

# Identification of a non-redundant set of 202 in silico SSR markers and applicability of a select set in chickpea (*Cicer arietinum* L.)

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Abstract The paucity of sequence information flanking the simple sequence repeat (SSR) motifs identified especially in the transcript sequences has been limiting factor in the development of SSR markers for plant genome analysis as well as breeding applications. To overcome this and enhance the genic SSR marker repertoire in chickpea, the draft genome sequence of kabuli chickpea (CDC Frontier) and publicly available transcript sequences consisting of in silico identified SSR motifs were deployed in the present study. In this direction, the 300 bp sequence flanking the SSR motifs were retrieved by aligning 566 SSR containing transcripts of ICCV 2 available in public domain on the reference chickpea genome. A set of 202 novel genic SSRs were developed from a set of

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507 primer pairs designed, based on in silico amplification of single locus and having no similarity to the publicly available SSR markers. Further, 40 genic SSRs equally distributed on chickpea genome were validated on a select set of 44 chickpea genotypes (including 41 Cicer arietinum and 3 Cicer reticulatum), out of which 25 were reported to be polymorphic. The polymorphism information content (PIC) value of 25 polymorphic genic SSRs ranged from 0.11 to 0.77 and number of alleles varied from 2 to 9. Clear demarcation among founder lines of multi-parent advanced generation inter-cross (MAGIC) population developed at ICRISAT and near-isogenic nature of JG 11 and JG11 + demonstrates the usefulness of these markers in chickpea diversity analysis and breeding studies. Further, genic polymorphic SSRs reported between parental lines of 16 different mapping populations along with the novel SSRs can be deployed for trait mapping and breeding applications in chickpea.

**Keywords** Genic SSR · In silico · Transcriptome · *Kabuli · Desi* · Wild

### Introduction

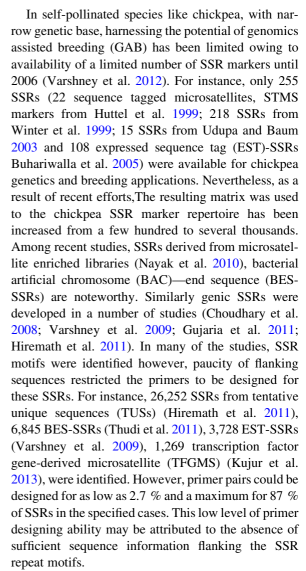
Chickpea (*Cicer arietinum* L.) is the second most important grain legume widely cultivated throughout tropical and sub-tropical regions of the world (Gaur et al. 2012). It is a self-pollinated crop with basic



chromosome number eight (2n = 2x = 16) and genome size of 738 Mb (Varshney et al. 2013b). Globally it is cultivated on 11.55 million ha with an annual production of 10.46 million tons and average productivity of 905 kg/ha (FAOSTAT data 2013). Based on the market class, two types of cultivated chickpeas viz. *desi* and *kabuli* are grown worldwide. *Desi* types are cultivated mainly in Indian subcontinent while *kabuli* types are grown in the Mediterranean region including Southern Europe, Western Asia and Northern Africa (Moreno and Cubero 1978).

Chickpea besides being rich and important source of protein, vitamins and essential minerals for human diet also enhances soil fertility by fixing atmospheric nitrogen through symbiotic association. Chickpea production is affected by several abiotic stresses like drought, heat, salinity and biotic stresses like Fusarium wilt, Ascochyta blight, Botrytis grey mould and dry root rot. Despite its economic importance, the climate changing scenarios in recent times are further aggravating impact of production constraints which kept the average productivity less than one ton per hectare for last six decades (Thudi et al. 2014). Nevertheless, the recent advances in genomics coupled with the availability of the genome sequence for several important crop plants (Michael and Jackson 2013) including chickpea (Varshney et al. 2013b) provide an opportunity to translate the knowledge to breed superior variety with enhanced resistance/tolerance to biotic and abiotic stresses in several crop species including chickpea (Gaur et al. 2012; Varshney and Tuberosa 2013a, b).

Molecular markers have become indispensable tools in genomics-assisted breeding (Varshney et al. 2007) for crop improvement. Among several kinds of molecular markers, the simple sequence repeat (SSR) markers are considered as the markers of choice owing to high reproducibility and co-dominant nature (Gupta and Varshney 2000). Further, genic SSRs (derived from cDNA/expressed sequence tags (ESTs) are, more robust for gene discovery and to study variation in transcribed regions and genes of known-function, very useful in plant genome analysis and crop improvement (Varshney et al. 2005, 2007). In addition, genic SSRs derived from ESTs with homology to candidate genes are good targets for genetic mapping and aligning genome linkage maps across distantly related species for comparative analysis (Holton et al. 2002).



In a recent study (Agarwal et al. 2012), transcript derived SSRs were identified, however, for 273 SSR motifs primers could not designed due to lack for sufficient flanking sequence information. Nevertheless, the constraint of designing the primers for SSR motifs without flanking sequence has been overcome in the present study because of the availability of reference chickpea genome sequence (Varshney et al. 2013b). Although in silico genome-wide survey is useful for the identification of large number of SSRs, identification of polymorphic markers is time consuming under laboratory conditions. In view of above, the present study demonstrates the utility of chickpea genome sequence for designing the primers for genic



SSRs. Furthermore, this study also validates a select set of SSRs in 44 chickpea genotypes.

### Materials and methods

### Plant material and DNA isolation

Forty-four chickpea genotypes representing the members of primary gene pool including 41 genotypes of *C. arietinum* (cultivated) and 3 genotypes of *C. reticulatum* (wild) were used in the present study. The origin, pedigree and salient features of the chickpea genotypes are listed in Table 1.

Genomic DNA from all 44 genotypes was extracted at ICRISAT from about 80 mg of fresh leaves collected from 15 to 20 day-old seedlings employing high-throughput DNA isolation method suggested by Cuc et al. (2008). DNA was normalized to 5 ng/µl after checking the quality on 0.8 % agarose gel.

# Primer designing

Previous transcriptome study by Agarwal et al. (2012) on *kabuli* cultivar of chickpea, ICCV 2 identified 43,389 transcripts which contained 566 transcripts having in silico identified genic polymorphic SSRs (here on regarded as in silico genic SSRs). For the present study the above transcripts available in the public domain (CTDB v 1.0, http://www.nipgr.res.in/ctdb.html) were utilized for designing primers and further validated a few of these markers on a set of 44 chickpea genotypes mentioned above.

The genomic loci of the in silico genic SSRs were identified by aligning the transcripts containing these SSRs on the publicly available chickpea genome (Varshney et al. 2013b). SSR motifs showing precise alignment with the genome with at least a 300 bp flanking sequences were selected. The SSR motifs along with respective flanking sequences were then used for designing primers employing batch Primer3web version 4.0.0 (http://probes.pw.usda.gov/ batchprimer3/). In order to remove the redundancy, the designed primers were subjected to Blast against publicly available SSR markers (Winter et al. 1999; Lichtenzveig et al. 2005; Gaur et al. 2011; Sethy et al. 2006; Buhariwalla et al. 2005; Choudhary et al. 2006, 2008; Varshney et al. 2009; Nayak et al. 2010; Hiremath et al. 2011; Thudi et al. 2011; Kujur et al. 2013). The primers showing hits with an expected product size and covering the SSR motif were considered to be redundant.

# Polymerase chain reaction (PCR)

For amplification of SSRs, PCR was carried out in a 10 µl reaction volume containing 10 ng DNA, 10X PCR buffer, 2.5 mM MgCl<sub>2</sub>, 2 mM dNTPs, 0.1 U *Taq* DNA polymerase (Sib-Enzyme, Novosibrisk, Russia), 2 pmol primer using a thermocycler (Bio-Rad, Berkeley, CA, USA). PCR amplification was carried out using touchdown cycles involving initial denaturation for 4 min at 94 °C; followed by 10 cycles of 94 °C for 20 s, 65 °C for 20 s and 72 °C for 30 s; (touchdown from 65 to 55 °C with 1 °C decrease in each cycle for 20 s)then by 35 cycles of 94 °C for 30 s; 55 °C for 50 s; and 72 °C for 30 s, a final extension of 20 min at 72 °C and left at 4 °C until further use.

# Capillary electrophoresis

Equal volumes of PCR products of four different markers labelled with different fluorescent dyes (FAM- VIC- NED- and PET) were pooled along with 7 μl of formamide (Applied Biosystems, CA, USA), 0.05 μl of the GeneScan<sup>TM</sup> 500 LIZ Size Standard (Applied Biosystems, CA, USA) and 2.95 μl of distilled water. The pooled PCR amplicons were denatured and size fractioned using capillary electrophoresis on ABI 3730 DNA Genetic Analyzer (Applied Biosystems, CA, USA) at ICRISAT. Allele calling was done using GeneMapper 4software (Applied Biosystems, CA, USA) by using the internal LIZ-500 size standard.

# Data analysis

To estimate the genetic diversity the amplified DNA fragments were either scored as presence (1) or absence (0) for each primer-genotype combination. The resulting matrix was used to compute Jaccard's similarity coefficients (Sneath and Sokal 1973) and UPGMA (Unweighted Pair Group Method with Arithmetic means) cluster analysis using computer package NTSYSpc (Numerical Taxonomy and Multivariate Analysis System) version 2.1 (Rohlf 2000) to determine the genetic relationship. Grouping of genotypes was further done by partitioning the



Table 1 Detailed information of 44 chickpea genotypes used in the present study

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Genotype	Pedigree	Origin	Type	Botanical name	Trait of importance
ICCV 05107	ICC 4958 $\times$ ICCV 92311	ICRISAT	Desi	C. arietinum	Ascochyta blight and Bortrytis grey mold resistant
ICCV 05530	Pant G-114 $\times$ ICC 3935	ICRISAT	Desi	C. arietinum	Ascochyta blight resistant
Pb 7	1	India	Desi	C. arietinum	Ascochyta blight susceptible
PI 489777	I	Turkey	Wild	C. reticulatum	Ascochyta blight susceptible
ICCV 04516	ICC $10169 \times NEC 138-2$	ICRISAT	Desi	C. arietinum	Ascochyta blight resistant
ICC 1496	P 1292-1	India	Desi	C. arietinum	Botrytis grey mould resistant
ICC 8261	NEC 2430	Turkey	Kabuli	C. arietinum	Drought tolerant
ICC 4958	JGC 1	India	Desi	C. arietinum	Drought tolerant (MAGIC parental line)
Arerti	ICC 14198 $\times$ FLIP 82-150C	Ethiopia	Desi	C. arietinum	Drought susceptible
ICC 1882	P 1506-4	India	Desi	C. arietinum	Drought susceptible
ICCV 10	P 1231 × P 1265	ICRISAT	Desi	C. arietinum	Drought tolerant (MAGIC parental line)
Pusa 256	$(JG 62 \times K 850) \times (L 550 \times H 208)$	India	Desi	C. arietinum	Fusarium wilt susceptible
Vijay	P 127 $\times$ Annigeri-1	India	Desi	C. arietinum	Fusarium wilt resistant
WR 315	I	India	Desi	C. arietinum	Fusariumwilt resistant
KWR 108	Selection from gerrmplasm (P 108)	India	Desi	C. arietinum	Fusarium wilt resistant
JG 62	Selection from gerrmplasm	India	Desi	C. arietinum	Fusarium wilt susceptible
JG 74	I	India	Desi	C. arietinum	Fusarium wilt resistant
Annigeri-1	I	India	Desi	C. arietinum	Fusarium wilt susceptible
C214	$(G24 \times IP58) \times G24$	India	Desi	C. arietinum	Fusarium wilt susceptible
ICC 3137	P 3659-2	Iran	Desi	C. arietinum	Helicoverpa susceptible
IG 72933	1	Turkey	Wild	C. reticulatum	Helicoverpa resistant
IG 72953	I	Turkey	Wild	C. reticulatum	Helicoverpa resistant
ICC 506	P 386 HR	India	Desi	C. arietinum	Helicoverpa resistant
ICCV 95423	$(ICC 7676 \times ICCC 32) \times [(ICCC 49)]$	ICRISAT	Kabuli	C. arietinum	High yielding
	$\times$ FLIP 82-1C) $\times$ ICCV 3]				
ICCV 97105	ICCV $10 \times GL$ 769	ICRISAT	Desi	C. arietinum	High yielding (MAGIC parental line)
JAKI 9218	$[(ICCC 37 \times GW5/7) \times ICCV 17]$	India	Desi	C. arietinum	High yielding (MAGIC parental line)
JG 11	(Phule G-5 $\times$ Narsinghpur bold) $\times$ ICCC 37	India	Desi	C. arietinum	High yielding (MAGIC parental line)



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Genotype	Pedigree	Origin	Type	Botanical name	Trait of importance
JG 11+	JG 11 × ICC 4958	ICRISAT	Desi	C. arietinum	High yielding with high root traits (MABC derived line)
JG 130	ICCC 42 $\times$ BG 256	India	Desi	C. arietinum	High yielding (MAGIC parental line)
JG 16	(ICCC 42 × ICCV 88506) $\times$ (KPG 59 × JG 74)	India	Desi	C. arietinum	High yielding (MAGIC parental line)
KAK 2	(ICCV 2 × Surutato 77) × ICC 7344	India	Kabuli	C. arietinum	High yielding
Pusa 362	BG $303 \times P$ 179	India	Desi	C. arietinum	High yielding and drought tolerant
Chefe	(ICCV 2 $\times$ Surutato 77) $\times$ ICC 7344	Ethiopia	Kabuli	C. arietinum	High yielding
DCP 92-3	ı	India	Desi	C. arietinum	High yielding
Ejere	X94TH71/FLIP87-59 C $\times$ UC15	Ethiopia	Desi	C. arietinum	High yielding
ICCC 37	$P 481 \times (JG 62 \times P 1630)$	ICRISAT	Desi	C. arietinum	High yielding and tolerant to dry root rot.
ICCV 00108	IG 9216 × ICCV 10	ICRISAT	Desi	C. arietinum	High yielding (MAGIC parental line)
ICCV 2	$[(K 850 \times GW 5/7) \times P 458] \times (L 550 \times Guamuchill)$	India	Kabuli	C. arietinum	Extra-short duration and salt sensitive
ICCV 03112	(ICCV 93001 $\times$ JAKI 9218) $\times$ BG 256	ICRISAT	Desi	C. arietinum	MARS population parent
ICCV 04112	ICCV 2 $\times$ PDG 84-16	ICRISAT	Desi	C. arietinum	MARS population parent
ICC 995	P 809	Mexico	Desi	C. arietinum	Parental line used to develop mapping population for protein content
ICC 1431	P 1260-1	India	Desi	C. arietinum	Salt tolerant
ICC 6263	NEC 142	Russia	Kabuli	C. arietinum	Salt sensitive genotype
ICC 283	P 223-1	India	Desi	C. arietinum	Tolerant to herbicide and drought



variance of the data-sets using principal component analysis (PCA). Polymorphism information content (PIC) was calculated using the following formula

$$PIC = 1 - \sum_{i} (Pi)^2$$

where Pi is the proportion of the population carrying the ith allele, calculated for each SSR locus.

### Results

Identification of novel genic SSRs

A total of 566 unique transcripts containing 623 in silico identified genic SSRs were downloaded from Chickpea Transcriptome Database (CTDB v 1.0, http://www.nipgr.res.in/ctdb.html). A total of 608 hits were obtained as a result of BlastN search of 566 unique transcripts against CDC Frontier genome of which 559 were unique hits. Further, the genome coordinates with a minimum of 300 bp flanking sequences on either side of genic SSR motifs were identified for 526 genic SSR motifs. These sequences

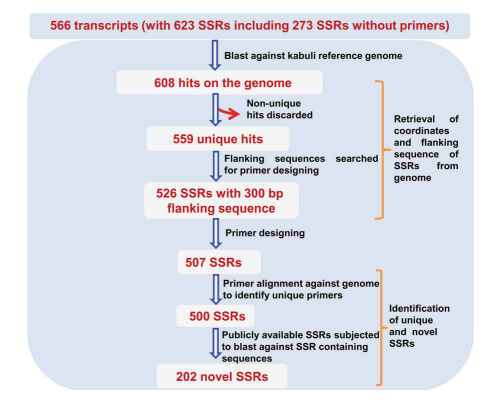
were extracted using customised perl script and then subjected to primer design using Batch Primer3. As a result primer pairs could be designed for 507 SSR motifs.

In order to determine the uniqueness and develop novel, robust genic SSR markers for applications in crop improvement the following in silico analysis was adopted:

- All 507 primers designed were subjected to BlastN analysis against the reference chickpea genome to identify primer pairs that amplify single genomic locus.
- 2. Primer pairs designed for SSR motifs with >10 dinucleotide repeats and >5 tri-nucleotide repeats were considered.
- 3. Primer pairs were blasted against available marker repertoire.

As a result of BlastN analysis, out of 507 primer pairs, 7 primer pairs showed multiple hits (>9 hits) on the reference genome and therefore 500 were considered for further studies. These 500 primer pairs included primers for 239 genic SSRs out of a total of 273 genic SSR motifs targeted, which were not

Fig. 1 Schematic workflow of identification of novel genic SSRs. The results of alignment of SSR containing transcripts on reference chickpea genome and identification of novel genic SSRs are depicted





reported in earlier study (Fig. 1). Although primer pairs were designed for all 273 genic SSR motifs targeted, 34 primer pairs did not flank the desired repeat motif. Hence, the primer pairs for 239 genic SSRs were tested for novelty by doing Blast against available marker repertoire. Flanking sequence of 300 bp on either side of SSR motif from chickpea genome were obtained and subjected to BlastN against the publicly available primers. Forward primers resulted in 142 hits while reverse primers gave 128 hits. However, there were only 32 unique hits that were shown by both forward and reverse primers. We therefore considered only 32 out of 239 SSRs as redundant. BAC-end sequences (46,270) reported by Thudi et al. (2011) were also used to further find out the redundant SSRs. These BAC-end sequences were also subjected to BlastN against the primers designed in this study and the hits were checked on the basis of alignment length including the SSR motif covered. Sequences showing hits with BAC-end sequences including the desired SSR motif including flanking region that could produce primers were discarded. We got 11 such genic SSRs containing sequences. However, only 5 out of these 11 were not a subset of 32 sequences which showed hits with the existing primers set. A total of only 37 primers (both forward and reverse primers) had unique hits with existing markers. Thus, primer pairs designed for 202 in silico genic SSRs were identified as novel. The details of all primer pairs designed including the genome coordinates of the repeat motifs are available in Supplementary Table 1.

### Validation and characterisation of SSRs

Among 202 novel in silico genic SSRs, 40 evenly distributed SSRs across eight pseudo molecules of chickpea were selected for validation. Of these 40 markers, 15 markers were monomorphic on a set of 44 chickpea genotypes that were genotyped. In total 95 alleles were produced by 25 genic polymorphic SSR markers. The number of alleles per locus ranged from 2 to 9 with an average of 3.8 alleles (Table 2). Polymorphism information content (PIC) ranged from 0.11 to 0.77 with an average of 0.42. The major allele frequency ranged from 0.34 to 0.93. The gene diversity ranged from 0.12 to 0.80.

A maximum of 9 (ICCV  $2 \times JG$  11 and ICCV  $2 \times JG$  62) and a minimum of 1 polymorphic marker

(C  $214 \times WR 315$ ) were identified for 16 different parental combinations used to generate segregating mapping populations for various traits (Supplementary Table 2).

Of the 25 SSR markers used in the study, 9 amplified genotype specific alleles for seven genotypes (Table 3). These SSRs could be used as molecular tags to identify the specific genotypes and these markers can be used to identify the seed lot for specific genotypes. The two wild chickpea species accessions (PI 489777 and IG 72953) could be identified using four different genic SSRs. Two genic SSRs (CakTSSR01526, CakTSSR00621) were specific to IG 72953, and the other two (CakTSSR01652, CakTSSR04407) were specific to PI 489777. Arerti (a desi variety of Ethiopian origin), Chefe (kabuli), ICC 1882 (a desi cultivar and parent of mapping population segregating for drought), ICCV 04112 (desi) and ICCV 97105 (desi) varieties were amplified for unique alleles using one SSR marker (CakTSSR00763, CakTSSR04052, CakTSSR00621, CakTSSR03039 and CakTSSR03970) for each genotype respectively.

### Genetic diversity analysis

In order to understand the genetic diversity among the 44 chickpea genotypes, a UPGMA (Unweighted Pair Group Method with Arithmetic Averages) dendrogram was constructed. Among 44 chickpea genotypes used in the present study, IG 72953 and PI 489777 had least similarity with rest of the genotypes analysed (Fig. 2). While remaining 42 genotyped were grouped into two major clusters namely Cluster I and Cluster II. Cluster I contained 18 genotypes and Cluster II contained 24 genotypes. The pairwise genetic similarity among the accessions are given in Supplementary Table 3.

Among 18 genotypes grouped in Cluster I, 12 were desi types (Annigeri-1, ICCV 04516, ICCV 05107, ICC 3137, ICCV 03112, ICC 506, ICCV 04112, ICC 995, ICCV 05530, Vijay, Arerti and Ejere) and 6 were kabuli (ICCV 95423, KAK 2, ICCV 2, Chefe and ICC 6263, ICC 8261) types. Further, among 24 genotypes clustered in Cluster II included wild type, IG 72933 (C. reticulatum accession) and the remaining 23 were cultivated desi type (C. arietinum accessions) (Fig. 2).

The diversity within eight *desi* founder parents (ICCV 10, ICCV 00108, JG 16, JAKI 9218, JG 130, ICC 4958, JG 11, ICCV 97105) used to develop first



Table 2 Summary of marker attributes of novel in silico genic pSSRs used in the study

SSR ID	Repeat motif	LGª	Primer sequences (5′–3′)	Fragment size (bp)	Major allele frequency	No. of alleles per locus	Gene diversity	PIC <sup>b</sup> value
CakTSSR00118	(AGA)7	7	F-GTTCACCACAGAATTCATCAT R-ACGATTTCCGATTCATCTTA	156–162	0.40	3	0.64	0.56
CakTSSR00392	(TC)11	5	F-CTTGCAAGTAAAAGTGTTTGG R-ATTTGGAAGGTTTATGGAGAA	167–169	0.81	2	0.29	0.25
CakTSSR00621	(CT)15	7	F-TTCTCTCTCTCTCTGGAGTT R-TCCCTCGCAACTAATATAACC	161–175	0.34	9	0.80	0.77
CakTSSR00729	(TA)14	8	F-TCGTAGACACGAAAATCTGTT R-AACCTTGATCAACATCTGGTA	164–172	0.43	5	0.68	0.62
CakTSSR00763	(GTT)8	3	F-TGCAACAGGGTTTAACTCTAA R-ACATAAGCACTC0CACATGAAC	167–170	0.86	3	0.24	0.22
CakTSSR01394	(TGA)15	1	F-GTTGGATTGAAGATTTCGAG R-TGCAACAACACTATTGAAGAA	174–183	0.81	4	0.31	0.30
CakTSSR01526	(AG)12	3	F-GATGCCATGTTACAACAACTT R-TAGTTCATTCAAAGCCTCTCA	154–164	0.47	5	0.59	0.51
CakTSSR01652	(ATTC)5	7	F-AAAATCATCTTTTGGACACCT R-ACCCATTTCTTCCTCTCTA	161–167	0.86	4	0.24	0.23
CakTSSR01996	(CT)17	1	F-AACCCATGAAAAATAACAGA R-GATGAAGTTCAACTGCCTATG	164–170	0.54	4	0.56	0.48
CakTSSR02234	(AG)10	1	F-TCATACACAGAAAAGCAAACA R-GGATGATATGGTATGGTTTGT	175–177	0.70	4	0.45	0.40
CakTSSR02460	(CAA)8	3	F-TGGGTCGTTTGTTAGTTCTTA R-ATACAAGTTTCTTCGGCTTTT	186–192	0.88	3	0.20	0.19
CakTSSR02655	(AAG)6	7	F-TTGTCAGAAGTTGATGGTTCT R-GAATCAAAATCTAGCAGCTCA	167–170	0.70	3	0.45	0.40
CakTSSR02667	(CTT)5	1	F-CTATGACAAAGTGGCATGATT R-ATCCACTTATCATTTGACGTG	174–177	0.45	3	0.64	0.57
CakTSSR03039	(ATG)6	2	F-ATGATATTGAAGGTGGTGATG R-ATTAGGACACCTTTGAAATCC	164–179	0.70	4	0.44	0.39
CakTSSR03099	(AG)10	4	F-GTTTGCGCTAAACAATATTAGA R-GTTTGCAGTGAAAAGAACAGT	169–181	0.63	5	0.50	0.43
CakTSSR03118	(AG)12	6	F-AATCCATGTCTTAATCTGCAA R-GAATTTGAAGAGCCCTAAGAG	165–167	0.93	2	0.12	0.11
CakTSSR03248	(CAC)8	4	F-CTAAAGAATGGAATTGGGATT R-CTCGTTTGTTTGCTCTATTGT	163–166	0.50	3	0.62	0.55
CakTSSR03332	(TCA)6	1	F-ACCCAACTTTGTACCTACCAT R-TAGAAACTTGTGGTGGAGATG	171–174	0.50	3	0.57	0.48
CakTSSR03899	(ATA)10	2	F-TTACACTATATGCGGTATCTGC R-ATCAAACTCAGTAGGCCAAA	148–151	0.81	3	0.30	0.27
CakTSSR03970	(TGA)9	6	F-TGAGGTTGAGAATTTGAGTGT R-CTCACTTCTCATCACCATCAT	160–172	0.47	4	0.59	0.51
CakTSSR04052	(ATTC)6	4	F-GCGTGAAGAAGAGAGAGAGAT R-TTGTTAGGCCTTAATCAATCA	163–167	0.56	4	0.52	0.42
CakTSSR04062	(TGG)6	4	F-CTCAAATATCTCTCCCAACCT R-GTCGTCGGAGAAATAGTCTTC	166–169	0.81	3	0.31	0.28
CakTSSR04214	(GAA)5	4	F-TAAAAGATGCTGCAAGAAGTG R-AAAGAGACAATGAAAGGGGTA	160–163	0.61	3	0.49	0.39
CakTSSR04255	(TTC)8	6	F-CTCATTCCATTATCCCCTTAC R-TGAGGATAGTGTTGAGAGGTG	177–180	0.59	3	0.53	0.44



Table 2 continued

SSR ID	Repeat motif	LGª	Primer sequences (5'-3')	Fragment size (bp)	Major allele frequency	No. of alleles per locus	Gene diversity	PIC <sup>b</sup> value
CakTSSR04407	(AG)6	1	F-TTGAACGATGATCGATAGAAG R-GGTGATCCAACCTAGAAGAAC	152–166	0.43	6	0.66	0.60

a Linkage group

**Table 3** SSR markers with their product sizes specific to seven chickpea genotypes

Genotype	Marker	Product size (bp)
Arerti	CakTSSR00763	170
Chefe	CakTSSR04052	169
ICC 1882	CakTSSR00621	169
ICCV 04112	CakTSSR03039	179
ICCV 97105	CakTSSR03970	172
IG 72953	CakTSSR01526	160
IG 72953	CakTSSR00621	151
PI 489777	CakTSSR01652	161
PI 489777	CakTSSR04407	152

multi-parent advanced generation inter-cross (MAGIC) population in chickpea was revealed using the markers developed in the present study. The MAGIC parents were uniformly distributed in cluster II and revealed considerable variation. The genetic similarity within MAGIC parents ranged from 0.40 (between ICCV 00108 and ICCV 10) to 0.80 (between JAKI 9218 and JG 130) (Supplementary Table 3).

The PCA analysis was performed to further validate the results reflected from dendrogram. PCA was utilized to derive a 2-dimensional scatter plot of individuals, such that the geometrical distances among individuals in the plot reflect the genetic distances among them with minimal distortion (Fig. 3). The 2D PCA plot broadly depicted two Clusters (Cluster I and II). However, 2 out of 3 wild chickpea species accessions, originated from Turkey, IG 72953 and PI 489777, did not fall into any of the two clusters and fall solitarily into two different places. Cluster I consisted of 26 genotypes, all of which are *desi* types and a wild accession (IG 72933). Cluster I also represented a leading variety JG 11 and it's MABC derived line JG 11 + for root trait QTL (Varshney et al. 2013a). It also represented all the 8 MAGIC founder parents (desi type) used to develop MAGIC population. The other Cluster (Cluster II) was represented by 16 genotypes, of which 10 genotypes are *desi* and 6 are of *kabuli* type. This group shows similarity to a part of Cluster I of dendrogram and consisted of two genotypes of Ethiopian origin namely, Ejere and Arerti.

### Discussion

Genic SSRs have been the marker of choice and are more robust as compared to non-genic SSRs in genetic and QTL mapping experiments. Genic SSRs are not prone to recombination between marker and trait associated with it as they lie within the gene of interest which also limits the identification of false positives in marker-assisted selection program (Gupta et al. 2010). Another important feature of the genic SSR markers is that, unlike genomic SSRs, they are transferable among related species and genera (Varshney et al. 2005). Recent advances in next generation sequencing technologies changed the sequencing scenarios which led to availability of draft genomes and transcriptome of several crop plants including chickpea (Varshney et al. 2013b; Varshney et al. 2009; Hiremath et al. 2011). The availability of chickpea genome made it possible to identify several thousands of SSRs in the genome using MISA (Thiel et al. 2003). However, classifying these into genic and genomic SSRs will be a difficult task. Nevertheless, several transcriptomic studies in recent past lead to identification of SSRs motifs in the transcriptome (Varshney et al. 2009; Hiremath et al. 2011; Agarwal et al. 2012). However, genic SSR markers could be developed for only a few SSR motifs. In the study done by Agarwal et al. (2012), although 623 genic SSRs were reported, primer pairs could be designed only for 350 SSRs and for the remaining 273 in silico genic SSRs primers could not be designed. In order to design the primers for SSR motifs lacking the flanking sequence in



<sup>&</sup>lt;sup>b</sup> Polymorphism Information Content

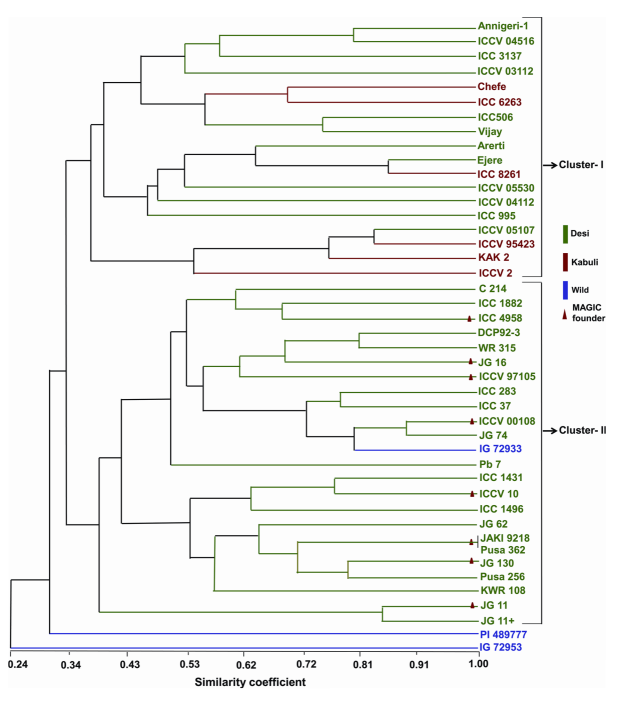


Fig. 2 Dendrogram showing hierarchical clustering (NTSYSpc) of 44 chickpea genotypes based on 95 alleles generated by 25 SSRs in chickpea

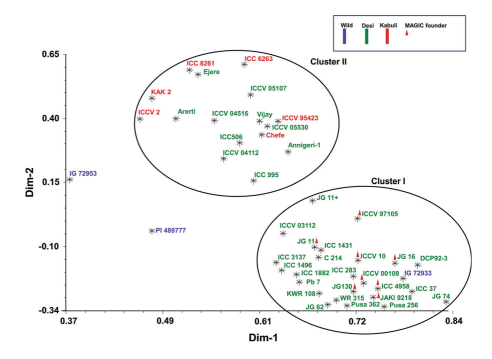
transcripts, 300 bp flanking sequence was obtained after aligning the transcripts on genome. The reason for considering such a long stretch of sequence for primer designing was to identify the novel SSRs) by

subjecting the publicly available primers to Blast against these flanking sequences.

In the present study, primers for the mentioned 273 genic SSR motifs were designed utilizing the available



Fig. 3 Two-dimensional plot of PCA analysis (NTSYSpc) showing genetic diversity of 44 genotypes using 95 alleles from 25 SSR loci



chickpea draft genome. Also, the novelty of these SSRs were confirmed. Further, a subset of uniformly distributed genic SSRs (40 markers) on the genome was validated for their usefulness in genetic diversity analysis. In silico identification of SSRs reduces the effort that goes in validation of number of markers for their polymorphism. These polymorphic markers are used to study the genetic diversity and relationship among the germplasm available, which will be helpful in chickpea improvement programs.

In this study, we report primer pairs for a set of 202 novel, non-redundant genic SSRs. The criterion adopted for removing the redundancy further avoids overestimation of the EST-SSRs in chickpea. We also evaluated and proved the authenticity of 25 in silico identified genic SSRs. Similar work was also reported in castor bean (Qiu et al. 2010), pepper (Kong et al. 2012) and date palm (Zhao et al. 2012), where the available EST sequences were used for identification of genic polymorphic SSRs followed by validation of some of these markers. In this study, 44 chickpea genotypes of kabuli, desi and wild type representing cultivated chickpea breeding lines, wild relatives and parents of mapping populations were exploited for diversity study using 25 genic SSR markers. In the present study we obtained PIC values in the range of 0.11–0.77 using only 25 SSRs in a set of 44 genotypes as compared to PIC values of 0.27 to 0.50, 0.37 to 0.91, 0.46 to 0.97 and 0.40 to 0.80 reported by Keneni et al. (2012), Sefera et al. (2011), Upadhyaya et al. (2008) and Choudhary et al. (2012) respectively using more number of genotypes with more number of markers. The polymorphic markers identified among parental genotypes of 16 different mapping populations segregating for different biotic (*Helicoverpa*, *Ascochyta* blight, *Fusarium* wilt) and abiotic stresses (drought etc.) can be used in marker assisted breeding programs.

Two of the reported markers (CakTSSR04062 and CakTSSR04214) were identified in the "QTL-hotspot" region located on CaLG04 (Varshney et al. 2014) which harbour QTLs for several drought tolerance related traits. The markers can be used in trait mapping and reducing the QTL region. As very few SSR markers are available for marker-assisted backcrossing (MABC; Varshney et al. 2012) and polymorphism of these markers in several genetic backgrounds is very low (Thudi et al. 2014), additional markers in this region will enable introgression of this region into different genic backgrounds. Other genic SSRs reported for the parental crosses can be useful in a marker-assisted selection program like background selection and trait mapping studies. Similar diversity studies were carried out by Keneni et al. (2012) in



Ethiopian chickpea accessions of different geographical origins using only 33 previously identified polymorphic SSR markers with a PIC value ranging from 0.27 to 0.50 and 3.36 bands per marker advocating the usefulness of those markers in germplasm characterization. SSR markers like CakTSSR00621 with high PIC value (0.77) and the markers signature alleles like CakTSSR01996 and CakTSSR03248 can be used to distinguish the accessions from each other signifying the relevance of these markers to generate specific genetic fingerprints for each genotype useful for genetic purity analysis.

High genetic distance between a wild and cultivated accession and a low genetic distance between two cultivated accessions was an expected outcome, for instance, molecular analysis revealed a high genetic distance (0.90) between parents IG 72953 (wild) and Annigeri-1 (parents of 2 different populations segregating for Helicoverpa resistance and drought avoidance root traits). Similar trend of genetic distance was observed in chickpea accession by Choudhary et al. (2012). Interestingly, the cluster analysis revealed that, Pusa 362 and JAKI 9218 genotypes are closely related (with almost 100 % similarity) with this set of markers followed by 90 % similarity (0.90) between ICCV 00108 and JG 74. Also, this study reconfirms that JG 11 + is a derived near isogenic line (NIL) of JG 11 with a genetic similarity of 89 %. This analysis also substantiates the diversity of selected chickpea MAGIC parents as revealed by dendrogram.

Cluster analysis separated all chickpea accessions into two major clusters with a wild accession IG 72953 not being part of either of the two clusters. PCA was in accordance with cluster analysis as IG 72953 out lied distantly with the other wild accession PI 489777 and not falling in either of the two clusters. The grouping of all the *kabuli* cultivars into Cluster II (Fig. 3) highlights the utility of these markers in discriminating kabuli genotypes with that of desi genotypes. Similarly, other *desi* accessions were intervened by breeding lines (desi) suggesting that breeding lines tend to cluster with kabuli and desi genotypes with the exception of a couple of desi genotypes grouped with the kabuli genotypes. Therefore, it can be concluded that kabuli and desi genotypes can be distinguished to a good extent exploiting SSR based diversity studies and this is in accordance with the findings of Upadhyaya et al. (2008), Sefera et al. (2011). However, the accessions originating from Turkey did not cluster together, which could be due to geographical separation. On the other hand, the accessions from Ethiopia were present in the same cluster despite having different pedigree and representing both *desi* (2) and *kabuli* (1) types.

The present study has resulted in identification of novel genic SSRs using the available chickpea genome. The strategy used here can also be deployed in other crops where there is scarcity of genic SSRs. The study has also avoided overestimation of chickpea genic SSRs by avoiding the duplicity with the already available SSRs in the public domain. Not only have these markers added to the existing repertoire of SSRs but also demonstrated their applicability in diversity studies, trait mapping, saturating genetic maps and in MABC programs for chickpea improvement.

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