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Identifying Transcription Factor Genes Associated with Yield Traits in Chickpea

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Abstract Identification of potential transcription factor (TF) gene-derived natural SNP allelic variants regulating pod and seed yield component traits by large-scale mining and genotyping of SNPs in natural germplasm accessions coupled with high-resolution association mapping is vital for understanding the complex genetic architecture of quantitative yield traits in chickpea. In these perspectives, the current study employed a genome-wide GBS (genotyping-by-sequencing) and targeted gene amplicon resequencing-based simultaneous SNP discovery and genotyping assays, which discovered 1611 novel SNPs from 736 TF genes physically mapped on eight chromosomes and unanchored scaffolds of kabuli chickpea genome. These SNPs were structurally and functionally annotated in diverse synonymous and non-synonymous coding as well as non-coding regulatory and intronic sequence

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components of chickpea TF genes. A high-resolution genetic association analysis was performed by correlating the genotyping information of 1611 TF gene-based SNPs with multi-location/years field phenotyping data of six major pod and seed yield traits evaluated in a constituted association panel (326 desi and kabuli germplasm accessions) of chickpea. This essentially identified 27 TF gene-derived SNPs exhibiting significant association with six major yield traits, namely days to 50% flowering (DF), plant height (PH), branch number (BN), pod number (PN), seed number (SN) and seed weight (SW) in chickpea. These trait-associated SNPs individually and in combination explained 10–23% and 32% phenotypic variation respectively for the studied yield component traits. Interestingly, novel non-synonymous coding SNP allelic variants in five potential candidate TF genes encoding SBP (squamosal promoter binding protein), SNF2 (sucrose non-fermenting 2), GRAS [Gibberellic acid insensitive (GAI)-Repressor of GAI (RGA)-SCARECROW (SCR)], bZIP (basic leucine zipper) and LOB (lateral organ boundaries)-domain proteins associated strongly with DF, PH, BN, PN, SN and SW traits respectively were found most promising in chickpea. The functionally relevant molecular signatures (TFs and natural SNP alleles) delineated by us have potential to accelerate marker-assisted genetic enhancement by developing high pod and seed yielding cultivars of chickpea.

Keywords Association mapping \cdot Chickpea \cdot Desi \cdot Kabuli \cdot SNP . Transcription factor

Introduction

Chickpea (Cicer arietinum), represented by diverse desi and kabuli cultivars, is a legume food crop of profound economic

and nutritional value (Kumar et al. [2011](#page-12-0); Varshney et al. [2013a\)](#page-12-0). Enhancing chickpea productivity by developing multiple high seed and pod-yielding desi and kabuli cultivars is the major current genomics-assisted breeding objectives to sustain global food and nutritional security amidst climate change and huge population growth scenario. However, most of the pod and seed yield-contributing traits like flowering time, plant height, branch number, seed/pod number and seed weight are polygenic in nature and exhibit complex quantitative genetic inheritance pattern in chickpea (Gaur et al. [2014](#page-11-0); Varshney [2016\)](#page-12-0). Significant efforts have been made by several global research groups to dissect these complex yield traits by employing diverse genomics-assisted breeding strategies including high-resolution QTL (quantitative trait loci) and association mapping in chickpea (Kujur et al. [2013](#page-11-0), [2014,](#page-11-0) [2015a,](#page-11-0) [b,](#page-11-0) [c,](#page-11-0) [2016;](#page-12-0) Saxena et al. [2014a](#page-12-0); Thudi et al. [2014](#page-12-0); Varshney et al. [2014;](#page-12-0) Bajaj et al. [2015a](#page-10-0), [b,](#page-11-0) [c](#page-11-0), [2016a](#page-11-0), [b;](#page-11-0) Das et al. [2015,](#page-11-0) [2016;](#page-11-0) Kale et al. [2015;](#page-11-0) Upadhyaya et al. [2015](#page-12-0); Singh et al. [2016\)](#page-12-0). This is primarily accomplished by high-throughput genotyping of gene/genome-based SSR (simple sequence repeat) and SNP (single nucleotide polymorphism) markers in phenotypically well-characterized natural core/mini-core germplasm accessions and advanced generation intra-/interspecific mapping populations of chickpea. Unfortunately, very limited number of informative sequence-based markers tightly linked to the major QTLs/genes governing aforesaid complex yield traits are identified till date to be exploited for marker-assisted genetic improvement of chickpea (Varshney et al. [2013a](#page-12-0), [b,](#page-12-0) [2014;](#page-12-0) Saxena et al. [2014a](#page-12-0); Upadhyaya et al. [2015\)](#page-12-0). To drive these prime objectives, the gene-based markers essentially appear much more informative and proficient for fast delineation of potential genes and alleles underlying major QTLs regulating diverse vital agronomic traits and deciphering the complex genetic architecture of seed and pod yield component quantitative traits in chickpea (Kujur et al. [2013,](#page-11-0) [2014;](#page-11-0) Saxena et al. [2014a;](#page-12-0) Bajaj et al. [2015a,](#page-10-0) [b,](#page-11-0) [2016a,](#page-11-0) [b;](#page-11-0) Upadhyaya et al. [2015](#page-12-0)).

Transcription factors (TFs) controlling the expression of genes are vital for regulation of diverse biological processes including growth, development and cellular differentiation as well as responses to various environmental cues (Udvardi et al. [2007](#page-12-0); Century et al. [2008](#page-11-0); Libault et al. [2009;](#page-12-0) Kujur et al. [2013,](#page-11-0) [2014](#page-11-0); Saxena et al. [2014a,b](#page-12-0); Yu et al. [2010](#page-12-0)). The significant role of TF genes as prime regulators in governing diverse complex cellular processes like growth, development and stress responses as well as various domestication traits is welldemonstrated in legumes including chickpea (Udvardi et al. [2007;](#page-12-0) Century et al. [2008](#page-11-0); Libault et al. [2009](#page-12-0); Kujur et al. [2013](#page-11-0), [2014](#page-11-0); Saxena et al. [2014b](#page-12-0); Yu et al. [2010](#page-12-0)). In chickpea, the draft whole genomes as well as genome/transcriptome resequencing for diverse growth/development and stressimposed tissues/stages of multiple cultivated (desi and kabuli) and wild Cicer accessions are underway (Garg et al. [2011,](#page-11-0)

[2014](#page-11-0); Gaur et al. [2012](#page-11-0), [2015;](#page-11-0) Hiremath et al. [2012](#page-11-0); Jain et al. [2013](#page-11-0); Singh et al. [2013](#page-12-0); Varshney et al. [2013c](#page-12-0); Deokar et al. [2014](#page-11-0); Kudapa et al. [2014](#page-11-0); Pradhan et al. [2014](#page-12-0); Parween et al. [2015;](#page-12-0) Gupta et al. [2016\)](#page-11-0). These currently serve as a useful genomic/transcriptomic resource for genome-wide identification aside structural/functional annotation of the TF-encoding genes in chickpea. Consequently, these exertions assisted in detection of diverse differentially expressed TF genes including tissue- and stage-specific TFs governing manifold stress and developmental responses in chickpea accessions with contrasting agronomic traits. Besides, these efforts are found much expedient to develop numerous genic SSR and SNP markers representing various coding and regulatory sequence components of TF genes (especially differentially expressed TFs) annotated from chickpea genome (Garg et al. [2011,](#page-11-0) [2014](#page-11-0); Gaur et al. [2012](#page-11-0), [2015](#page-11-0); Hiremath et al. [2012;](#page-11-0) Jain et al. [2013;](#page-11-0) Singh et al. [2013](#page-12-0); Varshney et al. [2013c;](#page-12-0) Deokar et al. [2014](#page-11-0); Kudapa et al. [2014](#page-11-0); Pradhan et al. [2014;](#page-12-0) Parween et al. [2015;](#page-12-0) Gupta et al. [2016](#page-11-0)). These TF gene-based markers thus have functional significance in establishing marker-trait linkages for driving genetic enhancement of chickpea. Essentially, this will expedite the identification and targeted mapping of potential TF genes underlying major QTLs associated with vital agronomic traits by employing QTL and association mapping as well as diverse integrated genomics-assisted breeding strategies in chickpea (Kujur et al. [2013,](#page-11-0) [2014;](#page-11-0) Saxena et al. [2014a;](#page-12-0) Upadhyaya et al. [2015](#page-12-0); Bajaj et al. [2016a](#page-11-0), [b](#page-11-0)). These informative genic markers, therefore, have been extensively deployed in various high-throughput genetic analysis and marker-aided genetic enhancement of chickpea (Kujur et al. [2013,](#page-11-0) [2014;](#page-11-0) Saxena et al. [2014a](#page-12-0), [b;](#page-12-0) Bajaj et al. [2015a,](#page-10-0) [b,](#page-11-0) [2016b](#page-11-0); Upadhyaya et al. [2015\)](#page-12-0).

Based on aforesaid perspectives, the functional significance of TF gene-derived markers in rapid genetic dissection of complex quantitative traits is apparent in chickpea. To accomplish these, large-scale genome-wide discovery and highthroughput genotyping of especially the TF gene-based markers (SNPs) among phenotypically well-characterized desi and kabuli core/mini-core germplasm accessions (association panel) is essential. Henceforth, this will be useful to delineate functionally relevant natural alleles of the TF genes governing diverse major pod and seed yield component traits through genetic association analysis in chickpea. Moreover, outcomes obtained from these exertions will essentially enrich our deep understanding on trait inheritance pattern and complex genetic architecture of quantitative yield traits for driving genomics-assisted crop improvement to develop high pod and seed yielding desi and kabuli cultivars of chickpea.

In view of above prospects, the current study was undertaken with a prime objective to delineate functionally relevant natural SNP allelic variants of TF genes regulating diverse major pod and seed yield component traits by deploying high-resolution association mapping in chickpea.

Materials and Methods

Phenotyping of a Constituted Association Panel

For mining and high-throughput genotyping of TF gene-based SNPs for genetic association analysis, a trait-specific association panel comprising of 326 diverse desi (206) and kabuli (120) accessions were selected from the available chickpea core/mini-core germplasm collections following Kujur et al. [\(2014,](#page-11-0) [2015a](#page-11-0), [b](#page-11-0)). These accessions were grown in the field as per alpha-lattice design with two replications during crop season for two consecutive years (2012 and 2013) at two diverse geographical locations [National Institute of Plant Genome Research (NIPGR), New Delhi; latitude/longitude 28.4° N/77.1° E and International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), Patancheru, Hyderabad; 17.1° N/78.9° E)] of India. These natural germplasm accessions were further phenotyped for six major pod and seed yieldcontributing quantitative traits including flowering time, plant height, branch number per plant, pod number per plant, seed number per plant and 100-seed weight in chickpea.

Aside field phenotyping, greenhouse trail was conducted under both long (16 h light/8 h dark) and short (10 h light/14 h dark) day conditions at 22 ± 2 °C for evaluating the flowering time trait inheritance pattern in 326 desi and kabuli chickpea germplasm accessions. All the aforesaid pod and seed yield component traits were measured by estimating the average value of 10–15 representative plants selected from each germplasm accession. The days to 50% flowering time (DF) of each germplasm accession was determined by counting the number of days from sowing (first irrigation) to the stage when 50% of their plants have begun to flower in chickpea. The plant height (PH) was estimated by measuring the mean canopy height (cm) of each germplasm accession from soil surface at the time of their flower ending/pod setting initiation stage. The branch number (BN) of each germplasm accession was estimated by counting the average number of productive pods/seeds-bearing branches emerging from each plant at time of harvest. The pod number (PN) and seed number (SN) of each accession were measured by counting the average number of fully formed pods and seeds per plant at maturity, respectively. The 100-seed weight (SW) of each accession was estimated by measuring the average weight of 100-matured seeds (at 10% moisture content) of chickpea. Diverse statistical parameters including coefficient of variation (CV), frequency distribution, broad-sense heritability (H^2) and Pearson's correlation coefficient (r) of six major pod and seed yield component traits were measured using SPSSv17.0 ([http://www.spss.com/statistics\)](http://www.spss.com/statistics) and the methods of Bajaj et al. [\(2015a](#page-10-0)) and Kujur et al. [\(2015b,](#page-11-0) [c](#page-11-0)).

Targeted TF Gene Amplicon Resequencing-Based SNP Mining and Genotyping

A total of 16,376 SNPs physically mapped on eight chromosomes and unanchored scaffolds of *kabuli* chickpea genome were mined from the sequenced 92 diverse desi and kabuli chickpea germplasm accessions by employing a simple, robust and economical as well as simultaneous genome-wide SNP discovery and genotyping strategy, GBS (genotypingby-sequencing) assay (Elshire et al. [2011;](#page-11-0) Kujur et al. [2015a,](#page-11-0) [b\)](#page-11-0). From these, the genotyping as well as structural/ functional annotation information of SNPs especially derived from TF genes mapped on chromosomes and scaffolds of kabuli genome were acquired. The TF genes with SNPs selected from GBS data were further resequenced using the genomic DNA of 326 diverse desi and kabuli chickpea germplasm accessions (association panel) employing the multiplexed amplicon resequencing method (TruSeq Custom Amplicon v1.5) of Illumina MiSeq next-generation sequencer (Illumina, USA). The custom oligo probes targeting the CDS (coding DNA sequences)/exons, introns, 2 kb-URRs (upstream regulatory regions) and 2 kb-DRRs (downstream regulatory regions) of TF genes were designed and synthesized using Illumina Design Studio. All the probes producing amplicons with an average size of 500 bp per reaction were pooled into a custom amplicon tube. The template libraries were constituted after adding of sample-specific indices to each library by PCR using common primers. The uniquely tagged pooled amplicon libraries were normalized and generated clusters were sequenced by Illumina MiSeq platform. The visualization/mapping of sequenced TF gene amplicons and discovery of high-quality sequence variants among accessions were performed as per Saxena et al. ([2014a](#page-12-0)) and Malik et al. [\(2016](#page-12-0)). The pseudomolecules of kabuli chickpea genome (Varshney et al. [2013c](#page-12-0)) were used as a reference to map the high-quality gene amplicon sequence reads of each chickpea accession. Accordingly, the TF gene-derived highquality SNPs were detected among chickpea accessions as per Saxena et al. [\(2014a](#page-12-0)) and Kujur et al. ([2015a,](#page-11-0) [b,](#page-11-0) [c\)](#page-11-0).

Genetic Association Analysis

For association mapping, we selected the chickpea germplasm accessions exhibiting high-heritability (H^2) for six major yield component traits (DF, PH, BN, PN, SN and SW) across two environments (geographical regions) and experimental years. The environment and/or replication-wise six major yield trait phenotyping data obtained from each of the selected accession was used for estimating the mean trait value in an individual accession to be employed in association analysis. For candidate gene-based association mapping, the genotyping information of TF gene-derived SNPs (\geq 5% minor allele frequency) was correlated with multi-location/years replicated

field phenotyping data of six major pod and seed yield component traits (PH, BN, DF, PN, SN and SW) and population structure (Q), kinship (K) matrix and PCA (principal component analysis) (P) data of 326 desi and kabuli accessions belonging to an association panel of chickpea. The PCA and K matrix among accessions were measured using GAPIT (Lipka et al. [2012](#page-12-0)) and SPAGeDi 1.2 (Hardy and Vekemans [2002\)](#page-11-0), respectively. To perform association analysis, the CMLM (compressed mixed linear model) $(P + K, K \text{ and } Q + K)$ along with P3D [population parameters previously determined (Zhang et al. [2010](#page-12-0); Kang et al. [2010\)](#page-11-0)] interfaces of GAPIT were employed following Thudi et al. ([2014\)](#page-12-0); Kujur et al. [\(2015b\)](#page-11-0) and Kumar et al. [\(2015\)](#page-12-0). To determine the accuracy and robustness of TF gene-derived SNP marker-trait association, the quantile-quantile (Q-Q) plot-based false discovery rate (FDR cut-off ≤ 0.05) corrections (Benjamini and Hochberg [1995](#page-11-0)) for multiple comparisons between observed/expected $-log_{10}(P)$ values and adjusted P value threshold of significance were performed as per Kujur et al. [\(2015b\)](#page-11-0). The TF gene-derived SNP loci exhibiting significant association with DF, PH, BN, PN, SN and SW traits at a lowest FDR adjusted P values (threshold $P < 1 \times 10^{-6}$) and highest R^2 were identified in chickpea. The R^2 representing the magnitude of SNP marker-trait association was estimated based on model with the SNPs and adjusted P values following FDR-controlling method.

Results and Discussion

Discovery, Genotyping and Annotation of TF Gene-Derived SNPs for Genomics-Assisted Breeding Applications in Chickpea

A diverse array of TF-encoding genes is known to regulate multiple growth, development and yield-related traits in crop plants including chickpea (Udvardi et al. [2007](#page-12-0); Century et al. [2008;](#page-11-0) Libault et al. [2009](#page-12-0); Kujur et al. [2013,](#page-11-0) [2014;](#page-11-0) Saxena et al. [2014a,](#page-12-0) [b](#page-12-0); Yu et al. [2010\)](#page-12-0). Therefore, it would be interesting to determine the association potential of these TF genes in governing diverse major pod and seed yield component traits in chickpea. This can be primarily achieved by large-scale genotyping of novel synonymous and non-synonymous coding and non-coding intronic and regulatory SNP allelic variants discovered from the TF genes in a diverse set of phenotypically well-characterized natural desi and kabuli germplasm accessions (association panel) of chickpea. In the current investigation, considering the pros and cons of GBS assay, a high sequencing-depth coverage targeted gene amplicon resequencing strategy coupled with GBS assay was utilized to discover and genotype TF gene-based highquality SNPs uniformly across 326 desi and kabuli germplasm accessions (association panel) of chickpea. Primarily, the genotyping and sequencing of 92 accessions selected from 326 desi and kabuli chickpea accessions using GBS assay discovered 1029 high-quality SNPs in 736 TF genes annotated on kabuli reference genome (Table S1). A NGS-led GBS assay is proficient enough in fast large-scale discovery and high-throughput genotyping of SNPs simultaneously at a genome-wide scale for genomics-assisted breeding applications including association mapping to scan potential genes/ QTLs regulating vital agronomic traits in chickpea (Deokar et al. [2014;](#page-11-0) Bajaj et al. [2015a,](#page-10-0) [b](#page-11-0), [c](#page-11-0), [2016a,](#page-11-0) [b](#page-11-0); Jaganathan et al. [2015;](#page-11-0) Kujur et al. [2015a](#page-11-0), [b](#page-11-0), [c,](#page-11-0) [2016](#page-12-0); Upadhyaya et al. [2015\)](#page-12-0). However, reduced potential of GBS assay to generate nonerroneous and high-quality homozygous SNP genotyping information uniformly across accessions genotyped with a high genome/gene coverage is quite apparent. This could restrain the use of GBS assay in high-resolution association mapping study in a large chickpea genome with narrow genetic base. Henceforth, revalidation of GBS-derived SNP genotyping information as well as discovery and large-scale genotyping of novel SNPs covering the whole genome/gene regions at a high-resolution scale using numerous germplasm accessions are essential prior to deployment of these markers in genomics-assisted breeding applications and genetic enhancement of chickpea.

To accomplish these prime objectives, NGS-based amplicon resequencing strategy targeting TF genes (scanned previously by GBS assay) with high sequencing-depth coverage was employed in the present study for large-scale discovery and genotyping of SNPs in 326 desi and kabuli germplasm accessions belonging to an association panel of chickpea. Primarily, this strategy successfully validated all 1029 GBSderived SNPs mined from 736 chickpea TF genes (Table S1). Further, the kabuli reference genome-based GBS and targeted gene amplicon resequencing-led high-throughput SNP genotyping in 326 desi and kabuli accessions (association panel) altogether discovered 1611 SNPs from 736 TF genes (representing 30 TF gene family) with an average density of 2.2 SNPs per TF (Fig. [1](#page-4-0)). Maximum of 551 and 484 SNPs were discovered from the 257 and 221 TF-encoding genes especially belonging to Zinc finger and DUF (domain of unknown function) TF gene families, respectively (Fig. [1\)](#page-4-0). Of these, 1497 and 114 SNPs derived from 683 and 53 TF genes were physically mapped across eight chromosomes and unanchored scaffolds of kabuli genome, respectively (Fig. [2a](#page-5-0), b). All eight chickpea chromosomes contained maximum frequency of SNPs from the TF genes belonging to Zinc finger and ARF (auxin responsive factor) TF gene families (Fig. [2](#page-5-0)a). The comprehensive structural annotation of 1611 SNPs in 736 TF genes exhibited presence of highest and lowest proportion of SNPs in the exons/CDS (58.3%, 939 SNPs) and DRRs (0.4%, 7) of 422 and 3 TFs, respectively (Fig. [2](#page-5-0)c). The coding SNPs contained 529 (56.3%) synonymous and 410 (43.7%) non-synonymous (missense and nonsense SNPs) SNPs from

Fig. 1 Proportionate distribution of 1611 SNPs discovered from 736 TF genes representing 30 different TF gene families by their large-scale genotyping in a constituted association panel (326 desi and kabuli germplasm accessions) employing GBS and targeted gene amplicon resequencing assays. Digits within the round and square parentheses indicate the (number of TF gene-derived SNPs) and [number of TF genes with SNPs], respectively

204 and 218 TF genes, respectively (Fig. [2c](#page-5-0)). The abundance of non-synonymous SNPs derived from the TF genes representing Zinc finger and ARF TF gene families was evident. The novel TF gene-based natural SNP allelic variants discovered from a diverse set of desi and kabuli chickpea germplasm accessions can be employed for multiple genomics-assisted breeding applications in chickpea. Especially, this involves marker-trait association and quick identification of functionally relevant molecular tags (markers, TFs and alleles) as well as regulatory signatures governing diverse traits of agronomic importance for marker-aided genetic enhancement of chickpea.

Association Mapping Delineates Natural SNP Allelic Variants of TF Genes Regulating Major Pod and Seed Yield Component Traits in Chickpea

The SNP markers with their desirable inherent genetic attribute of abundant genomic distribution seems to be much promising in candidate gene-based association analysis for efficient quantitative dissection of diverse complex yield component traits in chickpea (Thudi et al. [2014](#page-12-0); Kujur et al. [2015b](#page-11-0); Upadhyaya et al. [2015](#page-12-0); Bajaj et al. [2016a](#page-11-0), [b\)](#page-11-0). In this context, a high-resolution candidate gene-based association analysis targeting TF-encoding genes was deployed in our study to identify the potential TFs-derived natural alleles governing six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW) in chickpea. For association mapping, the genotyping information of 1611 informative SNPs (differentiating the 326 desi and kabuli chickpea

germplasm accessions) discovered from 736 TF genes using genome-wide GBS and targeted gene amplicon resequencing assays were utilized. The use of 1611 TF gene-based SNPs primarily in neighbour-joining phylogenetic tree and population genetic structure construction as well as principal component analysis (PCA) classified 326 desi and kabuli chickpea germplasm accessions (association panel) into two distinct population groups- POP I (173 desi and 29 kabuli accessions) and POP II (33 desi and 91 kabuli accessions), which corresponded well with our previous studies (Kujur et al. [2014,](#page-11-0) [2015a,](#page-11-0) [b\)](#page-11-0) (Fig. S1). A wider degree of significant population divergence (mean F_{ST} 0.47 at $P < 0.001$) was observed between POP I and POP II, while F_{ST} -led population differentiation was maximum in POP I (0.39). All the accessions with their 81% inferred ancestry were derived from one of the model-based population and rest 19% contained admixed ancestry. Maximum and minimum admixed ancestry was observed in POP II and POP I, respectively. We observed a significant deviation from population assignment of 326 germplasm accessions (representing 58 diverse geographical regions of the world) based on desi and kabuli cultivarspecific classification which was more pronounced in cultivated kabuli accessions (29%) belonging to POP I. This is possibly due to a greater effect of geographical origin and adaptive environment rather than cultivar-types of accessions on their assignment to a specific population group. The multiple domestication events (evolutionary bottlenecks) followed by a complex breeding history coupled with strong adaptive selection pressure might have influenced their population group assignment resulting in numerous admixtures

Fig. 2 a Frequency distribution of SNPs mined from the TF genes including two most abundant class of TF gene families [zinc finger and ARF (auxin response factor)] physically mapped on eight chromosomes and unanchored scaffolds of kabuli chickpea genome. Digits above the bars indicate the total number of TF gene-derived SNPs mapped. Digits within the parentheses represent the number of TF genes with SNPs. **b** A Circos circular ideogram depicting the genomic distribution of 1611 TF gene-derived SNPs (represented by innermost circle) physically mapped on eight kabuli chromosomes. The outermost circles represent the eight kabuli chromosomes coded with diverse colours while the inner circles denote the relative distribution of TF genes with SNPs. c Relative

among accessions especially within POP II (Saxena et al. [2014b;](#page-12-0) Bajaj et al. [2015c](#page-11-0); Das et al. [2015](#page-11-0); Kujur et al. [2015a;](#page-11-0) Upadhyaya et al. [2015](#page-12-0)).

The comprehensive analysis of multi-location/year field phenotyping data revealed a normal frequency distribution along with a broader phenotypic variation for six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW) in a constituted association panel comprising of 326 *desi* and kabuli chickpea germplasm accessions (Table [1](#page-6-0)). Maximum CV was observed for SN (57.2–68.2%) followed by PN (52–54%) and minimum for PH (10.2–13.9%) (Table [1](#page-6-0)). Heritability (H^2) was estimated to be $\geq 80\%$ for all six yield traits with highest in case of SW (85–86%). This is agreed well with H^2 (80–88%) measured earlier for diverse pod and seed yield component traits using multiple

distribution and frequency of SNPs in different coding (synonymous and non-synonymous) and non-coding (intron, URR and DRR) sequence components of TF genes annotated from kabuli chickpea genome. The gene annotation of kabuli genome (Varshney et al. [2013c\)](#page-12-0) was considered as a reference to deduce the CDS (coding DNA sequence)/exons, introns and DRR (downstream regulatory region) sequence components of TF genes. Digits within the round and square parentheses indicate the (number of TF gene-derived SNPs) and [number of TF genes with SNPs], respectively belonging to each class of coding and non-coding regions of TF genes

geographical locations as well as several years field phenotyping data of chickpea germplasm accessions (Kujur et al. [2013,](#page-11-0) [2014,](#page-11-0) [2015a,](#page-11-0) [b](#page-11-0), [c,](#page-11-0) [2016](#page-12-0); Bajaj et al. [2015a,](#page-10-0) [b,](#page-11-0) [c,](#page-11-0) [2016a,](#page-11-0) [b;](#page-11-0) Upadhyaya et al. [2015\)](#page-12-0). A higher significant positive correlation between PN and SN ($r = 0.96$ with $P < 0.0001$) followed by SW and PH ($r = 0.61$ with $P < 0.0001$), whereas a lower negative correlation between SW and SN ($r = -0.65$ with $P < 0.0001$) followed by SN and PH ($r = -0.56$ with $P < 0.0001$) among 326 chickpea accessions based on Pearson's coefficient (r) was evident. This indicates that the 326 desi and kabuli germplasm accessions representing 58 diverse geographical regions of the world selected by us are rich in natural phenotypic diversity for all six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW). Therefore, the screened germplasm lines included in a

Table 1 Diverse statistical measures estimated for six major pod and seed yield-contributing quantitative traits phenotyped in an association panel based on multi-location/years replicated field evaluation

constituted association panel can serve as a useful genetic resource for mining novel functional allelic variants to expedite trait association mapping at a whole genome and/or gene-level in chickpea. This will essentially accelerate the delineation of functionally relevant potential genes and natural alleles regulating diverse pod and seed yield-contributing traits in domesticated desi and kabuli chickpea.

The CMLM and P3D/EMMAX-based candidate genebased association analysis at a FDR cut-off ≤ 0.05 detected 27 TF gene-derived SNPs that were significantly associated with six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW) at a $P \le 10^{-6}$ (Fig. [3,](#page-7-0) Table [2](#page-8-0)). All these 27 trait-associated SNPs were physically mapped on seven chromosomes (except chromosome 2) and unanchored scaffolds of kabuli genome (Table [2\)](#page-8-0). A maximum of 10 trait-associated TF gene-derived SNP loci were mapped on chromosome 3. Twenty-three and four of 27 trait-associated genic SNP loci were represented from diverse coding (20 synonymous and three non-synonymous SNPs) and noncoding (four intronic) sequence components of 16 genes, respectively (Table [2](#page-8-0)). The phenotypic variation for six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW) explained by 27 TF gene-derived maximum effect SNP loci varied from 10 to 23% R^2 (P 1.7 × 10⁻⁶ to 0.3 × 10⁻⁷) among 326 desi and kabuli chickpea germplasm accessions belonging to an association panel (Table [2](#page-8-0)). The combined phenotypic variation for six major pod and seed yield traits explained by all significant 27 TF gene-based SNP loci was 32%. Notably, six TF gene-derived SNPs identified to be associated commonly with PN and SN traits in a constituted association panel of chickpea. One SNP derived from a TF gene encoding SNF2 (sucrose non-fermenting 2) was associated with both BN and PH traits, whereas another SNP mined from a TF gene encoding B3-domain protein was associated both SW and PH traits in chickpea. This could be due to high phenotypic correlation as observed in our study between PN vs. SN and SW vs. PH traits in a constituted association panel. Therefore, complex genetic architecture of these PH, PN, SN and SW traits was apparent, which were dissected efficiently in this study through high-resolution association mapping involving functionally relevant informative natural SNP allelic variants discovered from the TF genes of chickpea.

Association Mapping Identifies TF Gene-Based Natural Allelic Variants Governing Flowering Time in Chickpea

Four SNPs derived from the diverse coding (two nonsynonymous SNPs) and intronic (two SNPs) sequence components of four TF genes exhibited significant association (13–23% R^2 with P 2.0 × 10⁻⁶ to 0.3 × 10⁻⁷) with DF trait (Fig. [3a](#page-7-0), Table [2](#page-8-0)). Four DF-associated TF gene-based SNPs were physically mapped on two *kabuli* chickpea chromosomes (3 and 4) with a maximum of 3 SNPs on the chromosome 3. The proportion of DF phenotypic variation explained by four SNP loci derived from four TF genes [encoding bZIP (basic leucine zipper), SBP (squamosal promoter binding protein), bHLH (basic helix-loop-helix) and Myb (myeloblastosis) TFs] in an association panel (326 desi and *kabuli* accessions) varied from 1[3](#page-7-0) to 23% R^2 (Fig. 3a, Table [2\)](#page-8-0). All four significant SNP loci in combination explained 29% DF phenotypic variation. One intronic SNP (T/A) in a bZIP TF gene and another non-synonymous SNP (A/T) causing amino acid substitutions from cysteine (TGT) to serine (AGT) in a SBP TF gene (20–23% R^2 with P 0.3– 1.1 × 10−⁷) exhibited strong association with DF trait in chickpea (Fig. [3](#page-7-0)a, Table [2\)](#page-8-0).

Fig. 3 GWAS-led Manhattan plot depicting significant P values associated with six major pod and seed yield component traits including DF (a), PH (b), BN (c), PN/SN (d) and SW (e) using 1611 TF genederived SNPs in chickpea. The genomic distribution of SNPs physically mapped on eight chromosomes and unanchored scaffolds of kabuli

Association Mapping Identifies TF Gene-Based Natural Allelic Variants Governing Plant Height in Chickpea

Four non-synonymous SNPs derived from the diverse coding sequence components of four TF genes revealing significant association (10–20% R^2 with P 2.3 × 10⁻⁶ to 1.2×10^{-7}) with PH trait were detected (Fig. 3b, Table [2](#page-8-0)). The PH-associated four TF gene-based SNPs were physically mapped on two kabuli chickpea chromosomes (6 and 7) with a maximum of 2 SNPs on the chromosome 6. The proportion of PH phenotypic variation explained by four SNP loci derived from four TF genes [encoding SNF2, WD40 (Trp-Asp 40), B3 and Tify-domain proteins] in an association panel varied from 10 to 20% R^2 (Fig. 3b, Table [2\)](#page-8-0). All significant four SNP loci in combination explained 25% PH phenotypic variation. Strong association of one nonsynonymous SNP (A/C) causing amino acid substitutions from histidine (CAC) to proline (CCC) in a WD40-domain protein-coding TF gene $(20\% \; R^2 \; \text{with} \;$ P 1.2 \times 10⁻⁷) with PH trait was observed in chickpea (Fig. 3b, Table [2](#page-8-0)).

genome are denoted by the x-axis. The y-axis indicates the $-log_{10} (P)$ value for significant association of SNP loci with six major yield traits. The SNPs revealing significant association with target yield traits at a cutoff P value $\leq 1 \times 10^{-6}$ are demarcated with a *dotted line*

Association Mapping Identifies TF Gene-Based Natural Allelic Variants Governing Branch Number in Chickpea

Six SNPs derived from the diverse coding (five nonsynonymous SNPs) and intronic (one SNP) sequence components of six TF genes exhibited significant association (12–20% R^2 with P 2.4 × 10⁻⁶ to 0.7 × 10⁻⁷) with BN trait (Fig. 3c, Table [2\)](#page-8-0). Six BN-associated TF gene-based SNPs were physically mapped on five kabuli chickpea chromosomes (1, 3, 4, 5 and 7) with a maximum of 2 SNPs on the chromosome 5. The proportion of DF phenotypic variation explained by six SNP loci derived from six TF genes [encoding GRAS [Gibberellic acid insensitive (GAI)- Repressor of GAI (RGA)-SCARECROW (SCR)], RINGtype Zinc finger, DUF827 (domain of unknown function 827), SNF2, C2H2-Zinc finger and SANT [switching-defective protein 3 (Swi3)-adaptor 2 (Ada2)-nuclear receptor corepressor (N-CoR), transcription factor (TF)IIIB]-domain proteins] in an association panel varied from 12 to 20% R^2 (Fig. 3c, Table [2\)](#page-8-0). All significant six SNP loci in combination explained 26% BN phenotypic variation. Two nonsynonymous SNPs- (C/A) and (C/T) causing amino acid

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^aValidated by previous studies (Kujur et al. [2013](#page-11-0), [2015b](#page-11-0), [c](#page-11-0); Bajaj et al. [2015a](#page-10-0), [b](#page-11-0), [2016a](#page-11-0), b; Upadhyaya et al. [2015](#page-12-0)) using an integrated genomic strategy

substitutions from alanine (GCA) to serine (TCA) and leucine (CTT) to phenylalanine (TTT) in GRAS and SNF2 TF genes (20% R^2 with P 0.7–1.0 × 10⁻⁷), respectively, revealed strong association with BN trait in chickpea (Fig. [3c](#page-7-0), Table [2](#page-8-0)).

Association Mapping Identifies TF Gene-Based Natural Allelic Variants Governing Pod and Seed Number in Chickpea

Six SNPs derived from the diverse coding (four nonsynonymous and one synonymous SNPs) and intronic (one SNP) sequence components of six TF genes revealed significant association (10–21% R^2 with P 2.2 × 10⁻⁶ to 0.3 × 10⁻⁷) with both PN and SN traits (Fig. [3d](#page-7-0), Table [2\)](#page-8-0). Six PN/SNassociated TF gene-based SNPs were physically mapped on three kabuli chickpea chromosomes (3, 5 and 7) with a maximum of 4 SNPs on the chromosome 3. The proportion of PN and SN phenotypic variation explained by six SNP loci derived from six TF genes [encoding DUF3437, LOB (lateral organ boundaries)-domain protein, C2H2-Zinc finger, WD40, ZF (zinc finger)-HD (homeodomain) homeobox protein and homeobox TFs] in an association panel varied from 10 to 21% (Fig. [3d](#page-7-0), Table [2\)](#page-8-0). All significant six SNP loci in combination explained 25% PN and SN phenotypic variation. One nonsynonymous SNP (A/G) causing amino acid substitutions from asparagine (AAC) to aspartic acid (GAC) in a LOBdomain protein-coding TF gene (21% R^2 with P 0.3 × 10⁻⁷) exhibited strong association with PN and SN traits in chickpea (Fig. [3](#page-7-0)d, Table [2\)](#page-8-0).

Association Mapping Identifies TF Gene-Based Natural Allelic Variants Governing Seed Weight in Chickpea

Seven SNPs derived from the diverse coding (five nonsynonymous and two synonymous SNPs) sequence components of seven TF genes revealed significant association (10– 20% R^2 with P 1.7 × 10⁻⁶ to 0.8 × 10⁻⁷) with SW trait (Fig. [3](#page-7-0)e, Table [2\)](#page-8-0). Six SW-associated TF gene-based SNPs were physically mapped on four kabuli chickpea chromosomes (1, 3, 6 and 8) with a maximum of 2 SNPs each on the chromosomes 1 and 3. The proportion of SW phenotypic variation explained by seven SNP loci derived from seven TF genes [encoding bZIP, SBP, Jumonji, RING-type Zinc finger, Med12 (mediator complex), B3 and WRKY-domain protein] in an association panel varied from 10 to 20% (Fig. [3e](#page-7-0), Table [2](#page-8-0)). All significant seven SNP loci in combination explained 30% SW phenotypic variation. One non-synonymous SNP (G/T) causing amino acid substitutions from threonine (ACG) to lysine (AAG) in a bZIP TF gene (20% R^2 with P 0.8 × 10⁻⁷) exhibited strong association with SW trait in chickpea (Fig. [3e](#page-7-0), Table [2](#page-8-0)).

Especially, the non-synonymous coding SNPs in the TF genes associated with six major pod and seed yield component traits (DF, PH, BN, PN, SN and SW) delineated in our study,

using high-resolution candidate gene-based association mapping, have functional significance for quantitative dissection of these complex yield traits in chickpea. This information can be useful for establishing rapid marker-trait linkages and efficient identification of potential TFs and natural allelic variants governing diverse pod and seed yield component traits in chickpea. Among six major yield trait-associated 27 TF gene-derived SNPs, merely two, four and three TFs-based SNP allelic variants governing SW, BN and DF traits, respectively delineated by us, have also been documented in our earlier studies for similar pod and yield component traits through integrating association analysis with QTL mapping, expression profiling and gene-based molecular haplotyping in chickpea (Kujur et al. [2013](#page-11-0), [2014](#page-11-0), [2015a](#page-11-0), [b](#page-11-0), [c](#page-11-0), [2016;](#page-12-0) Bajaj et al. 2015a, [b](#page-11-0), [c](#page-11-0), [2016a,](#page-11-0) [b](#page-11-0); Upadhyaya et al. [2015](#page-12-0)). The validation of these molecular tags in two of our independent studies infers the functional relevance and robustness of the identified TF gene-based natural SNP allelic variants in governing major pod and seed yield traits in chickpea. The six major pod and seed yield trait-associated 27 TF genes with SNPs delineated by association mapping in chickpea are reported to be involved in transcriptional regulation of growth, development and yield traits in multiple crop plants (Manning et al. [2006](#page-12-0); Agarwal et al. 2007, 2011; Udvardi et al. [2007;](#page-12-0) Nijhawan et al. [2008](#page-12-0); Libault et al. [2009](#page-12-0); Wang et al. [2011;](#page-12-0) Heang and Sassa [2012](#page-11-0); Martínez-Andújar et al. [2012;](#page-12-0) Jones and Vodkin [2013;](#page-11-0) Hudson and Hudson [2015;](#page-11-0) Wang et al. [2015;](#page-12-0) Wang and Wang [2015;](#page-12-0) Zhang et al. [2015](#page-12-0)). Therefore, functionally relevant novel as well as earlier documented TF gene-based molecular signatures (TFs and natural SNP alleles) regulating six major yield traits delineated in our study, once comprehensively validated and characterized, will essentially be employed for marker-assisted genetic enhancement to develop high pod and seed yielding cultivars of chickpea.

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