Genetic mapping of fiber color genes on two brown cotton cultivars in Xinjiang

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Abstract
In the present study, genetic linkage analysis was carried out to map the fiber color loci \(Lc_1\) and \(Lc_2\) on two brown cotton cultivars with SSR and EST-SSR markers in the reference map by \(F_2\) segregation populations. The \(Lc_1\) locus carried by Xincaimian6 (\(Gossypium hirsutum\) L.) was flanked by the marker NAU2862 and NAU1043 on the long arm of Chromosome 07, with genetic distance 7.8 cM and 3.8 cM, respectively. The \(Lc_2\) carried by Xincaimian 5 (\(Gossypium hirsutum\) L.) was flanked by the marker NAU5433 and NAU2968 on the short arm of Chromosome 06, with genetic distance 4.4 cM and 7.4 cM respectively. Moreover, the marker NAU3735 and marker NAU5434 co-segregated with the \(Lc_1\) and the \(Lc_2\) locus, respectively. The results of marker association studies with these two loci provides the basic information for the final isolation of these important genes in colored cotton, and these linkage markers also could facilitate application of marker assisted selection in the future.

Keywords: Colored cotton; Genetic mapping; SSR

Background
Naturally colored cotton, or colored cotton, appears as brown or green colored fiber during the fiber development process. Requiring no or less dying in the textile processing, colored cotton reduces the pollution to the environment; moreover, free from residual chemical toxicant, garments made from colored cotton are more comfortable, softer and healthier for human bodies (Yuan et al. 2012). However, colored cotton is generally inferior to white cotton, especially with respect to fiber quality, including shorter, weaker fiber and lower micronaire. There is typically a negative correlation between fiber color and fiber quality traits mainly due to the pleiotropic effects of fiber color genes. For example, the deeper fiber color is, the lower the fiber quality. It is a challenge for researchers and breeders to break the negative correlation between the fiber color and fiber quality. Mapping and cloning of fiber color genes should substantially give a clue to resolve that problem mentioned above.

Kohel carried out traditional genetics research on some brown cotton materials collected from all over the world and discovered that brown fiber was controlled by six loci (\(Lc_1 \sim Lc_6\)). \(Lc_1\) and \(Lc_2\) control the brown color of lint, \(Lc_3\) controls dark brown, \(Lc_4\), \(Lc_5\) and \(Lc_6\) control the light brown. Furthermore, he assigned \(Lc_1\) to chromosome 7 and \(Lc_2\) to chromosome 6 by linkage to morphological markers (Kohel 1985). In this study, we aimed to map \(Lc_1\) and \(Lc_2\) loci to a detailed reference molecular map (Rong et al. 2004). Complete diallel cross between four white cotton cultivars (Xinluza13, Xinluza31, Zhongmiansuo41 and Zhongmiansuo45) (\(Gossypium hirsutum\) L.) and two elite brown cotton cultivars (Xincaimian5 and Xincaimian6) (\(Gossypium hirsutum\) L.) cultivated in Xinjiang were conducted, and sixteen populations were obtained. Furthermore, we used the mapping populations which were in accordance with the 3:1 Mendelian inheritance (\(\chi^2_{0.05} < 3.84, df = 1\)) to locate the \(Lc_1\) and \(Lc_2\) genes in the cotton reference map. This research is a start point for further defining a finer location of the fiber color genes that may result in the eventual cloning of these genes by map-based and candidate gene approaches.

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Materials and methods

Plant materials
In the summer of 2007, parental lines (Table 1) were crossed in all combinations including reciprocals. The F1 plants were self-pollinated to produce the F2 populations in the second year. In the summer of 2009, a total of sixteen F2 populations were grown, and field evaluation and genetic analysis were undertaken. All field experiments were carried out in Korla Breeding Base of China Colored-Cotton (Group) Co., Ltd.

Cotton DNA extraction
The genomic DNA from the parental cultivars or lines and their F2 segregating populations were extracted from the young leaves by the CTAB method (Paterson et al. 1993; Zhang et al. 2000).

SSR-PCR analysis and genetic mapping
Sixty-five and seventy-one pairs of cotton simple sequence repeat (SSR) primers on Chromosome 06 and Chromosome 07 were chosen according to primer information published on Cotton Marker Database (CMD) (http://www.cottongen.org/data/markers). PCRs were performed using a PTC-200 (Bio-Rad, Hercules, CA, USA) thermocycler. The genotype of the white parent was scored as “1”, the brown parent and the heterozygous (F1) genotype were scored as “2” and “3” in the parents, F1 and F2 populations. Missing data were noted as “-”. The χ² test for goodness of fit was used to assess the Mendelian dominant inheritance in the F2 segregating population. After eliminating the segregation distortion markers by χ² test, other SSR markers were detected by linkage analysis.

The volume of PCR reaction system was 20 μl, including 2 μl of 10× PCR buffer (containing Mg²⁺), 1 μl of dNTPs (10 mM), 0.3 U of Taq enzyme (2 U/μl), 0.5 μl of SSR upstream primers and downstream primers (5 μM) and 1 μl of template DNA (40 ng/μl), 14.7 μl of ddH₂O. Reaction procedures were 94°C for 4 min; 94°C for 40 s, 57°C for 45 s, 72°C for 50 s, 35 cycles; 72°C for 10 min, and the amplified products were preserved at 4°C. Amplified sample was mixed with 10 μl of loading buffer, denatured at 95°C for 10 min, the mixture was immediately transferred to ice-bath cooling for electrophoresis. Vertical slab denaturing polyacrylamide gel (7%) electrophoresis was used to separate the SSR amplified products, pre-electrophoresis at 120 V for 10 min, and 180 V constant voltage electrophoresis for 4 h. DNA fragments were detected with ethidium bromide staining.

Data analysis
SPSS13.0 software was used to test for segregation ratio of selected markers in F2 segregating population and Mapmaker 3.0 software was employed to construct the genetic linkage map, and all linkage groups were determined at LOD scores ≥6. Mapping was completed using MapDraw software (Lander et al. 1987).

Results

Inheritance of fiber color
In each population, all plants in the F1 generation displayed the same phenotype as the brown parent. This result indicates that the brown fiber trait was determined by nuclear inheritance and brown was dominant over white. Segregation analysis of F2 segregation populations was accomplished by visual inspection of the lint color of individual plants in each population. Five F2 populations were consistent with a 3:1 ratio at significant level of α = 0.05 (Table 2). Therefore, these five populations were used for mapping. Totally, 65 and 71 SSR markers on Chromosome 06 and Chromosome 07 respectively were employed to screen polymorphisms among all six parental lines. Sixteen polymorphic marker loci, 9 on Chromosome 06 and 7 on Chromosome 07, were identified. Then, the polymorphic SSR markers between the two parental lines were run on the corresponding F2 populations and the marker genotypes were recorded. The mapping results were shown as follows.

Genetic mapping of Lc1
Linkage analysis suggested that the fiber color gene carried by Xincaimian6, Lc1, is preliminarily located between NAU3654 and MS58 on the long arm of Chromosome 07, based on analysis of three F2 populations derived from Zhongmiansuo41 × Xincaimian6, Xincaimian6 × Xinzhuo31 and Xinzhuo31 × Xinzhuo41. To narrow

Table 1 The information of parental lines for constructing the F2 populations

| Lint phenotype | Cultivars       | Fiber color     | Material source                        |
|---------------|----------------|-----------------|----------------------------------------|
| Brown         | Xincaimian5    | Dark brown      | China Colored-Cotton Group             |
|               | Xincaimian6    | Light brown     | China Colored-Cotton Group             |
|               | Xinzhuo31      | White           | Xinjiang Kuitunwansi Seed Industry     |
| White         | Zhongmiansuo41 | White           | Cotton Research Institute of China     |
|               | Zhongmiansuo45 | White           | Cotton Research Institute of China     |
down the \( Lc_1 \) locus region further, three SSR markers, NAU2862, NAU3735 and NAU1043, were detected as DNA polymorphisms between Xincaimian6 and Xinluza31 (Figure 1A). \( Lc_1 \) co-segregated with the marker NAU3735, and flanked by markers NAU2862 and NAU1043, with genetic distance of 7.8 cM and 3.8 cM, respectively (Figure 2A).

**Genetic mapping of \( Lc_2 \)**

Linkage analysis also suggested that the fiber color gene carried by Xincaimian5, \( Lc_2 \), is preliminarily located between CIR329 and NAU905 in the short arm of Chromosome 06 by \( F_2 \) populations derived from Zhongmiansuo41 × Xincaimian5 and Xincaimian5 × Xinluza31. To narrow down the \( Lc_2 \) locus region further, four SSR markers, NAU5373, NAU5433, NAU5434 and NAU2968, were detected as DNA polymorphisms between Zhongmiansuo41 and Xincaimian5 (Figure 1B). \( Lc_2 \) co-segregated with the marker NAU5434, and flanked by NAU5433 and NAU2968, with genetic distance of 4.4 cM and 7.4 cM, respectively (Figure 2B).

**Discussion**

The inheritance of cotton fiber color trait has been studied in several reports and a few of genetic loci potentially involved in fiber color formation have been identified (Harland 1935; Симонгулян 1984; Kohel 1985; Shi et al. 2002). Harland (1935) found that the inter-barbadense cross Egyptian brown × Sea Island white gave \( F_1 \) intermediate and complicated segregation of the blending type in \( F_2 \) and the factor \( K^B \) (brown lint) of the brown parent was accompanied by a number of plus modifiers absent in the white parent. He also concluded that brown lint in barbadense and hirsutum was not determined by the same gene, but by a pair of duplicate genes. Симонгулян (1984) made a conclusion through

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**Table 2 White/brown cross combinations consisted with Mendel segregation ratio**

| Parental combination | \( F_2 \) | Brown: white | \( \chi^2 \) |
|----------------------|-----------|-------------|-------------|
| Xinluzao13 Xincaimian6 | 30 85 | 2.83:1 | 0.072 |
| Xinluzao31 Xincaimian6 | 27 82 | 3.07:1 | 0.012 |
| Zhongmiansuo41 Xincaimian6 | 27 90 | 3.33:1 | 0.231 |
| Zhongmiansuo45 Xincaimian6 | 36 84 | 2.33:1 | 1.600 |
| Xincaimian6 Xinluzao13 | 25 78 | 3.12:1 | 0.029 |
| Xincaimian6 Xinluzao31 | 26 76 | 2.92:1 | 0.013 |
| Xincaimian6 Zhongmiansuo41 | 17 100 | 6:1 | – |
| Xincaimian6 Zhongmiansuo45 | 23 90 | 4:1 | – |
| Xinluzao13 Xincaimian5 | 40 72 | 1:1 | – |
| Xinluzao31 Xincaimian5 | 20 79 | 3:95:1 | – |
| Zhongmiansuo41 Xincaimian5 | 34 93 | 2.74:1 | 0.08 |
| Zhongmiansuo45 Xincaimian5 | 11 88 | 8:1 | – |
| Xincaimian5 Xinluzao13 | 37 81 | 2:1:9 | – |
| Xincaimian5 Xinluzao31 | 28 92 | 3.32:1 | 0.223 |
| Xincaimian5 Zhongmiansuo41 | 17 100 | 5:88:1 | – |
| Xincaimian5 Zhongmiansuo45 | 23 96 | 4:17:1 | – |

Note: \( \chi^2_{0.05} = 3.84, \text{df} = 1 \). Bold numbers represent \( F_2 \) populations which were used for linkage analysis and genetic mapping of \( Lc_1 \) and \( Lc_2 \).

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**Figure 1** PCR amplification products generated by SSR primer pairs in \( F_2 \) segregation populations. A. Segregation of SSR marker NAU1043 in \( F_2 \) population from Xincaimian6 × Xinluzao31. Lane M is a 100-bp molecular weight marker. Lanes 2, 3, and 4 are parental lines Xinluzao31, Xincaimian6 and \( F_1 \). Lanes 5–25 are a subset of the \( F_2 \) individuals; B. Segregation of SSR marker NAU5433 in \( F_2 \) population from Zhongmiansuo41 × Xincaimian5. Lane M is a 100-bp molecular weight marker. Lanes 2, 3, and 4 are parental lines Zhongmiansuo41, Xincaimian5 and \( F_1 \). Lanes 5–32 are a subset of the \( F_2 \) individuals.
the analysis of a hybrid between white lint upland cotton and brown semi-wild Mexican species (*G. hirsutum*) that the brown fiber was controlled by two pairs of complementary major genes *Lc*<sub>1</sub> and *Lc*<sub>2</sub>, lack of any pair of dominant alleles lint became white in color, and another gene *Lc*<sub>3</sub> was a supplementary gene which might strengthen the function of these two pairs of genes. Shi et al. (2002) concluded that brown and green colored lint were controlled by one pair of major genes incomplete in dominance on non-homologous chromosomes, and that there were genetic interactions between lint and fuzz coloring genes. Interestingly, in our studies, we noticed that some F<sub>2</sub> populations fitted a 3:1 ratio regardless of cross or reverse cross, e.g. a combination of Xinluzao 31 and Xincaimian 6, but some do not, e.g. a combination of Zhongmiansuo 41 and Xincaimian 5. Therefore, we deduced that modification of minor genes resulted in the difference in proportion of F<sub>2</sub> in reciprocal crosses.

It has been previously demonstrated that brown cotton varieties in Xinjiang had nearer genetic relationships with upland cotton (*G. hirsutum*), but far from Sea-island cotton (*G. barbadense*) (Wang et al. 2012). For many years, breeders have tried to improve the fiber quality of colored cotton by introgressing fiber quality traits from sea island cotton, but the results were poor (Lacape et al. 2005). For map-based cloning, because chromosome variability of target genes can be ensured between upland cotton (*G. hirsutum*) and sea island cotton (*G. barbadense*), and genetic map of both cotton species can be fully utilized as well, *G. hirsutum × G. barbadense* populations are usually chosen in map-based cloning (Park et al. 2005). However, in fact, few progenies of F<sub>1</sub> were fertile, and severe segregation distortion was observed in F<sub>2</sub> populations by brown cotton crossing various sea island cultivars. Distant hybridization-sterility between current brown cotton cultivars and Sea-island cotton cultivar is a bottleneck not only for the improvement of fiber quality but also for map-based cloning fiber color genes in colored cotton (Zhang et al. 1994).

Another of the major limitations to map-based cloning for the genes of interest in cotton is the lack of polymorphism resulting from a narrow genetic base. Ling et al. reported that genetic similarities based on Jaccard’s similarity coefficient between brown cotton and white upland cotton was an average of 0.7 using sequence-related amplified polymorphism (SRAP) markers, which suggests that these cultivars are very closely related concerning their genetic background or they have a common ancestor (Ling et al. 2009; Guo et al. 2004). In China, three major cotton growing agro-ecological zones have been divided based on cotton type, distribution and growth environment, including the northwest inland cotton region, the Yellow River valley region and Yangtze River valley region. Presently, according to the pedigrees known, brown cotton cultivars had nearer
genetic relationships with native upland cotton varieties in Xinjiang than those cultivars in the other two regions. Therefore, the cotton varieties in the other two regions are preferred to choose as parental lines for constructing segregation populations; on the other hand, more markers, such as single nucleotide polymorphism (SNP), cleaved amplified polymorphic sequences (CAPS) markers, need to be developed for further fine mapping. Many studies covering physiology, biochemistry, cell genetics and conventional breeding have been reported using colored cotton as subject materials (Qiu 2004).

In conclusion, we fulfilled the primary purpose of mapping the Lc1 and Lc2 loci using SSR markers. These findings could be used for marker-assistant selection breeding. To the best of our knowledge, this is the first effort at mapping fiber color genes of brown cotton cultivars using SSR markers. Fine mapping will be further carried on by enlarging the F2 populations constructed by near-isogenic line and developing the new markers. With the more sequences release of tetraploid cotton, final cloning of the fiber color genes would help us to understand the complex molecular mechanism of color development in cotton fiber.

Abbreviations

SSR: Simple sequence repeats; CMD: Cotton marker database; SRAP: Sequence-related amplified polymorphism; CAPS: Cleaved amplified polymorphic sequences.

Competing interests

All authors declare that they have no competing interests in regard to this manuscript.

Authors’ contributions

LW, HL, CL, and XX carried out the experiments and participated in data analysis. HL, XYL, XA, and CL carried out the collection of materials and statistical analysis. LZ participated in the design of the study. XBL conceived of the study, and participated in its design, coordination and data analysis. LW, XBL and XX wrote the final version of the manuscript. All authors read and approved the final version of the manuscript.

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