Full Length Research Paper

Genome shotgun sequencing and development of microsatellite markers for gerbera \((Gerbera\ hybrida\ H.)\) by 454 GS-FLX

Kyoung-In Seo\(^1\), Gi-An Lee\(^1\), Sang-Kun Park\(^2\), Mun-Sup Yoon\(^1\), Kyung-Ho Ma\(^1\), Jung-Ro Lee\(^1\), Yu-Mi Choi\(^1\), Yeon-ju Jung\(^1\) and Myung-Chul Lee\(^1\)*

\(^1\)National Agrobiodiversity Center, National Academy of Agricultural Science, RDA, 88-20, Seodun-Dong, Suwon, Gyunggi-do, 441-707, Korea.

\(^2\)Floriculture Research Division, National Institute of Horticultural and Herbal Science, RDA, Suwon, Gyunggi-do, 440-706, Korea.

Accepted 26 January, 2012

The objective of this research was to develop and characterize microsatellite markers for gerbera. We used shotgun sequencing with Roche 454 GS-FLX Titanium technology to identify microsatellite loci in gerbera genomic DNA \((Gerbera\ hybrida)\). The total length of non-redundant sequences obtained was 22,527,019 bp, which consisted of 3,085 contigs and 28,249 singletons. We assembled 61,958 reads into 3,085 contigs, of which 114 (3.70%) contained microsatellite repeats. The average G+C content was 39.3%. Functional annotation to known sequences yielded 14.7% unigenes in the ‘Raon’ cultivar. Analysis of the gerbera genome DNA \((‘Raon’)\) general library showed that sequences of \((AT)\), \((AG)\), \((AAG)\) and \((AAT)\) repeats appeared most often, whereas \((AC)\), \((AAC)\) and \((ACC)\) were the least frequent. Primer pairs were designed for 80 loci. Only eight primer pairs produced reproducible polymorphic bands in the 28 gerbera accessions analyzed. A total of 30 alleles were identified from the eight polymorphic SSR loci, with two to eight alleles per locus (average level of 3.75). These markers will be useful for investigating genetic diversity and differentiation in gerbera.

Key words: Genetic diversity, genomics, microsatellite isolation, pyrosequencing, SSRs.

INTRODUCTION

Gerbera species, belonging to the family Asteraceae, are among the most popular decorative plants worldwide, with uses as garden plants, potted plants and cut flowers. Gerbera plants generally show variation in their stalks, and through hybrid breeding (they are polyploid) thin, soft and cut flower varieties have been produced (Kanwar and Kumar, 2008; Yu et al., 1999). Most of the gerbera cultivated in Korea have large flowers that are used as cut flowers. Various cut flower gerbera cultivars have different flower colors. In addition, year-round production is possible and is a common practice. Evaluations of the genetic diversity of gerbera have been conducted using random amplified polymorphic DNA (RAPD) and expressed sequence tag-simple sequence repeats (EST-SSR) (Mata et al., 2009; Gong and Deng, 2010). In addition, Bhatia et al. (2009) studied the effects of explant source on the genetic integrity of 45 in vitro-raised gerbera plants \((Gerbera\ jamesonii\ Bolus)\) using 32 ISSR markers. However, microsatellite (SSRs) markers were not developed and tested for these gerbera genetic studies. Simple sequence repeats (SSR) or microsatellites have been used to examine genotypic variation and for breeding research (Eujayl et al., 2001; Ruiz et al., 2000). Furthermore, these markers are widely used in cultivar fingerprinting and molecular mapping. When compared with other types of molecular markers, SSR markers are abundant, co-dominant and have high reproducibility. In the past, microsatellite markers were developed using the enrichment method and hybridization probes. However, this method had a disadvantage: it could not detect the correct repeat sequence in the

*Corresponding author. E-mail: mclee@rda.go.kr. Tel: 82-31-299-1887. Fax: 82-31-294-6029.
In this paper, we report the development of microsatellites and the characterization of gene content in gerbera (*Gerbera hybrida*) using cost-effective 454-pyrosequencing.

### MATERIALS AND METHODS

#### Plant material and DNA extraction

PCR and genotyping was performed on 28 gerbera (*Gerbera hybrida* H.) accessions collected from the National Horticultural Research Institute (NHRI), Rural Development Administration (RDA) of Korea (Table 1). The 'Raon' and 'Misty Red' cultivars were bred at the NHRI of Korea and the rest of the cultivars were bred abroad. The scoring of flower type was based on the number of petals, such as single, double and semi-double. Among 28 accessions, only 2 accessions have single type flower and 12 other accessions were semi-double type and 14 accessions have double type.

Total genomic DNA was extracted from young fresh leaves by the modified cetyltrimethylammonium bromide (CTAB) method described by Dellaporta et al. (1983). A total of 8 µg of 'Raon' (double petal type) DNA was used to perform shotgun sequencing using 454 GS-FLX Titanium pyrosequencing (Roche Applied Science, Mannheim, Germany).

#### Shotgun genomic sequencing

A small/medium plate (1/8 PicoTiterPlate; PTP) was used for 454 GS-FLX Titanium pyrosequencing for samples processed by the shotgun method according to the manufacturer’s instructions. In short, the samples prepared using the shotgun method was fractionated into smaller pieces (300 to 800 bp) by nebulization. Subsequently, the fragment length distribution of the library was assessed using a gerbera DNA 7500 LabChip (Agilent Technologies, Waldbronn, Germany) to ensure that the input material was adequate. The ends of the sheared gerbera genome DNA fragments were polished by the action of T4 DNA polymerase and T4 polynucleotide kinase. Short adapters were then ligated to the ends of each sample DNA fragment. These adapters provide priming regions to support both amplification and nucleotide sequencing. Finally, the quality of the library was assessed with a Bioanalyzer RNA 6000 Pico LabChip (Agilent Technologies).

Emulsion titration was performed with different amounts of input ssDNA. Medium-scale emPCR was conducted based on the titrated ssDNA amount. Enriched bead samples were then counted using a Z1 Coulter Counter (Beckman Coulter, Brea, CA, USA) to calculate the percent enrichment.

Samples prepared using the above procedure were loaded,
along with other reagents, into the wells of a 1/8 PTP device, and sequencing was performed with a GS FLX Titanium Reagents XLR70, a 70 x 75 mm Titanium PTP device and a Genome Sequencer FLX Instrument (454 Life Science, Roche Applied Science). Sequencing data were obtained after a 9-h run of the GS-FLX.

After sequencing, sequence assembly was performed using the GS Dasy Nova Assembler software to produce the contigs and singletions. All sequence alignments were conducted using the Basic Local Alignment Search Tool (BLAST; database: NT) (Altschul et al., 1997).

Selection of SSRs and primer design

Identification of SSR motifs within contig sequences and primer design were conducted using the ARGOS program (Kim, 2004). The contigs were screened for mononucleotide, dinucleotide, trinucleotide, tetranucleotide, pentanucleotide and hexanucleotide repeats. The minimum number of repeats was set to four. From these contigs, we designed 80 primer pairs to amplify microsatellite regions. The primer design included the following specifications: (1) melting temperature (Tm) between 54 and 64°C; (2) GC content above 30%; (3) amplicon size of 150 to 300 bp. For each locus, forward primers were synthesized to incorporate a 5' fluorescent label, M13 (−21)-Tag TGTAAAACGACGGCCAGT (Messing, 1983).

Polymorphism analysis

The M13F-tail (5′-TG TAA AAC GAC GGC CAGT-3′) PCR method of Schuelke (2000) was used to measure PCR product size, as described by Lee et al. (2008). PCR was performed in a 20 μl reaction mixture containing 2 μl of template genome DNA (20 ng/μl), 0.2 μl of locus-specific primer (10 pmol/μl), 0.4 μl of M13 universal primer (10 pmol/μl), 0.6 μl normal reverse primer, 2.0 μl of 10X PCR buffer (SolGent, Daejeon, Korea), 1.6 μl of dNTP (2.5 mM) and 0.2 μl h-Taq-polymerase (5 U/μl; SolGent). The 5′-M13 sequence was attached to a forward primer to incorporate a fluorescent dye [blue (FAM), green (NED) or yellow (HEX)] into the PCR product. PCR was performed as follows: initial denaturation at 94°C (3 min), then 30 cycles each of 94°C (30 s), 55°C (45 s) and 72°C (1 min), followed by 10 cycles of 94°C (30 s), 53°C (45 s) and 72°C (1 min), and a final extension at 72°C for 10 min (PTC-100 Thermocycler, MJ Research, Waltham, MA, USA). All denatured products were run on an ABI PRISM 3130xl Genetic Analyzer (Applied Biosystems, Carlsbad, CA, USA) using the molecular size standard (35 to 500 bp) GeneScan 500 ROX (6-carboxy-rhodamine). Genotypes were obtained using GeneMapper v4.0 software (Applied Biosystems).

Data analysis

PCR genotyping results were converted to binary form (0 = absence; 1 = presence), and similarity coefficients were calculated among all 28 accessions considered in this study with the program NTSYS, version 2.1 (Exeter Software, Setauket, NY, USA), using SIMQUAL with the Dice coefficient (Dice, 1945). The data were subjected to unweighted pair group method with arithmetic mean (UPGMA) cluster analysis using NTSYS 2.1. The number of distinct genotypes and genetic diversity were computed using GenoType/GenoDive software (Meirmans and Van Tienderen, 2004).

Sequence annotation and analysis of gene content

Functional annotation and major functional categories (biological process, molecular function and cellular component) were analyzed using the BLAST2GO software suite v2.3.1 (www.blast2go.de) (Conesa and Gotz, 2008). Homology searches (BLASTX and BLASTN) were performed remotely on the NCBI server through a query-friendly version of BLAST (QBLAST), following a sequential strategy.

RESULTS

Sequence analysis of the gerbera genome

The status yield of sequencing runs and statistics for ‘Raon’ are summarized in Table 2. We also provided sequence data for all contigs of ‘Raon’ as Supplementary File 1. The 454 GS-FLX Titanium sequencing run yielded 22.5 Mbp of a genome DNA library from ‘Raon’ (G. hybrida), including 22,527,019 reads, with an average read length of 363.6 bp and 114 contigs (3.69%) containing microsatellite repeats. The ‘Raon’ genome DNA was assembled into 3,085 contigs and 28,249 singletions. After the assembly (3,085 contigs + 28,249 singletions), the contigs had an average length of 344.03 bp and an average depth of coverage of 10.92 reads per base pair. A total of 430 large contigs (≥500 bp), with an average length of 1,046 bp, was detected for ‘Raon’. The longest contig was 25,066 bp.

To better understand the general mechanisms working in the gerbera, we performed the gene ontology (GO) analysis and carried out all aligned sequences (3,085 contigs and 28,249 singletions). Functional annotation was analyzed using BLAST2GO (Conesa and Götz, 2008), which led to consistent gene annotations and the assignment of gene names, gene products and gene ontology (GO) numbers. In total, 452 unigenes were assembled, and assembled contig sequences were available. All unigenes were investigated for their association with GO terms from level 4 biological processes, molecular function and cellular components; the number of associations was 137 (30.3%), 85 (18.8%) and 56 (12.4%), respectively. Eighty-five assignments were made to molecular function ontology, with a large proportion of these functioning in DNA binding (32.7%) and cation binding (27.3%) (Figure 1). Under biological process ontology, 137 assignments were made, with a large proportion of the assignments falling into cellular macromolecule metabolic process (24%) and nucleobase, nucleoside, nucleotide and nucleic acid metabolic process (23%) categories. The top 10 contigs by number of assembled reads were aligned to the NCBI database using BLASTN (Table 3). Four had significant (e-value ≤ 1 x 10⁻³ and a bit score >40) hits to different genes, while three aligned to Lactuca sativa cultivar Salinas chloroplast complete genome DNA and three others aligned to Guizotia abyssinica chloroplast complete genome DNA.

Characteristics of the genome SSR

SSR motifs found in ‘Raon’ are summarized in...
Of the 3,085 unique contigs, 114 (3.70%) contained microsatellite sequences. Among the 114 microsatellite regions, we found dinucleotide, trinucleotide and pentanucleotide repeat motifs. The frequency of the appearance of these SSRs was estimated to be one SSR for every 9.3 kb in the 1.06 Mbp sequences from the Gerbera genome. The repeat number for dinucleotides (62.3%) was higher than that for trinucleotides (36.8%). In addition, a pentamer repeat was found in one contig (0.9%). The number of repeats ranged from 4 to 33. Among the SSRs, AT/TA repeat motifs were the most common (28.9%) followed by AG/GA repeat motifs (21.1%). The most frequent trinucleotide motif was AAG/AGA/GAA (13.2%), which was closely followed by the AAT/ATA/TAA motif (7.9%).

### Development of SSRs markers

After screening all the contigs (n = 3,085) for SSR regions with the ARGOS PROGRAM (Kim, 2004), we identified a total of 114 SSR regions. A random set of 80 primer sets was designed which included SSR regions; these primers were used to amplify gerbera genomic regions. There were six perfect repeats having no interruptions, one imperfect repeat and one compound repeat sequence. Four of the loci: GB-GJ-002, GB-GJ-049, GB-GJ-059 and GB-GJ-063, which had (AAT)$_4$, (AT)$_4$, (ACA)$_6$, (AG)$_7$, (ATC)$_4$, and (CAT)$_5$ repeats, respectively, were labeled with FAM (blue). Two others, GB-GJ-011 and GB-GJ-074, with (CAC)$_5$ and (CAT)$_5$ repeats, respectively, were labeled with HEX (yellow). The last loci, GB-GJ-021 and GB-GJ-077, which had (AAAG)$_4$ and (TGA)$_6$ repeats, respectively, were labeled with NED (green). GB-GJ-049 had the widest allele size range of the eight loci (160 to 317 bp) (Table 4). When used with 28 accesses, eight primer pairs managed to amplify 30 distinct alleles, with an average of 3.75 alleles per locus.

### Genetic diversity based on SSR data

Genetic diversity among G. hybrida accessions was evaluated using an UPGMA model (Figure 3). Cluster analysis separated the three groups: Groups-I, II and III. Group-I accommodated the highest number of accesses (16 accesses, 57.1%) followed by Group-II (nine accesses, 32.1%). Group-III contained the lowest number of accesses (three accesses, 10.7%). As shown in Figure 3, most of the semi-double type accessions were clustered in Group-I. Group-II consisted of nine accesses, which were mainly double type; Group-III consisted of three accesses: one single type, one double type, and one semi-double type. Coefficients of similarity ranged from 0.68 to 0.96. Flower type did not differentiate clearly among gerbera accessions. The ‘Misty Red’ and ‘Raon’ accessions, which were originally bred in Korea (Choi et al., 2001), belonged to Groups-I and II, respectively. Our analyses of genetic distances (NTSYS) and calculations of clonal diversity (GenoType/GenoDive) divided the G. hybrida accessions in Group-I into 14 genotypes. The Groups-II and III populations were divided into eight and three genotypes, respectively. This is because SSR markers detected multilocus genotypes within the groups. Calculations of clonal diversity (Genotype/GenoDive) resulted in a Nei’s diversity index (corrected for sample size) of 0.98 for Group-I (Table 5). For Groups-II and III, clonal diversity was 0.97 and 1.00, respectively, as nine and three genotypes were identified.
respectively. Total diversity over all accessions was 0.99 and average diversity within populations was 0.98. The fraction of diversity among populations, corrected $G_{ST}$, was 0.01.

**DISCUSSION**

The 454 GS-FLX technology provided a very cost-effective method for obtaining SSR sequences. This technology will accelerate research on less-studied gerbera plants. The 454 GS-FLX has been broadly used for transcriptome sequencing, whole genome sequencing, metagenomics and microbial diversity (http://454.com/applications/index.asp). As next-generation sequencing, 454 GS-FLX Titanium possesses more powerful sequencing capabilities, with a read length of 400 bp and higher sequencing accuracy. In this study, *G. hybrida*
Table 3. The top ten contigs determined by the number of assembled reads and their respective alignments.

<table>
<thead>
<tr>
<th>Contig</th>
<th>BLASTN result</th>
<th>Contig Length (bp)</th>
<th>No. of assembled reads</th>
</tr>
</thead>
<tbody>
<tr>
<td>1931</td>
<td><em>Lactuca sativa</em> cultivar Salinas chloroplast, complete genome</td>
<td>25066</td>
<td>916</td>
</tr>
<tr>
<td>1954</td>
<td><em>Guizotia abyssinica</em> chloroplast, complete genome</td>
<td>18240</td>
<td>460</td>
</tr>
<tr>
<td>2020</td>
<td><em>Corokia cotoneaster</em> 26S ribosomal RNA gene, complete sequence</td>
<td>6606</td>
<td>319</td>
</tr>
<tr>
<td>28</td>
<td><em>Guizotia abyssinica</em> chloroplast, complete genome</td>
<td>14762</td>
<td>306</td>
</tr>
<tr>
<td>5</td>
<td><em>Guizotia abyssinica</em> chloroplast, complete genome</td>
<td>11466</td>
<td>253</td>
</tr>
<tr>
<td>56</td>
<td><em>Trachyium caeruleum</em> chloroplast, complete genome</td>
<td>10443</td>
<td>230</td>
</tr>
<tr>
<td>3831</td>
<td><em>Oryza punctata</em>, complete sequence</td>
<td>1296</td>
<td>226</td>
</tr>
<tr>
<td>1446</td>
<td><em>Oryza australiensis</em>, complete sequence</td>
<td>5698</td>
<td>218</td>
</tr>
<tr>
<td>2</td>
<td><em>Lactuca sativa</em> cultivar Salinas chloroplast, complete genome</td>
<td>6924</td>
<td>150</td>
</tr>
<tr>
<td>64</td>
<td><em>Lactuca sativa</em> cultivar Salinas chloroplast, complete genome</td>
<td>7569</td>
<td>134</td>
</tr>
</tbody>
</table>

Figure 2. Distribution pattern of SSRs found in sequence of the 'Raon' cultivar.

Genome DNA was sequenced using a 454 GS-FLX system, producing a total of 22,527,019 sequence reads before assembly (reads only). After assembly, 3,085 contig sequences were obtained. The GC content of all reads was 39.3%, which is within the range commonly observed for plant genomes [e.g., tomato (Zhu et al., 2008) and Arabidopsis (Arabidopsis Genome Initiative, 2000)]. Our sequencing runs yielded reads with an average length of 363.5 bp.

In the present study, 3,085 contigs (about 1.1 Mb) were used in an SSR search, which produced 114 (3.69%) SSR-containing sequences, including at least four dinucleotide, trinucleotide, tetranucleotide, penta-nucleotide and hexanucleotide repeats. This is a relatively low abundance of SSRs as compared to the numbers of ESTs containing SSRs found in an earlier report: *Citrus* spp. 6.09%, *Lactuca sativa* 4.90%, *Rosa* spp. 8.05% and *Prunus persica* 4.99% (Kumpatla and Mukhopadhyay, 2005). The occurrence of SSRs in ESTs from the gerbera transcriptome was at the rate of one SSR per 5.16 kb (Gong and Deng, 2010), whereas in this study, using genomic sequences (represented by 1.1 Mbp of gerbera genome DNA), the frequency was one SSR for every 9.3 kb.

The proportion of AT/TA motifs in the SSRs generated from the gerbera genome was much higher than the proportion of (CA)/(TG) repeat motifs (Figure 2). This result is consistent with surveys of microsatellites in *Jatropha curcas* L. (Sato et al., 2010) and also with studies on barley (Becker and Heun, 1995), *Cannabis sativa* (Algharim and Almirall, 2003) and citrus (Chen et al., 2006). However, studies of *Brassica napus* (Uzunova and Ecke, 1999) and *Phaseolus vulgaris* (Benchimol et al., 2007) found greater abundances of (CA)/(TG) motifs.
Malausa et al. (2011) reported that enriched libraries are able to design primers about three times more than shotgun library. But, the genomic library enriched method have the disadvantage of the difficulty encountered in hybridization with AT and GC probe, which makes it to easily construct the self-complementary structure than the other kinds of repeat types of DNA sequences (Ashkenazi et al., 2001; Lagercrantz et al., 1993). Malausa et al. (2011) analyzed *Apis mellifera* using enriched and shotgun libraries; the percentage of microsatellite motifs of AT nucleotides was 8 and 39%, respectively. These results were similar to that detected in our gerbera shotgun library.

The importance of microsatellites as genetic markers and their potential use in various applications, such as diversity analysis or genetic mapping for crop improvement, have been well verified in plants (Csencsics et al., 2010; Lee et al., 2009; Takundwa et al., 2010). Despite their many advantages, SSR markers had not been developed or used in gerbera. Therefore, we developed and used SSRs to assess genetic diversity in gerbera accessions. Of these SSR markers, 75% were perfect type, 12.5% were imperfect and 12.5% were compound type.

In this study, we reported the successful development of eight polymorphic markers using 454-pyrosequencing technology. We also tested the applicability of these markers in *G. hybrida*. Although, only eight markers have been tested here, these microsatellite markers provide a useful tool for gerbera breeding programs. The comprehensive dataset from this study will contribute to further genetic improvements, genomic information and gene discovery in gerbera. To our knowledge, this is the first report on microsatellite markers for gerbera (*G. hybrida*) and it confirms the usefulness of this kind of marker as a tool for cultivar identification and the breeding of

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### Table 4. Characteristics of polymorphic microsatellite markers isolated for gerbera.

<table>
<thead>
<tr>
<th>Locus</th>
<th>GenBank accession number</th>
<th>Repeat motif</th>
<th>Repeat status</th>
<th>Primer sequence (5’→3’)</th>
<th>Fluorescent dye</th>
<th>Tm (^{\circ\text{C}})</th>
<th>Expected Product size</th>
<th>Gerbera size range (bp)</th>
<th># Alleles (^{W})</th>
</tr>
</thead>
<tbody>
<tr>
<td>GB-GJ-002</td>
<td>JF343527</td>
<td>(AAT)(_4)</td>
<td>Perfect</td>
<td>F: TCAGTCTTCTTGGCGGTTC R: TGATCCACAGATTTCCCAG</td>
<td>FAM</td>
<td>58</td>
<td>188</td>
<td>203-206</td>
<td>2</td>
</tr>
<tr>
<td>GB-GJ-011</td>
<td>JF343528</td>
<td>(CAC)(_5)</td>
<td>Perfect</td>
<td>F: GTGCATTGGCTTTGGTCAA R: CCCAGCAAAATGACGCC</td>
<td>HEX</td>
<td>58</td>
<td>255</td>
<td>234-273</td>
<td>4</td>
</tr>
<tr>
<td>GB-GJ-021</td>
<td>JF343529</td>
<td>(AAAG)(_4)</td>
<td>Perfect</td>
<td>F: GCCTCCAAATTCTGTTTT R: TTCAAGCATGAGGGCAAG</td>
<td>NED</td>
<td>58</td>
<td>283</td>
<td>299-303</td>
<td>2</td>
</tr>
<tr>
<td>GB-GJ-049</td>
<td>JF343530</td>
<td>(AT)(_4), (ACA)(_6)</td>
<td>Compound</td>
<td>F: AATTGTCTCGACTCCGGTT R: ATCCATGTAGCAAGCCCAT</td>
<td>FAM</td>
<td>58</td>
<td>296</td>
<td>160-317</td>
<td>8</td>
</tr>
<tr>
<td>GB-GJ-059</td>
<td>JF343531</td>
<td>(AG)(_7)</td>
<td>Perfect</td>
<td>F: TGTCATCTTGAGATGTCGC R: GGTAGCAGATGAGGGCCAGA</td>
<td>FAM</td>
<td>58</td>
<td>189</td>
<td>182-211</td>
<td>4</td>
</tr>
<tr>
<td>GB-GJ-063</td>
<td>JF343532</td>
<td>(ATC)(_4), (CAT)(_5)</td>
<td>Imperfect</td>
<td>F: TCTCCCACATCTCCGT T: ATCCACGTGACGCC</td>
<td>FAM</td>
<td>58</td>
<td>192</td>
<td>194-235</td>
<td>4</td>
</tr>
<tr>
<td>GB-GJ-074</td>
<td>JF343533</td>
<td>(CAT)(_6)</td>
<td>Perfect</td>
<td>F: TCTCAATTGGCTTCCCCAG T: GGTCTATCAAGGGGGCCAG</td>
<td>HEX</td>
<td>58</td>
<td>283</td>
<td>296-299</td>
<td>2</td>
</tr>
<tr>
<td>GB-GJ-077</td>
<td>JF343534</td>
<td>(TGA)(_6)</td>
<td>Perfect</td>
<td>F: GAGCAAGCCATCATCACC T: CTTGCTTCTCCTGCCCT</td>
<td>NED</td>
<td>58</td>
<td>197</td>
<td>195-218</td>
<td>4</td>
</tr>
</tbody>
</table>

\(^{2}\text{Tm, annealing temperature; #Alleles}^{W}\), the total number of observed alleles among the 28 accessions.
Figure 3. Dendrogram showing relationships among 28 *G. hybrida* accessions based on UPGMA of SSR profiles.

Table 5. Calculation of clonal diversity within groups.

<table>
<thead>
<tr>
<th>Group</th>
<th>Number of samples</th>
<th>Number of genotypes</th>
<th>Nei's div$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group-I</td>
<td>16</td>
<td>14</td>
<td>0.98</td>
</tr>
<tr>
<td>Group-II</td>
<td>9</td>
<td>8</td>
<td>0.97</td>
</tr>
<tr>
<td>Group-III</td>
<td>3</td>
<td>3</td>
<td>1.00</td>
</tr>
<tr>
<td>Total</td>
<td>28</td>
<td>25</td>
<td>0.99</td>
</tr>
</tbody>
</table>

$^2$Nei’s div: Nei’s (1987) genetic diversity corrected for sample size.

ACKNOWLEDGEMENT

This study was supported by a grant (Code no. PJ006825) -> PJ008368 from the National Academy of Agricultural Science, RDA, Republic of Korea.

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