**Article**

**QTL Mapping for Gummy Stem Blight Resistance in Watermelon (Citrullus spp.)**

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**Abstract:** Watermelon (Citrullus lanatus) is an economically important fruit crop worldwide. Gummy stem blight (GSB) is one of the most damaging diseases encountered during watermelon cultivation. In the present study, we identified quantitative trait loci (QTLs) associated with GSB resistance in an F2 population derived from a cross between maternal-susceptible line '920533' (C. lanatus) and the paternal-resistant line 'PI 189225' (C. amarus). The resistance of 178 F2 plants was assessed by two different evaluation methods, including leaf lesion (LL) and stem blight (SB). To analyze the QTLs associated with GSB resistance, a linkage map was constructed covering a total genetic distance of 1070.2 cM. QTL analysis detected three QTLs associated with GSB resistance on chromosome 8 and 6. Among them, two QTLs, qLL8.1 and qSB8.1 on chromosome 8 identified as major QTLs, explaining 10.5 and 10.0% of the phenotypic variations localizing at same area and sharing the same top markers for both LL and SB traits, respectively. A minor QTL, qSB6.1, explains 9.7% of phenotypic variations detected on chromosome 6 only for the SB trait. High-throughput markers were developed and validated for the selection of resistant QTLs using watermelon accessions, and commercial cultivars. Four potential candidate genes were predicted associated with GSB resistance based on the physical location of flanking markers on chromosome 8. These findings will be helpful for the development of watermelon cultivars resistant to GSB.

**Keywords:** gummy stem blight; high-resolution melting; single nucleotide polymorphism; linkage map; quantitative trait loci

1. **Introduction**

Watermelon (Citrullus lanatus; 2n = 2x = 22) is one of the most important fruit crops, accounting for 11.4% of cultivation area among fruit around the world [1]. Watermelon consists of more than 91% of water and contains functional compounds such as lycopene, citrulline, and β-carotene that contribute in the uptake of a balanced diet and healthy nutrients [2,3].

Gummy stem blight (GSB), caused by Stagonosporopsis species pathogen Didymella bryonie (Auersw.) Rehm, causes severe damages to Cucurbit crops, including watermelon. GSB occurs mainly under high temperature and humidity [4,5]. GSB results in small brown spots on the leaves, yellow coloration of the leaves with round or irregular lesions, and greyish-brown stems followed by withering [6]. In particular, leaves and stems tend to dry
out and fall early [7], resulting in the rapid advancement of GSB after fruit setting, which leads to watermelon blight before the harvest [8]. The GSB can cause crown blight, severe defoliation, rotting of fruits, and deterioration of the fruits during the transportation and storage, resulting in heavy economic losses. The yield loss could be as high as 100% during the warm and humid season under high disease pressure [9].

In order to control GSB in watermelon, agricultural fungicides should be sprayed prior to the appearance of disease, and crop rotation ought to be performed in case of severe occurrence at the greenhouse or field [9,10]. However, excessive use of chemicals to control GSB can be hazardous to the ecosystem and agricultural safety. Cultural and agronomic controls are expensive and labor-intensive [10]. The development of disease-resistant cultivars can be the best alternative for controlling GSB [11,12]. The watermelon accessions ‘PI 189225’ and ‘PI 271778’ (C. amarus), from the US Germplasm Collection of USDA-ARS (United States Department of Agriculture-Agricultural Research Service) have been reported to exhibit GSB resistance [13,14]. Other accessions including, ‘PI 279461’, ‘PI 526233’, and ‘PI 482283’ have also been reported as new resources for GSB resistance [15]. According to Norton et al. [16], the GSB resistance in ‘PI 189225’ was controlled by a single gene, db; however, recent research by Gusmini et al. [17] suggested the involvement of multiple genes with environmental factors for GSB resistance. Further studies are required to determine the mode of inheritance associated with resistance to GSB.

Recently, QTL mapping analysis was carried out for resistance to bacterial fruit blotch (BFB), anthracnose race 1, fusarium wilt race 1, and papaya ringspot virus-watermelon strain (PRSV-W) using bi-parental mapping populations in watermelon. [18–22]. The quantitative trait loci (QTL) mapping for Fusarium oxysporum f. sp. niveum race 2 of watermelon has also been carried out [23]. The results of QTL mapping associated with GSB resistance have been reported in various cucurbits. In cucumber, major QTLs related to GSB resistance on leaves and stems were identified on chromosome 5 and 6, respectively [24,25]. Meanwhile, in melon, QTLs conferring resistance of GSB were detected on chromosome 9 [26,27]. The QTL identification linked to GSB (isolate: JS002; JAAS, Nanjing, China) resistance in watermelon were performed using bulked segregant analysis (BSA) [28]. This study detected a QTL (Qgsb8.1) on chromosome 8 with an LOD score ranged from 13.6 to 16.4, explaining 31.54 to 32.42% phenotypic variations in two different seasons [28]. Recently, GSB-resistant QTLs were mapped on chromosome 3, 5, and 7 explaining between 6.4 to 21.1% of phenotypic variation [29].

Molecular markers have been developed and deployed for marker-assisted selection in watermelon disease-resistant breeding. Cleaved amplified polymorphic sequence (CAPS) markers and high-resolution melting (HRM) based markers have been reported using ‘Arka Manik’ and ‘PI 254744’ resistant to powdery mildew race 1W in watermelon [30,31]. In addition, CAPS molecular markers related to zucchini yellow mosaic virus (ZYMV) resistance were developed in watermelon [32]. The HRM and KASP assays are relatively simple, rapid, and cost-effective approaches compared to other methods such as CAPS, sequence characterized amplified region (SCAR), and simple sequence repeat (SSR) analysis [33]. Recently, KASP assay were developed for GSB resistance in watermelon [28,29]. However, more high-throughput molecular markers are required for efficient prediction of GSB resistance to use in marker-assisted selection programs.

In the present study, QTL analysis was conducted to map resistant loci associated with GSB resistance in watermelon, using next-generation re-sequencing based SNPs. High-throughput molecular markers linked to GSB resistance were developed and validated using watermelon accessions and commercial cultivars. In addition, four potential candidate genes were predicted in QTL regions associated with GSB resistance in watermelon.

2. Results

2.1. Resistance to GSB in Parents and an F2 Population

The average of leaf lesion (LL) of ‘PI 189225’ and ‘920533’ to GSB for the 20 plants was 14.04% and 73.73%, respectively (Figure S1). Both the resistant and susceptible parental lines
showed significant difference in disease severity (Student’s t-test; \( p < 0.001 \)). According to the combined disease index (DI) scoring, a total of 178 plants in \( F_2 \) population segregated to 65 susceptible and 113 resistant plants, which do not fit into 3:1 (R:S) segregation ratio from Chi-square value of 12.59 at \( p < 0.001 \) (Table S1). The frequency distribution of LL of the \( F_2 \) population displayed a pattern with two peaks (Figure 1a). Plants of the left peak (resistant) were greater in number than those of the right peak (susceptible). Frequency distribution plot of SB of \( F_2 \) population was drawn to a right-skewed pattern (Figure 1b). The frequency distribution curves and segregation data supported the presence of a quantitative mode of inheritance of resistance to GSB in \( F_2 \) population. These results imply that multiple genes in this population control disease resistance.

Figure 1. Frequency distribution of disease severity. (a) Leaf lesion (LL) and (b) stem blight (SB) among parental lines and \( F_2 \) population. Green, orange and purple bar indicates resistant parent ‘PI 189225’, susceptible parent ‘920533’, and an \( F_2 \) population obtained from a cross between ‘920533’ \( \times \) ‘PI 189225’, respectively. Error bars indicates the standard error (SE).

2.2. SNP Genotyping Using Fluidigm® SNP Type™ Assays

The genotypes of 178 individuals in the \( F_2 \) population were analyzed using previously developed 438 Fluidigm® SNP Type™ assays [34]. Polymorphic SNPs were clustered into three groups in a graph according to the ratios of FEM to HEX fluorescence for individuals of \( F_2 \) population and two parental lines (Figure S2). Red and green-clustered dots indicate homozygous genotypes, and blue-clustered dots indicate heterozygous genotypes (Figure S2). Among these 438 Fluidigm® SNP Type™ assays, 113 polymorphic SNP between the parental lines were selected using Fluidigm® genotyping platform and used for the linkage analysis in \( F_2 \) population (Table 1).
### Table 1. Summary of the distribution of single nucleotide polymorphism (SNP) markers in watermelon genetic map.

| Linkage Group | Number of Markers | Map Length (cM) | Marker Density (cM/Marker) | Fluidigm® Markers | Previously Reported HRM Markers | Newly Designed HRM Markers | Previously Reported KASP™ Markers |
|---------------|-------------------|----------------|-----------------------------|-------------------|-------------------------------|----------------------------|----------------------------------|
| Chr.1a        | 6                 | 59.1           | 9.85                        | 1                 | 1                             | 4                         | -                                |
| Chr.1b        | 4                 | 10.0           | 2.5                         | 3                 | -                             | -                         | -                                |
| Chr.1c        | 7                 | 39.2           | 5.6                         | 3                 | 1                             | 3                         | -                                |
| Chr.2a        | 11                | 57.8           | 5.25                        | 10                | -                             | -                         | -                                |
| Chr.2b        | 7                 | 34.5           | 4.97                        | 6                 | -                             | -                         | -                                |
| Chr.3a        | 5                 | 21.5           | 4.3                         | 3                 | 1                             | -                         | -                                |
| Chr.3b        | 5                 | 30.2           | 6.04                        | 4                 | -                             | -                         | -                                |
| Chr.4         | 10                | 81.4           | 8.14                        | 4                 | -                             | 6                         | -                                |
| Chr.5a        | 5                 | 44.0           | 8.8                         | 4                 | -                             | 1                         | -                                |
| Chr.5b        | 6                 | 44.7           | 7.45                        | 6                 | -                             | -                         | -                                |
| Chr.6         | 22                | 97.0           | 4.41                        | 5                 | -                             | 17                        | -                                |
| Chr.7         | 20                | 98.8           | 4.94                        | 12                | -                             | 8                         | -                                |
| Chr.8         | 38                | 131.7          | 3.47                        | 10                | -                             | 21                        | 7                                |
| Chr.9         | 18                | 134.2          | 7.46                        | 15                | -                             | 3                         | -                                |
| Chr.10        | 14                | 89.5           | 6.39                        | 11                | 1                             | 2                         | -                                |
| Chr.11        | 10                | 96.3           | 9.63                        | 7                 | -                             | 3                         | -                                |
| Unmapped      | 23                | -              | -                           | 9                 | -                             | 11                        | 3                                |
| Total         | 211               | 1070.2         | 4.69                        | 113               | 4                             | 84                        | 10                               |

### 2.3. Identification of SNPs Using NGS

In order to develop SNP markers that cover the whole genome, a homozygous susceptible line (‘920533’) and a resistant line (‘PI 189225’) were sequenced with the next generation sequencing method using a Hiseq™ 4000 sequencer (Illumina Inc., San Diego, CA, USA). The raw reads for the parental lines were generated about 17.8 Gbp and 16.8 Gbp, respectively. By trimming the raw data, the sequences were shortened to 13.6 Gbp for ‘920533’ and 13.1 Gbp for ‘PI 189225’. The 320 Mbp (90.13%) and 305 Mbp (85.92%) trimmed data of the resistant and susceptible parental lines was mapped to the reference genome, respectively. The mapped sequences were used to detect SNPs between ‘920533’ and ‘PI 189225’. A total of 6 million raw SNPs were obtained between the parental lines (Table S2) [35].

### 2.4. Development of HRM Markers Using Identified SNPs

Among the total SNPs obtained, 5 million SNPs were selected as homozygous SNPs (Table S2) [35]. Subsequently, 4.3 million SNPs were filtered for A/G, A/C, T/G, and T/C combinations, excluding A/T and G/C. Furthermore, around 2 million SNPs whose flanking sequence (single copy) was found only once in the whole genome were chosen using the basic local alignment search tool nucleotide (BLASTN). Finally, 354,860 SNPs located on the genic region were further selected, and then 888 HRM markers that were evenly distributed on the watermelon genome were developed (Table S2) [35]. Polymorphisms of these HRM markers were tested using two parental lines, the F1 plants and 13 randomly selected plants of the F2 population. As a result, 84 HRM markers were identified showing polymorphic melting curves between the parents. A total of 88 HRM primer sets, including the 84 newly designed HRM markers and four previously developed HRM markers were used to analyze the genotypes of 178 F2 individuals (Table S3 and Figure S3) [36]. Newly designed HRM markers were abbreviated on watermelon gummy stem blight resistance SNPs (WGRS).

### 2.5. Construction of a Watermelon Linkage Map

A genetic linkage map was constructed using 178 F2 plants segregating for GSB resistance, and 211 SNP markers. The genetic map was generated using 211 SNP markers consisting of 88 high-resolution melting (HRM), 113 Fluidigm®, and 10 recently reported kompetitive allele specific PCR (KASP™) markers [28]. Among them, 188 SNP markers that included 104 Fluidigm® markers, 77 HRM markers, and 7 previously reported KASP™ markers were mapped on 16 linkage groups (LGs) whereas 23 markers remained unmapped (Figure 2 and Figure S4). LGs were named according to chromosomes, as there was a high identity between the genetic map and physical map. The total length of the genetic linkage
map was 1070.2 cM where the average length of the linkage groups was 66.89 cM, and the average interval between the markers was 5.69 cM with a mean of 11.8 markers per linkage group (Table 1 and Table S4). The lengths of linkage groups ranged from 10.0 cM to 134.2 cM (Table 1). The number of markers per linkage group ranged from 4 (Chr. 1b) to 38 (Chr. 8). In the genetic map, most of the markers’ orders were collinear to the physical map.

Figure 2. Genetic map of watermelon constructed by SNP markers. Numbers on the left side correspond to the genetic distance in cM from the top of each chromosome. Names on the right side indicate marker name that represents marker type according to color (blue, Fluidigm® SNP Type™ assays; red, previously reported HRM markers; green, newly designed HRM markers; purple, previously reported KASP™ markers).

2.6. Identification of QTLs Conferring Resistance to GSB in Watermelon

A total of three QTLs, qSB6.1, qLL8.1, and qSB8.1 were identified associated with GSB resistance in 178 individuals of the F2 population (Figure 3a,b). Two QTLs located on chromosome 8 were detected as major QTLs and one QTL located on chromosome 6 as a minor QTL (Figure 3a,b and Table 2). The QTLs, qLL8.1 and qSB8.1 explained 10.5 and 10.0% of phenotypic variations ($R^2$, %) for GSB resistance, with LOD scores of 4.28 and 4.02, respectively (Table 2). Interestingly, qLL8.1 and qSB8.1 both localized in the same area (85 cM) and sharing the same top markers (20.6 to 21.5 Mbp) on chromosome 8 (Figure 3a,b and Table 2). Whereas, qSB6.1 on chromosome 6 was detected only for SB trait at 57.1 cM and explained 9.7% of phenotypic variations in GSB resistance with a LOD score of 3.96.
Alleles conferring resistance to GSB were contributed by the resistant parent (‘PI 189225’) because additive effect of each QTL were positive (>0) (Table 2). Disease score associated with specific QTL (qSB8.1 and qSB6.1) genotypes were evaluated and compared to determine the independent and combined QTLs genetic effects. A clear trend related to resistance level for GSB was observed with the presence/absence of susceptible (S) and resistance (R) alleles of the QTLs (Figure 4a). The genotype carrying S alleles for both QTL showed reduced resistant level significantly for the SB trait with average score of 1.69 (Figure 4a). The genotypes carrying S allele for qSB8.1 and R allele qSB6.1 showed average disease score 1.38 whereas the genotypes carrying R allele for qSB8.1 and S allele qSB6.1 showed average disease score 0.94 (Figure 4a).

When the genotypes carrying R alleles for the both QTLs were compared the average disease score was significantly reduced as up to 0.38 (Figure 4a). These results revealed the additive effect of the two QTL when combine (Figure 4a). For LL trait, boxplot results revealed that the homozygous resistant genotype ‘B’ of qLL8.1 is associated with increased

### Table 2. QTL mapping results for leaf lesion (LL) and stem blight (SB) traits on chromosome 6 and 8 in the F2 population.

| Trait       | QTL   | Chr. | Position (cM) | Marker Interval | Location (bp) | LOD  | R² (%) | Gene Effect |
|-------------|-------|------|---------------|----------------|---------------|------|--------|-------------|
| Leaf lesion | qLL8.1 | 8    | 85.376        | chr6:WGRS512   | 20,663,001–21,535,005 | 4.28 | 10.5   | 9.21         |
| Stem blight | qSB6.1 | 6    | 57.139        | chr6:WGRS131   | 7,533,583–7,625,669 | 3.96 | 9.7    | 9.17         |
|             | qSB8.1 | 8    | 85.263        | chr8:WGRS102   | 20,663,001–21,535,005 | 4.02 | 10.0   | 10.50        |

1 Positions of the markers on the linkage map. 2 LOD, logarithm of odds. 3 Percent of the phenotypic variation explained by the QTL.
resistance compared to the homozygous susceptible genotype ‘A’ of qLL8.1 in the F2 population (Figure 4b).

![Box plots of tightly linked SNP markers to QTL for leaf lesion (LL) trait in the F2 population.](image)

**Figure 4.** Comparison of GSB resistance in F2 population according to genotypes for QTLs qSB6.1 and qSB8.1 (a) The mean of stem blight (SB) according to genotypes of two QTL-linked markers (chr8_WGRS(3)185 and chr6_WGRS(3)092) where R, resistant genotype and S, susceptible genotype. Bars on graphs indicate standard error and different small letters (a, ab, bc and c) refer to significant differences (p < 0.05) according to Duncan multiple range test. (b) Box plots of tightly linked SNP markers to QTL for leaf lesion (LL) trait in the F2 population. A, genotype of female parent (‘920533’); B, genotype of male parent (‘PI 189225’); H, genotype of heterozygote. The pink dots within the boxplots represent the mean of lines harboring respective alleles.

**2.7. Validation of Flanking Markers Using Watermelon Accessions and Cultivars**

A total four markers were developed and validated for GSB-resistant QTL selection. Among them, two markers were developed on chromosome 8 for QTLs (qLL8.1 and qSB8.1) and two on chromosome 6 for QTL (qSB6.1) (Table 3). For all these markers, disease severity for SB trait was significantly lower in the individuals having homozygous-resistant alleles R compared to the individuals carrying homozygous susceptible alleles S (Figure 4a). In order to validate QTLs effect of qLL8.1 and qSB8.1, two flanking markers (chr8_WGRS240 and chr8_WGRS(3)185) were converted into KASP™ markers and validated in 9 watermelon accessions and 13 commercial cultivars (Table 3). These two flanking markers tightly linked to qSB8.1 and qLL8.1 were found to be associated with resistance to GSB because 6 accessions (C. amarus) had high resistance to GSB and possessed resistant genotypes for (KASP_WGRS240 and KASP_WGRS(3)185) (Table 4). To test the utility of the developed KASP™ markers linked to qSB6.1, the markers (KASP_WGRS(3)089, KASP_WGRS(3)092) were also validated in 9 accessions and 13 commercial cultivars (Table 4). However, two flanking markers linked to qSB6.1 didn’t show a very strong correlation with resistance to
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GSB because ‘PI 279461’ and some commercial cultivars (‘Fair Fax’, ‘Crimson sweet’, and ‘Charleston Gray’) susceptible to GSB held resistant genotypes for (KASP_WGRS(3)089 and KASP_WGRS(3)092). However, these markers in combination can maximize the selection of QTL segments for GSB resistance.

Table 3. Primer sequence details of (KASP™) markers associated to GSB resistance.

| Marker Name          | Chromosome | Location (bp) | Allele       | Fluorescent Primer Sequence (5′-3′) | Sequence (5′-3′) |
|----------------------|------------|---------------|--------------|-------------------------------------|-----------------|
| KASP_WGRS(3)089      | 6          | 7,533,583     | T            | Allele-specific 1 (FAM)             | GAATCAACGATCCAGACCA |
|                      |            |               | C            | Allele-specific 2 (HEX)             | AATTCACCTGAATCCAGACCA |
|                      |            |               | -            | Common                              | GTAAAGGCGCTAAAAGGACCAA |
| KASP_WGRS(3)092      | 6          | 7,625,669     | A            | Allele-specific 1 (FAM)             | GAGGCCAACAGAAGAAGAGCACCA |
|                      |            |               | G            | Allele-specific 2 (HEX)             | GAGGCCAACAGAAGAAGAGCACCA |
|                      |            |               | -            | Common                              | GAGGCCAACAGAAGAAGAGCACCA |

Table 4. Validation of KASP™ markers developed for the gummy stem blight (GSB) resistance in QTL regions.

| Watermelon Accessions/Commercial Cultivars | Phenotype | Genotype |
|------------------------------------------|-----------|----------|
| Common Name                             | Origin    | Scientific Name 1 | Disease Index (DI) 2 | KASP_WGRS240 (Chr.8) | KASP_WGRS(3)185 (Chr.8) | KASP_WGRS(3)089 (Chr.6) | KASP_WGRS(3)092 (Chr.6) |
| PI 189225                               | Congo     | CA        | R               | R               | R               | R               | -               |
| ‘92/553’                                | South Korea | CL        | S               | S               | S               | S               | S               |
| PI 482283                               | Zambia    | CA        | R               | R               | R               | R               | R               |
| PI 164248                               | Liberia   | CM        | R               | S               | S               | R               | -               |
| PI 503334                               | Zambia    | CA        | R               | R               | R               | R               | R               |
| PI 244019                               | South Africa | CA    | R               | R               | R               | R               | R               |
| PI 482315                               | Zimbabwe  | CA        | R               | R               | R               | R               | R               |
| PI 379243                               | North Macedonia | CA | R               | R               | R               | R               | R               |
| PI 279461                               | Japan     | CL        | S               | S               | S               | S               | R               |
| PI 505960                               | Zambia    | CL        | S               | S               | S               | S               | S               |
| ‘Au-Jubilant’                           | USA       | CL        | S               | S               | S               | S               | S               |
| ‘Au-Producer’                           | USA       | CL        | S               | S               | S               | S               | S               |
| ‘Crimson Sweet’                         | USA       | CL        | S               | S               | S               | R               | S               |
| ‘Soupsipperlasoskul’                    | South Korea | CL    | S               | S               | S               | H               | H               |
| ‘Heugho’                                | South Korea | CL    | S               | S               | S               | -               | S               |
| ‘Norarmacindara’                        | South Korea | CL    | S               | S               | S               | S               | S               |
| ‘Daljona’                               | South Korea | CL    | S               | S               | S               | H               | H               |
| ‘Urickul’                               | South Korea | CL    | S               | S               | S               | S               | S               |
| ‘Orenjiking’                            | South Korea | CL    | S               | S               | S               | H               | H               |
| ‘Santakkul’                             | South Korea | CL    | S               | S               | S               | H               | H               |
| ‘Charleston Gray’                       | USA       | CL        | S               | S               | S               | R               | S               |
| ‘Seotaeja’                              | South Korea | CL    | S               | S               | S               | S               | S               |

1 CA, C. amarus Schrad.; CL, C. lanatus (Thunb.) Matsum. & Nakai; CM, C. mucosospermus (Fursa) Fursa. 2 Disease index (DI) was presented as resistant (R, 0–3) and susceptible (S, 4–7).

2.8. Identification of Candidate Genes for GSB Resistance

Two major QTLs (qLL8.1 and qSB8.1) on chromosome 8, flanked by two markers (chr8_WGRS240 and chr8_WGRS(3)185) (Figure 3b). The interval between these two flanking markers covered approximately 0.87 Mbp on chromosome 8, a region containing 83 genes in Cucurbit Genomics Database (http://cucurbitgenomics.org/; accessed on 15 June 2020) (Table S5). Among the 83 genes, three were encoding receptor-like kinase (RLK) domain-containing proteins and one leucine-rich repeat (LRR) receptor-like protein kinase were considered candidate genes for GSB resistance (Table 5). Three RLK genes (Clato22133, Clato22184, and Clato22195) were located at 825, 219, and 114 kb upstream of the most significant SNP (chr8_WGRS(3)185), respectively (Table 5). An LRR-RLK gene (Clato22196) was identified at 111 kb upstream of the highly significant SNP (Table 5). These genes could be potential candidate genes for GSB resistance. However, further studies would confirm their role in GSB resistance in watermelon.
Table 5. Candidate genes associated with gummy stem blight (GSB) resistance and their Gene Ontology (GO) description.

| Gene ID     | Location            | Annotation                        | SNP ('920533'/'PI 189225') |
|-------------|---------------------|-----------------------------------|-----------------------------|
|             | Chr. | Position       | Annotation                     | Nucleotide     | Position (bp) |
|             |      | (bp, Start–End) |                               | A/G            |               |
|             |      |                |                               | A/C            |               |
| Cla022133   | 8    | 20,710,211–20,711,725 | Receptor-like protein kinase | A/G            | 20,710,460   |
|             |      |                |                               | A/C            | 20,710,552   |
|             |      |                |                               | A/G            | 20,710,776   |
|             |      |                |                               | C/A            | 20,710,810   |
|             |      |                |                               | A/G            | 20,710,904   |
|             |      |                |                               | G/A            | 20,711,132   |
|             |      |                |                               | G/T            | 21,316,051   |
|             |      |                |                               | G/A            | 21,316,141   |
|             |      |                |                               | T/C            | 21,316,345   |
|             |      |                |                               | T/C            | 21,316,486   |
|             |      |                |                               | G/A            | 21,317,089   |
|             |      |                |                               | C/T            | 21,317,571   |
|             |      |                |                               | C/T            | 21,318,065   |
|             |      |                |                               | C/T            | 21,318,362   |
|             |      |                |                               | T/C            | 21,318,714   |
| Cla022184   | 8    | 21,315,878–21,318,792 | Receptor kinase | G/T            | 21,421,391   |
|             |      |                |                               | G/A            | 21,421,694   |
|             |      |                |                               | G/T            | 21,421,865   |
|             |      |                |                               | A/G            | 21,421,905   |
|             |      |                |                               | T/C            | 21,422,026   |
|             |      |                |                               | C/T            | 21,422,027   |
|             |      |                |                               | A/G            | 21,422,704   |
|             |      |                |                               | T/C            | 21,422,920   |
|             |      |                |                               | G/T            | 21,423,010   |
|             |      |                |                               | G/A            | 21,423,031   |
|             |      |                |                               | C/T            | 21,423,285   |
|             |      |                |                               | G/A            | 21,423,367   |
|             |      |                |                               | A/G            | 21,423,644   |
| Cla022195   | 8    | 21,421,183–21,423,645 | Receptor kinase | T/C            | 21,423,751   |
|             |      |                |                               | T/C            | 21,423,821   |
|             |      |                |                               | C/T            | 21,423,899   |
|             |      |                |                               | C/T            | 21,423,965   |
|             |      |                |                               | C/T            | 21,424,071   |
|             |      |                |                               | A/C            | 21,424,242   |
|             |      |                |                               | A/G            | 21,424,352   |
|             |      |                |                               | G/A            | 21,424,355   |
|             |      |                |                               | G/T            | 21,424,503   |

3. Discussion

Several different inheritance patterns have been described for controlling GSB resistance in cucurbits, such as a single recessive gene [16,37], single dominant [26,38], and multiple genes [11,17,28,29,37]. Inheritance of resistance to GSB in watermelon was previously described as a single recessive gene, db, from ‘PI 189225’ [16]. However, recent research has suggested that several loci might be involved in GSB resistance [15,17]. These discrepancies might be due to the use of different sources of resistance, mapping population, different pathogen isolates and inoculation methods [28]. In the present study, the segregation ratio of F2 plants supported that multiple genes are involved in resistance to GSB (Table S1). Two types of disease assessment methods, including leaf lesion (LL) and stem blight (SB), were used for the assessment of accurate phenotypes related to GSB resistance. Continuous distribution was observed in the present study for the F2 population (‘920533’ × ‘PI 189225’) in disease assessment results suggesting the quantitative control of GSB resistance.

D. bryoniae can infect the watermelon plants at any stage and on all parts of the plant, including the stem, foliar and fruits. Disease resistance to GSB normally assessed through spraying the inoculum with spore suspension at four to six leaf stage of watermelon seedlings [28,29,35]. Pathogenicity, aggressiveness and environmental factors such as temperature and humidity can influence the resistance evaluation [28]. Variation in the isolates’ aggressiveness, inoculation, post-inoculation disease management practices and disease scoring method can also affect the resistance evaluation and further mapping studies [28,29]. Recently, a research group observed less resistance level of ‘PI 189225’ when kept under 100% relative humidity after inoculation for 48 h compared to 24 h [28]. In the present study, for precise resistance evaluation we used a previously tested aggressive D. bryoniae isolate KACC 40937 and evaluated the resistance using two different criteria [8].
These methods could be useful for the separate resistance evaluation on different plant parts as well as comparison of the resistance on foliar and stem in the future studies.

The high-density genetic linkage maps are crucial to detect quantitative trait locus (QTL) for important traits such disease resistance [28,29]. With the advent of high-density molecular genetic linkage maps and advanced mapping technologies, it is achievable to assess the number of QTLs in the genome. Using modern and high-throughput sequencing technology, more genomes sequences are available, thus enabling the development of abundant molecular markers [39]. The earliest SNP markers based maps were developed and compared utilizing three populations in watermelon [40]. The spans of these linkage maps were 1144 cM, 1438 cM, and 1514 cM with average marker intervals of 3.4 cM, 3.8 cM, and 4.2 cM, respectively [40]. SNP markers have improved marker resolution in *Cucurbita pepo*, and melon [41,42]. In our study, we employed two different approaches to obtain SNP markers including NGS-based re-sequencing (HiSeq™ 4000) and Fluidigm® assays. We constructed a genetic linkage map with 188 SNP markers using 104 Fluidigm® assays, 77 HRM markers, and 7 previously reported KASP™ markers, which yielded a genetic linkage map spanning 1070.2 cM. Our linkage map was comparable with the previously developed linkage maps [29,39,40].

QTLs controlling resistance for GSB have been reported in cucurbits such as, cucumber and watermelon [24,25,28,29]. Two QTLs associated to GSB resistance on chromosome 4 (*GSB4*) and 6 (*GSB6b*) at genomic position 12 cM and 11 cM in cucumber (*Cucumis sativus*) were reported using introgression lines [11]. Another research group reported GSB-resistant QTLs using recombinant inbred lines (RIILs) and identified total six QTLs on chromosomes 3, 4, 5 and 6 in cucumber [24]. Among those QTLs, a locus *gsb5.1* was repeatedly detected in three seasons and explained 17.9% phenotypic variations [24]. Another QTL analysis reported five QTLs associated to GSB resistance in cucumber using RIILs population and major QTL *gsb-s6.2* was mapped on chromosome 6, which explained 22.7% phenotypic variations [25]. A QTL conferring resistance against GSB in watermelon was detected on chromosome 8 (*Qgsb8.1*) using an F2 segregating population developed by “PI 189225” as the resistance parents and “K3” as a susceptible parent [28]. *Qgsb8.1* was mapped between two SNP markers KASP_JS9168 and KASP_JS9383 spanning a 0.57 Mb region at physical location of 11,425,655–11,995,922 on chromosome 8 [28]. Another recent study detected three QTLs linked to GSB resistance in watermelon on chromosomes 3, 5 and 7 using F2:3 mapping population developed by crossing susceptible line Crimson Sweet and resistant accession PI 482276 [29]. We identified three QTLs on chromosome 6 (*qSB6.1*) and 8 (*qLB8.1* and *qSB8.1*) for LL and SB traits; among them, two QTLs for the SB traits were co-located and shared the common markers. However, the genomic and physical position of the recently detected QTL *Qgsb8.1* was different from the QTLs identified in present study (Table 2) [28]. Although the source of resistance was the same (PI 189225) in both studies but Ren et al. [28] used local *S. cucurbitacearum* (syn. *D. bryoniae*) isolate collected from east China for resistance assessment, whereas we used *D. bryoniae* Korean isolate. Furthermore, the physical position of the QTLs detected by Ren et al. [28] on chromosome 8 corresponded to 11.4 to 11.9 Mbp on the reference genome, whereas we detected QTLs at the genomic position of 20.6 to 21.5 Mbp on chromosome 8. It is yet to be discovered whether the QTLs conferring resistance against different geographic isolates of *Stagonosporopsis* spp. are carrying broad-spectrum resistance. Further fine mapping of the QTLs detected on chromosome 8 is required for a better understanding of the GSB resistance mechanism in watermelon.

Candidate genes for GSB resistance in cucumber and watermelon have been predicted in QTL regions [24,28,29]. One hundred and two potential candidate genes were suggested in the 0.5 cM QTL region in cucumber, and 7 genes associated to disease resistance were reported [24]. In another study of GSB resistance in cucumber, one hundred and seventeen candidate genes were predicted in a QTL region between 3.2 cM genomic distance; among them, 14 were associated to disease resistance [25]. A recently reported QTL, *Qgsb8.1* spans a 571.27 kb region and encompass approximately 19 annotated genes; among them two
were related to disease resistance [28]. Another GSB resistance study predicted several candidate genes in three QTL regions on different chromosomes among them CICCG07G013230, encoding an Avr9/Cf-9 rapidly elicited disease resistance protein, which comprises a non-synonymous point mutation in the DUF761 domain was predicted as a strong candidate gene for GSB resistance in watermelon [29].

In the present study, two major co-located QTLs (qLL8.1 and qSB8.1) spanned at 0.87 Mbp region on watermelon chromosome 8. A total 83 genes were predicted in QTLs region, out of which four were related to disease resistance. These candidate genes were included three RLK genes (Cla022133, Cla022184, and Cla022195) and one LRR-RLK (Cla022196) gene. An overview of reported works related to GSB resistance in cucurbits revealed that mostly candidate resistance genes belongs to NBS-LRR and RLK proteins [37]. We also predicted three RLK genes and one NBS-LRR gene as a candidate gene for GSB resistance in watermelon. However, additional fine mapping, expression analysis and cloning work will be required to pinpoint the candidate gene.

One of the main hurdles in the development of GSB-resistant cultivars is time-consuming difficult phenotyping process and discrepancies in phenotyping results [15,28]. Development and implementation of molecular markers can help in early and precise selection to breed GSB-resistant cultivar through marker-assisted selection in watermelon. In recent studies, high throughput KASP™ assays for GSB resistant has been developed for resistant QTLs selection on chromosome 5, 7 and chromosome 8 [28,29]. We developed and validated four high throughput KASP assays for efficient selection of GSB resistance on chromosome 6 and 8. Two flanking markers (chr8_WGRS240 and chr8_WGRS (3)185) on chromosome 8 linked to QTLs (qLL8.1 and qSB8.1) showed promising results for selection with high marker–trait association, whereas two flanking markers (chr6_WGRS (3)089 and chr6_WGRS (3)092) on chromosome 6 linked to QTL (qSB6.1) showed a resistant genotype with watermelon accessions but showed some contrasting genotype with commercial cultivars. However, as the results of QTL–QTL interaction of this study revealed, the presence of both alleles (qSB6.1 and qLL8.1–qSB8.1) significantly enhanced the GSB resistance (Figure 4a). These four markers in combination can be utilized to facilitate the marker-assisted selection to incorporate GSB resistance in elite watermelon cultivars.

4. Materials and Methods

4.1. Plant Materials and DNA Extraction

The GSB-resistant line (C. amarus Schrad.; ‘PI 189225’) and GSB-susceptible line (C. lanatus (Thunb.) Matsum. & Nakai; ‘920533’) were obtained from Fruit Vegetable Breeding Laboratory of National Institute of Horticultural & Herbal Science (NIHHS, Wanju, South Korea) in 2013. The parental lines were subjected to five rounds of self-pollination to obtain homozygosity. The F₁ plants were produced by crossing ‘920533’ as the maternal parent with ‘PI 189225’ (paternal parent) in 2014. In May 2017, the F₁ plants were self-pollinated to produce F₂ progeny. The F₂ seeds were sown in November 2017, and 178 F₂ individuals were used for DNA extraction and disease assessment. Genomic DNA was isolated from susceptible and resistant parents and an F₁ plant, 178 F₂ individuals, 9 watermelon accessions, and 13 commercial cultivars using the cetyltrimethylammonium bromide (CTAB) method [43]. Subsequently, 700 µL of 70% ethanol was added, centrifuged for 1 min. at 12,000 rpm, and then the supernatant was removed and the pellet was washed two times. Finally, the DNA was dissolved in 50 µL triple-distilled water (TDW). The concentration of DNA was quantified by NanoVue (GE Healthcare, Chicago, IL, USA) and DNA diluted to 20 ng·L⁻¹ was used for further analysis.

4.2. Pathogen Inoculation and Disease Assessment

The D. bryoniae ‘KACC 40937 isolate’ was collected from Korean Agricultural Culture Collection (KACC) [8]. Potato dextrose agar (PDA) was used to culture D. bryoniae. The agar plugs with mycelium were sub-cultured in PDA plates and incubated for five days at 25 °C under alternating periods of 12 h of fluorescent and 12 h of darkness for conidia.
formation. Spore suspension was prepared by flooding the culture plates with distilled water and filtering the liquid through fine cloth. Spore concentration was measured with a hemocytometer and adjusted to a four \( \times 10^5 \) spore \( \cdot \text{mL}^{-1} \) by adding sterile water. Seedlings with two to three fully expanded leaves were inoculated with \( D. \text{bryoniae} \) by spraying 10 mL spore suspension per plant. The infected plants were incubated in a humidity chamber at 25–27 °C for 48 h and then transferred to a growth chamber at 25 °C with 12 h light a day [4].

The disease severities on each parental lines, their \( F_1 \) progenies, 178 individuals of \( F_2 \) population, 9 watermelon accessions, and 13 commercial cultivars were assessed at five days after inoculation (DAI). Briefly, disease assessment was performed using two disease severity indexes. First, leaf lesion (LL) was measured for evaluation of disease severity (Figure S5). Five days after inoculation, LL of all true leaves on each plant was measured in multiples of five. LL percentage (%, 0–100) was scored by calculating the ratio of GSB lesion area to total leaf area. The average lesion of leaves on each plant was calculated. Second, symptom for evaluation of disease severity on stem and petiole were measured (Figure S6). Stem blight (SB) was rated on a scale of 0–3, as follows: 0, no symptoms; 1, moderate symptoms; 2, slight necrosis; 3, severe blight symptoms. Third, assessment using modified ordinal disease rating scale was also conducted [15]. To score the individuals as a resistant and susceptible a disease index (DI, 0–7) was defined by considering the disease severity of LL and SB together, with 0, no symptoms; 1, moderate symptoms (\( \leq 20\% \) necrosis) on leaves only; 2, slight symptoms (\( >45\% \) necrosis) on leaves only; 4, moderate symptoms (\( \leq 20\% \) necrosis) on leaves with moderate necrosis also on petiole and stem; 5, slight necrosis also on petiole and stem; 7, severe necrosis also on petiole and stem, death of plant (Table S6). Lastly, plants with a disease index (DI) 0–3 were considered resistant to GSB and plants with a disease index (DI) 4–7 were considered susceptible to GSB (Table S6).

4.3. SNP Genotyping Using Fluidigm® SNP Type™ Assays

Fluidigm® genotyping was performed by the Foundation of Agricultural Tech, Commercialization & Transfer (FACT) using the Fluidigm® EP1™ (Fluidigm, San Francisco, CA, USA) high-throughput (HT) genotyping platform. Previously developed 438 Fluidigm® SNP Type™ assays were used to analyze the genotype of \( F_2 \) population [34]. After performing polymorphic survey, genotyping analysis was conducted to 178 individuals of \( F_2 \) population using Fluidigm® SNP Genotyping Analysis version 4.1.3 software.

4.4. Next-Generation Re-Sequencing and SNP Detection

The homozygous susceptible ‘920533’ and resistant ‘PI 189225’ parental lines were sequenced through next-generation sequencing by the bioinformatics company Macrogen (Seoul, Korea) [35]. The sequencing libraries were prepared according to the manufacturer’s instructions using TruSeq DNA Nano Sample Preparation Kit (Illumina Inc., San Diego, CA, USA). The whole-genome resequencing was performed using a Hiseq™ 4000 sequencer (Illumina Inc., San Diego, CA, USA). After raw reads from re-sequenced data were aligned to the reference genome, reads were mapped to the reference genome [44] using Burrow-Wheeler Aligner (BWA) version 0.7.17. SNPs using SAMtools version 1.6 software were detected [45,46]. Next, 400-bp flanking sequences (amplicon sequences) of each SNP detected between ‘920533’ and ‘PI 189225’ were obtained. Among these, SNPs were filtered to select those with the following characteristics: (i) homozygous SNPs, (ii) SNPs with A/G, A/C, T/G, and T/C combinations, excluding A/T and G/C, (iii) SNPs with one copy number reference genome as a result of BLASTN version 2.7.1+, and (iv) SNPs on the genic region [35,47].

4.5. HRM Primer Design and Genotyping

A total of 888 uniformly distributed HRM primers that amplify sequences for each SNP from NGS data were designed [35,48]. Reactions were conducted in a 20 µL total volume containing 2.0 µL of genomic DNA, 2.0 µL of 10× PCR buffer, 1.0 µL of 2.5 mM
dNTP mixture, 0.1 µL of Taq DNA polymerase (Takara, Japan), 1.0 µL of SYTO® 9 green-fluorescent nucleic acid stain (Life Technologies™, Carlsbad, CA, USA), 1.0 µL of each primer at a concentration of 10 pmole · mL⁻¹ (0.5 µL of each forward and reverse primer) and the rest of total volume was adjusted with TDW. PCR was performed using a PCR machine (Eppendorf, Hamburg, Germany). The PCR cycles were as follows: pre-incubation at 98 °C for 2 min., then 40 cycles: 98 °C for 5 s (denaturation), 60 °C for 10 s (annealing & extension). A HRM analysis was conducted using a CFX96 Touch Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA) to select polymorphic SNP markers. After PCR, HRM melting curves were displayed by increasing the temperature from 70 °C to 94.6 °C at 0.3 °C intervals. After polymorphic HRM markers based on SNPs were selected using Precision Melt Analysis software (Bio-Rad), two parents, an F₁ plant, and 178 plants of the F₂ population were categorized for genotype by three letters (‘A’, ‘H’, ‘B’). Four previously developed HRM markers were also used to analyze genotype in the F₂ population (Table S3) [36].

4.6. Genotyping Using Previously Reported KASP™ Markers

Ten previously reported KASP™ assays (LGC Biosearch™ Technologies, Teddington, UK) were designed for genotyping of the F₂ population [28,49]. KASP™ assays were performed in a 10 µL reaction volume with 5 µL 2 × KASP™ master mix (LGC Biosearch™ Technologies), 0.14 µL assay mix, and 5 µL of 20 ng/µL genomic DNA. The PCR conditions used for the KASP™ assays were as follows: 15 min. at 94°C, followed by 10 cycles of touch down PCR with 20 s 94 °C, 1 min of primer annealing temperature 61 °C with 0.6 °C decrease each cycle, and 26 cycles of 20 s at 94 °C, 1 min at primer annealing temperature, and then 1 min at 37 °C. KASP™ fluorescent end-point readings were measured using a CFX96 Touch Real-Time PCR Detection System (Bio-Rad), and genotype calls were made using Bio-Rad CFX Maestro version 1.1 software (Bio-Rad).

4.7. Construction of Linkage Map and QTL Analysis

A genetic linkage map was constructed using JoinMap® version 4.1 software (Kyazma B.V., Wageningen, The Netherlands), with the Kosambi mapping function [50]. Logarithm of odds (LOD) scores from 2.0 to 10.0 and a maximum genetic distance of 5.0 cM were taken as thresholds for the determination of linkage groups (LGs) and genetic distance (GD; cM). Linkage maps were drawn using MapChart version 2.32 software (Wageningen University & Research, Wageningen, The Netherlands) [51]. QTL analysis was performed using the phenotypic data of GSB resistance and genotypic data through marker analysis by MapQTL® version 6.0 software (Kyazma B.V.) [52]. In order to estimate phenotypic variation of each marker, composite interval mapping (CIM) was conducted using the multiple-QTL model (MQM) mapping function. The genome-wide LOD threshold at the 5% significance level was fixed by 1000 permutation tests. QTL analysis was carried out separately using two type of phenotypic data: leaf lesion (LL) and stem blight (SB). Duncan test (p < 0.05) was performed to compare between ‘A’, ‘H’, and ‘B’ genotypes of F₂ individuals using the R agricolae package [53].

4.8. Validation of Flanking SNP Markers and Identification of Candidate Genes

Nine watermelon accessions (previously reported to have GSB resistance) and 13 commercial cultivars were used for the validation of SNP markers linked to detected QTL regions. The HRM markers within the QTL region were converted into KASP™ markers. KASP™ genotyping assays were designed using Kraken™ Primer Picker software (LGC Biosearch™ Technologies) [49]. The flanking sequences (0.87 Mbp) of QTLs were retrieved from watermelon ‘97103’ genome version 1 in Cucurbit Genomics Database (http://cucurbitgenomics.org/; accessed on 15 June 2020) and candidate genes were predicted in the QTL regions.
5. Conclusions

In summary, we performed genetic analyses for gummy stem blight (GSB) resistance using two different phenotyping methods (leaf lesion and stem bight) in F$_2$ segregating population of watermelon. We used Fluidigm®, HRM, and KASP™ genotyping platforms to analyze SNP markers and constructed a genetic linkage map covering 1070.2 cM with an average marker interval of 5.69 cM. QTL analysis revealed major QTLs, qLL8.1, and qSB8.1, for LL and SB on chromosome 8, and a minor QTL, qSB6.1 on chromosome 6, for SB trait only. We suggested four candidate genes associated to GSB resistance that may be key genes for candidate gene analysis in the future. In addition, we developed and validated four high throughput KASP™ assays to facilitate MAS in watermelon for GSB resistance. The finding of the present studies will help to incorporate GSB resistance in elite watermelon cultivars.

Supplementary Materials: The following are available online at https://www.mdpi.com/2223-7477/10/3/500/s1, Figure S1: Gummy stem blight (GSB) reactions on the leaves of resistant parent ‘PI 189225’, susceptible parent ‘920533’, and F$_1$, Figure S2: SNP genotyping using Fluidigm® SNP Type™ assays for two parents, F$_1$, and 178 individuals of the F$_2$ population, Figure S3: Melting curves of 88 polymorphic HRM markers for two parents, F$_1$, and 178 individuals of the F$_2$ population, Figure S4: SNP genotyping of KASP™ markers for two parents, F$_1$, 178 individuals of the F$_2$ population, Figure S5: Evaluation of disease severity using leaf lesion (LL) area, Figure S6: Symptoms caused by Didymella bryoniae on stem and petiole of test plants, Table S1: Inheritance of resistance to GSB in parental lines, F$_1$ progenies and F$_2$ population, Table S2: Summary of single nucleotide polymorphisms (SNPs) between watermelon lines ‘920533’ and ‘PI 189225’, Table S3: List of 88 polymorphic SNPs between ‘920533’ and ‘PI 189225’ developed into HRM marker, Table S4: Details of SNPs markers used for the genetic map construction in this study, Table S5: List of genes and their annotation in a QTL regions on chromosome 8, Table S6: Disease index (DI) scale for distinguishing resistance and susceptible plants to GSB.

Author Contributions: E.S.L., H.-E.L. and J.L. designed the study. E.S.L. performed the experiments. E.S.L. analyzed the data. Y.-C.H. and H.-E.L. constructed segregation populations derived from two inbred parental lines. C.-G.B. and S.G.K. participated in inoculating for disease assessment. E.S.L. wrote the manuscript. D.-S.K., Y.-R.L., M.I.S., K.H. and J.L. revised the manuscript. All authors have read and agreed to the published version of the manuscript.

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Abbreviations

BFB Bacterial Fruit Blotch
BLASTN Basic Local Alignment Search Tool Nucleotide
BSA Bulked Segregant Analysis
CAPS Cleaved Amplified Polymorphic Sequence
CIM Composite interval mapping
DAI Days After Inoculation
dCAPS derived Cleaved Amplified Polymorphic Sequence
DI Disease Index
GBS Genotyping-by-Sequencing
GO Gene Ontology
GSB Gummy Stem Blight
HRM High Resolution Melting
HT High-throughput
KASP Kompetitive Allele Specific PCR
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