

Mining of simple sequence repeats in the Genome of *Gentianaceae*

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ABSTRACT

Simple sequence repeats (SSRs) or short tandem repeats are short repeat motifs that show high level of length polymorphism due to insertion or deletion mutations of one or more repeat types. Here, we present the detection and abundance of microsatellites or SSRs in nucleotide sequences of *Gentianaceae* family. A total of 545 SSRs were mined in 4698 nucleotide sequences downloaded from the National Center for Biotechnology Information (NCBI). Among the SSR sequences, the frequency of repeat type was about 429 -mono repeats, 99 -di repeats, 15 -tri repeats, and 2 -hexa repeats. Mononucleotide repeats were found to be abundant repeat types, about 78%, followed by dinucleotide repeats (18.16%) among the SSR sequences. An attempt was made to design primer pairs for 545 identified SSRs but these were found only for 169 sequences.

Key words: *Gentianaceae*, nucleotide, simple sequence repeats

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INTRODUCTION

Gentianaceae, or the Gentian family, is a family of flowering plants of 87 genera and over 1650 species.^[1] Plants are usually rhizomatous. These are annuals or perennials, mostly upright though a few species lie on the ground and have upright branch tips. Leaves are opposite or whorled with entire edges and bases connately attached to the stem, mostly without a petiole. Flowers have four to five sepals, petals, and stamens, but only one pistil. Sepals and petals are fused at the base, with four to five free lobes above. Stamens alternate with the corolla lobes. Ovary is superior; fruit is a capsule. Stipules is absent. Plants usually accumulate bitter iridoid substances; bicollateral bundles are present. The fruits are dehiscent septicidal capsules splitting into two halves. The *Gentianaceae* contains many species with interesting phytochemical properties. They have been widely used in traditional medicine and also as constituents in bitters and similar concoctions. The family

consists of trees, shrubs, and herbs showing a wide range of colors and floral patterns.

Simple sequence repeats (SSRs),^[2] or microsatellites^[3] or short tandem repeats,^[4] are short (1–6 bp) repeat motifs that show a high level of length polymorphism due to insertion or deletion mutations of one or more repeat types.^[5] Studies suggest that both protein coding and noncoding regions of DNA sequences contain SSRs.^[6] SSRs present in coding sequences are less polymorphic than those in the genomic sequences. Moreover, different taxon varies in abundance of different types of SSRs and these are present in greater abundance in noncoding regions than coding SSRs.^[7] The SSRs are either developed conventionally^[8] or from sequence databases.^[9] PCR-based techniques such as AFLP and microsatellites or SSRs have also played important roles in plant DNA profiling. Primers are essential components of PCR-based systems as well as modern microarray systems which utilize appropriate probes for PCR amplification.^[10]

In genetics, a sequence motif is a nucleotide or amino acid sequence pattern that is widespread and is believed to have, a biological significance. When a sequence motif appears

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in the exon of a gene, it may encode the "structural motif" of a protein, that is, a stereotypical element of the overall structure of the protein. "Noncoding" sequences are not translated into proteins. Outside of gene exons, there exist regulatory sequence motifs and motifs within the "junk," such as satellite DNA.^[11] Robinson *et al.*^[12] developed a computer program to identify and design PCR primers for amplification of SSR loci based on available DNA sequence information. SSR primers have been designed using publicly available expressed sequence tags (ESTs) in barley,^[13] almond (*Prunus communis* Fritsch.), and peach (*P. persica* (L.) Batsch.),^[14] *T. aestivum*, and *O. Sativa*.^[15] These SSRs are useful as molecular markers because their development is inexpensive, they represent transcribed genes, and their putative function can often be deduced by a homology search.^[16] SSRs have been the backbone to creating molecular maps for a number of years.

The increasing number of genomic and expressed sequences in public databases provides a valuable source for bioinformatical data mining. However, there are a number of exciting application of these sequence data; used in comparative genome analysis – to trace the evolution among the related species, to study the genome structure and their gene functions. Comparative genome analysis requires the same sets of genes (i.e., cross-reference genes) to be mapped to chromosomes in the species compared. Thus, comparative maps with sets of EST-derived markers (i.e., cross-species markers) are essential for comparative genome analysis. Several studies have utilized publicly available ESTs to mine SSRs or microsatellites markers for plants,^[17–20] catfish,^[21] insects,^[22] animals,^[23] and human.^[24] The EST-derived SSR markers (EST-SSRs) have proved very useful for the construction of genetic and comparative maps.^[25] The software used here is MISA, a microsatellite identifying tool which has the advantage of detecting the mono- to decamer repeats and also compound repeats. But it has the disadvantage of inability to detect above decanucleotide repeats. Riju and Arunachalam,^[26] mined the SSRs in oil palm ESTs with five different software and have reported that MISA program has given maximum coverage of SSRs in both oil palm ESTs and Contigs.

PCR primer design in general

Understanding of primer properties is very important for primer design. The major aspects of primer properties include specificity, melting temperature (T_m), and intraprimer or interprimer homology. Primer specificity is mostly determined by the 3'-end sequences. It was reported that single internal mismatches had no significant effect on PCR product yield while the 3'-terminal mismatches, especially the A:A, A:G, G:A, and C:C mismatches, markedly reduced overall PCR product yield.^[27] Khabar *et al.*^[28] assessed the annealing specificity of primers in PCR

reactions under different annealing temperatures (35°C, 40°C, and 45°C) and found perfect matches between at least eight bases at the 3'-end of the 5'-primers and the target region, whereas mispriming occurred only toward the 5'-end. Therefore it is critical to include 8–10 unique bases at the 3'-end of the primer.

Ideally the primer has a T_m in the range of 50–65°C, random nucleotide composition with a 40–60% GC-content, and 18–30 bases long. The intraprimer or interprimer homology is kept as low as possible to avoid formation of hairpin structures or primer dimers (>3 bp complementarities between primers) which otherwise will interfere with annealing of primer to the DNA template.^[29]

ESTs, which represent the expressed part of genome, also serve as a source of SSRs.^[9] Detection of SSRs facilitates the development of SSR markers that are useful in the study of genetic variation, gene tagging, and linkage mapping,^[30] and are also useful across a number of related species.^[13] Microsatellites can be amplified for identification by the polymerase chain reaction (PCR) process, using the unique sequences of flanking regions as primers. Once the potentially useful microsatellites are determined (removing nonuseful ones such as those with random inserts within the repeat region), the flanking sequences can be used to design oligonucleotide primers which will amplify the specific microsatellite repeat in a PCR. Microsatellite loci are widely distributed throughout the genome and can be isolated from semidegraded DNA of older specimens, as all that is needed is a suitable substrate for amplification through PCR. Hence, the present study was to find out the distribution and abundance of SSRs for the development of markers and to annotate SSR-containing sequence in *Gentianaceae* family. Nucleotide database, which contains sequences of well-characterized genes as well as hundreds of thousands novel EST sequences, was retrieved to perform the analysis.

MATERIALS AND METHODS

Retrieval of nucleotide sequences and detection of SSRs

A total of 647 nucleotide sequences of *Gentianaceae* were downloaded from the NCBI (<http://www.ncbi.nlm.nih.gov/Nucleotide/?term=Gentianaceae>) and harvested for SSRs using a perl script. The minimum length of SSR was fixed at 14 bp according to the criteria used by Gupta *et al.*^[31] The SSRs were defined as 14-bp mononucleotide or dinucleotide repeats; 15-bp trinucleotide repeats; 16 tetranucleotide repeats; 20 pentanucleotide repeats; 18 hexanucleotide repeats. The poly A and poly T repeats were removed by using an inhouse developed perl script,

as these are not considered as SSRs due to their presence at 3'-end of mRNA/cDNA sequences.

Primer designing for SSRs

A pair of primer flanking each SSR was designed using FastPCR software available at www-genome.wi.mit.edu/cgi-bin/primer/primer3_www.cgi, which takes input according to user-defined conditions and pick primers according to these specified parameters. Default parameters of the FastPCR, viz, the optimum primer size of 20.0 (the range was 18–28), the optimum annealing temperature of 60.0 (the range was 57.0–63.0), and the range of % GC content of 44–60, were selected for primer designing.

Detection of SSR positions with respect to open reading frames

Open reading frames (ORFs) are predicted for all the SSR-containing sequences using ORF finder available at NCBI (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>) using standard genetic code. Sequence fragments with maximum length uninterrupted by stop codon were taken as the primary encoding segment (ORF) of the query sequences. In all the predicted ORFs, the relative positions of SSRs were detected, that is, whether the SSR was present within the ORF, in the 5' UTR untranslated region (UTR) or in the 3' UTR.

RESULTS

Screening of *Gentianaceae* sequences for SSRs

In the present study, 4698 nucleotide sequences of *Gentianaceae* available at NCBI (<http://www.ncbi.nlm.nih.gov>) were searched for SSRs with a minimum length of 18 bp. A total of 545 SSRs were detected from 2889 kb of data mined, excluding poly A and poly T. Depending upon the length of the repeat unit itself (1–6 bp), the lengths of the identified SSRs varied from 14 to 48 bp, respectively.

Frequencies of classified repeat types of *Gentianaceae*

From a number of 4698 sequences screened, only a subset of 461 sequences contained 545 SSRs, suggesting that merely 9.83% of sequences contained SSRs. The frequencies of SSRs with mono-, di-, tri-, tetra-, and hexanucleotide repeat units showed the frequent repeat type within the nucleotide sequences of *Gentiana* family that were found to be in mononucleotide (84.58%) followed by dinucleotide repeats (18.16%), trinucleotide (2.75%), and hexanucleotide (0.65%), respectively [Figure 1]. Whereas, no tetranucleotide and pentanucleotide repeat was detected during the analysis.

The observed frequency of different repeat types comprising the SSRs is presented in Figure 2a–d and summarized in Table 1. SSRs were comprised of four

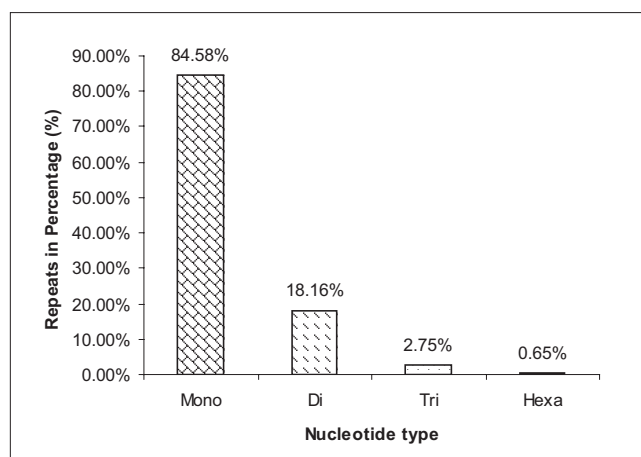


Figure 1: Frequency distribution of different repeat types identified in nucleotide sequences of *Gentianaceae*

different types of mononucleotide (A, T, C, and G), nine different types of dinucleotide (CA)_n, (TG)_n, (AC)_n, (GA)_n, (CT)_n, (TA)_n, (AT)_n, (GC)_n, (TC)_n, (AG)_n, (GT)_n repeats, seven different types of trinucleotide (GAG)_n, (ATG)_n, (CTT)_n, (TTA)_n, (CAA)_n, (AAC)_n, (ACA)_n repeats, and two types of hexanucleotide (CCACAC)_n, (GGTCAA)_n repeats.

Designing of primers for SSRs

Out of 545 SSRs detected, the primers could be designed only for 169 (31%) SSRs and the rest 376 (69%) sequences did not produce any acceptable primers. These 169 SSRs for which primers were designed include 133 mono-, 29 di-, 7 tri-, and no hexanucleotide repeats. The details of the accession numbers of nucleotide sequences of *Gentiana*, repeat motif of SSRs for which primer were designed, primer sequences, GC%, product size, and annealing temperature are given in Table 2.

Prediction of ORF in SSR-containing sequences

An attempt was made to predict the ORFs in SSR-containing sequences using ORF finder. Out of the 545 SSRs identified, the positions of 359 SSRs with respect to ORF were determined, while for the remaining 186 SSR-containing sequences, no ORF were predicted. Of these 359 SSRs, a large number of 161 (44.84%) were present in the 5' untranslated region, 129 (35.93%) SSRs occurred within ORF, and the remaining 69 (19.22%) occurred in the 3' untranslated region.

DISCUSSION

In the present study, a large number of nucleotide sequences (4698) of *Gentiana* retrieved from NCBI were mined for SSRs. In the sequences that were mined the SSRs were characterized, and a subset of these SSRs was

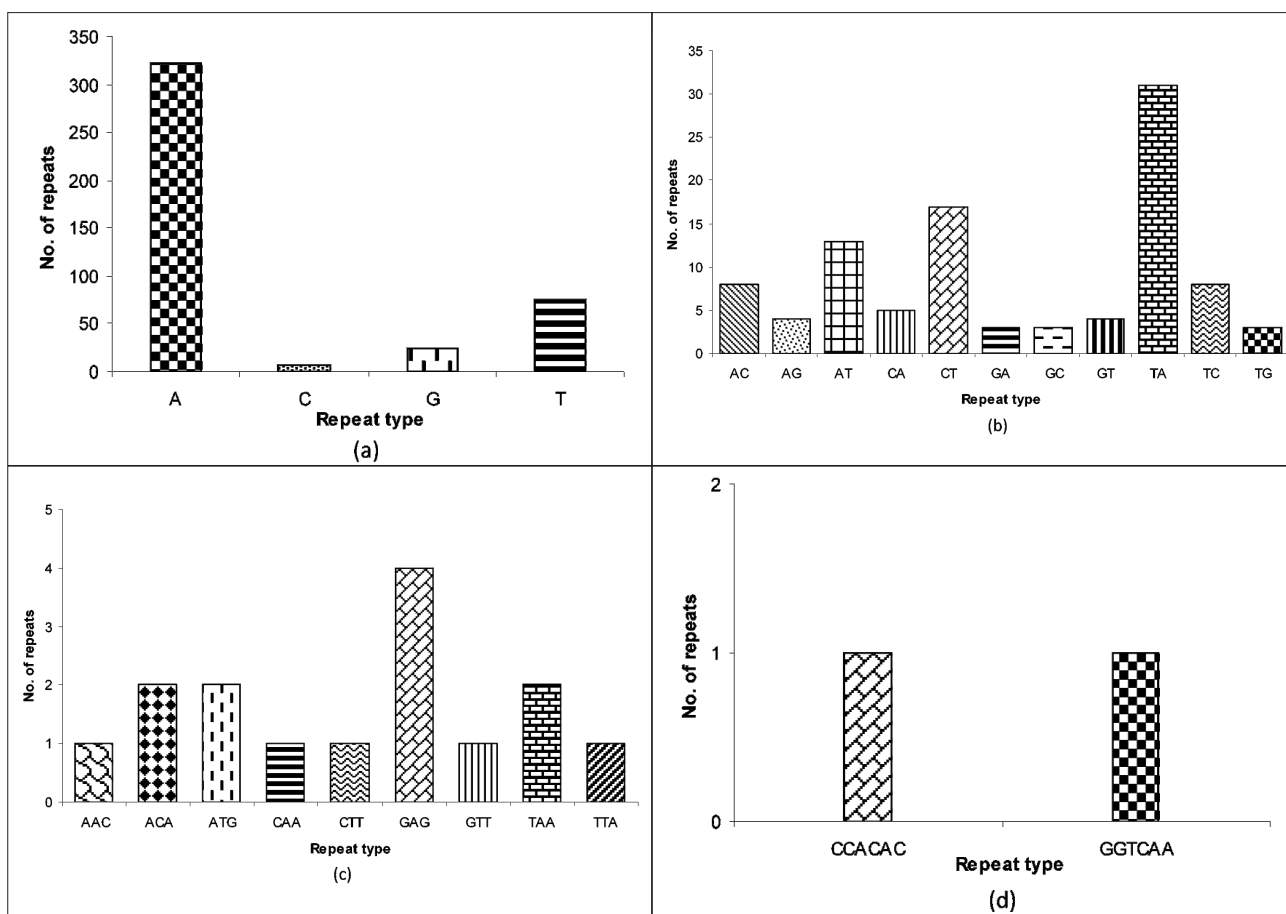


Figure 2: Frequency distribution of (a) mono-, (b) di-, (c) tri-, and (d) hexanucleotide repeat motifs in the genome of *Gentianaceae*

Table 1: Summary of in silico mining of Nucleotide sequences of *Gentianaceae*

Parameters	Values
Total number of sequences searched	4698
Total number of SSRs after removing poly A and poly T	545
Total size of examined sequences (bp)	2289303
Total number of sequences containing single SSRs	429
Number of sequences containing two SSRs	99
Number of sequences containing three SSRs	15
Number of sequences containing six SSRs	2
Number of sequences containing more than one SSR	57
Number of SSRs present in compound formation	47
Repeat type	
Mononucleotide	429 (84.58)
Dinucleotide	99 (18.16)
Trinucleotide	15(2.75)
Hexanucleotide	2 (0.65)

Data in parentheses is the percentage value of the repeat

used for designing the markers. A total of 545 SSRs was detected and this was in accordance to the findings of^[32] who reported that the abundance of different repeats varied broadly depending upon the species.

Microsatellites or SSRs are stretches of DNA containing tandem repeats of di-, tri-, tetra-, and above nucleotide

units ubiquitously distributed throughout the eukaryotic genome. They are found to be abundant in plant genomes and are thought to be the major sources of genetic variation in quantitative traits. The abundance of the different repeat motifs (1–6 bp) in the SSRs as detected in *Gentiana* family during the present study was variable so that the SSRs with different repeat motifs were not evenly distributed. The SSRs with dinucleotide repeats (18.16%) were abundant. This is in agreement with the results of earlier studies on *Arabidopsis* in which the dinucleotide repeats were also found to be abundant,^[33] perhaps because the genomic sequences of this species may include SSRs in noncoding regions too. The smaller repeat motifs were found to be predominant among SSRs identified and as the length of repeat unit increases, their occurrence decreases. We excluded poly A and poly T repeats due to which their number is under-represented. The abundance of trinucleotide SSRs may be attributed to the absence of frame shift mutations due to variation in trinucleotide repeats.^[34]

Molecular genetic markers can be used to examine a group of individuals or populations to estimate various diversity measures and genetic distances, intergenetic structure and

Table 2: Synthesis of primer

Accession number	Motif	Forward Primer		Reverse primer		Annealing temperature (C)	Product size (bp)
		Primer	GC %	Primer	GC %		
gi 261865195 gb GQ864096.1	(TA)7(AT)7	GGCAGAACTCAACGGGAAGC	59.1	GAGGCTAAAGGGTCTCGGA	58.8	57.9	709
gi 169798749 gb EU370947.1	(A)13	CGAGGCTGTAAGTGCCTGC	60	GACGGTGAGAGGCTGTGTATG	57.1	56.85	415
gi 24494572 gb EF371451.1	(TG)17	GAGGCTGTAAGTGCCTGC	50	GACGGTTAGAGGCTGTGTAGG	54.5	54.25	414
gi 124494568 gb ef371447.1	(AG)7(TG)8 (GA)12 (CA)29(GA)21(AG)6 (CT)21	GTCTGAGTGAGGGAGCCATC	60	GTGATGCTGTGTCTCCCAAGAG	54.5	56.8	80
gi 112293600 gb dq822583.1	(A)21	GGGATGGCACTCACCTACAGC	61.9	GGCATAGTCTGAGTTCCTCCAC	55	54.45	123
gi 112293594 gb dq822577.1	(A)21	GCGTTCTCCGACTGCCGAG	65	GCAGAGCATAGTGGACGC	61.1	62	656
gi 156787489 gb ef569230.2	(A)16	CTGTGTAGCTGAGTTGC	55	GGCTGAACCCAGGGACAACC	65	59	461
gi 89199712 gb dq398766.1	(A)10	GATAGTGTTCAGAGGACG	50	GGTTGACACTTTACAGCAGC	50	57	760
gi 89199711 gb dq398765.1	(A)16	GCGATGTGTTCAAGGACCG	55	GTGAACGCCTATCCAGTG	55.6	59	676
gi 89199709 gb dq398763.1	(A)16	CGCTGTTCCGCTCACGCTTC	65	GTTGACACTTTACAGCCCT	50	63	409
gi 89199708 gb dq398762.1	(A)12	GAGGCTGTAAGTGATAG	44.4	GCGTTAGGCTGACTTCGTG	57.9	56	349
gi 89199707 gb dq398761.1	(A)12	GTCGCTGAGTAGTGCCAAAG	57.9	GGCGGTTAGGGGCTGACACG	70	64	345
gi 89199706 gb dq398760.1	(A)12	GCTGCTGTAAGTGCCAAAG	55	GAGGCACGATTAGGGGCTGAC	61.9	65	414
gi 89199705 gb dq398759.1	(A)11	GAGCCTGTAAGTGCCAAAC	52.6	GTGCTGTAGAGCCTGAACCG	59.1	64	806
gi 89199704 gb dq398758.1	(A)11	GAGCTGTAAAGTGCCAAAC	57.1	GTAGCGGTAACTCTGACAC	52.6	58	228
gi 89199703 gb dq398757.1	(A)11	GAGTGCTGTAGAGTGCCATC	60	CGGCGGATAGCATAGAG	56.6	61	539
gi 89199702 gb dq398756.1	(A)11	GAGTCTGTAGTGCCGAACG	47.4	GCGAGATTAGAGCGTGAACAG	52.4	58	313
gi 89199701 gb dq398755.1	(A)12	GTCGCTGTAAGTGCTAAC	57.9	GTCATCAAGTCGCACAGGC	60	56.35	346
gi 89199700 gb dq398754.1	(A)12	CTGTTCAAGGACCCGTGC	63.2	GAGCCTTTTCTCGTCGGTGG	55	58	415
gi 89199699 gb dq398753.1	(A)11	GGGAGTAAAGTGCCAAAC	55.6	GCTTGCTAACGGATTCTGCG	60	56.55	409
gi 89199698 gb dq398752.1	(A)11	GGAGAGTAAAGTGCCAAAC	57.1	GAGCATTGCGAACACAGTG	55.6	56.55	349
gi 89199697 gb dq398751.1	(A)15	GGAGGCTGGCTAAAGTGCTAC	52.4	CTATGGTCAGGAACCGGTGG	57.9	57.05	345
gi 89199696 gb dq398750.1	(A)13	CAATGTTCAAGCGATGCCGTC	52.6	GATACCTTGCTACGAACGG	50	57.35	419
gi 89199695 gb dq398749.1	(A)21	CATCAGAGTTCAGGACCCG	57.1	GCACCTTGCTCGAACCGCGG	61.1	57.05	349
gi 89199694 gb dq398748.1	(A)12	CATCGTGTCAAGGACCCG	57.9	GAACCTCTGTCGGAACCG	61.1	57.05	141
gi 89199693 gb dq398747.1	(A)12	GTGCTGTATCAAGGACCCG	55.6	GGTCGTGCGGTAGGTGG	55.6	59.75	143
gi 89199692 gb dq398746.1	(A)14	GGAGGTCTAAGTGCCAAAC	57.9	GAAGTCCACCGTTAGGT	63.2	58	417
gi 89199691 gb dq398745.1	(A)15	GGAGGTCTAAGTGCCAAAC	52.4	GGCTAATGTCGTCCGGAGG	63.2	58	417
gi 89199690 gb dq398744.1	(A)15	GGAGGTCTAAGTGCCAAAC	55	GCCATTGTCGGAACCGAGG	63.2	58	415
gi 89199689 gb dq398743.1	(A)12	GGAGGTCTAAGTGCCAAAC	57.9	GCCTTGTCGGAACCGTGG	66.7	58	415
gi 89199688 gb dq398742.1	(A)14	GGAGGTCTAAGTGCCAAAC	57.1	GCCATTGTCGGAACCGAGG	63.2	58	417
gi 89199687 gb dq398741.1	(A)10	GGGTCTCTAAGTGCCAAAC	55.6	GCCAGTCTCGGAACAGTG	63.2	56.55	413
gi 89199686 gb dq398740.1	(A)12	GGAGGTCTAAGTGCCAAAC	55	GCGGATTGTGGAACGGCG	66.7	56.55	413
gi 89199685 gb dq398739.1	(A)12	GGAGGTCTAAGTGCCAAAC	57.9	GGCTATGCTCAGTCAGGG	61.1	56.55	413
gi 89199684 gb dq398738.1	(A)12	GGAGGTCTAAGTGCCAAAC	57.9	GACATTGCTCGGAACAGGG	57.9	56.55	418
gi 89199683 gb dq398737.1	(A)15	GGAGGTCTAAGTGCCAAAC	55.6	GAGCCTTCCGAACCGTGG	66.7	58.02	462
gi 89199682 gb dq398736.1	(T)10	GGAGGTCTAAGTGCCAAAC	50	GGATAAGTCGGAAGAGGC	55.6	58.02	462
gi 89199681 gb dq398735.1							
gi 89199680 gb dq398734.1							
gi 89199679 gb dq398733.1							
gi 89199678 gb dq398732.1							
gi 89199624 gb dq398678.1							
gi 89199623 gb dq398677.1							

Table 2 (contd....)

Accession number	Motif	Forward Primer		Reverse primer		Annealing temperature (C)	Product size (bp)
		Primer	GC %	Primer	GC %		
gi 89199624 gb dq398678.1	(A)21	CGTGTAGAGTGCCATCCG	61.1	GATGACTACGAGGATGGCG	57.9	58.05	462
gi 89199623 gb dq398677.1	(T)10	CGTGTAGTCGGTCCATCG	61.1	GTGATAGACAGAGGAGCGG	55	58.05	462
gi 89199612 gb dq398666.1	(T)10	GGATGAGCAGAGGAGAGCC	63.2	GATAGAGTCAGAGGAGGGC	57.9	58.05	461
gi 45738090 gb ay563392.1	(TA)8	GGTGCATAGACTCAACGG	57.9	GTATCGCTATCGCACAGTC	52.6	58.15	676
gi 45738088 gb ay563390.1	(TA)7(AT)7	GAGTCACAGTCGTCAGCG	61.1	GCGTGAGTATCGTAGCAGTC	55	58.15	639
gi 9994240 gb at102469.2	(A)14	GCTGCGTATGCGAGACAC	61.1	GCACGGGTATTTTCAGTCTCGC	56.5	58.1	648
gi 9994232 gb at102419.2	(A)10	CATAGTCGTCAGGATGCG	57.9	GCACGAGTCAGTCTCGC	66.7	57.5	786
gi 9994224 gb at102376.2	(A)10	CACGCCAATCCTGACGCAC	63.2	GCACGGGTTTCAGTCTCGC	65	58.85	779
gi 224985956 gb fj014139.1	(A)10(A)14	CTGGATGGAACCCCTGAGTG	57.9	GCTTGACGCAGAACGGTG	61.1	55.45	356
gi 46403206 gb ay596976.1	(GAG)5	GGAGACGATTGGAGTTGGTG	55	CTCGTTAGATACTCGCC	55.6	56	669
gi 7578882 gb at240764.1	(A)20	GGTGAGGGCATAGAGGC	66.7	GGTGAGGGCATAGAGGC	57.9	55.85	691
gi 6685069 gb at205859.1	(A)19	GAAGCCACAGGAAGCAG	61.1	GTCAAATCACTTCGCGCCAG	52.6	55.85	555
gi 6110321 gb ah008318.1	(A)12	GTATCGGCGTATGTGGC	61.1	GCCAAACCCATTCGTAAGTCC	55	57.8	550
gi 260079916 gb lgq245007.1	(A)10	GGACACACAGCGAGCAG	66.7	GCGACGGTATTCACCTCAC	55	57.95	320
gi 1644387 gb u72654.1	(A)23	GGCTGTTGGTAGATGGCTG	57.9	GGCGAACTCCTATGAACGG	57.9	57.8	690
gi 209483591 gb fj232569.1	(GC)7	CACCTGCGATAGCGGACGAC	63.2	GCAGCATCTCTCGTGGGAC	61.9	61.35	602
gi 94317216 gb dq449916.1	(C)10	CGAATCCAGCGGAAAGCAGAGG	61.9	CGCTTACAGTCGCGGAGTCTC	61.9	62.75	608
gi 57634567 gb ay858677.1	(A)10	GGGAAACCCAGCGAGCGATG	63.2	GTGGCTTCGAGCGCAACTG	63.2	60.1	307
gi 259435649 emb hb880950.1	(A)18	GAGAAGCCATAGGAGGTC	55.6	CGCAATACCTCTCGTAGCTCG	55	63.1	822
gi 205289952 gb fj010824.1	(TA)7	GCGCAGAACTCAACGAC	61.1	GCGAGGCTATCCACCACTC	65	58.15	671
gi 205289938 gb fj010810.1	(TA)6	GTGGCAGGACTCAACGGC	66.7	GCACGAGCATACGCCAGTC	65	58.15	669
gi 257693471 emb hb769727.1	(GAG)5	CTCGTCGTTGGCGTGAATC	60	CAGCCATAACCTCAGCGATAG	52.4	55.6	549
gi 241661579 db jab453155.1	(T)13(T)10	GGGAGTATCTTATCGGAGCG	52.4	GCTGCTATTGATTGCCCGTC	55	54.95	312
gi 241661523 db jab453127.1	(T)10(T)10	GGGTTACTTATCGGGAATCG	50	CGATAGGCATTTTGAGCGGC	57.1	58.55	331
gi 241661499 db jab453115.1	(T)11	GGGAGTTTATCGGGAATCG	52.6	CGTTAGGCGTTTGGGCTG	57.9	58.3	331
gi 89511875 db jab222605.1	(T)10	CACGAGACTTGGTTACGC	57.9	GAATCCCCCAAAACCGAGG	63.2	56.25	337
gi 89511875 db jab222605.1	(T)11	GAATAAGAGGACGCCACG	55.6	GCAGAGCAAGGCCCAATG	66.1	56.4	714
gi 62183686 gb ay879942.1	(A)13	GAATAAGAGGACGCCACG	55.6	GCTTGACGGCAGAACGGC	66.7	56.25	299
gi 166407456 gb eu326062.1	(T)12(GT)7	GAGCAGCAGACGAGTAGC	61.1	GACGACGCACATCTCCAC	61.1	57.25	320
gi 219929423 emb fb699668.1	(GAG)5	GGAGACGATGGAGTTGGTG	57.9	CATAGGTGACATACGCCG	55.6	56	669
gi 218478034 db jab459662.1	(T)12(GT)7	GCGACTATGGCTGCTGCTGC	65	CATAGGTGACATACGCCG	55.6	57.5	
gi 218478034 db jab459662.1	(T)17	GCACCGAAGGCCAGCACCT	65	GAACATACTCTGCCACCG	57.9	58.6	
gi 164454772 db jab289445.1	(ATG)5	CTTCTTCTACTCCGCAGC	55.6	GCAGAAGATGACTCCTCCAG	55	54.9	559
gi 164454770 db jab289444.1	(CTT)7	CCAGAAGTGAGGAAAGCG	55.6	GCAGTGACCAGAGACCCC	66.7	56.95	549
gi 193795409 gb ef203258.1	(TA)7 (T)15 (A)29	CGTGAGGATTGGCTGTCGGC	65	CTATGCGACCAGCGATTAC	55	58.45	703
gi 193795407 gb ef203257.1	(G)14 (A)86	CAGACCAAGGAAACACCCG	57.9	GGATGAGGCACCCACCAAC	63.2	55.3	513
gi 113735515 db jab271691.1	(AT)6	GTAGCAGCAATGTGGTCCGC	65	GATTCAGCAACACCGGTG	50	55	677
gi 170673145 gb eu541812.1	(G)10	CGACCTTTGTAGCAGCCG	61.1	CGCTTTGTGTGCTTCG	55	55.15	499
gi 170673141 gb eu541808.1	(A)10	CACGACTCTCCAGCAGCG	68.4	GTCTCTGCTGTGCTATCG	57.1	55.4	614
gi 170673139 gb eu541806.1	(T)10	CACGAATCATCTCAGTCTCTC	52.4	GTCTTTGCTGTGCTTCG	55.6	55.4	611
gi 170672874 gb eu528047.1	(TAA)6 (T)10	GGGTAATCTGAGCCAAATCC	50	GCGAGGCTATCCGACAC	68.4	57.9	735
gi 4092183 gb at102463.1	(T)10 (A)10	CGGGTCGCAATCCTGAGCC	68.4	GCTTGACAGGCAGAACGGG	63.2	56.7	290

Table 2 (contd....)

Accession number	Motif	Forward Primer		Reverse primer		Annealing temperature (C)		Product size (bp)
		Primer	GC %	Primer	GC %	temperature (C)	GC %	
gi 4092141 gb af102421.1	(A)11	GGGTCCGCATCCTGAGCC	72.2	GCTTGACGGCAGAACGGG	66.7	56.7	290	
gi 4092124 gb af102404.1	(A)11	GCGGGTCGGATGTGAGCC	72.2	GCTTGACAGTGCGAACGGG	63.2	56.25	290	
gi 133874211 dbj ab190181.1	(A)24	CGGGTCTTGGCATGCCTGGG	71.4	CGTCCCTCTTCTCCACTGCC	65	58.3	592	
gi 124388815 gb ef069436.1	(A)21	GGCTCAATCGCTCGGTAAC	57.9	GCCAGTCCAGTGAGTTCCG	63.2	57.7	137	
gi83758482dbj AB19627.1	(TA)6	GGTTACGGTGAAGAGTGACAGG	54.5	GGATGGGAAGAGTGACAGG	57.9	58.25	374	
gi 83758480 dbj AB219625.1	(TA)7	GGTTACGGGAAGAGTGACAGG	55	CCATACCAAGGCTCAATCC	52.6	56.2	228	
gi 156787487 gb ef569229.2	(AT)6 (A)20	CTTCTCCACGGTCGCCCTTAC	60	GTGACTGAAGCATCCTACC	52.6	60.15	539	
gi 156787485 gb ef569228.2	(A)18	GGATAGAGGCTGTGGGATGC	60	CACCAGTCTCAACACCTC	55.6	56.75	313	
gi 146272406 dbj ab281494.1	(CAA)7	CGAGGATTCAAGTTCACGGC	55	GAGTTCAGGGACCGCATAGC	60	58.15	371	
gi 147743640 gb ef571643.1	(CT)10	CCGTAGTGTGGTCAGAAACAGG	56.5	GCGACCTGTAGAACCGATGATG	54.5	57.85	350	
gi 147743624 gb ef571635.1	(TC)6	GCAAGGGGAGCACCCCAAG	66.7	GTTAGCCAGGATGCGGAAGC	57.9	57.75	712	
gi 147743612 gb ef571629.1	(TC)7	GCATTCCGGTCAGCCAAAG	61.1	GGTGGTATTCACTTCCGCCGG	57.1	58.55	825	
gi 147743609 gb ef571627.1	(CT)8	GCAACACAGCGGGACTAAC	57.9	GCACGACAGAACGAGCGGG	66.7	60.25	617	
gi 147743597 gb ef571621.1	(CT)11	CCTCGGACCACCAATCAGC	63.2	GTGTGAACGACTCCGCTTG	57.9	56.35	486	
gi 147743595 gb ef571620.1	(CT)6	CAGTTTCGGTCAGAAAGC	55.6	GTGAGAACAAACCCCGCTG	63.2	57.35	440	
gi 147743553 gb ef571599.1	(TC)7	GCAAGAGGAGGACCCAAAG	55.6	GCAGGTATTCACTCCGCCGG	60	59.05	705	
gi 44829182 gb ay466118.1	(A)28	GCCTACCCAAAACGCTGACC	60	CACCTGTCTTCTGTAGCGGG	55	59.2	645	
gi 124295133 gb ef203260.1	(A)18	GTTGTGAGGTGGCTTCG	61.1	CTGCGTAACACTCATCAAGCC	52.4	56.02	603	
gi 118145133 dbj dd357421.1	(A)18	CCAAATGACGGACGCTACCCC	60	CTATGCCACCAATGTCCC	55.6	57.35	791	
gi 117935906 gb ef062505.1	(A)18	GGAGCCTATCGGAACGGG	66.7	GCGGTTCACTTTCACGAGCG	52.4	58.85	700	
gi 95118583 gb dq497593.1	(A)11	CGTGATCAAGGACCCGCC	61.1	GGAGCCTATCGGAACGGG	66.7	56.35	345	
gi 95118580 gb dq497590.1	(A)11	CTGTGTTCAAAGGACCCGTGC	57.1	GAGCCTATGTCGGAACGGG	61.1	56.35	345	
gi 13018492 dbj je31220.1	(A)18	CTCTGTGCTTTGTGATGCTG	50	GTGTAITTTGAGCGGCAGC	52.6	54.8	650	
gi 92242664 dbj bd340504.1	(G)14(A)86	GGCTGGTGACTACTGCTG	61.1	CCGCACTCCAGCACGAAGC	68.4	58.95	561	
gi 92128002 dbj bd289870.1	(A)15	CCTTGGTTGTGCTGTCTGCTGGT	59.1	GAAGGCAACAGAGGTGACGC	60	58.6	447	
gi 90959871 dbj ab027191.1	(T)14	GAGGATACGACCCACGGACG	63.2	CAGAGCGTAGAAGCCATCAG	55	55.7	704	
gi 90959867 dbj ab027189.1	(A)23	GATACCCACAGAGGAACC	57.9	GACAACTCCCAAGCAAGAC	57.9	55.45	680	
gi 90959865 dbj ab027188.1	(A)27	GCGAACTGCTACTCAAAACCACC	54.5	CATTGAGCGACATCCCTGC	57.9	55.35	693	
gi 90959863 dbj ab027187.1	(A)18	CGACATCAACCCCTACGCTG	60	GCCATAAGGGACGCCATTG	57.9	57.1	730	
gi 6681691 dbj ab017370.1	(TA)7(T)15	CCTGTAACCTCCACCTCAACC	57.1	GCTGTCAATGTAGTCTGCTTCG	52.2	57.9	679	
gi 6681687 dbj ab017368.1	(TA)8	GGTAGTAAGTCCAGGTGCTCG	57.1	CGCTTCTTCACTCCCTTTGC	55	56.95	590	
gi 89276224 gb dq402068.1	(T)14	CGCAGCAATAAAGCCACAG	55	GGTGAGAAGGAGACGGTGG	63.2	58.15	320	
gi 86451166 gb dq358898.1	(T)10	GCCGTCAAATCTCTTCTCCTCG	54.5	CGGGTCTTTGTTGGCTGGCTATG	56.5	60.1	658	
gi 76443559 dbj ab011014.1	(A)18	GTTGTGGAGGTGGCTTCGG	63.2	GGCTGAATGTTGAGAGACG	55	56.6	446	
gi 75812155 dbj d38043.1	(T)12	GCGAGGCATAAGACTGGCTG	60	CAACCGCAACAGGACACCG	63.2	57.2	681	
gi 74267409 dbj ab208689.1	(A)10	GAATGGCAAGTGGTCGCTGG	60	GTATGGCTCGGCAAAAGTTTC	55	54.8	460	
gi 18146806 dbj ab028666.1	(G)13	GCAAGCAAGGGAATCAGG	55.6	CCGCACTCCAGCACGAAGC	68.4	58.6	629	
gi 6687482 emb aj236195.1	(A)10	GTGACTTCGTGAGGACAGC	57.9	GTCGCAATGTCGTCGACGG	63.2	56	453	
gi 62433122 dbj d38168.1	(CT)8	CAGTCTCCTCCGTACCCGAAG	61.9	GAATGGAATCACGGGAG	52.9	56.75	450	
gi 62086548 dbj ab193314.1	(A)28	GAATCATCGCCATCTCCACC	55	GTCCATAACCCCTTTCAGCC	52.6	55.5	362	
gi 62086546 dbj ab193313.1	(A)12(A)12	GGTGTAGCGGGCAATTCG	61.1	CAGACGCTCCACCTTCCC	66.7	57.25	524	
gi 62086540 dbj ab193310.1	(A)19	GAATGGCAAGTGGTCGCTGG	60	GTATGGCTCGGCAAAAGTTTC	55	54.8	620	
gi 38568130 emb aj489906.1	(C)10	GATGATTGCCGCTGCTGG	63.2	GGACTCGCATTTAGCCAGC	57.9	59.8	349	

Table 2 (contd....)

Accession number	Motif	Forward Primer		Reverse primer		Annealing temperature (C)		Product size (bp)
		Primer	GC %	Primer	GC %	GC %	temperature (C)	
gi 4455871 emb aj010509.1	(T)12	TGGTAAAGATGCTCCGTC	50	CCATAGTAAGTCCGAAAGTCC	50	55.15	55.15	317
gi 37954874 gb ay251732.1	(A)19	GGTGCCAAATCCTGAGCCG	66.7	GCTTGACGGCAGAACCGG	66.7	56.4	56.4	307
gi 37954873 gb ay251731.1	(A)16	GGATTGAGCCTGGTATGG	55.6	GCCTTGACAGCAGAACCGG	63.2	56.5	56.5	375
gi 37991673 dbj jab124878.1	(A)16	GCAGTAAGGGAGCGTGAC	61.1	GTCGTAAGTCAGCCAAC	55.6	54.6	54.6	460
gi 22773822 dbj jab080739.1	(A)13	CATAGGAAGCGAAGAACG	52.6	GATTCAGGTAGCAACGGAGTGG	54.5	58.9	58.9	528
gi 33945370 emb aj490190.1	(A)10	GGTGGCAGGACTCAACGG	66.7	GCGATTGACAGCAGAACGGG	60	56.95	56.95	204
gi 15149940 emb aj315324.1	(A)11	GTCGCTGCTGCCGATTGGG	63.2	CTACACAGGTTCCGAGGG	61.1	62.4	62.4	75
gi 27530874 dbj jab076697.1	(A)18	GATTGAGCAGCCGAGGAG	61.1	GTTATGCCCTCCGCTCAGTG	57.9	55.35	55.35	458
gi 22796301 emb aj430909.1	(AT)7	GATGGCAGTCTCCGCTTC	63.2	CCGTTGATGTCTCTGTACC	55	55.95	55.95	597
gi 18146804 dbj jab028865.1	(G)14(A)86	CAGTTCCATCAAGCACACGCGG	59.1	CTTCGGTGGATGTACAG	61.1	56.7	56.7	622
gi 1785485 dbj d14589.1	(A)15	GGCTGTTGAAATGGCGG	61.1	GAGCGTTTATCTGGGCG	55.6	56	56	672
gi 3808127 emb aj011983.1	(A)12	CCGTAGGGTTGGGCTTTC	61.1	GCCAAACCATTCGTAAGTCC	52.6	56.05	56.05	527
gi 10189942 emb aj028840.1	(A)17	GGTCATTACTCGGGGTGTG	57.9	GCACCTCACATCACATTGGGC	54.5	56.6	56.6	797
gi 7415596 dbj jab026494.1	(TA)6(A)21	CTTGAGAAATGCCGTGTTG	50	CTACAAATGGCGGCTCTAC	55	54.8	54.8	410
gi 221150824 gb gh694656.1	(C)12	GGAGGAGTGAATCGGAACCC	60	GTGAGGTGGAGGACTGG	66.7	58.55	58.55	181
gi 62086548 dbj jab193314.1	(A)28	GAATCATCGCCATCTCCAC	55	GTCCATAACCCCTTCAGCC	52.6	55.5	55.5	362
gi 62086546 dbj jab193313.1	(A)12(A)12	GGGTAGCGGGCATTCG	61.1	CAGACGCTCCACCTTCCC	66.7	57.25	57.25	424
gi 62086540 dbj jab193310.1	(A)19	GAATGGCAAGTGGTCGCTGG	60	GTATGGCTCGGCAAGGTTTC	55	54.8	54.8	620
gi 3858130 emb aj489906.1	(C)10	GATGATTGCCGCTGCTGG	63.2	GGACTCGCATTTAGCCAGC	57.9	59.8	59.8	349
gi 33945346 emb aj489882.1	(C)10	CACAAACGGCGACGAGAAG	57.9	CAGCACACGAGTTGAGGC	61.1	61.25	61.25	445
gi 4455871 emb aj010509.1	(T)12	TGGTAAAGATGCTCCGTC	50	CCATAGTAAGTCCGAAAGTCC	50	55.15	55.15	317
gi 37954874 gb ay251732.1	(A)19	GGTGCCAAATCCTGAGCCG	66.7	GCTTGACGGCAGAACCGG	66.7	56.4	56.4	307
gi 37991673 dbj jab124878.1	(A)16	GGATTGAGCCTGGTATGG	55.6	GCCTTGACAGCAGAACCGG	63.2	56.5	56.5	375
gi 22773822 dbj jab080739.1	(A)16	GCAGTAAGGGAGCGTGAC	61.1	GTCGTAAGTCAGCCAAC	55.6	54.6	54.6	460
gi 33945370 emb aj490190.1	(A)13	CATAGGAAGCGAAGAACG	52.6	GATTCAGGTAGCAACGGAGTGG	54.5	58.9	58.9	728
gi 15149940 emb aj315324.1	(A)10	GGTGGCAGGACTCAACGG	66.7	GCGATTGACAGCAGAACCGG	60	56.95	56.95	204
gi 27530874 dbj jab076697.1	(A)11	GTCGCTGCTGCCGATTGGG	63.2	CTACACAGGTTCCGAGGG	61.1	62.4	62.4	75
gi 10189940 emb aj028838.1	(A)18	GATTGAGCAGCCGAGGAG	61.1	GTTATGCCCTCCGCTCAGTG	57.9	55.35	55.35	458
gi 22796301 emb aj430909.1	(A)17	CGGGCTAAGAGACACGGC	66.7	GGTGTGGCTCGTTGAATGGG	60	59.3	59.3	285
gi 18146804 dbj jab028865.1	(AT)7	GATGGCAGTCTCCGCTTC	63.2	CCGTTGATGTCTCTGTACC	55	55.95	55.95	597
gi 1785485 dbj d14589.1	(G)14(A)86	CAGTTCCATCAAGCACACGCGG	59.1	CTTCGGTGGATGTACAG	61.1	56.7	56.7	522
gi 3808127 emb aj011983.1	(A)15	GGCTGTTGAAATGGCGG	61.1	GAGCGTTTATCTGGGCG	55.6	56	56	672
gi 10189942 emb aj028840.1	(A)12	CCGTAGGGTTGGGCTTTC	61.1	GCCAAACCATTCGTAAGTCC	52.6	56.05	56.05	427
gi 7415596 dbj jab026494.1	(TA)6(A)21	GGTGAAGTGAATCGGAACCC	50	CTACAAATGGCGGCTCTAC	55	54.8	54.8	410
gi 221150824 gb gh694656.1	(C)12	GGAGGAGTGAATCGGAACCC	60	GTGAGGTGGAGGACTGG	66.7	58.55	58.55	181

clustering patterns, test for Hardy-Weinberg equilibrium and multilocus equilibrium, and to test polymorphic loci for the evidence of selective neutrality. This can be useful to plant breeders, germplasm managers, or others who are interested in population genetic properties of materials that they are working with. The three most common types of markers used today are RFLP, RAPD, and microsatellites. A wide variety of methods for the construction of libraries enriched for microsatellite sequences have been reported, the most popular among those being the ones based on vectorette PCR using anchored primers. But this method is highly time-consuming and expensive, and the alternative is to use bioinformatics, that is, computational tools to screen the public database and find SSR. EST-derived molecular markers, especially SSR and SNP, are highly useful in developing linkage maps and markers assisted breeding programs. These markers are also transferable to related genera.

Molecular marker techniques are advantageous as they directly reflect variations in the DNA sequences and therefore of independence of environment. Among many molecular marker techniques currently available, microsatellites and SSRs^[35] provide an improved technology in assessing genetic diversity and genetic relationships in plants as they are highly polymorphic, codominants, very informative, and PCR based. EST-SSRs offer the following advantages over other genome DNA-based markers: (1) they should detect variation in the expressed portion of the genome so that gene tagging should give “perfect” marker–trait associations; (2) they can be developed at no cost from the EST databases; and (3) once developed, these markers, unlike genomic SSRs, may be used across a number of related species. With the growth of sequence databases, several authors have reported an abundance of SSRs in different genomes. The Distribution of SSRs in the rice genome has also been studied on the basis of the two whole genome draft sequences released, respectively, by Syngenta and by the Beijing Genome Institute (BGI). In the draft sequence released by Syngenta, for instance, 48,351 SSRs (including di-, tri-, and tetranucleotide repeats) were available, giving a density of 8 kb per SSR in the whole genome; SSRs represented by di-, tri-, and tetranucleotide repeats accounted respectively for 24%, 59%, and 17% of the total SSRs.

SSRs are very polymorphic due to the high mutation rate affecting the number of repeat units. Such length-polymorphisms can be easily detected on high-resolution gels (e.g., sequencing gels), by running PCR-amplified fragments obtained using a unique pair of primers flanking the repeat.^[36] Chung and Staub^[37] developed a set of consensus chloroplast primer pairs for ccSSRs from *N. tabacum* chloroplast sequences. All primer pairs produced

amplicons after PCR employing chloroplast DNA from members of the *Cucurbitaceae* (six species) and *Solanaceae* (four species). Sixteen, 22, and 19 of the initial 23 primer pairs were successively amplified by PCR using template DNA from species of the *Apiaceae* (two species), *Brassicaceae* (one species), and *Fabaceae* (two species), respectively. Twenty of the 23 primer pairs were also functional in three monocot species of the *Liliaceae* (onion and garlic), and the *Poaceae* (oat). ccSSR primers were strategically “recombined” and referred to correctly as recombined consensus chloroplast primers (RCCP) for PCR analysis of cucumber DNA such that the primers designed for the SSR-containing genus of Gentiana family would be utilized for the production of amplicons from different members of family.

Kijas *et al.*^[38] tested two primer sets in 10 different *Citrus* species and two related genera and found conservation of the sequences. Cross-species amplification has also been reported between cultivated rice and related wild species^[39] and between *Vitis* species.^[40] Provan *et al.*^[41] could show successful amplification of two tomato SSR primer pairs tested on potato cultivars. Weising *et al.*^[42] reported conservation of SSR flanking sites in different species of kiwifruit (*Actinidia chinensis*). Usually, a low percentage of markers also amplified fragments from species belonging to other genera from the same family. Within the *Poaceae* family, primers worked even across different genera,^[43] but only 50% of microsatellite loci identified in wheat were also polymorphic in rye and barley cultivars. Whitton *et al.*^[44] tested 13 SSR loci in 25 representatives of the *Asteraceae*, where it was demonstrated that the regions flanking in the repeats are not highly conserved, neither in the nucleotide sequence nor in the relative position.

Indeed, in general, transferability of polymorphic markers in plants is likely to be successful mainly within genera (success rate close to 60% in eudicots and close to 40% in the reviewed monocots) rather than between genera (transfer rates are approximately 10% for eudicots) within the same family.^[45] This transferability of polymorphic markers nature in plant generally enhances the utilization of the primers in random way. Comparative genome analysis facilitates high-throughput comparative mapping with the assistance of cross-species markers, and further facilitates gene cloning by identifying cross-reference genes. Seventeen SSR primer sets developed for *Quercus petraea* were tested on eight different members of the *Fagaceae* family.^[46] In total 66% resulted in interpretable amplification products and most of them were really homologous to the originally cloned SSR fragment from *Q. petraea*. The primers could be designed successfully for a very large number (169, 31%) of SSRs [Table 2]. However, it was not possible to design the primers for remaining SSRs (376, 69%) because

the sequence flanking at both ends of the SSRs was inadequate in size to design the primers. The large number of primer pairs for the SSRs that have been designed during the present study may be utilized for a variety of purposes, for example, gene tagging, genetic mapping, population studies, etc. Due to a high level of potential for length polymorphisms, SSRs have become a valuable source of genetic markers and have been broadly applied to various areas of genetic research including studies of genome variation, establishment of genetic maps, integration of physical and genetic maps, determination of evolutionary relationships, and comparative genome analyses.

CONCLUSIONS

Nucleotide sequences of *Gentiana* family were systematically searched for SSRs using the “ssr_finder.pl” perl program for the development of SSR markers. This is a valuable approach for both costs and time, given a sufficient amount of available *Gentiana* family sequences. The use of SSRs in genetic diversity studies is a novel tool that reveals variation in genomes.

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