Fine mapping of the QTL cqSPDA2 for chlorophyll content in Brassica napus L.

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Abstract

**Background:** Chlorophyll is the most important factor enabling plants to absorb, transfer and transform light energy and plays an important role in yield formation. *Brassica napus* is one of the most important oil crops. Breeding *Brassica napus* for high light efficiency by improving photosynthetic efficiency has considerable social and economic value. In *Brassica napus*, there have been studies of the initial location of chlorophyll in seed embryos and pericarps, but there are few reports on the fine mapping of chlorophyll QTLs. We constructed near-isogenic lines (NIL), fine-mapped a chlorophyll locus, and evaluated the effect of this dominant locus on agronomic traits.

**Results:** The *cqSPDA2* locus was mapped to an interval of 21.87-22.91 Mb on the chromosome A02 of *Brassica napus* using doubled haploid (DH) lines. To fine-map *cqSPDA2*, we built NIL and designed Indel primers covering the mapping interval. The 469 individuals in the BC\(_3\)F\(_2\) population were analyzed using these indel primers. Among the indel primers, 15 could narrow the mapping interval to 188 kb between Indel3 and Indel15. Next, 16 indel primers and 19 SSR primers were designed within the new narrower mapping interval, and 5 of the primer-amplified fragments were found to be polymorphic and tightly linked to the *cqSPDA2* locus in the BC\(_4\)F\(_2\) population. The mapping interval was narrowed to 152 kb on A02 between SSR2 and Indel15. By gene expression analysis, we found three annotated genes in the mapping interval, including *BnaA02g30260D*, *BnaA02g30290D* and *BnaA02g30310D*, which may be responsible for chlorophyll synthesis.

**Conclusions:** The locus *cqSPDA2*, a dominant QTL for chlorophyll content in *Brassica napus*, was fine-mapped to a 21.89-22.04 Mb interval on A02. Three annotated genes (*BnaA02g30260D*, *BnaA02g30290D* and *BnaA02g30310D*) that may be responsible for chlorophyll synthesis were found.

Background

The material basis of crop yield formation is derived from photosynthesis, and high yield based on photosynthesis has long been a hot topic in crop breeding [1]. Chlorophyll is the most important factor enabling plants to absorb, transfer and transform light energy and plays an important role in the growth and development of plants [2]. Maintaining a high level of chlorophyll content in leaves is an important factor in increasing photosynthetic activity [3]. Within a certain range, there is a positive correlation between chlorophyll content and photosynthetic rate, which directly determines the yield [4, 5]. Therefore, chlorophyll content plays an important role in yield formation [6, 7]. Robust seedling development in *Brassica napus* also leads to higher yield stability and has a high importance for plant breeders.

Chlorophyll content is a quantitative characteristic that is primarily controlled by nuclear genes and has high heritability[8,9]. Previous studies have suggested that at least 27 genes are involved in 15 steps of chlorophyll synthesis[10], and biosynthetic defects are considered to be one of the main reasons for low chlorophyll content. Other reasons for chlorophyll deficiency include deficient signal transduction,
restrained heme feedback, harmful photooxidation and so on. Overall, the molecular mechanisms of chlorophyll synthesis are very complex[11]. In recent years, researchers have analyzed QTLs for chlorophyll content in the seedling leaves of different populations of multiple crops from different perspectives, making considerable progress and establishing a foundation for future research attempting to elucidate the molecular genetic mechanisms that determine of chlorophyll content [12-18].

The completion of the whole genome sequencing of *Brassica napus* indicates that research on the *Brassica napus* genome has entered a new era. In recent years, with the rapid development of molecular marker technology, it has become possible to construct a high-density molecular marker genetic map of *Brassica napus*. Therefore, the study of chlorophyll QTLs that will allow efficient breeding of *Brassica napus* for greater light efficiency will provide important information for improving the yield potential and direction of high-yield *Brassica napus* breeding in the future. Breeding *Brassica napus* for high light efficiency by improving photosynthetic efficiency is socially and economically important. At present, studies of the initial location of chlorophyll content in the seed embryos [19] and pericarps [20] of winter *Brassica napus* have been conducted, QTLs have been identified under drought and salt stress, and candidate genes related to salt tolerance have even been predicted [21-23]. However, there are few reports of the fine mapping of chlorophyll content QTLs in *Brassica napus*[11].

In a previous study, we discovered a dominant major QTL named *cqSPDA2* located in 21.87-22.91 Mb on A02 using DH lines derived from a cross between Zhongshuang11 (ZS11—a semi-winter variety) and QU (a spring variety), which explained 15.72% variance of phenotype and was detected in six environments stably (under review). In this study, a near-isogenic line (NIL) population for *cqSPDA2* was constructed to further narrow the interval of *cqSPDA2*. This study established a foundation for the cloning of chlorophyll genes controlling photosynthetic function and provided a theoretical basis for improving germplasm resources and selecting new high-yield varieties with molecular markers.

**Results**

*Phenotypic and genetic analysis*

The first fully developed leaves counting from the tops of the 2061 individuals in the BC*$_4$F$_2$* population at the six-leaf stage were measured by SPAD (SPAD 502, Japan). The distribution of chlorophyll content in the BC*$_4$F$_2$* population was bimodal, with the lowest point of the column chart as the dividing line (SPAD=43). SPAD$\geq 43$ was regarded as the high chlorophyll content (n=1514), and SPAD$\leq 43$ was regarded as the low chlorophyll content (n=547). A chi-square test showed that the segregation pattern of the chlorophyll content trait was in keeping with the expected Mendelian segregation ratio of 3:1 ($\chi^2=2.53$) (high-chlorophyll content vs. low-chlorophyll content) (Fig. 1). Among the random selection of 198 individuals in the BC$_6$F$_1$ population, the markers SSR2 and Indel100 were used to validate the effect of *cqSPDA2*. The result of a chi-square test was in keeping with a 1:1 ($\chi^2=1.46; \chi^2=1.82$) (AA:Aa) Mendelian ratio (Additional file 1: Table S1).
**Fine mapping of cqSPDA2**

To fine-map the cqSPDA2 locus and identify the candidate genes, 87 primer pairs of indel markers were designed to uniformly cover the preliminary mapping interval (A02 21.87-22.91Mb). As a result, 28 polymorphic markers were detected in the two parental lines and selected DH lines. Twenty-three of these markers were found to cosegregate in the DH lines. The 469 individuals in the BC3F2 population were then analyzed using these indel primers. The linkage map constructed using the indel data and corresponding chlorophyll content phenotypes showed that 15 of the indel primers were tightly linked to the cqSPDA2 locus (Additional file 2: Table S2). The cqSPDA2 locus was delimited to an interval of 5.2 cM between Indel3 and Indel15 (Additional file 3: Fig. S1). The fragments amplified by the primers Indel3, Indel6, Indel15 and Indel 17 near the QTL were recovered. TA cloning was performed with the PMD18T vector, and the physical location of the region was found to be within the 188-kb range of 21.88-22.07 Mb (Additional file 4: Fig. S2).

Next, 16 indel and 19 SSR primers were designed within the new, narrow mapping interval, and the fragments amplified from 5 primers were polymorphic and tightly linked to the cqSPDA2 locus (Additional file 5: Table S3). These new primers helped to narrow the interval for the 250 individuals in the BC4F2 population. As a result, the cqSPDA2 locus was mapped to a 152-kb interval between SSR2 and Indel15 (Fig. 2). BSNP88 and BSNP90 were developed competitive allele-specific PCR (KASP) markers based on SNP analysis in this interval (Additional file 6: Table S4). SSR2 is a codominant marker closely linked to cqSPDA2. Twenty BC4F2 plants with low-, medium- and high-chlorophyll phenotypes were selected, and their SSR2 genotypes were determined. The results showed that the three groups of different phenotypes can be divided into three genotypes: AA (dominant homozygous, high-chlorophyll phenotype), Aa (heterozygote, medium-chlorophyll phenotype) and aa (recessive homozygous, low-chlorophyll phenotype). These results suggested that SSR2 was closely linked to cqSPDA2 and could be effectively used in marker-assisted selection (MAS) (Fig. 3).

**Quantitative RT-PCR of genes in the mapping interval**

According to the Brassica napus genome annotation database (http://www.genoscope.cns.fr/brassicanapus/), twenty-seven genes were identified in the targeted mapping interval of 21.89-22.04 Mb on A02 (Additional file 7: Table S5). Quantitative real-time PCR (qRT-PCR) was used to identify the expression levels of the genes in the targeted mapping interval between the leaves of different genotypes. The melting and amplification curves of twenty-seven genes were analyzed, and the results showed that 24 primer pairs could be used to analyze gene expression (Additional file 8: Table S6). Thus, twenty-four genes in the mapping interval and the housekeeping gene Actin7 were quantified by qRT-PCR (Additional file 9: Table S7). The results showed that the expression levels of the three genes (BnaA02g30260D, BnaA02g30290D and BnaA02g30310D) were all higher in ZS11 and BC4F2:3 (aa) than in QU and BC4F2:3 (AA) at three stages. Student’s t-test was used to compare QU with ZS11 and BC4F2:3 (AA) with...
BC₄F₂:3 (aa). \textit{BnaA02g30290D} and \textit{BnaA02g30310D} showed significant differences in expression between BC₄F₂:3 (AA) and BC₄F₂:3 (aa) at the 6-leaf stage (p<0.05), and the difference in \textit{BnaA02g30260D} was highly significant (p<0.01) (Fig. 4). There was no consistent expression difference in the other genes tested among QU, BC₄F₂:3 (AA), ZS11 and BC₄F₂:3 (aa) at the three stages. Therefore, \textit{BnaA02g30260D}, \textit{BnaA02g30290D} and \textit{BnaA02g30310D} were likely candidate genes for \textit{cqSPDA2}.

\textbf{Agronomic traits analysis}

To investigate the effect of \textit{cqSPDA2} on agronomic traits, 50 plants with the AA genotype (high-chlorophyll content) and 50 plants with the aa genotype (low-chlorophyll content) were selected from the BC₄F₂ population by molecular marker and SPAD analysis. We investigated plant height, silique length, number of seeds per silique, number of siliques per plant, 1000-grain weight, and individual plant yield. The results showed that plant height, number of seeds per silique, number of siliques per plant and individual plant yield were highly significantly different between the AA-genotype plants and the aa-genotype plants (P<0.01). There was a difference in 1000-grain weight but no difference in silique length between groups (P<0.05) (Table 1).

\textbf{Discussion}

\textbf{Significance and QTL analysis of chlorophyll content}

The leaf is the main photosynthetic organ in plants, and its chlorophyll content is an important agronomic trait for crop yield. Ninety to ninety-five percent of plant dry matter is produced by photosynthesis, and crop yield is primarily derived from the photosynthetic products of leaves [24]. Chlorophyll is an important pigment involved in photosynthesis in chloroplasts, which can absorb and transform light energy, and an important index used to evaluate leaf photosynthetic capacity [25]. Increasing crop yield by increasing chlorophyll content is one of the important objectives of high-light-efficiency breeding [26]. The emergence of high-throughput sequencing technology provides a new method for the development of molecular markers. Molecular marker technology promotes rapid genetic map construction, and provides convenient data for fine mapping. Chlorophyll content is a quantitative characteristic that is primarily controlled by nuclear genes and has high heritability. QTL mapping has been proved to be an efficient way to reveal the genetic mechanisms underlying the chlorophyll content trait. At present, research on chlorophyll QTLs has been performed in different populations of various crops, such as rice [12, 14, 16, 17], wheat [13, 15], beans [27, 28], and cabbage [18], and especially rice [17]. Previous studies have examined the location of chlorophyll in the embryo [19] and pericarp [20] of winter \textit{Brassica napus}. In addition, QTLs were detected under drought and salt stress conditions, and a candidate gene related to salt tolerance was predicted [21-23]. Wang et al.[11] detected a QTL on chromosome C08 in \textit{Brassica napus} using chlorophyll deficient mutants by measuring the absolute chlorophyll content in leaves.

\textbf{Fine mapping using NILs and MAS}
The development of NILs is a productive strategy for fine mapping and evaluating genetic effects during chlorophyll QTL studies[29]. In this study, a NIL population for the cqSPDA2 regions was constructed using ZS11 as the recurrent parent with flanking markers. The flanking markers' Indel1, 3, 15, 87 were used to carry out foreground selection for construction the population of BC₃F₂ and BC₄F₂ populations, and the mapping interval was reduced to 152 kb (Fig. 2). This strategy for mapping the target genes is reasonable, inexpensive, and highly efficient. Based on BC₄F₂ phenotype and BC₆F₁ genotype analysis, the segregation of chlorophyll content was consistent with chi-square test (Fig. 1 and Additional file 1: Table S1). These results indicated that cqSPDA2 is the dominant QTL controlling chlorophyll content in Brassica napus. The NIL analysis also showed positive correlations between cqSPDA2 and agronomic traits such as yield, plant height, seeds per silique and siliques per plant (Table 1).

Molecular marker SSR2 with close linkage was developed through fine mapping (Fig. 3). Indel 15 was also next to cqSPDA2, these markers could accelerate the breeding process for molecular marker assisted selection in the future.

**Candidate genes prediction**

According to qRT-PCR analysis, BnaA02g30260D, BnaA02g30290D and BnaA02g30310D were suitable candidate genes for cqSPDA2 among the twenty-nine genes annotated in the Brassica napus genome in the mapping interval (Additional file 9: Table S7). BnaA02g30260D, which is part of a disease-resistance protein family, has transmembrane receptor activity, nucleoside-triphosphatase activity, nucleotide binding function, and ATP binding function and is involved in signal transduction, defense response, apoptosis, and innate immune response according to its annotations. Further study is needed to determine whether BnaA02g30260D affects chlorophyll synthesis. BnaA02g30290D is FK506- and rapamycin-binding protein 15 kD-2 (FKBP15-2) and has peptidyl-prolyl cis-trans isomerase activity related to protein folding. Luan et al. [30] found that AtFKBP15-1 and AtFKBP15-2 had the highest homology to FKBP13 and encoded functional homologs of FKBP13. AtFKBP13 was reported to be associated with Rieske protein both before and after the import of proteins into the chloroplast stroma, and AtFKBP13 can play a role in the downregulation of Rieske protein accumulation. Rieske is a subunit of the cytochrome b₆f complex, which is one of the four complexes of the photosynthetic electron transport chain [31]. It was also reported that when ScFKBP12 was transferred into Arabidopsis, chloroplast formation and the expression of genes related to chloroplast formation were inhibited [32]. In this study, the expression levels of BnaA02g30290D (AtFKBP15-2) in the NIL (aa) and ZS11 plants were all higher than those in the NIL (AA) and QU at the three stages; if this gene inhibited the formation of chlorophyll, this finding would be consistent with the abovementioned results. BnaA02g30310D is homologous to GCH-1 in Arabidopsis thaliana. GCH-1 is the first enzyme in tetrahydrobiopterin (BH4) biosynthesis [33]. BH4 is an essential coenzyme for all three kinds of nitric oxide synthase (NOS) [34]. AtNOA1 (AtNOS1) is located in Arabidopsis chloroplasts, and OsNOA1 (OsNOS1) is also located in rice chloroplasts [35-37]. Yang et al. [37] found that the chlorophyll content decreased with increasing OsNOA1 at a low temperature (22 °C). He [38] suggested that OsNOA1 directly regulates chloroplast-encoded proteins by affecting the function of the chloroplast ribosome and then transmits a signal to the nucleus through the chloroplast retrograde
signaling pathway mediated by Mg protoporphyrin IX, further affecting the expression of chloroplast proteins encoded by the nuclear genes. The location of this study, Qinghai Province, is on the Qinghai-Tibet Plateau, where the average temperature during crop growth is low. In this location, the expression level of \textit{BnaA02g30310D} (\textit{GCH-1}) in NIL (aa) and ZS11 was high at the three stages observed, especially at the six-leaf stage, which is consistent with the results obtained by He [37, 38].

Additional experiments, such as transgenic complementation test, CRISPR/Cas9, VIGS and RNAi, are warranted to investigate whether \textit{BnaA02g30260D}, \textit{BnaA02g30290D} and \textit{BnaA02g30310D} are the genes underlying \textit{cqSPDA2}. Analysis of the regulatory network controlling chlorophyll synthesis will facilitate molecular breeding of \textit{Brassica napus} for high yield.

**Conclusions**

In this study, we constructed NILs and narrowed the interval of \textit{cqSPDA2} to 152 kb on A02 between SSR2 and Indel15. According to the \textit{Brassica napus} genome annotation database, there were twenty-seven genes in this mapping interval. \textit{BnaA02g30260D}, \textit{BnaA02g30290D} and \textit{BnaA02g30310D} were identified as suitable candidate genes for \textit{cqSPDA2} according to a qRT-PCR analysis, and thus, these genes may be responsible for chlorophyll synthesis. In addition, the dominant locus \textit{cqSPDA2} has positive effects on agronomic traits.

**Methods**

**Plant materials**

The leaves of ZS11 have low chlorophyll content, and the leaves of QU have high chlorophyll content (Fig. 5). To obtain a relatively simple genetic background and to fine-map \textit{cqSPDA2}, we constructed an NIL population. An F\textsubscript{1} line with the QU genotype in the \textit{cqSPDA2} region was selected and backcrossed to ZS11 for three generations. BC\textsubscript{3}F\textsubscript{1} individuals were selfed to generate a BC\textsubscript{3}F\textsubscript{2} mapping population backcrossed to ZS11. The flanking markers Indel1 and Indel87 were used to construct the NIL population with foreground selection (Additional file 2: Table S2).

The BC\textsubscript{4}F\textsubscript{1} individuals with a QU genetic background in the \textit{cqSPDA2} region as selected by the flanking markers Indel3 and Indel15 (Additional file 3: Fig. S1) were selfed to generate a BC\textsubscript{4}F\textsubscript{2} population, which was used for fine mapping of the \textit{cqSPDA2} locus. The detailed process of population development is illustrated in Additional file 12: Fig. S4. BC\textsubscript{4}F\textsubscript{2:3} individuals with the AA genotype (homozygous for \textit{cqSPDA2}) and aa genotype (without \textit{cqSPDA2}) were detected by the flanking markers Indel3 and Indel15 and subjected to qRT-PCR analysis. In addition, each population was grown in the experimental plots with a spacing of 30 cm (between rows) × 15 cm (within rows). The BC\textsubscript{3}F\textsubscript{2} and BC\textsubscript{4}F\textsubscript{2} populations were grown at the same density in fields in Yuanmou, Yunnan (altitude 898 m, 101°52′N, 25°42′E) and Xining, Qinghai (altitude 2225 m, 101°49′N, 36°34′E), respectively. The BC\textsubscript{4}F\textsubscript{2:3} and BC\textsubscript{6}F\textsubscript{1} populations were grown in a
greenhouse at the Academy of Agricultural and Forestry Sciences, Qinghai University (Xining, Qinghai, China). Standard crop management practices were followed.

**Phenotypic trait and data analysis**

The testing targets included every plant of the population that was lacking diseases and insect pests. According to the previous method of chlorophyll determination, we measured three positions in the distal third of the first fully developed leaf from the top of each plant at the six-leaf stage by SPAD (SPAD 502, Japan) during the 9:00-11:00 am interval [39]. Each measurement was repeated three times per leaf, and the veins were avoided. Statistical analysis was performed with Excel. Chi-square tests were performed on the segregation data to determine the genetic regulation of the chlorophyll content.

**DNA extraction and development of molecular markers development**

Total DNA was extracted from fresh leaves using the CTAB method [40]. PCR was performed in a 20-μL reaction solution containing 2 μL DNA, 2 μL 2 mM dNTPs, 2 μL 10× PCR buffer, 1 μL Taq, 1 μL of 2 μM forward and reverse primers and 12 μL of ddH₂O. The PCR program was carried out according to Yang's method with minor modifications [41]. The PCR products were separated on 6% non-denatured polyacrylamide gels and detected by silver staining [42]. Indel (insertion/deletion) markers were developed from the resequencing data of the parents according to the *B. napus* ‘Darmor-bzh’ reference genome sequence in the primary mapping interval. SSR markers were developed based on the *B. napus* ‘Darmor-bzh’ reference genome sequence corresponding to the interval. The sequences of the SSR markers were designed using SSR Hunter 1.3 and Primer Premier 5.0 [43,44].

**Mapping of the cqSPDA2 locus**

The BC$_3$F$_2$ and BC$_4$F$_2$ family populations were used to fine map the cqSPDA2 locus using indel and SSR markers. First, we designed 87 indel markers in the primer mapping interval (21.87-22.91 Mb on chromosome A02) to map the cqSPDA2 locus. A linkage map for the cqSPDA2 locus was constructed using JoinMap 4.0 [45]. The mapping interval for the cqSPDA2 locus was gradually reduced using the mapping results of the BC$_3$F$_2$ population. Finally, additional indel and SSR markers within the new narrow mapping interval were designed to fine-map the cqSPDA2 locus based on the BC$_4$F$_2$ population with WinQTLCart 2.5. The physical location was obtained by blasting the *Brassica napus* genome database using the indel and SSR sequences. The physical linkage map was produced with MapDraw 2.1 [46].
TA cloning

The specific markers closely linked to \( cqSPDA2 \) were sequenced by NIL population scanning. Specific fragments were collected according to Yi et al. [47]. The product was ligated into the pDM18-T vector (Takara), and the transformed clone was detected with M13 primers. Six positive clones were randomly selected and sequenced by Sangon Biotech (Shanghai) Co., Ltd. [48].

Genes in the mapping interval

All genes within the targeted mapping interval on A02 were identified using annotations from the \textit{Brassica napus} genomes (http://www.genoscope.cns.fr/brassicanapus/) and annotated according to the BRAD annotations (http://brassicadb.org/brad/blastPage.php). The homologous sequences were aligned using BLASTN (http://blast.ncbi.nlm.nih.gov/).

RNA extraction and qRT-PCR analysis

\( \text{BC}_4F_{2:3} \) (91 AA-genotype plants homozygous for \( cqSPDA2 \) and 104 aa-genotype plants without \( cqSPDA2 \)) were grown in a greenhouse in September 2019. Total RNA was isolated from the first fully developed leaves counted from the top of each plant (at the 4-leaf stage, 6-leaf stage and squaring stage) of the \( \text{BC}_4F_{2:3} \) population and the parental lines using TRNzol-A+ Total RNA Reagent (Takara, Dalian, China) according to the manufacturer’s protocol. RNA integrity was monitored using 1% agarose gel electrophoresis. cDNA was obtained via reverse transcription of total RNA using the PrimeScript RT Reagent Kit (Takara, Dalian, China) and following the manufacturer’s instructions.

We performed a qRT-PCR analysis to identify the genes in the mapping interval. Real-time PCR was conducted using LightCycler 480 II 96-Well PCR Plates (Roche, Rotkreuz, Switzerland). The utilized reaction system contained 10 \( \mu \)L of 2×SG Fast qPCR Master Mix (B639271, BBI), 2 \( \mu \)L cDNA, and 10 \( \mu \)M gene-specific primers in a final volume of 20 \( \mu \)L. The thermal cycling conditions used were 95 °C for 3 min, followed by 45 cycles at 95 °C for 5 s and 60 °C for 30 s followed by a final extension stage. The relative expression levels of all the genes in the mapping interval were calculated by the \( 2^{-\Delta\Delta Ct} \) method based on the QU samples and three replicates were performed for each sample [49]. The housekeeping gene Actin7 was used as the internal control to calculate the relative expression levels of each gene. Three biological replicates were performed in this experiment and t-test was used for statistical analysis.

Phenotyping for agronomic traits

To evaluate the agronomic efficiency of \( cqSPDA2 \), 100 individuals (50 AA- and 50 aa- genotype plants) from the \( \text{BC}_4F_2 \) population were sampled using the markers Indel3 and Indel15, and their chlorophyll content was characterized. The agronomic traits investigated were as follows: plant height (cm), total
siliques per plant, silique length (cm), seeds per silique, 1000-grain weight (g), and yield per plant (g). The mean values, standard deviations and significance analyses of all the agronomic traits were compared between the AA- and aa-genotype plants by Minitab16 and Excel2010.

**Abbreviations**

QTL: Quantitative trait locus; NIL: Near-isogenic line; DH: Doubled haploid; qRT-PCR: Quantitative real-time PCR; SNP: Single nucleotide polymorphism; Indel: Insertion-deletion; SSR: Simple sequence repeat; KASP: kompetitive allele-specific PCR; SPAD: Soil and plant analyzer development; FKBP: FK506- and rapamycin-binding protein; GCH-1: GTP cyclohydrolase I; BH4: Tetrahydrobiopterin; NOS/NOA: nitric oxide synthase.

**Declarations**

*Ethics approval and consent to participate*

This study does not contain any research requiring ethical consent or approval.

*Consent to publish*

Not applicable.

*Availability of data and materials*

All data used during the study are included in this published article and its additional files.

*Competing interests*

The authors declare no competing financial interests.

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Authors’ contributions

YJX, LHD and DDZ conceived and designed the research; YJX and LHD performed the experiments; YJX, LHD and DDZ discussed the results and strategies; YJX analyzed the data and wrote the manuscript. ZZ, XL and LKX corrected the manuscript. All authors read and approved the manuscript.

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**Table**

Due to technical limitations, Table 1 can be found in the Supplementary Files.

**Figures**
Figure 1

Phenotypic frequency distribution of chlorophyll SPAD in the BC4F2 population. The lowest point of the column chart represents the dividing value (SPAD=43). Leaves were considered to have high chlorophyll content (n=1514) with a SPAD value $\geq$ 43 and a low chlorophyll content (n=547) with a SPAD value $\leq$ 43. A chi-square test showed that the segregation pattern of the chlorophyll content trait was in keeping with the expected Mendelian segregation ratio of 3:1 ($\chi^2=2.53$).
Figure 2

Genetic and physical maps of the cqSPDA2 gene locus and candidate gene analysis. a Genetic linkage map of the cqSPDA2 region on chromosome A02. The numbers between the markers indicate genetic distance in centimorgans. b Fine mapping of the cqSPDA2 locus in the BC4F2 population. The cqSPDA2 was narrowed down to a 152-kb interval between the markers SSR2 and Indel15. The numbers between the markers indicate physical distance. c Genetic and physical maps of the candidate genes in the targeted interval and the annotated genes in the Brassica napus genome annotation database (http://www.genoscope.cns.fr/brassicanapus/)
Figure 3

Codominant marker and closely linked to cqSPDA2. The aa group (low-chlorophyll phenotype): the mean of SPAD=40.0±0.27; the Aa group (medium-chlorophyll phenotype): the mean of SPAD=44.8±1.44; the AA group (high chlorophyll phenotype): the mean of SPAD=53.3±0.80

Figure 4

Differential expression of 3 genes in the mapping interval in the parents and NILs. The relative expression levels were calculated by the $2^{-\Delta\Delta Ct}$ method based on the QU samples, and three replicates were performed for each sample. The housekeeping gene Actin7 was used as the internal control. Values shown are means ± SD (n=3). Student’s t-test was used to compare QU with ZS11 and BC4F2:3 (AA) with BC4F2:3 (aa). * denotes significance at the probability level of 0.05. ** denotes significance at the probability level of 0.01

Figure 5
Leaf color phenotypes of the parents QU and ZS11. QU: deep green leaves, high-chlorophyll content, SPAD value=50.4; ZS11: light green leaves, low-chlorophyll content, SPAD value=40.6

**Supplementary Files**

This is a list of supplementary files associated with this preprint. Click to download.

- Table1.xlsx
- Additionalfiles.pdf
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