Quantitative trait loci analysis of *Verticillium* wilt resistance in interspecific backcross populations of *Gossypium hirsutum* × *Gossypium barbadense*

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**Abstract**

**Background:** *Verticillium* wilt (VW) caused by *Verticillium dahliae* (Kleb) is one of the most destructive diseases of cotton. The identification of highly resistant QTLs or genes in the whole cotton genome is quite important for developing a VW-resistant variety and for further molecular design breeding.

**Results:** In the present study, BC₁F₁, BC₁S₁, and BC₂F₁ populations derived from an interspecific backcross between the highly resistant line Hai1 (*Gossypium barbadense* L.) and the susceptible variety CCRI36 (*G. hirsutum* L.) as the recurrent parent were constructed. Quantitative trait loci (QTL) related to VW resistance were detected in the whole cotton genome using a high-density simple sequence repeat (SSR) genetic linkage map from the BC₁F₁ population, with 2292 loci covering 5115.16 centiMorgan (cM) of the cotton (AD) genome, and the data concerning VW resistance that were obtained from four dates of BC₂F₁ in the artificial disease nursery and one date of BC₁S₁ and BC₂F₁ in the field. A total of 48 QTLs for VW resistance were identified, and 37 of these QTLs had positive additive effects, which indicated that the *G. barbadense* alleles increased resistance to VW and decreased the disease index (DI) by about 2.2–10.7. These QTLs were located on 19 chromosomes, in which 33 in the A subgenome and 15 QTLs in the D subgenome. The 6 QTLs were found to be stable. The 6 QTLs were consistent with those identified previously, and another 42 were new, unreported QTLs, of which 31 QTLs were from *G. barbadense*. By meta-analysis, 17 QTL hotspot regions were identified and 10 of them were new, unreported hotspot regions. 29 QTLs in this paper were in 12 hotspot regions and were all from *G. barbadense*.

**Conclusions:** These stable or consensus QTL regions warrant further investigation to better understand the genetics and molecular mechanisms underlying VW resistance. This study provides useful information for further comparative analysis and marker-assisted selection in the breeding of disease-resistant cotton. It may also lay an important foundation for gene cloning and further molecular design breeding for the entire cotton genome.

**Keywords:** Cotton, *Verticillium* wilt (VW), Quantitative trait loci (QTL), Interspecific backcross population

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Background
Cotton is an important economic crop worldwide. It provides important natural fibers for the textile industry. Of the two most economically important tetraploid cultivated species, *Gossypium hirsutum* (Upland cotton), which has high yield and wide adaptability, makes up about 95% of the total cotton yield worldwide [1]. *G. barbadense* (sea-island cotton), however, is grown in only limited areas because of its relatively low yield and limited adaptability, despite its strong resistance to *Verticillium* disease and extra-long, fine, strong, fibers. Cultivated forms of both these species show very different traits regarding yield, fiber quality, disease resistance, environmental adaptation, and other traits [2].

*Verticillium* wilt (VW) is one of the important diseases of cotton (*G. hirsutum* L) worldwide. It is caused by *Verticillium dahliae* (Kleb.), a soil-borne fungal pathogen that has a broad range of hosts and moves among them using a variety of mechanisms. It has high pathogenicity and can survive for long periods [3]. Infected plants usually exhibit symptoms of marginal chlorosis or necrosis in leaves, discoloration of the stem vascular bundles, and even full defoliation and plant death. Severe infection results in significant reduction of fiber yield and quality [2, 4]. Worldwide, the disease was first reported in Virginia in 1914 [5]. Today, it is found in almost all cotton growing areas worldwide. This disease broke out in China in 1993, resulting in the infection of approximately 80% of the cotton crop in the seriously infected region of cotton in North China and lint yield losses of 100 million kg in the whole China in that year [6, 7]. Another VW outbreak occurred in major cotton-producing regions in China, including the Yellow River and the Northwest in 2002 and 2003. In 2010, the yield loss caused by VW was 0.31% of the total cotton yield in the U.S. [8], and losses have reached as high as 3–5% in New Mexico [4]. The VW has not been effectively controlled mainly due to its biology characteristics, the indeterminacy of the genetic mechanism of resistance to VW and the lack of highly resistant commercial Upland cotton varieties, except for some modern Acala cotton cultivars developed in California and New Mexico [4].

Planting a resistant cultivar has long been considered the most practical, economic, and effective means of decreasing losses from VW. However, sources of VW resistance in cotton are very limited and no source of heritable immunity has been found in Upland cotton [4, 9]. Many *G. barbadense* genotypes are known to carry high levels of resistance to VW [4, 10, 11], but its resistance has not been successfully transferred into commercial Upland cotton due to hybrid breakdown except for introgressed breeding lines [4, 12, 13]. Using conventional breeding, breeders have made interspecific crosses between Upland cotton and sea-island cotton to improve the VW resistance of Upland cotton for many years, but no breakthrough has been reported because of the negative genetic correlation between lint yield and fiber quality, between lint yield and disease resistance, as well as linkage drag and hybrid breakdown [14]. This makes it challenging for breeders to realize the synchronous improvement of fiber quality, yield, and disease resistance. However, the development of molecular quantitative genetics has made it possible to locate the quantitative trait loci (QTL) for yield, fiber quality, and disease resistance, thus facilitating the use of marker-assisted selection (MAS) for genetic improvement. With the assistance of tightly linked markers to VW resistance, it is possible to transfer the resistance genes from sea-island cotton to Upland cotton.

In recent years, a great deal of progress has been made in mapping VW resistance genes and QTLs in cotton. Many genetic linkage maps have been constructed, and QTLs have been identified for VW resistance in cotton from interspecific populations of *G. hirsutum* × *G. barbadense* [2, 15–21] and from *G. hirsutum* intraspecific populations [22–27]. Some QTLs have also been detected by association mapping using a natural population, chromosome segment introgression lines and molecular mapping based on restriction-site associated DNA (RAD) sequencing technology using a RIL population [17, 28, 29]. These provided some information for further study of the QTLs/genes of VW resistance. However, most of genetic linkage maps used to identify QTLs for VW resistance in all these studies offered no more than 60% coverage of the cotton genome. Although the map reported by Wang et al. covered 3745.9 centi–Morgan (cM), an estimated 74.92% of the tetraploid cotton genome, there were only 430 marker loci on the map [18]. Constructing a high-density linkage map is of great significance for identifying and studying the genes and QTLs of VW resistance.

To introgress the good fiber quality and VW resistance from *G. barbadense* into a commercial Upland cotton variety, we have developed the advanced backcrossing populations with the commercial Upland cotton cultivars (CCRI36) as the recipient parents, and sea-island cotton (Hai1) as the donor parent [30–32], and developed a high-density simple sequence repeat (SSR) genetic linkage map from a BC1F1 population of *G. hirsutum* × *G. barbadense*, which comprised of 2292 loci and covered 5115.16 cM of the cotton AD genome with an average marker interval of 2.23 cM [33]. To the best of our knowledge, no QTL for VW resistance were identified using any map with more than 2000 SSR loci which covered almost the whole cotton AD genome and data for VW resistance from the different the dates, generations, or environments. In this study, a whole-genome screening strategy was used to map QTLs
related to VW resistance using data from different populations related to BC₁F₁, which is quite useful for further fine gene mapping, gene cloning, and marker assisted selection (MAS) in cotton and for further molecular design breeding.

Results

Differences between the parents and phenotypic variation in the populations

In 2005, BC₂F₁ family lines and their parents were evaluated for VW resistance both in the artificial disease nursery and in the field, BC₁S₁ family lines and their parents were also evaluated for VW resistance in the field. The recurrent parent CCRI 36 was susceptible to VW disease, and the average disease index (DI) was 30.6. Hai1 was highly resistant to VW disease with an average DI of 0.4. There was a significant difference in resistance to VW disease between the two parents. The F₁ was also highly resistant to VW disease with an average DI of 4.3. The DI of BC₂F₁ family lines in the artificial disease nursery and in the field and the DI of BC₁S₁ family lines in field exhibited a continuous and normal distribution consistent with multi-gene inheritance for VW resistance (Figs. 1 and 2). Transgressive segregation towards to low DI value was observed (Fig. 1).

Construction of the genetic linkage map

The high-density SSR genetic linkage map was developed from a BC₁F₁ population of G. hirsutum × G. barbadense [33]. Briefly, the map comprised 2292 loci and covered 5115.16 cM of the cotton AD genome with an average marker interval of 2.23 cM. The map was used to detect QTL in the BC₁S₁ population in the field (BC₁S₁-FD), BC₂F₁ population in the field (BC₂F₁-FD), and BC₂F₁ population in the artificial disease nursery (BC₂F₁-NY).

QTL mapping in BC₂F₁ population in the artificial disease nursery

Based on composite interval mapping, a total of 28 QTLs of VW resistance were detected in the BC₂F₁ population during four different dates (June 23, July 19, August 15, and August 25) in the artificial disease nursery, with 5.48–16.66 % of the total phenotypic variation explained. The 28 QTLs were located on 15 chromosomes. Chromosome (C) 5 contained 7 QTLs, and C1 and C15 contained 3 QTLs each, C10, C11, and C26 contained 2 QTLs each, and C6, C9, C13, C14, C17, C19, C20, C21, and C22 contained 1 QTL each. Of these, 19 QTLs had positive additive effects, which indicated that the G. barbadense alleles increased VW resistance and decreased the DI by about 2.2–7.3 (Fig. 3, Table 1).

QTLs of VW resistance for June 23

Six QTLs, qVW-9-1, qVW-10-1, qVW-10-2, qVW-13-1, qVW-14-3 and qVW-19-1, were identified in the BC₂F₁ population in the artificial disease nursery and found on 5 chromosomes, each explaining 5.71–10.16 % of the total phenotypic variation. All QTLs except qVW-9-1 had positive additive effects, which indicated that the Hai1 alleles increased VW resistance and decreased VW DI by 3.2–4.3.

QTLs of VW resistance for July 19

Nine QTLs, qVW-1-2, qVW-1-3, qVW-5-1, qVW-5-2, qVW-5-3, qVW-5-4, qVW-15-4, qVW-17-1 and qVW-21-1, were detected in the BC₂F₁ population in the
artificial disease nursery, each explaining 5.48–12.01 % of the total phenotypic variation. These were located on 5 chromosomes, four QTLs were found on Chr5 alone. All QTLs except qVW-15-4 had positive additive effects, which indicated that the Hai1 alleles increased VW resistance and decreased VW DI by 2.2–3.4.

**QTLs of VW resistance for August 15**
Eleven QTLs, qVW-1-1, qVW-5-2, qVW-5-3, qVW-5-4, qVW-5-5, qVW-6-1, qVW-11-1, qVW-11-2, qVW-15-3, qVW-26-1 and qVW-26-2, were identified in the BC2F1 population in the artificial disease nursery, each explaining 6.92–16.66 % of the total phenotypic variation. The QTLs were located on 6 chromosomes, four on Chr5. All QTLs except qVW-6-1, qVW-15-3, qVW-26-1 and qVW-26-2 had positive additive effects, which indicated that the Hai1 alleles increased VW resistance and decreased VW DI by 5.0–7.3. In addition, qVW-C26-1 and qVW-C26-2 were both located on C26 and explained 11.12 and 7.95 % of the phenotypic variation, respectively. The CCRI36 alleles increased VW resistance and decreased VW DI by 5.0 and 5.9, respectively.

**QTLs of VW resistance for August 25**
Five QTLs, including qVW-5-6, qVW-5-7, qVW-15-1, qVW-20-1 and qVW-22-1, were detected in the BC2F1 population in the artificial disease nursery and located on 4 chromosomes, each explaining 6.93–9.50 % of the total phenotypic variation. Two QTLs (qVW-5-6 and qVW-5-7) had positive additive effects, and they were located on the same chromosome, C5. They explained 9.22 and 7.09 % of the phenotypic variation, respectively. The Hai1 alleles increased VW resistance and decreased VW DI by 5.8 and 5.1, respectively.

qVW-C5-2, qVW-C5-3 and qVW-C5-4 on C5 were detected in the two dates on July 19 and August 15, and were localized in the marker intervals PGML03048-DPL0063, DPL0063-DPL0724, and CGR6708a-MGHE506, respectively. qVW-5-2 explained 9.76–11.74 % of the phenotypic variation, and the G. barbadense allele decreased the DI by about 3.0–6.0. qVW-C5-3 explained 9.8–13.52 % of the phenotypic variation, and the G. barbadense allele decreased the DI by about 3.0–6.5. qVW-C5-4 explained 12.01–16.66 % of the phenotypic variation, and the G. barbadense allele decreased the DI by about 3.2–7.3.

**QTL mapping in BC2F1 and BC1S1 populations in the field**
Based on composite interval mapping, a total of 13 QTLs associated with VW resistance, qVW-5-2, qVW-5-3, qVW-5-4, qVW-5-7, qVW-7-1, qVW-7-2, qVW-7-3, qVW-7-4, qVW-12-1, qVW-12-2, qVW-12-3, qVW-12-5, qVW-15-3 and qVW-26-3, were detected in the BC1S1 population on September 17, explaining 5.87–13.59 % of the total phenotypic variation. These 13 QTLs were located on C5, C7, C12, C15 and C26. C7 contained 4 QTLs, C5 and C12 contained 3 QTLs each, C15 contained 2 QTLs and C26 contained 1 QTL. All QTLs except qVW-15-2, qVW-15-3, and qVW-26-3 had positive additive effects, which indicated that the G. barbadense allele decreased the DI by about 7.2–10.7 (Fig. 3, Table 1).

A total of 16 QTLs for VW resistance, qVW-3-1, qVW-3-2, qVW-3-3, qVW-3-4, qVW-3-5, qVW-3-6, qVW-5-1, qVW-5-2, qVW-5-3, qVW-5-4, qVW-5-5, qVW-8-1, qVW-8-2, qVW-9-2, qVW-14-1 and qVW-14-2, were detected in the BC2F1 population on September 17, explaining 5.76–13.46 % of the total phenotypic variation. The 16 QTLs were located on C3, C5, C8, C9 and C14. C3 contained 6 QTLs, C5 contained 5 QTLs, C8 and C14 each contained 2 QTLs and C9 contained 1.
Fig. 3 Chromosomal locations of QTLs for Verticillium wilt (VW) resistance in the BC$_2$F$_1$-FD, BC$_1$S$_1$-FD, and BC$_2$F$_1$-NY populations derived from an interspecific cross of *G. hirsutum* (CCRI36) × *G. barbadense* (Hai1)
| QTL      | Gen.     | Date | Env. | C Position (cM) | Nearest Marker                     | Marker interval | LOD  | Additive effect | PV% |
|----------|----------|------|------|-----------------|------------------------------------|-----------------|------|----------------|-----|
| qVW-C1-1 | BC2F1    | 8.15 | DN C1 | 42.7            | DPL0692                            | DPL0692-NAU3744b | 3.5  | 5              | 7.88 |
| qVW-C1-2 | BC2F1    | 7.19 | DN C1 | 89.5            | CGR6870                            | BNL2921-CGR6870 | 4.2  | 3.4            | 10.43|
| qVW-C1-3 | BC2F1    | 7.19 | DN C1 | 141.6           | PGML0280                           | NAU5163-CGR6803a | 2.6  | 2.8            | 7.11 |
| qVW-C3-1 | BC2F1    | 9.17 | FD C3 | 94.2            | MUS50087                           | MUS50087-PGML0376 | 3.3  | 5.7            | 7.32 |
| qVW-C3-2 | BC2F1    | 9.17 | FD C3 | 98              | NAU1286-DPL0232                    | DPL0232-PGML03195 | 3.3  | 6.2            | 8.76 |
| qVW-C3-3 | BC2F1    | 9.17 | FD C3 | 101.3           | DPL0232                            | DPL0232-PGML03195 | 3.3  | 6.2            | 8.76 |
| qVW-C3-4 | BC2F1    | 9.17 | FD C3 | 105.3           | HAU1022                            | TMB1898-DPL0224 | 5.6  | 7.2            | 11.33|
| qVW-C3-5 | BC2F1    | 9.17 | FD C3 | 109.6           | NAU2297-DPL0321                    | DPL0232-PGML03195 | 3.3  | 6.2            | 8.76 |
| qVW-C3-6 | BC2F1    | 9.17 | FD C3 | 113.8           | NAU2297                           | DPL0232-PGML03195 | 3.3  | 6.2            | 8.76 |
| qVW-C5-1 | BC2F1    | 7.19 | DN C5 | 30.5            | CIR224b                            | DPL0274a-PGML03048 | 3.3  | 2.7            | 8.1  |
| qVW-C5-2 | BC2F1    | 7.19 | DN C5 | 30.5            | CIR224b                            | DPL0274a-PGML03048 | 3.3  | 2.7            | 8.1  |
| qVW-C5-3 | BC2F1    | 9.17 | FD C5 | 38.2            | HAU0746                            | DPL0063-DPL0724 | 5.5  | 6.5            | 13.52|
| qVW-C5-4 | BC2F1    | 9.17 | FD C5 | 38.2            | HAU0746                            | HAU0746          | 6    | 10.6           | 13.13|
| qVW-C5-5 | BC2F1    | 9.17 | FD C5 | 38.2            | HAU0746                            | DPL0063-DPL0724 | 5.5  | 6.5            | 13.52|
| qVW-C5-6 | BC2F1    | 8.15 | DN C5 | 45              | DPL0138                            | MGHE506-DPL03048 | 4.6  | 6.1            | 11.63|
| qVW-C5-7 | BC2F1    | 8.15 | DN C5 | 45              | DPL0138                            | MGHE506-DPL03048 | 4.6  | 6.1            | 11.63|
| qVW-C5-8 | BC2F1    | 8.15 | DN C5 | 45              | DPL0138                            | MGHE506-DPL03048 | 4.6  | 6.1            | 11.63|
| qVW-C5-9 | BC2F1    | 8.15 | DN C5 | 45              | DPL0138                            | MGHE506-DPL03048 | 4.6  | 6.1            | 11.63|
| qVW-C6-1 | BC2F1    | 8.25 | DN C6 | 5.0             | NAU5038                            | NAU5038-CGR5128a | 2.9  | -4.9           | 7.13 |
| qVW-C6-2 | BC2F1    | 8.25 | DN C6 | 5.0             | NAU5038                            | NAU5038-CGR5128a | 2.9  | -4.9           | 7.13 |
| qVW-C6-3 | BC2F1    | 8.25 | DN C6 | 5.0             | NAU5038                            | NAU5038-CGR5128a | 2.9  | -4.9           | 7.13 |
| qVW-C6-4 | BC2F1    | 8.25 | DN C6 | 5.0             | NAU5038                            | NAU5038-CGR5128a | 2.9  | -4.9           | 7.13 |
| qVW-C6-5 | BC2F1    | 8.25 | DN C6 | 5.0             | NAU5038                            | NAU5038-CGR5128a | 2.9  | -4.9           | 7.13 |
| qVW-C7-1 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-2 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-3 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-4 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-5 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-6 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-7 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-8 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
| qVW-C7-9 | BC2F1    | 8.15 | DN C7 | 88.1            | NG6381                             | NG6381-NAU1763 | 3.4  | 7.8            | 7.08 |
All 16 QTLs had positive additive effects, which indicated that the \textit{G. barbadense} allele decreased the DI by about 5.1–7.9 (Fig. 3, Table 1).

Stability of QTLs over multiple generations, environments and dates

A total of 48 QTLs of VW resistance were detected in the BC\textsubscript{2}F\textsubscript{1} and BC\textsubscript{1}S\textsubscript{1} populations and the five dates. These were located on 19 chromosomes. C5 contained 7 QTLs, C3 contained 6 QTLs, C7 and C15 each contained 4 QTLs, C1, C12, C14 and C26 each contained 3 QTLs, C8, C9, C10 and C11 each contained 2 QTLs, and C6, C13, C17, C19, C20, C21 and C22 each contained 1 QTL. Of these, 37 QTLs (77.08 \%) had positive additive effects, which indicated that the \textit{G. barbadense} allele decreased the DI (Fig. 3, Table 1).

Of these 48 QTLs, 3 QTLs (qVW-C5-2, qVW-C5-3, qVW-C5-4) were detected in two generations (BC\textsubscript{2}F\textsubscript{1}, BC\textsubscript{1}S\textsubscript{1}), in two environments (the field, the artificial disease nursery) and in the three dates (July 19, August 15 and September 17), explaining 9.76–16.66 \% of the phenotypic variation. The Hai1 allele decreased the DI by approximately 3.0–10.7, increasing VW disease resistance. One QTL (qVW-C5-1) was simultaneously detected in both environments in the artificial disease nursery and in the field for BC\textsubscript{2}F\textsubscript{1}, explaining 7.72–11.63 \% of the phenotypic variation. The Hai1 allele decreased the DI by about 5.1–8.7. One QTL (qVW-C15-1) was simultaneously detected in two environments or dates on August 15 in the artificial disease nursery and on September 17 in the field for BC\textsubscript{2}F\textsubscript{1}, explaining 7.22–16.3 % of the phenotypic variation. The Hai1 allele decreased the DI by approximately 5.8–6.1 (Figs. 3 and 4, Table 1). One QTL (qVW-C15-3) was simultaneously detected on September 17 in BC\textsubscript{1}S\textsubscript{1} in the field and in BC\textsubscript{2}F\textsubscript{1} on August 15 in the artificial disease nursery, explaining 6.05–6.92 \% of the phenotypic variation. And the CCRI36 allele decreased the DI by about 4.8–7.1. A total of six QTLs were stably detected in different generations, environments, and/or different dates. These stable QTLs could be used for MAS breeding.

Meta-analysis of QTLs

In the Meta-analysis, a total of 17 QTL hotspot regions were identified on 13 chromosomes of the consensus map, including C3, C5, C7, C8, C9, C12, C14, C15, C17, C19, C20, C21 and C26 (Fig. 5, Table 2). 7 QTL hotspot regions (c5-VW-Hotspot-2, c7-VW-Hotspot-1, c7-VW-Hotspot-2, c8-VW-Hotspot-1, c19-VW-Hotspot-1, c21-VW-Hotspot-1 and c26-VW-Hotspot-1) were consistent with those identified previously by Zhang et al. and Said et al. [13, 34] (Table 2), and the other ten were new, unreported hotspot regions. Two QTL hotspot regions were on C5, C7, C9 and C19, respectively. Of 17 QTL hotspot regions, c3-VW-Hotspot-1, c5-VW-Hotspot-1, c8-VW-Hotspot-1 and c26-VW-Hotspot-1 had more QTLs. 29 QTLs from this paper were distributed in 15 hotspot regions, 29 of them were in 12 hotspot regions and were all from \textit{G. barbadense} (Table 3). It showed that QTL-rich regions on the map of CCRI36 × Hai1 BC\textsubscript{2}F\textsubscript{1} were consistent with the QTL hotspot regions (Table 3, Additional file 1). These consensus QTL regions (QTL-Table 1 QTLs of \textit{Verticillium} wilt (VW) resistance detected during the five dates across two generations (Continued)

| QTL   | Generation | Date | Chromosome | Genomic Location | Additive Effect |
|-------|------------|------|------------|------------------|----------------|
| qVW-C14-1 | BC\textsubscript{2}F\textsubscript{1} | 9.17 | C14 | 8.16 | C2-0079 NAU3499-CGR6550 | 4.4 | 6.4 | 8.7 |
| qVW-C14-2 | BC\textsubscript{2}F\textsubscript{1} | 9.17 | C14 | 88.9 | PGM02953b CGR6550-CICR0377 | 2.9 | 5.5 | 6.43 |
| qVW-C14-3 | BC\textsubscript{2}F\textsubscript{1} | 6.23 | C14 | 184.6 | NAU5465 SHIN-0659b-CICR0052 | 2.6 | 3.2 | 5.71 |
| qVW-C15-1 | BC\textsubscript{2}F\textsubscript{1} | 8.25 | C15 | 43.9 | CGR5834 NAU3177-HAU3050a | 3.8 | 5.8 | 9.5 |
| qVW-C15-2 | BC\textsubscript{2}F\textsubscript{1} | 9.17 | C15 | 141.9 | NAU0861 NAU3680-DPL0437 | 2.6 | 7 | 5.87 |
| qVW-C15-3 | BC\textsubscript{2}F\textsubscript{1} | 9.17 | C15 | 152 | TMB1664 COT059-PGM02824a | 2.9 | 7.1 | 6.05 |
| qVW-C15-4 | BC\textsubscript{2}F\textsubscript{1} | 7.19 | C15 | 161.2 | CGR5056a PGM02824a-CER0013 | 3.4 | 2.9 | 9.02 |
| qVW-C17-1 | BC\textsubscript{2}F\textsubscript{1} | 7.19 | C17 | 5.3 | HAU0800a CER0560-DC4029 | 2.6 | 2.2 | 5.48 |
| qVW-C19-1 | BC\textsubscript{2}F\textsubscript{1} | 6.23 | C19 | 185.6 | HAU1385b DPL0192-NAU5489 | 4.5 | 4.3 | 10.16 |
| qVW-C20-1 | BC\textsubscript{2}F\textsubscript{1} | 8.25 | C20 | 22.2 | CICR0254 | 3.6 | 5.6 | 9.01 |
| qVW-C21-1 | BC\textsubscript{2}F\textsubscript{1} | 7.19 | C21 | 111.3 | Gh074a CICR0046-DC40250 | 4.2 | 3.1 | 9.99 |
| qVW-C22-1 | BC\textsubscript{2}F\textsubscript{1} | 8.25 | C22 | 149.7 | CER0139b TMB0206-CICR0438 | 2.8 | 5 | 6.93 |
| qVW-C26-1 | BC\textsubscript{2}F\textsubscript{1} | 8.15 | C26 | 164.5 | DPL0915 | 3.9 | 5.9 | 11.12 |
| qVW-C26-2 | BC\textsubscript{2}F\textsubscript{1} | 8.15 | C26 | 169.4 | NAU3905 NAU4914-PGM04182 | 3.5 | 5 | 7.95 |
| qVW-C26-3 | BC\textsubscript{2}F\textsubscript{1} | 7.19 | C26 | 172.4 | CGR6759 DPL0890-PGM02118 | 3 | 7.4 | 6.48 |

Note: 8.15 means that the DI was investigated at August 15; DN the artificial disease nursery, FD the field, Gen. generation, Env. environment, C chromosome, PV\% phenotypic variation explained; positive additive effect indicates that the locus derived from Hai1 decreased the value of DI.
rich regions and QTL hotspot regions) could be worthy of further research and used for MAS.

Discussion
Map used to identify QTLs for Verticillium wilt (VW) resistance
To identify QTL associated with VW resistance from *G. hirsutum* and *G. barbadense* or *G. hirsutum* and an introgressed line. For example, Zhang et al. detected QTLs for VW resistance based on a 392 SSR polymorphic loci linkage map covering 2895 cM or about 57.90 % of the cotton genome using the BIL population of SG 747 × Giza 75 [13]. Fang et al. constructed a 292-marker genetic linkage map for the BIL population of SG 747 × Pima S-7 [15]. It covered 1226 cM or about
27% of the cotton genome. Yang et al. constructed two genetic linkage maps [19]. One of these included 35 linkage groups with an average distance between two markers of 8.7 cM with 219 SSR loci. They covered 1772.5 cM or approximately 31.93% of the tetraploid cotton genome from BC1 of Hai 7124 × Junmian 1. Another genetic linkage map with 420 genome-wide loci using F2 of Hai 7124 × Junmian 1, with an average distance of 7.3 cM per marker covered 2726.9 cM or approximately 49.10% of the tetraploid cotton genome.

Wang et al. constructed a genetic linkage map with 430 marker loci using F2 population of XinLuZao1 × Hai7124, with an average distance of 8.71 cM per marker and covered 3745.9 cM, which is an estimated 74.92% of the total recombination length of the tetraploid cotton genome [18]. Bolek et al. constructed eleven linkage groups using an F2 population derived from the interspecific cross of Pima S-7 and Acala 44, consisting of 35 markers and spanning 531 cM (approximately 10.62% of the tetraploid cotton genome) with an average distance of 15.17 cM [2]. Gao et al. constructed a genetic linkage map with 99 marker loci using an F2 population of Handan208 × Pima90 [20]. It had an average distance of 18.61 cM per marker and covered 1842.8 cM or approximately 36.8% of the tetraploid cotton genome. Fang et al. constructed a genetic linkage map with 882 marker loci using the RIL population of TM-1 × NM24016 [16]. It had an average distance of 7.06 cM per marker and covered 2267 cM, which is an estimated 55.7% of the total recombination length of the tetraploid cotton genome. NM24016 is an inbred Upland cotton line with considerable but stable introgression from several G. barbadense lines. Ning et al. constructed a genetic linkage map with 279 marker loci using a RIL population of Prema × 86–1 in which Prema was an introgressed line from G. thurberi and G. barbadense [23]. This map covered 1576.25 cM, which is an estimated 35.42% of the total recombination length of the tetraploid cotton genome. Using the same population, Wang et al. constructed a molecular map based on restriction-site associated DNA (RAD) sequencing technology, covering 3499.69 cM, which is an estimated 69.99% of the total recombination length of the tetraploid cotton genome [29].

In addition, some genetic maps have also been developed using intraspecific populations of G. hirsutum to identify QTL for VW resistance. Zhang et al. constructed a genetic linkage map with 141 marker loci using an F2 population of LHB22 × JM11 [22]. It had an average distance of 8.11 cM per marker and covered 1143.1 cM, which is an estimated 22.86% of the tetraploid cotton genome. Jiang et al. constructed a genetic linkage map using F2 segregating population of 60182 × Junmian 1, with 139 loci, covering 1165 cM [24]. It had an average distance of 8.38 cM between two markers, or 25.89% of the length of the cotton genome. Ge et al. constructed a genetic linkage map with 122 marker loci using an F2 population of Chang96 × Junmian 1 [25]. It had 33 linkages and covered 1172.7 cM or approximately 23.5% of the tetraploid cotton genome. Wang et al. constructed a genetic linkage map with 10 marker loci using F2 population of Lumianyan22 × Luyuan343

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Table 2 QTL hotspots identified for *Verticillium* wilt (VW) resistance on the consensus map by meta-analysis

| Hotspot name     | C   | Position (cM) | No. of QTLs | No. of QTLs in this paper | Reported previously                                      |
|------------------|-----|---------------|-------------|--------------------------|----------------------------------------------------------|
| c3-VW-Hotspot-1  | C3  | 16–28 cM      | 7           | 6                        | Zhang et al. 2015 [13] and Said et al. 2015 [34]         |
| c5-VW-Hotspot-1  | C5  | 28–43 cM      | 7           | 5                        | Zhang et al. 2015 [13]                                   |
| c5-VW-Hotspot-2  | C5  | 47–68 cM      | 5           | 2                        | Zhang et al. 2015 [13]                                   |
| c7-VW-Hotspot-1  | C7  | 43–59 cM      | 4           | 1                        | Zhang et al. 2015 [13]                                   |
| c7-VW-Hotspot-2  | C7  | 64–85 cM      | 5           | 3                        | Zhang et al. 2015 [13]                                   |
| c8-VW-Hotspot-1  | C8  | 17–34 cM      | 8           | 2                        | Zhang et al. 2015 [13]                                   |
| c9-VW-Hotspot-1  | C9  | 55–85 cM      | 6           | 1                        | Zhang et al. 2015 [13]                                   |
| c9-VW-Hotspot-2  | C9  | 124–147 cM    | 5           | 0                        | Zhang et al. 2015 [13]                                   |
| c12-VW-Hotspot-1 | C12 | 17–26 cM      | 4           | 3                        | Zhang et al. 2015 [13]                                   |
| c14-VW-Hotspot-1 | C14 | 26–43 cM      | 5           | 3                        | Zhang et al. 2015 [13]                                   |
| c15-VW-Hotspot-1 | C15 | 42–63 cM      | 6           | 3                        | Zhang et al. 2015 [13]                                   |
| c17-VW-Hotspot-1 | C17 | 11–36 cM      | 4           | 1                        | Zhang et al. 2015 [13]                                   |
| c19-VW-Hotspot-1 | C19 | 1–26 cM       | 5           | 0                        | Zhang et al. 2015 [13]                                   |
| c19-VW-Hotspot-2 | C19 | 120–132 cM    | 4           | 1                        | Zhang et al. 2015 [13]                                   |
| c20-VW-Hotspot-1 | C20 | 1–26 cM       | 4           | 1                        | Zhang et al. 2015 [13]                                   |
| c21-VW-Hotspot-1 | C21 | 30–44 cM      | 6           | 1                        | Zhang et al. 2015 [13]                                   |
| c26-VW-Hotspot-1 | C26 | 54–79 cM      | 9           | 3                        | Zhang et al. 2015 [13]                                   |
It had three linkages and covered 108.9 cM. Yang et al. constructed a genetic linkage map with 115 marker loci using an RIL population of 5026 × Li 8 [27]. It had 20 linkage groups and covered 560.1 cM or approximately 11.2 % of the tetraploid cotton genome.

Only 1–18 QTLs for VW resistance were detected using these maps in previous reports. In the previous reports, most of these linkage maps used to identify QTLs associated with VW resistance had no more than 2000 SSR markers and covered less than 60 % of the cotton genome [2, 15, 16, 19, 22–24]. It is difficult to identify more QTLs for VW resistance from G. barbadense using the limited number of markers and low coverage of the whole cotton genome.

In this study, the genetic map comprised 2292 SSR marker loci and covered 5115.16 cM of the cotton genome [33]. A total of 48 QTLs for VW resistance were detected and distributed on 19 chromosomes. This showed that more QTLs for VW were found and located in more chromosomes and were mapped to a narrower interval with tightly linked markers using our map than maps of the traits reported in previous papers.

Distribution of QTLs through the whole genome

QTL clusters of different fiber quality and disease resistance have also been observed in previous studies [1, 35–41]. The current results showed some QTL-rich regions on C3, C5, C7, C12, C15 and C26 for VW resistance. Six QTLs were detected within a C3 region (93–114 cM) between MUSB0087 and DPL0321. A C5 region (27–47 cM) between DPL0274a and DPL0241 harbored 5 QTLs and a C7 region (75–100 cM) between DPL0136 and DC40253 carried 4 QTLs. Three QTLs each were found within a specific range on C12 (93–112 cM) between DPL01491 and NAU3713, within a specific range on C15 (141–165 cM) between NAU3680 and CER0013, and within a specific

Table 3 QTLs identified using interspecific backcross populations on the map of CCRI36 × Hai1 BC1F1 and the corresponding hotspots

| QTL-rich regions     | QTL   | Direction | Hotspot name and range |
|----------------------|-------|-----------|------------------------|
| On the map of CCRI36 x Hai1 BC1F1 |       |           |                        |
| C3-VW-QTL-rich-1: 93–114 cM | qVW-C3-1 | Hai1       | c3-VW-Hotspot-1: 16–28 cM |
|                      | qVW-C3-2 | Hai1       |                        |
|                      | qVW-C3-3 | Hai1       |                        |
|                      | qVW-C3-4 | Hai1       |                        |
|                      | qVW-C3-5 | Hai1       |                        |
|                      | qVW-C3-6 | Hai1       |                        |
| C5-VW-QTL-rich-1: 27–47 cM | qVW-C5-1 | Hai1       | c5-VW-Hotspot-1: 28–43 cM |
|                      | qVW-C5-2 | Hai1       |                        |
|                      | qVW-C5-3 | Hai1       |                        |
|                      | qVW-C5-4 | Hai1       |                        |
|                      | qVW-C5-5 | Hai1       |                        |
|                      | qVW-C5-6 | Hai1       | c5-VW-Hotspot-2: 47–68 cM |
|                      | qVW-C5-7 | Hai1       |                        |
|                      | qVW-C6-1 | CCRI36     |                        |
| C7-VW-QTL-rich-1: 75–100 cM | qVW-C7-1 | Hai1       | c7-VW-Hotspot-1: 43–59 cM |
|                      | qVW-C7-2 | Hai1       | c7-VW-Hotspot-2: 64–85 cM |
|                      | qVW-C7-3 | Hai1       |                        |
|                      | qVW-C7-4 | Hai1       |                        |
|                      | qVW-C8-1 | Hai1       | c8-VW-Hotspot-1: 17–34 cM |
|                      | qVW-C8-2 | Hai1       |                        |
|                      | qVW-C9-1 | CCRI36     | c9-VW-Hotspot-1: 55–85 cM |
|                      | qVW-C9-2 | Hai1       |                        |
|                      | qVW-C10-1| Hai1       |                        |
|                      | qVW-C10-2| Hai1       |                        |
|                      | qVW-C11-1| Hai1       |                        |
|                      | qVW-C11-2| Hai1       |                        |
| C12-VW-QTL-rich-1: 93–112 cM | qVW-C12-1| Hai1       | c12-VW-Hotspot-1: 17–26 cM |
|                      | qVW-C12-2| Hai1       |                        |
|                      | qVW-C12-3| Hai1       |                        |
|                      | qVW-C13-1| Hai1       |                        |
|                      | qVW-C14-1| Hai1       | c14-VW-Hotspot-1: 26–43 cM |
|                      | qVW-C14-2| Hai1       |                        |
|                      | qVW-C14-3| Hai1       |                        |
|                      | qVW-C15-1| CCRI36     |                        |
| C15-VW-QTL-rich-1: 141–165 cM | qVW-C15-2| CCRI36     | c15-VW-Hotspot-1: 42–63 cM |
|                      | qVW-C15-3| CCRI36     |                        |

Table 3 QTLs identified using interspecific backcross populations on the map of CCRI36 × Hai1 BC1F1 and the corresponding hotspots (Continued)

| QTL     | Direction | Hotspot name and range |
|---------|-----------|------------------------|
| qVW-C17-1 | Hai1       | c17-VW-Hotspot-1: 11–36 cM |
| qVW-C19-1 | Hai1       | c19-VW-Hotspot-2: 120–132 cM |
| qVW-C20-1 | CCRI36     | c20-VW-Hotspot-1: 1–26 cM |
| qVW-C21-1 | Hai1       | c21-VW-Hotspot-1: 30–44 cM |
| qVW-C22-1 | CCRI36     |                        |
| qVW-C26-1 | CCRI36     | c26-VW-Hotspot-1: 54–79 cM |
| qVW-C26-2 | CCRI36     |                        |
| qVW-C26-3 | CCRI36     |                        |
range on C26 (158–177 cM) between PGML04562 and PGML02118 (Table 1, 3, Fig. 3). Importantly, the 5 QTLs for VW resistance that clustered on C5 were stably detected in different generations, different environments and/or different dates.

In this paper, a total of 48 QTLs for VW resistance were detected in BC1F1 and BC2F1 populations using the high-density linkage map and they were distributed on 19 chromosomes. Here, 33 (68.75 %) of them were located in the A subgenome, involving 11 chromosomes (C1, C3, C5, C6, C7, C8, C9, C10, C11, C12 and C13), and 15 QTLs were located on the D subgenome, involving 8 chromosomes (C14, C15, C17, C19, C20, C21, C22 and C26). This indicated that the A subgenome harbored more QTLs or genes for VW resistance than the D subgenome. This was consistent with results reported by Yang et al. [19], Ning et al. [23], and Bolek et al. [2]. However, they are not consistent with the idea that the D subgenome makes a greater contribution to genetic control of resistance traits than the A subgenome does [42, 43]. This could be relevant to the conclusion that more QTLs cluster on the A subgenome in this study. It may be because different materials were used in different studies.

**QTLs for Verticillium wilt (VW) resistance from G. barbadense**

In this paper, a total of 48 QTLs for VW resistance were detected in two generations (BC1F1, BC2F1) using a high-density linkage map. Of these, 37 (77.08 %) QTLs had positive additive effects, which indicated that the G. barbadense allele increased VW resistance and decreased the DI by about 2.2–10.7. Six QTLs were stably detected in two different generations and different environments and dates. Of these, 5 QTLs clustered on C5 and 1 was located on C15.

The six QTLs detected were found to be on the same chromosomes or subgenomes reported in previous studies, and they had shared SSR markers [17, 19, 21–23]. qVW-C7-1 located on C7 for VW resistance was the same as Yang's qVL-A7-1 F2 [19], based on shared markers of NAU1048. qVW-C19-1 may be the same as qVL-D5-1BC1S2592 in Yang et al. [19]. These were linked to the common marker BNL1878. qVW-C5-2 was same as QTL (qVW-A5-1) reported by Ning et al. with the linked common marker DC20067 [23]. The qVW-C21-1 here detected for VW resistance may be the same as the qVW-c21-1 reported in Zhang et al. with the linked common markers HAU0423 and CGRS5602 [22]. qVW-C9-2 was assigned to the same chromosome region as the qFD1711-11-4.01 reported by Wu et al. with the linked common marker BNL3031 [21]. qVW-C3-5 may be the same as qVW991-A3-1 in Wang et al. [17]. These were linked to the common marker NAU3479. We could not correlate the other QTLs for VW resistance to those found in other studies due to the lack of common markers, although some QTLs were on the same chromosomes [15, 18, 20, 22, 23, 26, 27, 29, 36]. The more frequently informative molecular markers are used in the cotton community, the greater the likelihood that QTL conditioning VW resistance among various cotton germplasm lines could be tagged. In this way, the other 42 QTLs for VW resistance could be considered newly identified by this map.

In summary, 6 stable QTLs (qVW-C5-1, qVW-C5-2, qVW-C5-3, qVW-C5-4, qVW-C5-5, qVW-C15-3) were detected in two or more populations or dates in this map and 6 consensus QTLs (qVW-C7-1, qVW-C19-1, qVW-C5-2, qVW-C21-1, qVW-C9-2, qVW-C3-5) were also reported in the previous studies. These 11 stable or consensus QTLs (qVW-C5-1, qVW-C5-2, qVW-C5-3, qVW-C5-4, qVW-C5-5, qVW-C15-3, qVW-C7-1, qVW-C19-1, qVW-C21-1, qVW-C9-2, qVW-C3-5) could be used for MAS. Among 48 QTLs, there were 42 new unreported QTLs, of which 31 QTLs were from G. barbadense.

By meta-analysis of QTLs, 17 QTL hotspot regions were identified on 13 chromosomes of the consensus map (Fig. 5, Table 2). Of them, 7 QTL hotspot regions were consistent with those identified previously by Zhang et al. and Said et al. [13, 34] (Table 2) and another 10 were new, unreported hotspot regions. 36 (75 %) QTLs from this paper were distributed in 15 hotspot regions, of which 29 QTLs were in 12 hotspot regions and were all from G. barbadense (Table 3). All QTL-rich regions on the map of CCR136 × Hai1 BC1F1 were consistent with the QTL hotspot regions of the consensus map (Table 3, Additional file 1). These consensus QTL regions (QTL-rich regions or QTL hotspot regions) could be worthy of further research and used for MAS.

**Further application of QTLs for Verticillium wilt (VW) resistance**

Chromosome segment substitution lines (CSSLs) with the two commercial Upland cotton cultivars (CCRI36 and CCRI45) were developed as the recipient parents to introgress genes and QTLs associated with good fiber quality and VW resistance from G. barbadense into a commercial Upland cotton variety, G. barbadense (Hai1) served as a donor parent. The fiber yield and quality traits of these CSSLs have been evaluated in previous works [30–32, 44–46]. VW resistance in these CSSLs is being evaluated. The minor difference between CSSLs and the recurrent parent lies in the donor chromosome segment from G. barbadense in G. hirsutum genetic background, reducing the interference of the genetic background. In this way, CSSLs are an ideal material for genetic studies on quantitative traits. The present study provides molecular information for fine gene mapping, gene cloning, gene pyramiding, and marker-assisted breeding for improving VW resistance and lays a solid
foundation for further molecular design breeding throughout the cotton genome.

Conclusions
In the present study, a high-density SSR genetic linkage map covering the whole cotton genome and data associated with VW resistance from different dates in populations of different generations in the field and in the artificial disease nursery were used to identify QTLs associated with VW resistance and never-before-reported QTLs were detected. Here, 48 VW resistance QTLs were mapped based on a 5115.16 cM linkage map with 2292 SSR marker loci. This work provides useful information for further comparative analysis and improving understanding of the genetic basis of VW resistance in cotton. Here, six QTLs were found to be stable, of which 5 QTLs clustered on C5 and 1 on C15. In this paper, six QTLs were the same or similar to previously reported QTLs and 42 QTLs were new. Of these 42, 31 QTLs were from G. barbadense. 17 QTL hotspot regions for VW resistance were identified and 10 of them were new, unreported hotspot regions. 29 QTLs in 12 hotspot regions were all from G. barbadense. These stable and consensus QTLs provide important molecular information for MAS in Upland cotton breeding aimed at improving the resistance to VW and may lay a better foundation for gene cloning and further study.

Methods
Plant materials
The mapping population with 135 BC1F1 plants was derived from an interspecific cross of G. hirsutum (CCRI36) × G. barbadense (Hai1), in which CCRI36 is a susceptible cultivar (G. hirsutum L.) and Hai1 is a highly resistant line (G. barbadense L.) with super fiber quality and the dominant glandless gene [33, 47].

The CCRI36× Hai1 F1 was made in Henan Province during the summer of 2003 and F1 plants, which served as the male parent, were backcrossed with the recurrent parent CCRI36 in Hainan Province. BC1F1 seeds were harvested during the spring of 2003. In 2004, the grafted plants in the BC1F1 population were pulled out at the seedling stage, and BC1F1 plants, which served as the male parent, were backcrossed with CCRI36, and the hybrid seeds originating from the same BC1F1 plants were harvested together (BC2F1 seeds were obtained). BC2F1 plants were self-pollinated and BC1S1 seeds were harvested.

Field evaluation of Verticillium wilt (VW) disease resistance for BC2F1 and BC1S1 populations
A field in the Anyang experiment farm at the Institute of Cotton Research of Chinese Academy of Agricultural Sciences, Anyang, Henan Province, China, had been consecutively planted for more than 30 years with cotton. So it had a relatively high and uniform density of V. dahliae with both of the defoliating and nondefoliating populations and could be used as a natural nursery to screen for VW resistance breeding lines. In April 2005, 133 BC2F1 and 121 BC1S1 families (BC2F1-FD and BC1S1-FD) and the two parental lines (CCRI36 and Hai1) and its F1 were planted in this field in single-row plots at one replication for per every family and in two-row plots for the two parents and F1. Each row was 8 m long and 0.8 m wide with 32 plants. The grafted plants in BC2F1 and BC1S1 were pulled out at the seedling stage.

Preparation of inoculum and the artificial disease nursery evaluation of Verticillium wilt (VW) for BC2F1 population
A defoliating, moderate pathogenic Anyang strain of Verticillium dahliae, was a dominant pathogenic strain, isolated from the seriously infected field of cotton at Anyang, Henan Province, China. Before sowing cotton, the Anyang strain was cultured with cotton seeds with fuzz at 25 °C for 10 days. And the dry culture was ground and placed uniformly in the artificial disease nursery, with the inoculum concentration of 45 g of dry culture per 1 square meter of the disease nursery with conidial content of 1 × 10^8 per 1 g of the dry culture. This inoculation procedure yielded a mean DI of ca. 50 on the susceptible check variety in the normal conditions. The VW resistance for the 133 BC2F1 families and their parents were evaluated on June 23, July 19, August 15, and August 25 in 2005 in the artificial disease nursery.

The artificial disease nursery consists of a series of cement pools used for the artificial inoculation. Every pool is 20 m long and 2.5 m wide. The test was arranged in a randomized complete block design (RCBD) with 3 replicates and one row (plot) for every family. Each row was 2.5 m long and 0.6 m apart with 15 plants.

Phenotyping
The leaf tissue damage in the seedling and the maturity stages were classified into five grades which is a national standard in Verticillium resistant scoring in China [19, 48]. The grades scored as 1 and 2 were considered as resistant, and grades 3 and 4 as susceptible to VW.

<25 % chlorotic/necrotic leaves, grades 1
25–50 % chlorotic/necrotic leaves, grades 2
50–75 % chlorotic/necrotic leaves, grades 3
>75 % chlorotic/necrotic leaves, grades 4

The disease grade of each plant in BC2F1 was recorded by the leaf symptoms on June 23, July 19, August 15, and August 25 in 2005 in the artificial disease nursery. The disease grade of each plant was recorded by the leaf symptoms for BC2F1 and BC1S1 on 17 September in 2005 in the experiment field in Anyang. The DI was
calculated using all plants scores within the family in each plot, and then the disease index (DI) for per each family was calculated to get the average value using the DI of the three replicates (the phenotype datasets in the Additional file 2). The DI was calculated as follows:

\[
\text{DI} = \left[ \sum (N_i \times i) / (N \times 4) \right] \times 100, \quad i = 1 \sim 4, \quad N_i = \text{number of plants with the grade}, \quad N = \text{the total number of plants in each family}
\]

DNA extraction, PCR amplification, electrophoresis and map construction
Cotton genomic DNA was extracted from young leaves of the 135 BC1F1 plants and the two parents as described by Paterson et al. [49]. SSR-PCR was conducted as described by Sun et al. [1] and the PCR products were electrophoresed and silver-stained according to the protocol of Zhang et al. [50]. The genotype datasets are in the Additional file 3. SSR markers were used to construct a genetic map for BC1F1 population using JoinMap 4.0 [51] and the linkage map was published in 2015 [33].

QTL analysis
We used the Linkage maps with the phenotypic data of four dates of BC2F1-NY populations and one date of BC3S1-FD and BC2F1-FD populations to identify QTLs for VW resistance. QTLs were analyzed using the composite interval mapping method [52] and Windows QTL Cartographer 2.5 [53], with a walk speed of 1 cM, a window size of 10 cM and 5 background control markers. The statistical significance of the QTLs identified for the trait was determined by running a permutation procedure 1,000 times [54]. Positive additive effects indicated that Hai1 alleles decreased the DI value, increasing the phenotypic values of VW resistance, and negative scores indicated that CCRI 36 decreased the DI value, increasing the values of VW resistance. The QTLs were named as follows: (q + trait abbreviation) + chromosome/linkage group + QTL number. QTLs for the same trait across different generations and dates were considered stable when their confidence intervals overlapped.

Meta-analysis of QTLs
Recently, Said et al. reported a meta QTL analyses and established CottonQTLdb database containing 2274 QTL for 66 different QTL trait types [34], including 139 QTLs of VW resistance from nine published papers [15, 16, 18, 19, 24–28]. We got the VW resistance QTL mapping information including their names and CI provided by Said et al. [34]. The other 5 studies for VW resistance QTL mapping were also lately reported [13, 17, 21–23], including 65 QTLs of VW resistance. Totally, we used 252 QTLs of VW resistance for Meta-analysis, including 48 QTLs identified in our study.

A meta-analysis of QTLs for VW resistance was performed using Biomercator V3 software [55]. The detailed description for the meta-analysis of QTLs could be found in Said et al. [35, 36]. Hotspots were determined manually. Four or more QTLs in an interval of 25 cM were considered as a consistent QTL region. If there was more than one trait involved in the QTLs, the region is called a QTL cluster. Otherwise, it is called a QTL hotspot for the region involving only one single trait [13].

Additional files

| Additional file 1: The details of QTLs identified using interspecific backcross populations on the map of CCRI36 × Hai1 BC2F1, and their corresponding hotspots. (XLSX 16 kb) |
| Additional file 2: The phenotype datasets of BC2F1 on June 23, July 19, August 15, and August 25 in 2005 in the artificial disease nursery and of BC2F1 and BC3S1 on 17 September in 2005 in the field. (XLSX 27 kb) |
| Additional file 3: The genotype datasets of 135 BC1F1 plants of CCRI36× Hai1. (XLSX 1247 kb) |

Abbreviations
C: chromosome; cM: CentiMorgan; CSSLs: Chromosome substitution segment substitution lines; DI: Disease index; MAS: Marker assisted selection; QTL: Quantitative trait loci; SSR: Simple sequence repeat; VW: Verticillium wilt;

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Authors’ contributions
YLY conceived the study. And BCZ and YZS performed the experiments, YZS conducted the data analysis and drafted the manuscript. YLY revised the manuscript. AYL and JWL helped with the field experiments. WTL, QWL, ZZ and SQL helped with data analyses. YLY, YZS, AYL, JWL, WKG, HHS, JWT, TT, TW, SQ, HQZ and LZ contributed reagents and materials. All authors read and approved the manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
This article does not contain any studies with human participants or animals performed by any of the authors.

Informed consent
Informed consent was obtained from all individual participants included in the study.
References
1. Sun FD, Zhang JH, Wang SF, Gong WK, Shi YZ, Liu AY, Li JW, Gong JW, Shang HH, Yuan YL. QTL mapping for fiber quality traits across multiple upland cotton breeding populations. World Cotton Res Conf. 2003;15(5):457–65.
2. Zhang JF, Percy RG, McCarty Jr JC. Introgression genetics and breeding improvement of cotton against Verticillium wilt. Mol Plant Breed. 2010;8:680–86.
3. Wang T, Shang HH, Gong WK, Yuan YL. Phenotyping traits related to yield and quality of BC_{F2} substitution lines in cotton (Gossypium hirsutum). Mol Plant Breed. 2015;290(5):450–67.
4. Said JI, Song MZ, Wang HT, Lin ZX, Zhang XL, Song MZ, Zhang JF. A comprehensive comparative meta-analysis of QTL between intraspecific and interspecific Gossypium barbadense populations. Mol Genet Genomic. 2015;290(3):1003–10.
5. Said JI, Knapka JA, Song MZ, Zhang JF. Cotton QTLdb: a cotton QTL database for QTL analysis, visualization, and comparison between Gossypium hirsutum and G. barbadense populations. Mol Gen Genomic. 2015;290(3):1615–25.
6. Said JI, Zhongyu Lin ZX, Zhang XL, Song MZ, Zhang JF. A comprehensive meta-QTL analysis for fiber quality, yield, related and morphological traits, drought tolerance, and disease resistance in tetraploid cotton. BMC Genomics. 2013;14:776. doi:10.1186/1471-2164-14-776.
7. Said JI, Song MZ, Wang HT, Lin ZX, Zhang XL, Fang DD, Zhang JF. A comparative meta-analysis of QTL between intraspecific Gossypium hirsutum and interspecific G. hirsutum × G. barbadense populations. Mol Genet Genomic. 2014;290(3):1003–25.
8. Ulloa M, Meredith WR. Genetic linkage map and QTL analysis of agronomic and fiber quality traits in an intraspecific population. J Cotton Sci. 2000;4:161–70.
9. Paterson AH, Saranga Y, Mentz M, Jiang CX, Wright RJ. QTL analysis of genetic variation by environment interaction affecting cotton fiber quality. Theor Appl Genet. 2003;106(3):384–96.
10. Shen XL, Guo WZ, Zhu XF, Yuan YL, Yu JZ, Kohel RJ, Zhang TZ. Molecular mapping of QTLs for fiber length in a trifoliate line. J Cotton Sci. 2005;9(2):69–77.
41. Chen L, Zhang ZS, Hu MC, Wang W, Zhang J, Liu DJ, Zheng J, Zheng FM, Ma J. Genetic linkage map construction and QTL mapping for yield and fiber quality in Upland Cotton (Gossypium hirsutum L.). Acta Agron Sin. 2008;34:1199–205.
42. Wright RP, El-Zik TK, Paterson AH. D-subgenome bias of Xcm resistance genes in tetraploid Gossypium (Cotton) suggests that polyploid formation has created novel avenues for evolution. Genetics. 1998;149:1987–96.
43. Rong J, Feltus EA, Waghmare VN, Pierce GJ, Chee PW, Draye X, Saranga Y, Wright RJ, Willkins TA, May OL, Smith CW, Gannaway JR, Wendel JF, Paterson AH. Meta-analysis of polyploid cotton QTL shows unequal contributions of subgenomes to a complex network of genes and gene clusters implicated in lint fiber development. Genetics. 2007;176(6):2577–88.
44. Yang ZM, Li JZ, Li AG, Zhang BC, Liu GP, Li JW, Shi YZ, Liu Y, Jiang JX, Wang T, Yuan YL. Developing chromosome segment substitution lines (CSSLs) in Cotton (Gossypium) using advanced backcross and MAS. Mol Plant Breed. 2009;7:233–41.
45. Lan MJ, Yang ZM, Shi YZ, Ge RH, Li AG, Zhang BC, Li JW, Shang HH, Liu Y, Wang T, Yuan YL. Assessment of substitution lines and identification of QTL related to fiber yield and quality traits in BcF2 and BcF3 populations from Gossypium hirsutum × Gossypium barbadense. Sci Agric Sin. 2011;44:3086–97.
46. Ma LJ, Shi YZ, Lan MJ, Yang ZM, Zhang JF, Zhang BC, Li JW, Wang T, Gong JW, Liu Y, Shang HH, Gong WK, Yuan YL. Evaluation of chromosome segment substitution lines related to fiber yield and quality traits from Gossypium hirsutum × Gossypium barbadense. Cotton Sci. 2013;25(6):486–95.
47. Jing SR, Zhan XH. Breeding of new types of dominant glandless Upland cotton (G. hirsutum L). Agric Sci Chin (in Chinese). 1990;23:22–7.
48. Wu ZB, Li J, Feng CD, Zhang JF. Studies on the identification techniques of cotton resistance to Verticillium wilt. Hubei Agric Sci. 1999;5:16–9.
49. Paterson AH, Brubaker CL, Wendel JF. A rapid method for extraction of cotton (Gossypium spp.) genomic DNA suitable for RFLP or PCR analysis. Plant Mol Bio Reporter. 1993;11:122–7.
50. Zhang J, Guo W, Zhang T. Molecular linkage map of allotetraploid cotton (Gossypium hirsutum L. × Gossypium barbadense L.) with a haploid population. Theor Appl Genet. 2002;105:1166–74.
51. Van Ooijen JW. JoinMap 4.0, Software for the calculation of genetic linkage maps in experimental populations. Kyazma BV, Wageningen. 2006.
52. Zeng ZB. Precision mapping of quantitative trait loci. Genetics. 1994;136:1457–68.
53. Wang S, Basten CJ, Zeng ZB. Windows QTL Cartographer 2.5, Department of Statistics, Raleigh: North Carolina State University; 2006. http://statgen.ncsu.edu/~shchwang/WQTLCart.htm.
54. Churchill GA, Doerge RW. Empirical threshold values for quantitative trait mapping. Genetics. 1994;138:963–71.
55. Arcade A, Labourdette A, Falque M, Mangin B, Chardon F, Charcosset A, Joets J. BioMercator: Integrating genetic maps and QTL towards discovery of candidate genes. Bioinformatics. 2004;20(14):2324–26.