Development of a high-density genetic linkage map and identification of flowering time QTLs in adzuki bean (*Vigna angularis*)

Changyou Liu, Baojie Fan, Zhimin Cao, Qiuzhu Su, Yan Wang, Zhixiao Zhang & Jing Tian

A high-density linkage map is crucial for the identification of quantitative trait loci (QTLs), positional cloning, and physical map assembly. Here, we report the development of a high-density linkage map based on specific length amplified fragment sequencing (SLAF-seq) for adzuki bean and the identification of flowering time-related QTLs. Through SLAF library construction and Illumina sequencing of a recombinant inbred line (RIL) population, a total of 4425 SLAF markers were developed and assigned to 11 linkage groups (LGs). After binning the SLAF markers that represented the same genotype, the final linkage map of 1628.15 cM contained 2032 markers, with an average marker density of 0.80 cM. Comparative analysis showed high collinearity with two adzuki bean physical maps and a high degree of synteny with the reference genome of common bean (*Phaseolus vulgaris*). Using this map, one major QTL on LG03 and two minor QTLs on LG05 associated with first flowering time (FLD) were consistently identified in tests over a two-year period. These results provide a foundation that will be useful for future genomic research, such as identifying QTLs for other important traits, positional cloning, and comparative mapping in legumes.

Key Laboratory of Crop Genetics and Breeding of Hebei Province; Institute of Cereal and Oil Crops, Hebei Academy of Agricultural and Forestry Sciences, Shijiazhuang 050035, China. Correspondence and requests for materials should be addressed to J.T. (email: nkytianjing@163.com)
To construct a high-density genetic map of adzuki bean, additional molecular markers need to be developed. Single-nucleotide polymorphisms (SNPs) are the most abundant class of polymorphisms in most genomes and are one of the most efficient markers for identifying candidate genes associated with QTLs. Based on the development of next-generation sequencing technology, several high-throughput methods for SNP and insertion/deletion polymorphism (InDel) marker identification and genotyping have been developed. These methods include restriction site-associated (RAD) sequencing (RADseq), genotyping-by-sequencing (GBS), and specific length amplified fragment sequencing (SLAF-seq). Among these methods, SLAF-seq combines pre-designed reduced representation library (RRL) schemes, high-throughput paired-end sequencing technology, and a double barcode system, which allows it to simultaneously genotype large populations with a considerable number of loci at a lower cost. Importantly, reference genome sequences and polymorphism information are not necessary when this method is used. This method has been applied in many species for genetic map construction as a rapid and cost-effective strategy for high-throughput SNP and InDel discovery and genotyping.

Flowering time is a very important target in adzuki bean breeding programs because it is critical for adapting cultivars to different cultivation areas or growing seasons. A series of genes or QTLs related to flowering time have been detected in Arabidopsis, rice, wheat, soybean, common bean, and other plants. However, few studies have focused on candidate genes or QTLs for flowering time in adzuki bean. Two studies that focused on the genetics of domestication in adzuki bean described 1 to 5 QTLs related to first flowering time (FLD). For example, Isemura et al. detected a QTL for FLD (Fld2.4.1, phenotypic variance explained (PVE): 23.9%) on LG 4 using an F2 population. Kaga et al. identified one major QTL of FLD (Fld3.4a.1, PVE: 43.7%) on LG 4a and four QTLs with smaller effects on LGs2 (Fld3.2.1, PVE: 6%), 3 (Fld3.3.1, PVE: 5.4%), 5 (Fld3.5.1, PVE: 8.8%), and 11 (Fld3.11.1, PVE: 5.8%) also using an F2 population. However, the precise genomic positions of these QTLs remain unclear, and the genes underlying these flowering time QTLs in adzuki bean are not known.

The main objective of this research was to analyze a recombinant inbred line (RIL) mapping population from a cross between wild and cultivated adzuki bean for SNP and InDel polymorphisms and for QTLs associated with flowering time during different years. The specific objectives were (a) to construct a high-density genetic map of adzuki bean based on the SLAF-seq high-throughput method and (b) to identify QTLs associated with FLD over two years.

### Results

#### SLAF sequencing and genotyping

SLAF library construction and Illumina sequencing generated a total of 47.8 Gb of raw data containing 240,904,338 paired-end reads with a length of 100 bp. The Q20 ratio (a quality score of 20, indicating a 1% chance of an error) was 81.12%, and the GC (guanine-cytosine) content is 33%. Of the loci were 190.14-fold for the male parent, 122.63-fold for the female parent, with an average of 4.25-fold for each RIL individual. Of the two years.

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#### Genetic linkage map construction

After discarding SLAF markers with a pair-wise independence LOD < 5, a total of 4425 markers could be assigned to 11 LGs. The coverage of the markers was 379.19-fold for the male parent, 243.04-fold for the female parent, with an average of 4.46-fold for each RIL individual. Of the assigned SLAF markers, 1938 markers did not segregate in the expected 1:1 ratio, as based on a chi-square test where the P-value of a marker segregating 1:1 was < 1%. The markers with distorted segregation were initially excluded from the map construction but were added later as accessory markers. To more reasonably calculate the average distance between adjacent markers, 3253 markers with the same genotype across the entire RIL population were merged into 861 bins. The bin information is supplied in Supplementary Table S1. The final map is 1628.15 cM in length with an average marker density of 0.80 cM (Fig. 1 and Table 1). The number of SLAF markers in each LG ranges from 108 (LG06) to 622 (LG07). The length of each LG ranges from 61.05 cM (LG06) to 270.83 cM (LG07), and the average distance between adjacent markers is 0.61 cM (LG05) to 0.95 cM (LG09). Except for one > 6-cM gap between adjacent markers on LG09, all other gaps are < 4.1 cM.

#### Comparative analysis

To evaluate the quality of the genetic map, the sequences of the 4425 mapped SLAF markers were aligned to the two draft genome sequences of adzuki bean using the stringent threshold described in the materials and methods section.

Based on the draft adzuki bean genome reported by Kang et al., the BLAST result indicated that 76.14% (3369/4425) of the SLAF markers map to pseudo-chromosomes or scaffolds (Supplementary Table S2). The SLAF markers on LG01, LG04, LG09, LG10, and LG11 primarily map to a single pseudo-chromosome (Table 2). The high Spearman's rank correlation coefficient (> 0.9) indicated high collinearity between the LGs and pseudo-chromosomes. The SLAF markers on LG02, LG03, LG05, LG06, LG07, and LG08 predominantly map to two to four pseudo-chromosomes or long scaffolds.

A similar analysis was performed for the adzuki bean genome reported by Yang et al. The BLAST results showed that 75.16% (3326/4425) of the SLAF markers map to pseudo-chromosomes or scaffolds (Supplementary Table S3). Except for LG01, LG03, LG05 and LG09, all other LGs are primarily homologous with a single pseudo-chromosome, with high syntenic (Table 2).
Figure 1. The high-density linkage map of adzuki bean. The high-density linkage map of adzuki bean was generated using JoinMap version 4.0. The name of the linkage groups is mentioned at the top of each LG. Distances between the loci (cM) are shown to the left, and the names of the loci are shown to the right of the linkage groups.

Table 1. Description of the basic characteristics of the 11 linkage groups.

| Linkage group ID | Length   | SLAF markers | Bin | Maker number after merging | Average distance between adjacent markers | Largest gap (cM) |
|------------------|----------|--------------|-----|----------------------------|------------------------------------------|-----------------|
| LG01             | 202.53   | 475          | 110 | 252                        | 0.80                                     | 3.73            |
| LG02             | 151.92   | 276          | 50  | 179                        | 0.85                                     | 3.75            |
| LG03             | 178.76   | 356          | 59  | 203                        | 0.88                                     | 3.73            |
| LG04             | 156.75   | 491          | 94  | 196                        | 0.80                                     | 3.02            |
| LG05             | 71.82    | 563          | 79  | 118                        | 0.61                                     | 2.34            |
| LG06             | 61.85    | 108          | 23  | 70                         | 0.87                                     | 3.38            |
| LG07             | 270.83   | 622          | 134 | 344                        | 0.79                                     | 4.08            |
| LG08             | 120.77   | 444          | 90  | 184                        | 0.66                                     | 3.03            |
| LG09             | 143.75   | 265          | 57  | 152                        | 0.95                                     | 6.63            |
| LG10             | 138.60   | 417          | 87  | 172                        | 0.81                                     | 3.03            |
| LG11             | 131.39   | 408          | 77  | 162                        | 0.81                                     | 4.08            |
| Total            | 1628.15  | 4425         | 860 | 2032                       | 0.80                                     |                 |

To align the genetic map with the reference genome of common bean [31], macrosynteny and microsynteny were detected between adzuki bean and common bean. All 11 LGs are syntenic with 10 common bean chromosomes (Fig. 2 and Supplementary Table S4). LG01, LG03, LG04, LG06 and LG09 correspond to only one common bean chromosome. The locus order of LG09 is most similar to common bean chromosome 6 (Pchr6). LG02, LG05, LG07, LG08, LG10 and LG11 mainly correspond to two common bean chromosomes. Interestingly, LG02 and LG07 are syntenic with common bean chromosomes 2 (Pchr2) and 3 (Pchr3) according to a ‘sharing’ model, which suggests that these two chromosomes in mungbean and common bean have recombined.

QTLs for first flowering time. FLD was evaluated over two years. As expected, the FLD of the wild parent (Yesheng10) occurred much later than did that of the cultivated parent (fjlong9218), at 78 days compared to 43 days. Transgressive segregation lines were observed in the RIL population. The distribution of FLD among RIL
population was nearly binomial (Fig. 3). A total of 8 QTLs were detected for FLD during the 2 years of the study (Fig. 4 and Table 3). Four QTLs were identified each year. Among them, two major QTLs (Fld3.2 and Fld3.3) for FLD with high LOD values (39.17 in 2013 and 35.63 in 2014) were consistently identified at the same map position as LG03 during both years, and the PVE by the QTLs was 70.9% (2013) and 66% (2014). Another 4 minor QTLs (Fld5.1 vs Fld5.3, and Fld5.2 vs Fld5.5) for FLD were detected at a similar map position on LG05 during both years. However, Fld3.1 on LG03 and Fld5.4 on LG05 were identified only during one year. The alleles of all QTLs from the wild parent delayed flowering time.

Discussion

The genetic map developed in this study contains 4425 SLAF markers, a majority of which are anchored to the adzuki bean draft genome scaffolds. The map consists of 11 LGs corresponding to the haploid chromosome number of the Vigna genus. Compared to the genetic map constructed by Han et al., the number of mapped loci (486 vs 4425), marker density (1.85 cM vs 0.8 cM), and total map length (1628.15 cM vs 832.1 cM) are significantly improved in this dense genetic linkage map. For our high-density genetic linkage map, the maximum genetic distance between flanking markers is 6.63 cM, which is much less than the 18.5 cM for the map of Han et al. Compared with the two SNP maps constructed by Kang et al. and Yang et al. to assemble the adzuki bean draft genome, the marker number in this individual map is substantially higher. However, the proportion of markers showing segregation distortion (43.8%) is higher than for the previous interspecific mapping populations of V. nepalensis x V. angularis (3.9%), V. angularis x V. nakashimae (19.7%) and V. umbellata x V. angularis (29.8%). One reason for these differences in segregation distortion may be the use of an RIL mapping population; in contrast, the above mentioned interspecific mapping populations were BC1F1 and F2 populations. A similar phenomenon was observed during the construction of maps for rice and chickpea. Another possible reason maybe that from generations F2 to F8 during RIL population generation, 19 lines were lost due to disease. However, compared to using BC or F2 as mapping populations, using an RIL mapping population can effectively reduce the influence of the dominant effect and reveal the additive effect of QTLs. Most importantly, an RIL mapping population as a permanent population can be planted in different years or environments, which would improve the accuracy of QTL mapping. Because discarding the markers with segregated distortion may not only remove biologically interesting segments of the genome but also dramatically reduce the marker density of the genetic map and because markers with skewed segregation can be successfully used for QTL mapping, we chose to include the

| Linkage group ID | SLAF markers | Adzuki bean draft genomea | Adzuki bean draft genomeb |
|-----------------|--------------|--------------------------|--------------------------|
|                 | SLAF markers | Homologous Chromosome/ Scaffold ID | Identified SLAF markers | Spearman | Homologous Chromosome ID | Identified SLAF markers | Spearman |
| LG01            | 475          | Chr6                     | 343                      | 0.9919   | Chr2                     | 297                      | 0.9668 |
| LG03            | 356          | Chr5                     | 66                       | 0.9737   | Chr4                     | 206                      | 0.8985 |
| LG04            | 491          | Chr8                     | 152                      | 0.9767   | Chr8                     | 225                      | 0.9685 |
| LG06            | 108          | Chr5                     | 20                       | 0.9727   | Chr2                     | 131                      | 0.5619 |
| LG07            | 622          | Chr11                    | 194                      | 0.9741   | Chr1                     | 341                      | 0.9900 |
| LG08            | 444          | SuperScaf_22             | 153                      | 0.2576   | Chr5                     | 321                      | 0.9832 |
| LG09            | 265          | Chr3                     | 146                      | 0.9360   | Chr9                     | 142                      | 0.9517 |
| LG10            | 417          | Chr2                     | 190                      | 0.9031   | Chr10                    | 36                       | 0.8759 |
| LG11            | 408          | Chr1                     | 259                      | 0.9755   | Chr7                     | 272                      | 0.9901 |

Table 2. Summary of the high-density linkage map aligned with two draft genomes of adzuki bean. Spearman: Spearman’s rank correlation coefficient; the closer the value is to 1, the better the synteny is. Only homologous chromosomes or scaffolds with the number of “Identified SLAF markers ≥ 20” are included in this table. aAdzuki bean draft genome reported by Kang et al. bAdzuki bean draft genome reported by Yang et al.
Figure 2. A map of synteny between the high-density map and the reference genome of common bean. Chromosomes of common bean are marked with PChr, and adzuki bean linkage groups are marked with LG.

Figure 3. Population distributions for FLD among the RIL mapping population from the cross between *V. nipponensis* and adzuki bean. Jihong9218 and Yesheng10 are the male and female parents, respectively.
markers with distorted segregation as accessory markers after map construction. Markers with the same genotype across the mapping population should be binned when constructing a high-density map; otherwise, the average inter-marker distance may not reflect the real distribution of the markers in the genome. A series of dense maps have been constructed according to this principle, such as the ultra-dense genetic maps of potato\textsuperscript{35}, rice\textsuperscript{36}, sorghum\textsuperscript{37}, and maize\textsuperscript{38}. In this study, the average inter-marker distance was 0.37 cM before we binned the SLAF markers from RIL with the same genotype but 0.80 cM after merging. Although this principle doubled the average inter-marker distance, it was obviously more reasonable.

A comparison of the dense genetic linkage map and two physical maps showed that the marker order along chromosomes was mostly collinear (Table 2), which confirmed the high quality of our map. However, we also found that several LGs mapped to more than one chromosome. For example, LG02, LG03, LG05, LG06, LG07, and LG08 mapped to 3, 2, 3, 3, 4, and 3 chromosomes, respectively, when aligned with the physical map of Kang \textit{et al.}\textsuperscript{29}. Four LGs (LG01, LG03, LG05 and LG09) mapped to 2 chromosomes in the physical map of Yang \textit{et al.}\textsuperscript{30}. To investigate these contradicting results and to determine the relationship between our map and that of Han \textit{et al.}\textsuperscript{8}, we
also aligned 186 pairs of SSR primer sequences from the map of Han et al. with the two reference genomes, and similar results were obtained (Supplementary Table S5). For example, LG 1 of Han's map mainly mapped to chromosomes 10 and 11 and scaffold 33 when aligned with the physical map by Kang et al.29, and it primarily mapped to chromosomes 1, 7, and 11 when aligned with the physical map of Yang et al.30. The mapping of LG07 from our high-density genetic linkage map produced similar results. Indeed, the results from mapping our genetic map as well as that of Han et al. to the two physical maps are in strong agreement for all 11 LGs, which will facilitate comparisons among these LGs. The mapping of LGs to more than one chromosome may be explained by genotyping errors, misassembled scaffolds or reciprocal translocations between chromosomes. We suggest that the mapping of LG05 to chromosomes 3 and 8 on the physical map of Yang et al.30 is due to a reciprocal translocation because it corresponds to an “LG 4 + 6” reciprocal translocation model in some wild adzuki bean accessions39. However, further efforts are needed to confirm this inference.

Based on the number of SLAF markers mapped to pseudo-chromosomes and the collinearity between LGs and pseudo-chromosomes, we determined that the assembly of the adzuki bean draft genome reported by Yang et al.30 is more accurate than that reported by Kang et al.29. This is in agreement with the genome assembly coverage (79.9% vs 75% of adzuki bean genome) results of the two physical maps. However, the assembly of pseudo-chromosome 1 on the physical map of Kang et al. may be better than the corresponding pseudo-chromosome on that of Yang et al., as it has more identified SLAF markers and better collinearity. This conclusion may be helpful when selecting a suitable reference genome for fine-mapping genes. Moreover, a high-resolution linkage map is useful for physical map assembly using next-generation sequencing. The mapping of SLAF markers to scaffolds will improve the ordering and orientation of the remaining unplaced sequences in the two draft genome sequence databases of adzuki bean.

Genomic synteny analysis is useful for comparative genomics. Given the close genetic relationship between common bean and adzuki bean, we aligned the genetic map with the reference genome of common bean. Our results demonstrated high synteny between the two genomes (Fig. 2), which suggests that candidate genes could be identified through comparative mapping. In addition, the extensive synteny and collinearity observed between the common bean genome and the current map provide additional support for the mapping accuracy of our high-density genetic map.

Flowering time is critical for adapting adzuki bean cultivars to different cultivation areas or growing seasons. Isemura et al. detected a QTL (Fld2.4.1) of FLD on LG 4 (map of Han et al.) after phenotyping an F2 population derived from a cross between cultivated adzuki bean (Vigna angularis) and wild relative (Vigna nepalensis) in a single year41. Kaga et al. identified one major QTL of FLD (Fld3.4a.1) on LG 4a and four QTLs with smaller effects on LGs2 (Fld3.2.1, 3(Fld3.3.1), 5(Fld3.5.1), and 11(Fld3.11.1) using an F2 population derived from a cross between cultivated adzuki bean (Vigna angularis) and wild adzuki bean (Vigna nipponensis) accession42. In the present study, a two-year phenotypic evaluation of flowering time in an RIL population produced from an early-flowering adzuki bean cultivar and a late-flowering wild adzuki bean accession indicated the presence of one major QTL on LG03 and two minor QTLs on LG05. Because both studies mentioned above were based on the genetic map of Han et al. and we had aligned both our map and that of Han et al. to the two physical maps of adzuki bean (Supplementary Table S5), it is possible to compare the QTLs from all three studies. Based on the alignment with the physical map of Yang et al., Fld2.4.1 (the nearest SSR markers were CEDG103 and CEDG011) is located on chromosome 3, and its physical position is between 32.8 and 39.1 Mb. Similarly, Fld3.4a.1 (the nearest SSR markers were CEDG036 and CEDG127) is also located on chromosome 3, and the physical positions between 36.2 and 40.0 Mb. Because these two QTLs are located in the same genomic region, they may be influenced by the same gene or genes. Fld3.2.1 maps to chromosome 4 in the physical map of Yang et al.; in this study, Fld3.2 and Fld3.3 were detected on the same chromosome. A neighbouring SSR marker (CEDG026) showed that its physical position is approximately 1.5 Mb on chromosome 4, and the distance from the associated marker (Marker56693) of Fld3.2 and Fld3.3 is approximately 2.2 Mb, indicating that they may be located at the same QTL. However, similar to Fld2.4.1 (PVE: 23.9%) vs Fld3.4a.1 (PVE: 43.7%), the PVEs by Fld3.2.1 (PVE: 6.0%) and Fld3.2 (PVE: 70.9%) differed greatly. These differences may be due to allelic variation or interactions between the QTL and the genetic background. Further research is needed to verify this inference.

Considering that LG03 shows high synteny with chromosome 9 (Pchr9) of the common bean genome (Fig. 2), we considered QTLs for flowering time on this chromosome. As expected, we found flowering time-related genes (i.e., PvZTL) on LGB9 (corresponding to Pchr9 in the common bean reference genome)28, which are located near the GH locus identified by Tar’an et al.40. Using the ZTL protein sequence of Arabidopsis, the adzuki bean homologues in GenBank were identified by a BLAST search as located near 0.2 Mb on chromosome 4; the distance from the associated marker (Marker56693) of Fld3.2 and Fld3.3 is approximately 3.5 Mb. Further studies are needed to confirm the role of this homologue as a potential flowering time gene.

In summary, this study is the first attempt to conduct QTL analysis using an NGS-derived dense genetic map in adzuki bean. The results provide a foundation that will be useful for future genomic research, such as the identification of QTLs for other important traits, positional cloning, comparative mapping in legumes, ordering and orienting the remaining unplaced scaffolds in the two draft genomes of adzuki bean and marker-assisted selection in adzuki bean breeding.

Materials and Methods
Mapping population. An F2RIL population was developed from the cross between a wild adzuki bean accession (Vigna nipponensis: Yesheng10) collected in Dandong, China (39.75N, 123.74E), and an adzuki bean cultivar (Jihong9218) that is widely grown in northern China. The cultivated parent had an early flowering time and was the male in the cross. The wild adzuki bean accession had a late flowering time and was the female in the cross. The RIL population consisted of 153 lines generated from a single seed descent from generations F2 to F8.
The parental accessions used in the cross were obtained from the Hebei Academy of Agricultural and Forestry Sciences (HAAFS).

**Phenotypic evaluation for variation in flowering time.** The RIL population of 153 lines and 20 plants of each parent was grown in the field at HAAFS, Shijiazhuang, China (37.95N, 114.73E), from June to November in 2013 and 2014. The soil at the site was a sandy loam with no major fertility problems (PH = 7.3). Experimental units consisted of two-row plots 3.5 m long and 1 m wide. Each planting was a randomized block design experiment that was repeated twice. The FLD (days from sowing to the first flower) was evaluated using the mean value of each RIL line and parent over a two-year period (2013 and 2014).

**DNA extraction.** Young leaves from 153 RIL lines and two parents were collected, immediately frozen in liquid nitrogen, and transferred to a −80 °C freezer. To obtain high-quality DNA for SLAF library construction, a plant DNAzol kit (ThermoFisher, Waltham, MA, USA) was used to extract the total genomic DNA according to the manufacturer’s instructions.

**SLAF library construction and high-throughput sequencing.** The procedure used for SLAF library construction was conducted as described by Sun et al. with minor modifications. In brief, a draft reference genome of adzuki bean was used to design SLAF marker discovery experiments by simulating in silico the number of markers produced by restriction digest with two different enzyme combinations. Accordingly, an SLAF pilot experiment was performed, and the SLAF libraries were constructed. Two enzymes (HaeIII and SspI-HF; NEB, Ipswich, MA, USA) were used to digest the genomic DNA of the parents and RIL population. Subsequently, a single nucleotide (A) overhang was added to the digested fragments using Klenow Fragment enzyme (NEB, Ipswich, MA, USA) and ATP at 37 °C. Next, duplex tag-labelled sequencing adapters (Life Technologies, Carlsbad, California, USA) were ligated to the A-tailed fragments using T4 DNA ligase (NEB, Ipswich, MA, USA). Then, the diluted and ligated DNA samples were used as a template for polymerase chain reaction (PCR). Each PCR reaction also contained High-Fidelity DNA Polymerase (NEB, Ipswich, MA, USA), dNTPs and PCR primers (Forward sequence: 5′-CAAGCAGAAGACGGCATACG-3′, reverse sequence: 5′-CGAGTACGGGACCGATACG-3′) (PAGE-purified, Life Technologies). PCR products were purified using Quick Spin columns (Qiagen, Venlo, Netherlands) and separated by 2% agarose gel electrophoresis. SLAFs of 264–414 bp (with adapter sequence indexes and adapters) in size were excised and purified using a QIAquick Quick Spin columns (Qiagen, Venlo, Netherlands) and separated by 2% agarose gel electrophoresis. SLAFs of 264–414 bp (with adapter sequence indexes and adapters) in size were excised and purified using a QIAquick Gel Extraction kit (Qiagen, Hilden, Germany). Finally, 100-bp paired-end sequencing was performed on an Illumina HiSeq 2500 sequencing platform (Illumina, Inc., San Diego, CA, USA) according to the manufacturer’s instructions.

**Sequence data grouping and genotype definition.** The procedures used for SLAF marker identification and genotyping were performed as described by Sun et al. with minor modifications. Briefly, low-quality reads (quality score < 20) were filtered out, and the barcodes were trimmed from each high-quality read. All of the clean reads were clustered based on sequence similarity as determined by BLAST (−tileSize = 10, −stepSize = 5). Sequences with >95% identity were grouped into one SLAF locus. Allele tags of each SLAF locus with a sequencing depth >10-fold for parental reads and >70% integrity in the offspring were collected. Both SNP and InDel loci were detected between parents, and SLAFs with >3 SNPs or InDels were filtered out. Because adzuki bean is diploid species and one locus contains at most four SNP tags, groups containing more than four tags were discarded. Only SLAFs with 2, 3, or 4 tags were identified as polymorphic and considered potential markers. All polymorphic markers were classified into eight segregation patterns (aa × bb, ab × cd, ef × eg, hk × hh, lm × ll, nn × np, ab × cc and cc × ab). Because the mapping population was composed of RILs, only the aa × bb segmentation pattern was used for genetic linkage map construction.

**Genetic linkage map construction.** JoinMap ver. 4.0 was used to construct a linkage map. Marker segregation ratios were calculated using the chi-square test. Markers showing significant (P < 0.01) segregation distortion were initially excluded from the map construction but were added later as accessory markers. Markers with the same genotype across the entire RIL population were binned. The grouping and ordering of the markers were established using a maximum likelihood algorithm. Pairwise marker loci that showed a likelihood-ratio statistic (LOD) value larger than 5.0 were used to create LGs, and the recombination frequencies were converted into map distances (cM) using the Kosambi mapping function.

**Comparative analysis of the high-density linkage map.** Colinearity between the high-density linkage map and the adzuki bean genome was determined by comparing the assigned SLAF sequences to two adzuki bean draft genomes with BLASTN (version 2.2.30) and a cut-off value of e−30. SLAF markers with query sequence lengths ≥80 and sequence identity >98% were selected to calculate the Spearman rank correlation according to their order on LGs and physical position on chromosomes using R (version 3.1.2) software. Synteny between adzuki bean and common bean (Phaseolus vulgaris) was determined with BLASTN searches against the genome of Phaseolus vulgaris (http://phytozome.jgi.doe.gov/pz/portal.html) using the source sequences of mapped SLAF markers as queries. The significance cut-off value was e−20 for an overlap of at least 70 bp. Synteny was visualized using MapChart (version 2.3) software.

**QTL analysis.** QTL analysis was conducted with MapQTL ver.6 as described by Van et al. Briefly, the entire genome was scanned for QTLs of FLD using a general interval mapping (IM) method. A regression algorithm was used to calculate the maximum likelihood, and a 1.0 mapping step size was used. The significance of each
QTL interval was tested with an LOD. The threshold of the LOD score for significance (P = 0.05) was determined using 10,000 permutations. The PVE by each QTL was estimated based on the population variance found within the segregating population.

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Author Contributions
Jing Tian and Changyou Liu designed the experiment. Changyou Liu carried out the DNA isolation and sequence data analyses and drafted the manuscript. Jing Tian assisted in manuscript preparation. Baojie Fan, Zhimin Cao, Quizhu Su, Yan Wang, and Zhixiao Zhang prepared the plant materials and participated in DNA extraction. All of the authors read and approved the final manuscript.

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