were all same as Js. Table 1 showed the economic characters of the NIL and the donor parent and recurrent parent.

The detection results of each backcrossing generation

The 3 male moths which had been demonstrated to harbor nsd-Z and had been selected to generation next generation were detected their genotype of the SSR markers. Only 5 pairs of BC6 moths whose 1/4 offspring were nonsusceptible to DNV-Z were detected for the genotype too. The results showed that linkage groups from the recurrent parent in the NILs increased rapidly along with the increase of the backcrossing generations (Figure 2). The average linkage groups which had been replaced by Js were 12, 19.3, 22.7, 24.3 and 26.7 in BC1, BC2, BC3, BC4 and BC6 generation respectively. The linkage groups of 9 of the 5 BC6 pairs, whose 1/4 offspring were non-susceptible to DNV-Z, were all replaced by the recurrent parent except the linkage group that held nsd-Z, and only part of one linkage group from the other one individual had not been replaced, as one marker showed the genotype of L10, while another showed the genotype of Js. According to the markers linked to nsd-Z, Fl0316 had not been replaced in all of the 5 pair moths, the genotype of Fl0568 showed Js type in 2 individuals, indicating crossover happened between Fl0568 and nsd-Z (Figure 2) in them.

DISCUSSION

NIL populations have been used extensively in genetic studies due to the advantages derived from their homozygosity. NILs were important material to locate and clone new genes, in a NIL population, the phenotypic variation
observed between pairs of lines can be assigned directly to the distinct genomic regions introgressed in another similar genetic background. Depending on the desired resolution one can minimize the number of lines by analyzing lines carrying large introgressions or even chromosome substitution strains (Nadeau et al., 2000).

Theoretically, if there were $n$ genes difference between recurrent parent and donor parent, after $m$ times backcrossing, the proportion of homozygotes will account for $[1-(1/2)^m]^n$ (Xu and Zhu, 1994). In this research, the velocity of homozygosity was accord with it but was a little lower than the theoretic value. During the backcrossing, the genotypes of the SSR markers were not same in all of the detected individuals in one generation; their genotypes were stochastic but accorded with Mendel's law.

SSRs are widely used in population genetics and linkage map construction because of their high levels of polymorphism and reproducibility (Tautz, 1989), as well as their genome-wide distribution. SSRs are inherited in a Mendelian fashion and show co-dominant alleles, and had been widely used in constructing linkage map, studying genetic diversity, gene mapping, map-based cloning and marker-assisted selection (Miao et al., 2005; Li et al., 2005; Li et al., 2006; Ren et al., 2005; Danson et al., 2006). In this research, there was no crossover had been detected between nsd-Z and the nearest marker (Fl0316, 4.4cM), but crossover had been detected between nsd-Z and another marker, Fl0568, which located on the other side of nsd-Z, indicated that marker assisted selection could be actualized to breed new silkworm races nonsusceptible to DNV-Z.

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