QTL-seq for rapid identification of candidate genes for 100-seed weight and root/total plant dry weight ratio under rainfed conditions in chickpea

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Summary
Terminal drought is a major constraint to chickpea productivity. Two component traits responsible for reduction in yield under drought stress include reduction in seeds size and root length/root density. QTL-seq approach, therefore, was used to identify candidate genomic regions for 100-seed weight (100SDW) and total dry root weight to total plant dry weight ratio (RTR) under rainfed conditions. Genomewide SNP profiling of extreme phenotypic bulks from the ICC 4958 × ICC 1882 population identified two significant genomic regions, one on CaLG01 (1.08 Mb) and another on CaLG04 (2.7 Mb) linkage groups for 100SDW. Similarly, one significant genomic region on CaLG04 (1.10 Mb) was identified for RTR. Comprehensive analysis revealed four and five putative candidate genes associated with 100SDW and RTR, respectively. Subsequently, two genes (Ca_04364 and Ca_04607) for 100SDW and one gene (Ca_04586) for RTR were validated using CAPS/dCAPS markers. Identified candidate genomic regions and genes may be useful for molecular breeding for chickpea improvement.

Introduction
Chickpea is the second most important grain legume crop, cultivated predominantly by resource poor farmers in arid and semi-arid regions of the world. Annual global production of chickpea is around 13.1 million tons from 13.5 Mha (FAOSTAT, 2013). India is the major chickpea producing country, producing 67% of the world’s chickpea in 2013 (FAOSTAT, 2013). Besides being an important source of protein for millions of people in developing countries, particularly in South Asia, it is a rich source of fibre, minerals (phosphorus, calcium, magnesium, iron and zinc) and β-carotene. In addition, chickpea can obtain over 70% of its nitrogen requirement through symbiotic nitrogen fixation (SNF) by fixation of 140 kg/ha of atmospheric nitrogen to the soil (Flowers et al., 2010; Gaur et al., 2012), although chickpea has high calorific value (364 kcal/100 g after soybean 446 kcal/100 g; Akibode and Maredia, 2011) and high protein content (18%–25%). However, the average per capita availability of chickpea decreased by 6% across the world and 14% especially in India between 1990 and 2008 due to increase in population growth. In addition, several biotic and abiotic stresses have also been limiting the chickpea productivity especially in India, which happens to be the largest producer and consumer as well. Nevertheless, with the expansion in international trade for chickpea, several developing countries (e.g. Ethiopia), as well as developed countries (e.g. Australia, Canada), have been emerging as major exporters of chickpea to India.

Development and adoption of improved varieties with higher yield will enhance per capita consumption, thereby reducing the number of malnourished people across the world, especially in developing countries. In this direction, in addition to conventional breeding strategies, genomics approaches are being deployed in recent years to identify genomic regions that confer resistance/tolerance for different stresses. Both biparental mapping and association mapping approaches have been utilized to dissect complex traits in chickpea. However, to date, the limited number of trait mapping studies has mainly focused on mapping biotic stresses such as Fusarium wilt (Gowda et al., 2009; Sabbavarapu et al., 2013), ascochyta blight (Udupa and Baum, 2003; Kottapalli et al., 2008) and botrytis grey mould (Anuradha et al., 2011), as well as agronomically important traits (Das et al., 2015; Gowda et al., 2011). Efforts have also been made to dissect complex abiotic stresses such as salinity (Vadez et al., 2011) and drought tolerance (Hamwieh et al., 2013; Jaganathan et al., 2015; Kale et al., 2015; Rehman et al., 2011; Thudi et al., 2014; Varshney et al., 2014a) with an emphasis on understanding the performance of agronomic or physiological traits.

Recently, Varshney et al. (2014a) reported a genomic region on linkage group 4 in chickpea (CaLG04) referred as ‘QTL-hotspot’ for drought responsive traits. This genomic region explains highest phenotypic variation (PVE) of 58.20% for 100SDW, an important trait that has relevance to yield and 16.67% PVE in case of root dry weight/total plant dry weight ratio (RTR) that has relevance to drought tolerance (Kashiwagi et al., 2006). Owing to the large size of this genomic region, the
region was refined using a genotyping-by-sequencing (GBS) approach (Jaganathan et al., 2015). As a result, the ‘QTL-hotspot’ region was narrowed to 14 cM (~3 Mb) and candidate genes prioritized for analysis (Kale et al., 2015).

In recent years, advances in next generation sequencing technologies (NGS) have seen the rapid decrease in cost of sequencing, enabling the cost-effective application of genomics in crop improvement possible (Varshney et al., 2014b). The advantage of identifying large numbers of SNPs through resequencing has made it possible to locate and refine candidate genomic regions more efficiently compared to traditional QTL mapping approaches (Chen et al., 2014; Kale et al., 2015; Qi et al., 2014; Xu et al., 2013). Bulk segregant analysis (BSA) is a rapid method to identify markers linked to the trait of interest (Michelmore et al., 1991) and when combined with NGS technologies can be used as a fast track approach to locate candidate genomic regions more rapidly. This approach is known as QTL-seq and involves the selection of 20–50 lines with extreme phenotypic values, pooling in equivalent concentration followed by sequencing of pools for downstream sequence analysis (Takagi et al., 2013). The QTL-seq approach has been used successfully to map candidate genomic regions for early flowering in cucumber (Lu et al., 2014), fruit weight and locule number in tomato (Illa-Berenguer et al., 2015) and 100-seed weight in chickpea (Das et al., 2015).

With an objective of identifying candidate genomic regions responsible for 100SDW and RTR, the QTL-seq approach was adopted. We were able to precisely localize the genomic regions for the two target traits and identify five genic SNPs in four candidate genes for 100SDW and six genic SNPs in five candidate genes for RTR. The involvement of candidate genes was further validated through co-segregation analysis in the entire RIL population derived from ICC 4958 × ICC 1882.

Results

Extreme bulks for 100SDW and RTR

Based on phenotyping data generated earlier for the ICC 4958 × ICC 1882 mapping population (Varshney et al., 2014a), two extreme bulks each for 100SDW and RTR were prepared and subjected to the QTL-seq pipeline as shown in Figure S1 for supplementary figures. In brief, phenotypic data for 100SDW showed a skewed segregation towards the tolerant parent ICC 4948 (Figure S2), with trait values of RILs ranging between 12.24 and 30.80 g [values for parental lines: 31.83%–0.02 (ICC 4958); 26.38%–13.59 g] were selected to prepare the two extreme bulks. The phenotyping data for RTR among RILs showed a significant deviation from SNP-index of 0.5 could indicate the contribution of equal genome contribution from both parents. A significant deviation from SNP-index of 0.5 could indicate the contribution of that SNP to the phenotypic difference observed in the bulks (Abe et al., 2012).

SNP-index values across the genome were calculated and sliding window averages of 2 Mb interval with 10 kb increments were plotted for high and low bulks. To facilitate detection

| Genotypes | Number of bulked lines | Number of reads | Unmapped read (%) | Genome coverage (%) | Average depth (X) |
|-----------|------------------------|-----------------|-------------------|----------------------|------------------|
| ICC 4958* | –                      | 29299428        | 0.39              | 96.35                | 6.52             |
| High 100SDW bulk† | 15          | 17795370        | 0.29              | 81.73                | 3.18             |
| Low 100SDW bulk† | 15          | 20670924        | 0.29              | 81.53                | 3.17             |
| High RTRbulk† | 15          | 17954432        | 0.28              | 81.94                | 3.21             |
| Low RTRbulk† | 15          | 17574432        | 0.29              | 81.05                | 3.17             |

*ICC 4958 short reads were aligned to the publicly available chickpea genome of CDC Frontier (Varshney et al., 2013), a kabuli chickpea variety.
†The short reads of bulks were aligned to the ICC 4958 assembly developed by replacement of SNPs between ICC 4958 and CDC Frontier.
differences in SNP-indices between high and low bulks, we plotted ΔSNP-index with a statistical confidence interval. With a statistical significance of \( P < 0.05 \), significant genomic positions were identified.

Sequence analysis for 100SDW revealed two genomic regions on two linkage groups (CaLG01 and CaLG04) (Figures 1 and S5–S7). In both regions, alleles from ICC 4958 contributed to higher trait values and those from ICC 1882 to lower trait values. A genomic region spanning 1.08 Mb (3.07–4.15 Mb) on CaLG01 showing significant \( P < 0.05 \) deviation from equal inheritance of the two parental genomes had five SNPs with ΔSNP-index = −1 (Table S3). However, none of the SNPs was from predicted genic regions. The lower number of SNPs may be due to the narrow candidate region and/or low coverage. Further, the second 2.70 Mb region (11.12–13.82 Mb) on CaLG04 harboured 21 SNPs with ΔSNP-index = −1. Of these 21 SNPs, five SNPs were present in four putative genes (Ca_04364, Ca_04600, Ca_04602 and Ca_04607; Table 2). Of these five genic SNPs, four were in intronic region and one SNP (Ca_04607_13822383) in the exon of a predicted gene Ca_04607, associated with transmembrane protein, causing a nonsynonymous substitution from valine (ICC 4958) to isoleucine (ICC 1882).

Similarly, sequence analysis of low and high RTR bulks for the identified 1.10 Mb region (12.73–13.83 Mb) on linkage group 4 (CaLG04) revealed 26 SNPs with ΔSNP-index = −1, suggesting a biased inheritance of parental genomes in the two populations (Figures 2 and S8–S10). The high RTR bulk showed SNP-index = 0 indicating that high trait value alleles were inherited from the tolerant parent ICC 4958. By contrast, low RTR bulk at these 26 positions possesses SNP-index = 1, indicating that their alleles were derived from the susceptible parent ICC 1882 (Table S4). Of 26 SNPs, six SNPs were found to occur in five predicted genes (Ca_04493, Ca_04586, Ca_04592, Ca_04595 and Ca_04602). Interestingly, two SNPs (at 13666705 bp for synonymous change and 13666728 bp for T to I amino acid change) were identified in the exon of a putative gene, Ca_04586 predicted to be coding for cytochrome P450 monoxygenase (Table 3).

Validation of identified genomic regions
CAPS/dCAPS markers represent cost-effective assays to genotype SNPs in a segregating population. Therefore, to validate the identified five genic SNPs for 100SDW and six genic SNPs for RTR regions, a total of 11 CAPS/dCAPS primers were designed (Table S5). Of 11, eight primers amplified prominent fragments of...
Table 2 Identification of SNPs in putative candidate genes for 100-seed weight (100SDW)

| Linkage group | Gene | Position | ICC 4958 allele | High 100SDW bulk allele | SNP-index (high 100SDW bulk)* | Low 100SDW bulk allele | SNP-index (low 100SDW bulk)† | Δ SNP-index‡ | SNP effect | Function |
|---------------|------|----------|----------------|-------------------------|-------------------------------|------------------------|----------------------------|-----------------|------------|----------|
| CaLG04       | Ca_04364 | 11311944 | A              | A                       | 0                             | C                      | 1                         | –1              | Intron     | Cell division protein kinase |
| CaLG04       | Ca_04600 | 13760326 | C              | C                       | 0                             | T                      | 1                         | –1              | Intron     | Uncharacterized protein     |
| CaLG04       | Ca_04602 | 13780146 | C              | C                       | 0                             | G                      | 1                         | –1              | Intron     | Random slug protein         |
| CaLG04       | Ca_04607 | 13822383 | G (Gtc/V)§     | G (Gtc/V)§              | 0                             | A (Atc/I)§             | 1                         | –1              | Exon       | Transmembrane protein       |
| CaLG04       | Ca_04607 | 13822453 | A              | A                       | 0                             | G                      | 1                         | –1              | Intron     | Transmembrane protein       |

*SNP-index of high 100SDW bulk was calculated based on the allele calls and read depth in comparison with ICC 4958 reference assembly.
†SNP-index of low 100SDW bulk was calculated based on the allele calls and read depth in comparison with ICC 4958 reference assembly.
‡Δ SNP-index of each SNP positions was calculated using following formula: Δ SNP-index = SNP-index of high 100SDW bulk – SNP-index of low 100SDW bulk.
§Value in parenthesis indicates the codon change due to SNP/Code for changed amino acids V, valine; I, isoleucine.

Figure 2 QTL-seq approach adopted for mapping genomic regions responsible for RTR. (a) Morphological differences between drought tolerant parent ICC 4958 (with larger root system) and ICC 1882 (sensitive parent with smaller root system) (b) Frequency distribution of RTR of 262 RILs based on 2 years mean data. The DNA of 15 RILs with extreme phenotypes (high and low RTR) was used to develop high and low RTR % bulks. (c) SNP-index plot of high RTR bulk (top), low RTR bulk (middle) and ΔSNP-index plot (bottom) of chromosome 4 with statistical confidence interval under the null hypothesis of no QTLs (orange, $P < 0.01$ and green $P < 0.05$). The significant genomic regions are highlighted in orange shaded colour (12.73–13.83 Mb).
expected size in two different parental combinations (ICC 4958 × ICC 1882; ICC 283 × ICC 8261). Of these eight primer pairs, six (three for each 100SDW and RTR trait) were found polymorphic using their respective restriction enzymes. Of six polymorphic markers, four Ca_04364_11311944 and Ca_04607_13822453 for 100SDW and Ca_04586_13666705 and Ca_04586_13666728 for RTR were validated in the high and low DNA pools. The fragment size of high DNA pools of 100SDW and RTR followed the similar pattern of the tolerant line ICC 4958 and the low DNA pool of 100SDW and RTR bulks followed the similar pattern of the sensitive line ICC 1882. The details on amplified fragment size and digested product size of each primer pairs are presented in Figures 3 and 4 and Table S6. Of 11 primer pairs tested, four CAPS markers follow the similar pattern in high trait parent and high bulk and similarly to low trait parent and low bulk. This ensures the utilization of these CAPS/dCAPS markers in marker-assisted breeding programme.

**Confirmation of marker–trait associations**

The genotyping data of four candidate CAPS/dCAPS markers (Ca_04364_11311944 and Ca_04607_13822453, Ca_04586_13666705 and Ca_04586_13666728) were used to develop genetic linkage map. As a result, a map length of 25.18 cM was obtained for linkage group 4. Single-marker QTL analysis showed a high significance (\( P < 0.01 \)) for all markers. The candidate markers for 100SDW, Ca_04364_11311944 and Ca_04607_13822453, explained high phenotypic variation of 28.61% with LOD score of 64.58 and 19.25% with LOD score of 45.61, respectively (Table 4). Similarly, the candidate markers for RTR, Ca_04586_13666728 and Ca_04586_13666705 explained a phenotypic variation of 23.29% with LOD score of 51.11 and 26.46% with LOD score of 56.33, respectively (Table 4).

**Discussion**

Genetic mapping of QTLs for economically important traits is an important component of the development of new and elite lines in plant breeding programmes. Classic methods of QTL mapping involve genotyping of segregating mapping population with polymorphic markers identified between parents and use of genotyping data with phenotyping data sets for defining significant genomic regions for the target traits. However, identification of polymorphic markers between contrasting parents is time-consuming and tedious task (Schneeberger and Weigel, 2011). For instance, during earlier studies, a total of 2717 (between ICC 4958 × ICC 1882) and 2410 (between ICC 283 × ICC 8261) markers including simple sequence repeats (SSRs), genic molecular markers (GMMs) and EST-SSRs were screened on the parental lines of these two mapping populations. The markers showed polymorphism between parents (between ICC 4958 and ICC 1882 after digestion with AluI and Msel restriction enzyme, respectively. The PCR amplicons also correspond to the high and low 100SDW bulk along with other two parental lines (ICC 8261—high seed weight parent and ICC 283—low seed weight parent) of different mapping population.

![Figure 3](Image)

**Figure 3** Validation of candidate gene-based markers for 100SDW. Two gene-based markers Ca_04364_11311944 (CAPS) and Ca_04607_13822453 (dCAPS) associated with 100SDW showed clear polymorphism between ICC 4958 and ICC 1882 after digestion with AluI and Msel restriction enzyme, respectively. The PCR amplicons also correspond to the high and low 100SDW bulk along with other two parental lines (ICC 8261—high seed weight parent and ICC 283—low seed weight parent) of different mapping population.
of NGS technologies and availability of the chickpea genome ensures that the number of genomewide markers is not a limiting factor for trait mapping (Varshney et al., 2014b). Therefore, the recently proposed NGS-based QTL-seq approach, which is a cost-effective and rapid method of trait mapping, was adopted in this study as it is a proven approach to identify the target genomic region in rice (Takagi et al., 2013). This approach avoids the tedious genotyping of a large population, and it has the capacity to identify shorter genomic regions associated with a trait of interest than classical QTL mapping approaches. QTL-seq also has an advantage that the markers located within the identified candidate regions can be further used for fine mapping and cloning experiments. These markers can be converted into cost-effective CAPS/KASP markers for deployment in genomics-assisted breeding programmes.

In this study, sequencing of both bulks (high RTR and low RTR bulk) identified a comparatively large number of SNPs (7318 total SNPs and 3792 homozygous SNPs), which were used for calculating genomewide SNP-index. Analysis of SNPs located in this region revealed a candidate gene, $Ca_{04586}$ coding for cytochrome P450 monooxygenase, an abscisic acid (ABA) 8'-hydrodrolase which involved in ABA catabolism (Krochko et al., 1998), reducing ABA during seed imbibition. Cytochrome P450 monooxygenase gene has been shown up-regulated during drought stress in Arabidopsis and found to play an important role in the maintenance of ABA levels in plants (Kushiro et al., 2004). The possible role of $Ca_{04586}$ in drought tolerance in chickpea needs to be investigated further.

Similarly for 100SDW, two genomic regions on CaLG01 (1.08 Mb) and CaLG04 (2.70 Mb) were identified. It is interesting to note that several studies reported the presence of QTL for 100SDW on CaLG04 in chickpea (Abbo et al., 2005; Cho et al., 2002; Cobos et al., 2007), showing the importance of this linkage group in yield trait improvement for chickpea. These results highlighted the significance of the QTL-seq approach in identifying refined and reliable candidate regions for the trait of interest. Detailed analysis of SNP frequencies identified the gene $Ca_{04607}$ on CaLG04, coding for a ‘transmembrane protein’. Role of transmembrane proteins in controlling grain weight, grain length, grain width and thickness has been earlier reported in rice (Fan et al., 2006; Shomura et al., 2008; Song et al., 2007). Based on these findings, it is possible to target $Ca_{04607}$ for fine mapping and cloning of seed weight-related genes in chickpea. However, it is important to note that this gene was not present in the prioritized list of genes in our recent study despite the mapping of 100SDW responsive QTLs in the same genomic region from 11.12 to 13.82 Mb on CcLG04 (Kale et al., 2015; 13.23–13.37 Mb for ‘QTL-hotspot-a’ and 13.39–13.54 for ‘QTL-hotspot-b’ on CaLG04). This might be due to low-coverage sequencing of RIL population (average 0.72 X per RILs) or because of errors in genotyping of RILs or resequencing of parental genotypes.

Recently, Das et al. (2015) utilized QTL-seq approach utilizing RILs population (ICC 7184 × ICC 15061) and reported major seed weight QTL on CaLG01 (CaqSW1.1: 0.83–0.87 Mb). This QTL region was further narrowed down using an integrated approach and reported CSN8 as a possible candidate gene for controlling seed weight in chickpea. Similarly, five robust QTLs on five different linkage groups (CaLG 1, 2, 5, 6 and 7) with PVE ranged from 10.07% to 22.31% using GBS approach in SBD377 × BGDL12 mapping population (Verma et al., 2015). Thus, the genomic regions reported using similar approaches in earlier study are different from that we reported in this study, which are novel.

To understand the robustness and precision of identification of the genomic regions responsible for 100SDW and RTR over the classical QTL mapping studies, we compared the results of this work with our earlier studies (Jaganathan et al., 2015 and Varshney et al., 2014a). In the case of 100SDW, Varshney et al. (2014a) reported two major QTLs each on CaLG01 and CaLG04 that explained 10.31% and 58.20% of phenotypic variation, respectively. Based on the physical position of flanking markers, the QTLs for 100SDW spanned 6.57 Mb (2.93–9.51 Mb) on CaLG01 and 6.75 Mb (10.07–16.83 Mb) on CaLG04, while Jaganathan et al. (2015) reported two major QTLs each on CaLG01 and CaLG04 that explained 16.23% and 60.41% of phenotypic variation, respectively. Based on the physical position of flanking markers, the QTLs for 100SDW spanned 8.43 Mb (1.08–9.51 Mb) on CaLG01 and 0.45 Mb (13.68–14.14 Mb) on CaLG04. While comparing this study with previous studies, the genomic region identified on CaLG01 was narrowed down from 6.57 Mb (Varshney et al., 2014a) and 8.43 Mb (Jaganathan et al., 2015) to 1.08 Mb (3.07 to 4.15 Mb) in the current study (Table S7 and Figure 5a), whereas genomic

![Figure 4 Validation of candidate gene-based markers for RTR. Two gene-based markers.](Image)
region identified on CalLG04 in the present study was 2.70 Mb in comparison with 6.75 Mb (Varshney et al., 2014a) and 0.45 Mb (Jaganathan et al., 2015) identified earlier.

Similarly, in the case of RTR, a major QTL (Qr3tr01) explaining 16.67% of phenotypic variance was reported between TAA170 and NCPGR21 on CalLG04 (Varshney et al., 2014a). Based on the physical position of the flanking SSR markers, the identified QTL was found to be located at 10.07–14.14 Mb (4.06 Mb region) on the physical map. With a denser genetic map (1007 loci), the same genomic region was further delimited to 3.76 Mb (10.07–13.84 Mb; Jaganathan et al., 2015). Nevertheless, the QTL-seq approach deployed in the present study narrowed down the region further (1.1 Mb) and identified the genomic region responsible for RTR (Table S7 and Figure 5b). In this region, a promising candidate gene, cytochrome P450 monoxygenase responsible for RTR (Table S7 and Figure 5b). In this region, a promising candidate gene, cytochrome P450 monoxygenase involved in ABA catabolism (Kushiro et al., 2004).

In addition to the identification of candidate genes located in genomic regions associated with 100SDW and RTR, SNP markers were developed that could be directly implemented in selection strategies in breeding programmes. Of 11 tested, four CAPS/dCAPS markers follow the similar pattern in the high trait parent and the high bulk and similarly to low trait parent and low bulk. Single-marker analysis of these markers shows significant correlation with the traits measured, confirming their suitability for application in marker-assisted selection. The current study suggests that WGRS-based BSA mapping methods can be adopted for other traits of relevance in crop improvement, potentially avoiding cost and time constraints associated with traditional QTL mapping and providing a powerful means to rapidly refine genomic regions containing candidate genes.

**Experimental procedures**

**Plant materials**

Chickpea recombinant inbred line (RIL) population (ICCRIL03) comprising 262 lines developed from a drought tolerant genotype, ICC 4958 (with high 100SDW and high RTR) and a drought-sensitive genotype, ICC 1882 (low 100SDW and low RTR) was used in the present study. Extensive phenotyping data under rainfed conditions for several drought component traits (20 traits) in 1–7 seasons and 1–5 locations along with marker data enabled by Varshney et al. (2014a) to provide greater insights into the drought tolerance and genomic regions responsible for the traits studied.

**Construction of pools**

Extreme bulks were prepared for 100SDW and RTR traits based on precise phenotyping data obtained for 5 years (2005, 2006, 2007, 2008 and 2009) and 2 years (2005 and 2007), respectively, under rainfed conditions. For developing the extreme bulks for each trait, 15 RILs with high mean phenotypic values and 15 RILs with low mean phenotypic values were selected. The equimolar concentration of DNA from 15 RILs with high phenotypic values was pooled together as one bulk, and similarly DNA from low mean phenotypic values was pooled together as another bulk. Thus, four extreme bulks two for each the traits were prepared for sequencing.

**Construction of libraries and Illumina sequencing**

A total of five Illumina libraries (four from extreme bulks mentioned above and one from ICC 4958 the drought tolerant parent) were prepared using TruSeq DNA Sample Prep kit LT, (set A) FC-121-2001. Two microgram DNA from each sample was sheared using diagenone Bioruptor® NGS (Diogenode, Liege, Belgium), end repaired and adapter ligated. Size selection of libraries was performed using 2% agarose gel to get a target insert size of 500–600 bp and purified for further analysis. Further, the libraries were enriched using adaptor compatible PCR primers. The size distribution of amplified DNA libraries was checked on an Agilent Technologies 2100 Bioanalyzer using a High Sensitivity chip. The DNA libraries were sequenced on Illumina MiSeq platform with MiSeq Reagent Kit v2 (500-cycles) (Illumina Inc., San Diego, CA) to generate 250-base pair-end reads.

**Construction of reference-based assembly**

The statistics of generated sequencing reads was estimated using raspberry tool of NGS-QCbox (Katta et al., 2015). Further, QTL-seq pipeline (http://genome-e.ibrc.or.jp/home/bioinformatics-team/matmap, developed by Ioate Biotechnology Research Center, Japan) was used for calculating SNP-index. Briefly, the cleaned reads of donor parent (ICC 4958) were first aligned to the reference genome (CDC Frontier; Varshney et al., 2013) using inbuilt BWA aligner (Li and Durbin, 2009). Coval was used for postprocessing and filtering of the alignment files (Kosugi et al., 2013). The variants called were then used to develop reference-based assembly of the donor parent (ICC 4958) by substituting the bases with confidence variants calls in the genome. The reads from high and low bulks for both the traits were then aligned, and variants were called for both the bulks against the developed assembly.

**Calculation of SNP-index**

SNP-index for each SNP position was calculated for both the bulks as per Abe et al. (2012) using the formula:

\[ \text{SNP-index (at a position)} = \text{count of alternate base(count of reads aligned)} \]

The positions with read depth <7 in both the bulks and SNP-index <0.3 in either of the bulks were filtered out for ΔSNP-index calculation. ΔSNP-index can be calculated by subtracting SNP-index of low bulk from SNP-index of high bulk. Only SNP positions with ΔSNP-index = −1 (i.e. the allele called in high bulk was same as that of ICC 4958 while contrastingly different in low bulk) were considered as the causal SNPs responsible for the trait of interest. The possible effects of the identified SNPs were inferred using SnpEff v3.0 open source program (Cingolani et al., 2012).

**Validation of identified candidate SNPs**

Five genic SNPs for 100SDW and six genic SNPs for RTR identified in the present study were validated on four genotypes (ICC 4958, ICC 1882, ICC 283 and ICC 8261) and four bulks (High and low bulks of RTR and 100SDW). For this purpose, the SNPs were converted into CAPS/dCAPS markers using dCAPS Finder 2.0 (Neff et al., 2002) for the development of cost-effective gel-based markers. Of 11 SNPs, seven SNPs were converted into CAPS and four SNPs were converted into dCAPS. The predicted CAPS and dCAPS candidates were amplified on parental genotypes (ICC 4958 and ICC 1882) and also amplified on the resistant and susceptible bulks of respective pools from 100SDW and RTR. PCR amplifications of each CAPS and dCAPS marker were subjected to digestion with their respective restriction enzymes. The restricted samples were checked on 2% agarose gel electrophoresis.

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Figure 5 Co-localization of QTLs from traditional and QTL-seq approach for 100SDW and RTR. (a) Co-localization of QTLs mapped for 100SDW through traditional and QTL-seq method. (i) Pseudomolecules of reference genome CDC Frontier (Varshney et al., 2013), (ii) upper probability values at 99% confidence ($P < 0.01$), (iii) upper probability values at 95% confidence ($P < 0.05$), (iv) genomewide $\Delta$SNP-index (red dots denote $\Delta$SNP-index ranged from 0 to $-1$ and contributed by high trait parent (ICC 4958) and green dots denote $\Delta$SNP-index ranged from 0 to 1 and contributed by low trait parent (ICC 1882)), (v) lower probability values at 95% confidence ($P < 0.05$), (vi) lower probability values at 99% confidence ($P < 0.01$), (vii) physical position of earlier mapped QTL (Varshney et al., 2014a) for 100SDW through traditional mapping approach. The physical positions of QTL were estimated through blast of the flanking primers to the chickpea genome. (viii) Common genomic positions on linkage group 1 (CaLG01) and linkage group 4 (CaLG04) were observed through both the approaches. (b) Co-localization of QTLs mapped for RTR through traditional and QTL-seq method. (i) Pseudomolecules of reference genome CDC Frontier (Varshney et al., 2013), (ii) upper probability values at 99% confidence ($P < 0.01$) for declaring significant $\Delta$SNP-index, (iii) upper probability values at 95% confidence ($P < 0.05$) for declaring significant $\Delta$SNP-index, (iv) genomewide $\Delta$SNP-index (red dots denote $\Delta$SNP-index ranged from 0 to $-1$ and contributed by high trait parent (ICC 4958) and green dots denote $\Delta$SNP-index ranged from 0 to 1 and contributed by low trait parent (ICC 1882)), (v) lower probability values at 95% confidence ($P < 0.05$), (vi) lower probability values at 99% confidence ($P < 0.01$), (vii) physical position of earlier mapped QTL (Varshney et al., 2014a) for RTR through traditional mapping approach. The physical position of QTL was estimated through blast the flanking primers into the chickpea genome. (viii) Common genomic positions on CaLG04 were observed through both the approaches.
Single-marker analysis
To study the effect of identified candidate markers on the population, the RIL population (ICC 4958 × ICC 1882) was genotyped with all the four candidate CAPS/dCAPS markers. The genetic map was constructed using JoinMap V4.0 with the default parameters. Single-marker analysis (SMA) was carried out using the map along with the phenotypic data by QTL Cartographer. The mean phenotyping data generated under rainfed conditions used for making the pool as described earlier for 100SDW and RTR were used for this analysis. Based on the p-value, the marker was determined to control the trait of interest.

Author's contributions
V.K.S., A.W.K. and D.J. performed most of the experiments; V.K. and A.C. generated sequence data; A.W.K., H.T. and R.T. performed QTL-seq analysis; P.M.G. contributed genetic material; V.K.S., A.W.K., D.J., M.T., M.R., V.G., T.S., R.T. and R.K.V. analysed and interpreted the QTL-seq data; D.J., V.K.S. and R.K.V. performed validation of candidate SNPs; V.K.S., D.J., M.T., M.R., T.S., R.T. and R.K.V. wrote the manuscript; R.K.V. conceived, designed and supervised the study and finalized the manuscript. All authors read and approved the manuscript.

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Conflict of interest
The authors declare that they have no competing interests.

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Supporting information

Additional Supporting Information may be found online in the supporting information tab for this article:

Figure S1 QTL-seq approach used in chickpea.

Figure S2 Frequency distribution of 100SDW of the 262 RILs derived from a cross between ICC 4958 and ICC 1882 for different years (2005, 2006, 2007, 2008, 2009 and average) in field conditions.

Figure S3 Frequency distribution of RTR levels of the 262 RILs derived from a cross between ICC 4958 and ICC 1882 over two different years (2005 and 2007) with mean values in controlled conditions.

Figure S4 (a) Based on the 2 year phenotyping of 262 RILs for RTR, a total of 15 RILs with high RTR and 15 with low RTR were used to develop high RTR and low RTR bulks, respectively. (b) Similarly, based on 5 years data of 100SDW a total of 15 RILs with high 100SDW and 15 with low 100SDW were used to develop high 100SDW and low 100SDW bulks.

Figure S5 SNP-index plots for eight chromosomes of high 100SDW bulked DNA.

Figure S6 SNP-index plots for eight chromosomes of low 100SDW bulked DNA.

Figure S7 The Δ(SNP-index) plot obtained by subtraction of high 100SDW SNP-index from low 100SDW SNP-index for RILs obtained from a cross between ICC 4958 and ICC 1882.

Figure S8 SNP-index plots for eight chromosomes of high RTR bulked DNA.

Figure S9 SNP-index plots for eight chromosomes of low RTR bulked DNA.

Figure S10 The Δ(SNP-index) plot obtained by subtraction of high RTR SNP-index from low RTR SNP-index for RILs obtained from a cross between ICC 4958 and ICC 1882.

Table S1 Frequency distribution of 100SDW of the 262 RILs derived from a cross between ICC 4958 and ICC 1882 for different years (2005, 2006, 2007, 2008, 2009 and average) in field conditions.

Table S2 Frequency distribution of RTR levels of the 262 RILs derived from a cross between ICC 4958 and ICC 1882 over two different years (2005 and 2007) with mean values in controlled conditions.

Table S3 List of putative SNPs identified for 100SDW based on SNP-index values.

Table S4 List of putative SNPs identified for RTR based on SNP-index values.

Table S5 List of primers successfully validated for 100SDW and RTR.

Table S6 List of primers used for validation of candidate genes for RTR and 100SDW.

Table S7 Comparison of the identified QTLs from QTL-seq with earlier studies.