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The 'PI 438489B' by 'Hamilton' SNP-Based Genetic Linkage Map of Soybean [*Glycine max* (L.) Merr.] Identified Quantitative Trait Loci that Underlie Seedling SDS Resistance

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Abstract

Soybeans [Glycine max (L.) Merr.] are susceptible to many diseases including fungal diseases such as soybean sudden death syndrome (SDS). Several studies reported SDS resistance quantitative trait loci (QTL) on the soybean genome using different recombinant inbred line (RIL) populations and low density genetic linkage maps. High density exclusively single nucleotide polymorphisms-based (SNP-based) maps were not yet reported in soybean. The objectives of this study were (1) to construct a high density SNP-based genetic linkage map of soybean using the 'PI438489B' by 'Hamilton' (PIxH, n=50) recombinant inbred line population, and (2) to map QTL for SDS resistance using this high-density reliable genetic SNP-based map. The PI438489B by Hamilton high-density SNP-based genetic map was a high density map composed of 31 LGs, 648 SNPs, and covered 1,524.7 cM with an average of 2.37 cM between two adjacent SNP markers. Fourteen significant QTL were identified for SDS resistance using interval mapping (IM) and composite interval mapping (CIM) with LOD scores that ranged between 2.6 and 5.0. Twelve QTL were identified for foliar disease severity (FDS) and three QTL for root rot severity (RRS) of which one QTL underlain both FDS and RRS. The fourteen QTL were mapped onto ten separate chromosomes of the soybean genome. Seven of the intervals encompassing the QTL had been identified previously (on LGs C1, C2, D1b, G, L, N and O) associated with resistance to SDS but seven were novel (LGs A2 (2), B1, C2, D1a, D1b and O). We constructed the first PI438489B by Hamilton exclusively SNP-Based map and identified fourteen QTL that underlie SDS resistance including both resistances to foliar and root rot symptoms caused by Fusarium virguliforme infection. The QTL discovered here for SDS resistance could be useful to include in breeding programs in developing soybean cultivars resistant to SDS.

Introduction

Soybean [*Glycine max* (L.) Merr.] sudden death syndrome (SDS) is caused by the fungus Fusarium virguliforme (Aoki et al., 2005). SDS is capable of causing major yield losses in soybean (Luo et al., 2000). Host resistance in soybean to SDS has been found to be polygenic and controlled by several quantitative trait loci (QTL) (Meksem et al., 1999; Iqbal et al., 2001).

In the 'Essex' by 'Forrest' recombinant inbred line (RIL) population (ExF, n=100) nine QTL have been discovered (Kassem et al., 2006, 2007). More specifically, two were found on chromosome 13 (Linkage Group [LG]) F); one QTL each on chromosomes 16 (LG J), 6 (LG C2), and 20 (LG I); and four QTL on chromosome 18 (LG G) (Kassem et al., 2007). To date 3 of these QTL were confirmed in near isogeninic lines (NILs); two on LG

G and one on LGC2. Three additional QTL for SDS resistance have been found on chromosomes 3 (LG N), 6 (LG C2), and 18 (LG G) using the 'Pyramid' by 'Douglas' RIL population (PxD, n=90) (Kassem et al., 2007; Njiti et al., 2002). Prabhu et al. (1999) tested the markers Satt038 (chromosome 18 - LG G) and BLT65 (chromosome 8 - LG A2) and found QTL for SDS resistance associated with the LG G markers in the 'Hartwig' by 'Flyer' RIL population (FxH, n=50) (Prabhu et al., 1999). Farias et al. (2007) mapped QTL for SDS resistance (DX) using two RIL populations 'Ripley' by 'Spencer' (RxS, n=91) and PI 567374 by 'Omaha' (PI567374xO, n=96) (Farias et al. 2007). These QTL were located on chromosomes 4, 17, and 19 (LGs C1, D2, and L, respectively) in the first population and on chromosomes 17 and 20 (LGs D2 and I) in the latter one (Farias et al., 2007). It is worth mentioning that the D2 QTL was confirmed in NILs too. In a recent study, Kazi et al. (2007) identified QTL on LG C2, D2 and G in FxH (n=94) and confirmed the D2 QTL in NILs (Kazi et al., 2007; Kazi et al., 2008). However, the ExF, HxF, PxD, RxS, and PI567374xO genetic linkage maps that were used to map SDS resistance QTL were not high density genetic maps and also the populations that used were limited. The ExF genetic linkage map was based on 237 markers (Kassem et al., 2006), the FxH genetic linkage map was composed of 144 markers (Kazi et al., 2007, 2008), the PxD genetic linkage map was composed of 112 markers (Njiti et al., 2002), the RxS map was composed of 112 simple sequence repeats (SSRs) with only 68 of the markers linked (Farias et al. 2007), and the PI567374xO map was composed of 104 SSRs with only 41 of the markers linked (Farias et al., 2007).

During the last two decades, soybean genetic maps have been increasing in the number of markers used to create the maps. The first genetic map in soybean was based on 150 markers (Keim et al., 1990). Subsequently, the number of markers used to create genetic maps has varied greatly and many different types of markers such as restriction fragment length polymorphisms (RFLPs), amplified fragment length polymorphisms (AFLPs), random amplifications of polymorphic DNA (RAPDs), and SSRs have used.. The number of markers has varied from 110 markers (Shoemaker et al., 1995), to 112 markers (Njiti et al., 2002), to 132 markers (Lark et al., 1995), to 237 markers (Kassem et al., 2006), to high density composite genetic maps (Song et al., 2004).

Since the discovery of single nucleotide polymorphism (SNP) markers in mice and humans (Collins et al., 1998; Lindblad-Toh et al., 2002), several methods of SNP discovery have been described in different species including plants (Nicod et al., 2003; Barbazuk et al., 2007; Hyten et al., 2010a). SNPs are the most abundant genetic marker available for the creation of genetic maps and can be used with high-throughput genotyping technology making them highly desirable for QTL mapping studies (Brookes et al., 1999; Nicod et al., 2003; Barbazuk et al., 2007; Hsu et al., 2008; Fan et al., 2003). On this basis, SNP-based maps have been constructed in many plant species including Arabidopsis thaliana (Li et al., 2000), almond (Wu et al., 2009), sugar beet (Mohring et al., 2004), soybean (Hyten et al., 2010b), and others. However, the genetic maps in soybean have not been exclusively SNP-based but contain SNPs anchored with

other markers such as RFLPs, AFLPs, RAPDs, and SSRs except of a recent high density soybean SNP-based map containing 1,790 SNPs (Hyten et al., 2010a).

Genetic linkage maps have been used to identify and map QTL for many important agronomic traits in soybean (SoyBase, 2011). High density maps, the marker-type, and population size, are among important factors for mapping QTL. The objectives of this study were (1) to construct a high density SNP-based genetic linkage map of soybean using the PI 438489B by 'Hamilton' (PIxH, n=50) RIL population, and (2) to map QTL for SDS resistance using this high-density genetic SNP-based map.

Materials and Methods

Plant Material

In this study, we used the PI 438489B by Hamilton recombinant inbred line (RIL) population (PIxH, n=50). PI 438489B is a plant introduction from China and is resistant to SDS. Hamilton was developed at the Illinois Agricultural Experiment Station and was released for its high yield (Nickell et al., 1990) although it is susceptible to SDS. The PI 438489B by Hamilton RIL population was developed at the University of Missouri Agronomy Research Center (Yue et al., 2001) and advanced to the $F_{6:13}$ generation by Dr. Silvia Cianzio at the ISU research site at the Isabela Substation, Univ. of Puerto Rico, Isabela, Puerto Rico.

DNA Isolation

DNA was extracted using DNeasy 96 Tissue Kit (QIAGEN, Inc., Valencia, CA, USA) from young leaf tissue. DNA quantification was performed using fluorescent nucleic acid stain with Hoechst 33258 dye (Cat no. H21491 Invitrogen, Carlsbad, CA, USA). The DNA samples were read in a Synergy 2 plate reader (Biotek , Winooski, VT, USA) and then diluted to 100 ng μ l⁻¹ at final concentration.

SNP Genotyping

The 1,536 Universal Soy Linkage Panel 1.0 from Hyten et al. (2010b) was used to screen the 50 RILs and the GoldenGate assay was performed as per the manufactures protocols and as described previously (Fan et al., 2003; Hyten et al., 2008). The Illumina BeadStation 500G (Illumina Inc., San Diego, CA) was used for genotyping the GoldenGate assay. The automatic allele calling for each locus was accomplished with the BeadStudio version 3.2 software (Illumina Inc., San Diego, CA). All BeadStudio data for the 1,536 SNPs were visually inspected and rescored if any errors in calling the homozygous or heterozygous clusters were evident.

SDS Phenotypic Scoring

The RIL population was phenotyped for disease severity in the plant pathology greenhouse at Iowa State University, in Ames, Iowa. The experiment was established as a randomized complete block design with three replications per RIL. The

experimental unit was a 3" styrofoam cup containing three plants growing in soil infested with F. virguliforme. Inoculum was prepared according to a protocol by Munkvold and O'Mara (2002). One kilogram of a sterile 5:1 sand:cornmeal mixture was infested with 0.6 ml of a 106 spore/ml suspension of the Mont 1 isolate of F. virguliforme (Li et al., 2000), resulting in an initial spore concentration of 600 spores g⁻¹ of inoculum. The inoculum was incubated for 10 days at room temperature (approximately 24°C), after which it was homogeneously mixed in a 1:5 ratio with pasteurized soil and sand mix (1:1 by vol). Three seeds were then planted in styrofoam cup filled with the infested soil mix. The cups were maintained on a greenhouse bench at 23±2°C, and a 16 h photoperiod for 30 days. Foliar diseases severity (FDS) was rated 15 (FDS 1), 20 (FDS 2), 25 (FDS 3), and 30 (FDS 4) days after planting (DAP) as a percentage of total leaf area with typical SDS symptoms. Root rot severity (RRS) was evaluated 30 DAP by thoroughly washing the roots in running tap water and visually evaluating the percentage of root area showing brown or black discoloration. This highly controlled assay has been shown to produce results in seedlings that correlate closely with field performances (Njiti, 2001). The experiment was conducted twice.

Statistical Data Analysis

Means, ranges and standard deviations were calculated for the RIL lines and their parents from raw data. Analyses were performed on JMP 8.02 (SAS Institute Inc., Cary, NC, USA).

QTL Mapping

For mapping of QTL and estimation of their effects, composite interval mapping (CIM) was performed using the Win-QTL Cartographer version 2.5 software (Wang et al., 2005). The Model 6 and its default settings were adopted. To establish experimental-wise LOD cutoff values for declaring QTL significant at $P \le 0.05$ 1,000 permutations were performed. In addition, a value of $P \le 0.0002$ or LOD 3.7 was suggested by an approximate Bonferroni correction (P<0.05/250) for the set of about 250 independent (unlinked or >10 cM apart) DNA markers (from the 679 mapped). However, at genomic regions where gaps between adjacent markers were greater than 10 cM in the map associations 0.005 > P > 0.0005 (2.3 < LOD < 3.3) were accepted as a potentially significant association. If the interval was large or was flanking a single marker the uncorrected P value of <0.05 was accepted. Precedents with first-pass mapping of other quantitative traits (lqbal et al., 2001; Kassem et al., 2006, 2007; Njiti et al., 2002; Prabhu et al., 1999; Farias et al., 2007; Kazi et al., 2007) have shown these criteria to be valid during the later NIL based maps of the intervals that were inferred at marginal P values (Meksem et al., 1999).

Genetic Map Construction

The current PI 438489B by Hamilton genetic map is exclusively SNP-based and was constructed in several steps using Join-Map 4.0 software (Feltus et al., 2010). The SNP markers were initially grouped and assigned to the soybean chromosomes based on their mapped position on the soybean Consensus 4.0 map (Hyten et al., 2010b). We used the regression mapping algorithm with the default parameters and Kosambi's mapping function to determine map order and genetic distances.

Finding Potential Genes that Underlie SDS QTL

Each QTL was bounded by two single nucleotide polymorphisms (SNPs). The DNA sequence of the SNPs was obtained by searching the NCBI Single Nucleotide Polymorphism database (http://www.ncbi.nlm.nih.gov/projects/SNP/). The SNP DNA sequences were used as a query in a BLAST search of the Glycine max genome, version Williams 82 genomic sequence (http:// soybase.org/gbrowse/cgi-bin/gbrowse/gmax1.01/).The BLAST search revealed the nucleotide position of each SNP on the chromosome of interest. The nucleotide positions were used to query the soybean whole genome sequence (Glycine max version 1.01) (Schmutz et al., 2010) that included gene names and gene descriptions (http://soybase.org/gbrowse/cgi-bin/ gbrowse/gmax1.01/). The results included the genes and gene descriptions of all genes bounded by the SNPs, and the data was pasted into Microsoft Excel version 2007 spreadsheet for subsequent analysis. To determine common genes within all QTL intervals, a single excel file was created with three columns: QTL, gene name and gene description. The QTL and gene name column were concatenated to form the QTL/gene column and a delimiter (semicolon) was appended at the end. The QTL and gene name columns were deleted leaving two columns: QTL/gene and gene description. The excel function Remove Duplicates was used to remove duplicate QTL/genes in a given QTL based on the gene description. The columns were sorted alphabetically based on gene description. The following function identified genes that were common to more than one QTL and created a new column (=IF(OR(B2=B3, B2=B1), A2 &" "& B2) where A is the QTL/ gene column and B is the gene description column. Genes that occurred only in one QTL were labeled as false. The column was sorted alphabetically and the false values deleted. The excel function Text to columns was used to separate the column into two columns using the semicolon delimiter. The first column was the QTL/gene data and the second column was the gene description. The columns were sorted alphabetically based on the gene description column. The function (=COUNTIF A:A, A1) where A is the column with gene description was used to count the number of repeating gene descriptions, and a new column gene frequency was created. The three columns QTL/gene, gene description and gene frequency were sorted based on gene frequency. The results are presented in Table 4 (Supplementary Data).

Results

Trait Distributions

SDS resistance was determined in the RIL population using foliar disease severity (FDS) and root rot severity (RRS) which are symptoms caused by *F. virguliforme* infection of plants in the greenhouse. FDS 1, 2, 3, and 4 were scored 15, 20, 25, and 30 days after planting (DAP), respectively and RRS was estimated 30 DAP. Means and standard deviations of FDS 1-4 and RRS are shown in Table 1. The RIL population demonstrated a broad range of variation for the FDS and RRS traits that were evaluated (Table 1). The variation of FDS 1 ranged from 0 to 30 (st. dev. 2.437), FDS 2-4 ranged from 0-100 (st. dev. 31.189-35.929) and RRS showed range from 0 to 100 (st. dev. 14.255) (Table 1).

Table 1. Means, ranges, skewness and test for nor-
mal distribution within the PI438489B by HamiltonRIL population.

Traits	Mean (±St. Dev.)	Range
FDS 1	0.406 (± 2.437)	0-30
FDS 2	17.461 (± 31.189)	0-100
FDS 3	47.677 (±35.929)	0-100
FDS 4	62.239 (± 32.450)	0-100
RRS	89.072 (±14.255)	0-100

SNP-Based Genetic Map

The PI 4384898 by Hamilton SNP-based genetic map was constructed in several steps using the JoinMap 4.0 software (Feltus et al. 2010). First, the 1536 SNP Universal Soy Linkage Panel 1.0 from Hyten et al. (2010b) was used to screen the parents PI 4384898 and Hamilton, and the 50 RILs. Out of the 1,536 SNPs there were a total of 679 SNPs which were polymorphic between the two parents and segregated within the RIL population. These 679 SNPs were used to construct the genetic linkage map. The final genetic map contained 31 LGs and 648 linked SNP markers (Figure 1). Thirty-one markers were unlinked. The map coverage was 1,524.7 cM (Figure 1). The average distance between markers was 2.35 cM (Table 3). This map was used for genetic mapping of the SDS resistance QTL.

SDS Resistance QTL

Both interval mapping (IM) and composite interval mapping (CIM) were used to identify SDS resistance QTL. Using CIM, eleven QTL were identified for SDS resistance and mapped on 10 separate chromosomes of the soybean genome. Eight QTL

Table 2. The fourteen QTL that underlie SDS resistance found in the soybean PI 438489B by Hamilton RIL population. The QTL were found using the CIM of WinQTL Cart., and reported with LOD scores equal or greater than 2.5. Traits measured were foliar disease severity (FDS 1, 2, 3, or 4 weeks after infection by F. virguliform, respectively) and root rot severity (RRS) symptoms. QTL were named according to the Soybean Genetics Committee recommendations as revised in March 2007 (http://soybase.org/resources/QTL.php).

Trait	No.	QTL	Chr./LG	Marker/Interval	Position (cM)	LOD	R²(%)
CIM							
FDS 2	1	qFDS002-01	1/D1a	ss107927723-ss107913849	27.2–42.8	3.0	7.5
	2	qFDS002-02	10/0	ss107930838-ss107912519	13.5-15.2	2.6	19.3
	3	qFDS002-03	19/L	ss107913933-ss107929955	42.0–49.9	4.0	17.7
FDS 3	4	qFDS003-01	2/D1b	ss107927695-ss107913858	19.4-21.8	3.6	9.0
	5	qFDS003-02	2/D1b	ss107920774-ss107912689	30.0–36.0	2.8	5.2
	6	qFDS003-03	6/C2	ss107929602-ss107925487	32.8–39.2	4.2	3.5
	7	qFDS003-04	8/A2	ss107919498-ss107915722	2.0-13.0	4.6	9.6
FDS 4	8	qFDS004-01	11b/B1	ss107912672-ss107924081	5.5-17.8	2.7	3.4
	9	qFDS004-02	18/G	ss4969823-ss107924619	24.4-28.1	2.9	8.8
RRS	10	qRRS001-01	3/N	ss107912585-ss107920575	38.3–42.6	4.2	9.9
	11	qRRS001-02	4/C1	ss107929213-ss107929551	51.5-57.3	5.0	8.6
	(9)	qRRS001-03	18/G	ss4969823-ss107924619	24.4–27.7	2.9	33.3
IM							
FDS 2	(3)	qFDS002-03	19/L	ss107924889-ss107921208	48.0-51.3	2.6	6.0
FDS 3	12	qFDS003-05	6/C2	ss107917031-ss107912977	16.9–32.8	4.2	2.4
	(6)	qFDS003-03	6/C2	ss107930961-ss107912561	34.5–39.8	4.6	4.7
	13	qFDS003-06	8/A2	ss107915722-ss107918074	15.0-28.0	2.7	17.4
FDS 4	14	qFDS004-03	4/C1	ss107924445-ss107918378	57.3-83.9	2.6	4.8
	(10)	qFDS004-03	6/C2	ss107917031-ss107912977	16.9–32.8	3.4	2.1
	(6)	qFDS004-04	6/C2	ss107930961-ss107912561	34.5–39.8	3.6	3.2
RRS	(3-9)	qRRS001-03	18/G	ss107924669-ss107921695	26.5-28.4	2.6	2.3

 Table 3. Statistics of the linkage groups (LGs) of the 'PI

 438489B' by 'Hamilton' SNP-based genetic linkage map.

LG. Chr. No	Coverage	No. of	cM/Marker
	(cM)	Markers	
1. Chr_1	93.0	32	2.91
2. Chr_2	60.2	31	1.94
3. Chr_3	51.7	24	2.15
4. Chr_4	98.4	31	3.17
5.Chr_5a	49.8	18	2.76
6. Chr_5b	34.1	8	4.26
7. Chr_6	68.4	44	1.55
8. Chr_7	79.9	35	2.28
9. Chr_8	119.4	50	2.39
10. Chr_9	73.0	53	1.38
11.Chr_10	118.5	26	4.55
12. Chr_11a	4.5	12	0.37
13.Chr_11b	18.7	8	2.34
14.Chr_11c	18.2	4	4.55
15. Chr_11d	6.9	4	1.72
16.Chr_12	58.5	25	2.34
17. Chr_13a	35.0	18	1.94
18.Chr_13b	65.9	21	3.14
19. Chr_14	99.5	33	3.01
20. Chr_15a	37.1	10	3.71
21. Chr_15b	37.3	25	1.49
22. Chr_16a	28.2	7	4.03
23. Chr_16b	37.9	12	3.16
24. Chr_17a	23.8	16	1.48
25. Chr_17b	27.7	8	3.46
26. Chr_17c	16.8	15	1.12
27. Chr_17d	11.9	3	3.96
28. Chr_18	35.1	30	1.17
29. Chr_19	54.2	16	3.38
30. Chr_20a	22.5	12	1.87
31. Chr_20b	38.6	11	3.51
Total	1524.7	642	2.37

were found for foliar (FDS) resistance only, two QTL for root rot (RRS) resistance only, and one QTL which conferred resistance on both leaves and roots. The first QTL was associated with the FDS 2 trait (qFDS002-01) was identified in the marker interval ss107927723–ss107913849 on chromosome 1 (LG D1a). The QTL interval spanned approximately 15.6 cM, had a peak LOD score of 3.0, and an R2 of 7.5% (Table 2, Figure 1). The second QTL associated with the FDS 2 trait (qFDS002-02) was identified in the marker interval ss107930838–ss107912519 on chromosome 10 (LG O). The QTL interval spanned approximately 1.7 cM, had a peak LOD score of 2.6, and an R2 of 19.3% (Table 2, Figure 1). The third QTL which was associated with the FDS 2 trait (qFDS002-03) was identified by the marker

interval ss107913933-ss107929955 on chromosome 19 (LG L). The QTL interval spanned approximately 7.9 cM, had a peak LOD score of 4.0, and an R2 of 17.7% (Table 2, Figure 1). The three QTL explained approximately 50% of the total variation in FDS 2. The fourth QTL which was associated with the FDS 3 trait (qFDS003-01) was identified by the marker interval ss107927695-ss107913858 on chromosome 2 (LG D1b). The QTL interval spanned approximately 2.4 cM, had a peak LOD score of 3.6, and an R2 of 9.0% (Table 2, Figure 1). The fifth QTL that was associated with the FDS 3 trait (gFDS003-02) was identified by the marker interval ss107920774-ss107912689 on the same chromosome 2 (LG D1b). The QTL interval spanned approximately 6 cM, had a peak LOD score of 2.8, and an R2 of 5.2% (Table 2, Figure 1). The sixth QTL that was associated with the FDS 3 trait (qFDS003-03) was identified by the marker interval ss107929602-ss107925487 on chromosome 6 (LG C2). The QTL interval spanned approximately 6.4 cM, had a peak LOD score of 4.2, and an R2 of 3.5% (Table 2, Figure 1). The seventh QTL that was associated with the FDS 3 trait (gFDS003-04) was identified by the marker interval ss107919498-ss107915722 on the same chromosome 8 (LG A2). The QTL interval spanned approximately 11 cM, had a peak LOD score of 4.6, and an R2 of 9.6% (Table 2, Figure 1). The four QTL explained approximately 43.7% of the total variation in FDS 3. The eighth QTL that was associated with the FDS 4 trait (qFDS004-01) was identified by the marker interval ss107912672-ss107924081 on chromosome 11 (LG B1). The QTL interval spanned approximately 12.3 cM, had a peak LOD score of 2.7, and an R2 of 3.4% (Table 2, Figure 1). The ninth QTL that was associated with the FDS 4 trait (qFDS004-02) was identified by the marker interval ss4969823-ss107924619 on chromosome 18 (LG G). The QTL interval spanned approximately 3.7 cM, had a peak LOD score of 2.9, and an R2 of 8.8% (Table 2, Figure 1). This same interval which was significantly associated with FDS 4 also was significant for RRS (gRRS001-03). The QTL interval spanned approximately 3.3 cM, had a peak LOD score of 2.9, and an R2 of 33.3% (Table 2, Figure 1). The two QTL explained approximately 26.4% of the total variation in FDS 4. The tenth QTL that was associated with the RRS trait (qRRS001-01) was identified by the marker interval ss107912585-ss107920575 on chromosome 3 (LG N). The QTL interval spanned approximately 4.3 cM, had a peak LOD score of 4.2, and an R2 of 9.9% (Table 2, Figure 1). The eleventh QTL that was associated with the RRS trait (qRRS001-02) was identified by the marker interval ss107929213-ss107929551 on chromosome 4 (LG C1). The QTL interval spanned approximately 5.8 cM, had a peak LOD score of 5.0, and an R2 of 8.6% (Table 3, Figure 1). Another QTL associated with the RRS trait (qRRS001-03) was identified by the marker interval ss4969823-ss107924619 on chromosome 18 (LG G). The QTL interval spanned approximately 3.3 cM, had a peak LOD score of 2.9, and an R2 of 33.3% (Table 2, Figure 1). However, this QTL is the same as gFDS004-02 described above. The three QTL explained approximately 53.5% of the total variation in RRS.

Using IM, three additional QTL were identified for SDS foliar resistance (FDS). The twelfth QTL that was associated with the FDS 3 trait (qFDS003-05) was identified by the marker interval



Figure 1. The 'PI 438489B' by 'Hamilton' SNP-based genetic linkage map and the positions of the fifteen QTL that underlie SDS resistance found in this RIL population (n=50).



Figure 1. Continued.



Figure 1. Continued.

Chr_17b

Chr_17c



Chr 17d

Chr_18

Figure 1. Continued.

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ss107917031–ss107912977 on chromosome 6 (LG C2). The QTL interval spanned approximately 15.9 cM, had a peak LOD score of 2.9, and an R2 of 2.4% (Table 2, Figure 1). The thirteenth QTL that was associated with the FDS 3 trait (qFDS003-06) was identified by the marker interval ss107915722–ss107918074 on chromosome 8 (LG A2). The QTL interval spanned approximately 13 cM, had a peak LOD score of 2.7, and an R2 of 17.4% (Table 2, Figure 1). The fourteenth QTL that was associated with the FDS 4 trait (qFDS004-03) was identified by the marker interval ss107924445–ss107918378 on chromosome 4 (LG C1). The QTL spanned approximately 26.6 cM, had a peak LOD score of 2.6, and an R2 of 4.8% (Table 2, Figure 1).

QTL that underlie both FDS 3 (qFDS003-03 and qFDS003-05), and FDS 4 (qFDS004-03 and qFDS004-04) were identified on chromosome 6 (LG C2) (Table 2, Figure 1). Similarly, QTL that underlie both FDS 4 (qFDS004-02) and RRS (qRRS001-03) were identified on chromosome 18 (LG G) (Table 2, Figure 1).

Common Genes within the QTL Regions

Using CIM, approximately 1,459 genes were identified within the genomic regions containing QTL for FDS 2, FDS 3, FDS 4, and RSS (Table 4-Supplementary data). The most frequently occurring genes (> 5 copies) within these regions were those encoding for plant-type serine-threonine protein kinase (Van Bentem et al., 2008), pentatricopeptide repeat-containing protein, MYB-related protein, RNA binding protein, ATP binding cassette transporter, protein phosphatase, ologopeptide transporter, ascorbate oxidase, cytochrome P450, copper transporter protein, CDC2-related kinase, amino acid transporter, zinc finger protein, mitochondrial carrier protein, membrane associated finger protein, homeobox protein, ATP dependent helicase, ATP independent helicase, subtilisin-related serine protease, sensor histidine kinase, RAS-related GTPase, NAD dependent epimerase, glucosyl transferase, F-Box/LRR protein, centurin, aspartyl protease, aquaporin transporter, thioredoxin-related protein, synthaxin, solute carrier protein, and many others (Table 4-Supplementary data).

Discussion

The first 'PI438489' by 'Hamilton' high density SNP-based genetic map presented here is among the few high density maps published in soybean to date (Choi et al., 2007; Hyten et al., 2010b; Vuong et al., 2010) and among the few in crops and other plant species (Feltus et al., 2010). The map can be used to discover new QTL for other agronomic traits and to decipher the candidate genes that underlie these traits.

Several chromosomes are fragmented in to more than one LG (5a and 5b; 11a, 11b, 11c, and 11d; 13a and 13b; 15a and 15b; 16a and 16b; 17a, 17b, 17c, and 17d; 20a and 20b) which increased the number of LGs into 31 compared to 20 chromosomes may be due to the relatively small population size used in this study (50 RILs). We are in the process of increasing this population size to >100 RILs in order to construct a robust genetic linkage map based on this population and to

accurately detect QTL for important agronomic traits.

Previously, Kassem et al. (2007) reported nine QTL for SDS resistance. In the same publication, a decade (1996-2006) of SDS resistance QTL mapping in ExF was also summarized. QTL reported in Kassem et al. (2007) were identified using the 'Essex' by 'Forrest' RIL population and the updated ExF genetic map based mainly on SSR markers (Kassem et al., 2006). Later reports increased the number of QTL recognized to 14 (Prabhu et al., 1999; Farias et al., 2007; Kazi et al., 2007, 2008) in diverse germplasm. In the study reported herein, fourteen QTL for SDS resistance were identified, twelve QTL for foliar (FDS) and three QTL for root (RRS) symptoms caused by F. virguliforme infection. At the same position of the QTL (gFDS002-01) identified on chromosome 1 (LG D1a), QTL for soybean cyst nematode (SCN) resistance were identified using this same population (Yue et al. 2001). QTL for several other agronomic traits such as javanese root-knot nematode resistance, protein concentration, oil content, and leaf length, were also identified in this region (SoyBase 2010). At the same region of qFDS002-02 identified on chromosome 10 (LG O), only one QTL for Southern root-knot nematode resistance was identified (SoyBase 2011; Li et al., 2001). Similarly, on the same region containing qFDS002-03 identified on chromosome 19 (LG L), QTL for lodging, seed linoleic acid content, plant height, and sucrose concentration were identified and mapped (SoyBase, 2011). Interestingly, a QTL for sclerotinia stem rot resistance was identified and mapped 5 cM downstream from this region containing qFDS002-03. The two QTL qFDS003-01 and qFDS003-02 identified on chromosome 2 (LG D1b) were only 8 cM apart and QTL for several morphological traits such as plant height, yield, leaf width, and leaf length were identified within this region (SoyBase, 2011). On chromosome 6 (LG C2), the genomic region containing gFDS003-03, qFDS003-05, qFDS004-03, and qFDS004-04 spanned from 17 cM to 40 cM (approx. 23 cM) and could be designated as one region that underlie resistance to both FDS 3 and FDS 4. Approximately 10 cM upstream of this region, a QTL for SCN resistance was identified (SoyBase, 2011; Wang et al., 2001).

Using the same RIL population (PI 438489BxH), QTL for SCN resistance were mapped 10 cM upstream and 60 cM downstream from the region containing qFDS003-03, qFDS003-05, gFDS004-03, and gFDS004-04 (Yue et al., 2001) on chromosome 6 (LG C2). Approximately 80 cM downstream of this region, QTL for SDS resistance were identified and mapped in different genetic backgrounds (lqbal et al., 2001; Kassem et al., 2006, 2007). Recently, QTL for SDS resistance (cgRfs4; leaf DX) were identified approximately 60 cM from the first region containing qFDS003-03, qFDS003-05, qFDS004-03, and qFDS004-04 (Kazi et al., 2008) on LG C2 which indicates the existence of several SDS resistance QTL on this chromosome. On chromosome 8 (LG A2), a cluster of SDS resistance QTL containing both qFDS003-04 and qRRS001-03 was identified from 2 to 28 cM (26 cM span). Interestingly, a QTL for SCN resistance (Concibido et al., 2004), and SDS resistance (SoyBase, 2011) were identified 4 cM, and 20 cM from this region, respectively. On chromosome 11 (LG B1), QTL for hypocotyls length and aluminum tolerance were identified on the same region containing qFDS004-01 and a cluster of QTL for SCN resistance was identified approximately 40 cM from this region (Yue et al., 2001; Concibido et al., 2004). Chromosome 18 (LG G) is of particular interest since it contains a cluster of SDS and SCN resistance QTL identified in different genetic backgrounds (Kassem et al. 2006, 2007; Kazi et al., 2008; Concibido et al., 2004). In this study, the region identified on this chromosome contains both qFDS004-02 and qRRS001-03 which indicates that this region confers resistance to both foliar (FDS) and root (RRS) symptoms. This agreed with earlier results (Meksem et al., 1999; Iqbal et al., 2001; Kassem et al., 2006, 2007) was in contrast to what has been previously reported in 'Flyer' by 'Hartwig' RIL population (Kazi et al., 2008).

Interestingly, a QTL for SDS resistance was mapped only 2 cM from the region containing qRRS001-01 identified on chromosome 3 (LG N) (Njiti et al. 2002). On chromosome 4 (LG C1) containing qRRS001-02 and qFDS004-03, no QTL for disease resistance have been mapped in this region; however, QTL for several morphological traits such as seed weight, pod maturity, and seed yield have been mapped at this location (SoyBase, 2011).

The most frequently occurring genes within these QTLs were plant-type serine-threonine protein kinase, pentatricopeptide repeat-containing protein and MYB-related protein. The four QTLs collectively had ten copies of each gene. QTL qFDS003-01 had four copies of each of the three genes which is the greatest number of copies. The serine-threonine protein kinase gene is a stress-induced gene which helps plants to cope with environmental stresses such as drought, salinity, and cold (Xinguo et al., 2010). The pentatricopeptide repeat-containing protein is essential for RNA editing in chloroplasts and mitochondrial transcripts (Kotera et al., 2005; Bentolila et al., 2010), and the MYB-related protein is a member of a class of transcription factors identified to be involved in plant stress responses (Martin and Paz-Ares, 1997, Ge et al., 2011, Uehara et al., 2010).

Conclusions

In this study, we constructed the first 'PI438489B' by 'Hamilton' SNP-based map and identified fourteen QTL that underlie SDS resistance including both resistance to foliar and root rot symptoms caused by *F. virguliforme* infection. The map presented here is among the few high density map published in soybean (Choi et al., 2007; Hyten et al. 2010b; Vuong et al., 2010) and other plant species (Feltus et al., 2010). The map can be used to discover new QTL for other agronomic traits and to decipher the candidate genes that underlie these traits.

Seven of the intervals encompassing the QTL had been identified previously (on LGs C1, C2, D1b, G, L, N and O) associated with resistance to SDS and this provided strong evidence that the map and trait scoring methods used were valid. Seven QTL were identified in novel locations (LGs A2 (2), B1, C2, D1a, D1b and O). Some of these QTL might be confirmed in NILs or by further mapping in a larger portion of the population. Equally some QTL may prove to have been type II errors by additional experiments.

Several studies reported SNP markers associated with QTL for several traits in soybean (Vuong et al., 2007). However, only

one of these studies reported an exclusively SNP-based genetic map (Hyten et al., 2010a). However, SNPs are being integrated with SSRs, RFLPs, AFLPs, and other markers to construct high density genetic linkage maps (Hyten et al. 2010a; Hyten et al. 2010b) that are being used by the soybean research community and biotechnology industry to help improve soybean (Vuong et al., 2007). Therefore, the SNP-based map and the QTL regions presented here can be used in breeding programs to develop soybean cultivars with increased resistance to SDS.

List of Abbreviations

SDS: Sudden death syndrome; SNP: Single nucleotide polymorphism; RFLP: Restriction fragment length polymorphism; AFLP: Amplified fragment length polymorphism; RAPD: Random amplified polymorphic DNA; DAP: Days after planting; RIL: Recombinant inbred line.

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