Intersimple Sequence Repeats Distinguish Genetic Differences in Easter Lily ‘Nellie White’ Clonal Ramets within and among Bulb Growers over Years

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ABSTRACT. The large genome size of easter lily [Lilium longiflorum (77.1 pg/2C nucleus)], coupled with repetitive DNA sequences, makes it difficult to use molecular techniques to identify or fingerprint lily (Lilium) species, hybrids, and clones. Previous research demonstrated that amplified fragment length polymorphisms could not be optimized for consistency and repeatability to obtain reliable genetic variation assessments of lily species and clones. The objective of this research was to analyze the effectiveness and stringency of intersimple sequence repeats (ISSRs) to determine genetic differences between L. longiflorum ‘Nellie White’ clonal ramet populations from bulb growers over years. DNA from closely related clones of L. longiflorum ‘Nellie White’ included 2002 (n = 11 bulb lots) and 2003 (n = 12 bulb lots). Comparison cultivars and species were also included. Five University of British Columbia (UBC) primers (P808, P810, P811, P814, and P818) that were used produced 56 polymorphic loci. ISSR banding patterns were consistent among three replications within ‘Nellie White’ clonal genotypes. ‘Nellie White’ clones differed significantly within (82%) and among (18%) growers in 2002 and 2003. ‘Nellie White’ clones are not uniform or part of a single ramet population. Principal clades within years separated at Nei’s genetic distances (GDs) of GD = 0.6 (growers 2, 4, and 12) to GD = 0.82 (grower 6) in 2002 and GD = 0.51 (grower 4) to GD = 0.78 (grower 14). The most closely related ‘Nellie White’ clones within growers ranged from GD = 0.8 to 0.95 in 2002 and GD = 0.7 to 0.91 for 2003. Five top-performing growers (1, and 3–6) from previous morphological studies and, particularly growers 3 and 5, were in similar clades, cosegregating with phenotypic traits of stem emergence and flowering dates. The lack of a meiotic sieve (Muller’s ratchet) may be responsible for the high level of mutational differences present in the ‘Nellie White’ clones and significantly affects the ability of commercial greenhouse growers to produce a uniform easter lily crop, particularly in years when the Easter holiday is early.

Lilium contains ≥100 species across the Northern Hemisphere between 10.00°N and 60.00°N. lat. (Anderson, 1986; Asano, 1986, 1989; De Jong, 1974). Many species are important commercial floricultural crops for use as flowering potted plants, cut flowers, and garden perennials (Dole and Wilkins, 2005). Of these, easter lily ranks among the top five potted flowering plants in sales for the United States (U.S. Department of Agriculture, 2009). The primary easter lily cultivar in North America is L. longiflorum ‘Nellie White’, a vegetatively propagated clone that is >60 years old (Zlesak and Anderson, 2003, 2007). A lily bulb grower, J. White, named ‘Nellie White’ after his wife (Texas A&M University, n.d.). The original seedling may have originated from crosses performed at Colorado State University (D. Hartley, personal communication).

Easter lily is the only major floral crop that is propagated independently by each of the ≈12 bulb growers in the Pacific Bulb Growers’ Association (Zlesak et al., 2007). Four to five of these 12 growers supply 75% of the ‘Nellie White’ clones for the market (Zlesak, 2006). Each bulb grower selects and propagates their own ‘Nellie White’ variants as a result of periodic screens of new, favorable lines (Zlesak and Anderson, 2003). This has the potential to allow for differing ‘Nellie White’ intraclonal mutational lines (ramets) to be selected, propagated, and produced by each wholesale bulb grower. Continuous asexual propagation of ‘Nellie White’ without periodic sexual cycles and meiotic purging of deleterious recessive alleles has allowed for operation of Muller’s ratchet (Anderson and Ascher, 1994; Muller, 1964). This has led to significant clonal decline [increases in sterility, lily symptomless virus titer (Anderson and Ascher, 1994; Zlesak and Anderson, 2007)]. Previous widely used easter lily clones (e.g., ‘Croft’ and ‘Ace’).

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also experienced clonal decline and were dropped from production (Zlesak and Anderson, 2003).

The potential for mutations within ‘Nellie White’ due to clonal propagation allows for morphological variation to surface in the clone (Veilleux and Johnson, 1998). This has often been dismissed with the assertion that most variation is caused by environmental factors rather than genetic ones (Fisher and Lieth, 2000). Environmental factors, such as the bulb grower’s field conditions, have caused greenhouse bulb forcers to routinely purchase from at least two bulb growers annually to minimize wide forcing variation for early Easter dates (Zlesak and Anderson, 2003). Populations of ‘Nellie White’ exhibited variability in days to stem emergence, visible flower bud, flower anthesis, plant and inflorescence height and number of leaves, bulb weight and circumference, and number of stems and flowers (Zlesak and Anderson, 2007). While the ‘Nellie White’ phenotype varied widely between bulb growers within years (2002 and 2003), this variation did not disappear when the environmental component was standardized in subsequent forcing cycles (Zlesak and Anderson, 2007). Thus, it would appear that a significant portion of the ‘Nellie White’ phenotypic variation may also arise from the genetic component because, by definition, phenotype = genotype × environment (G × E).

Objectively determining genetic variability in asexual crops (i.e., easter lily) can be performed irrespective of environmental variation, such as bulb growers’ field conditions (Veilleux and Johnson, 1998). Many protein and molecular DNA techniques are available for such research endeavors (Beckman and Soller, 1986). Polymorphisms of bulb scale proteins using isoelectric focusing have been used in Lilium for cultivar identification (Booy et al., 1998). Randomly amplified polymorphic DNA (RAPD) markers have been successfully used to detect genetic variation in many horticultural species and clonal cultivars; for example, in Camellia (Devarumath et al., 2002), Rosa (Debener et al., 1996; Torres et al., 1993), Lilium (Dai et al., 2007; Lee et al., 1993; Persson et al., 1998), and Scaevola (Swoboda and Bhalla, 1997). RAPD markers were used to infer phylogenetic relations and resolve paternity issues in wide hybrid lily crosses (Wen and Hsiao, 2001; Yamagishi, 1995). RAPDs, however, are often not reproducible or may lack sufficient markers and limit inferences about closely related clonal ramets (Yamagishi, 1995).

Polymerase chain reaction (PCR)-based approaches are more commonly used due to their simplicity, requiring only small DNA quantities (Devarumath et al., 2002). Intersimple sequence repeats (ISSRs) or intermicrosatellites are arbitrary multiloci markers produced by PCR amplification with a microsatellite primer. ISSR markers exhibit high polymorphism due to variability in the number of tandem repeats present in these sequences (Levinson and Gutman, 1987). ISSR fingerprinting is more reproducible than RAPD amplification due to the longer SSR-based primers, thus enabling higher-stringency DNA amplifications (Williams et al., 1990). It is a powerful tool for investigating genetic variation of closely related individuals that exhibit low levels of polymorphism (Godwin et al., 1997; Hale et al., 2005; Zietkiewicz et al., 1994). ISSRs have been used in fingerprinting and genetic analyses of several horticultural crops [e.g., Olea europaea (Hess et al., 2000), Magnifera indica (Liu et al., 2007), Chrysanthemum (Wolf et al., 1995), Leucadendron (Pharmawati et al., 2005), asiatic Lilium (Abe et al., 2002; Yamagishi et al., 2002), L. longiflorum hybrids (Wang et al., 2009), and interspecific (L. ×formolongi) × L. martagon hybrids (Anderson et al., 2009)].

ISSRs have also been used in the construction of genetic maps (Wu and Tanksley, 1993) and to detect clonal diversity by fingerprinting closely related genotypes (Camacho and Liston, 2001; Esselman et al., 1999; Mayes et al., 1998; Rongwen et al., 1995; Zietkiewicz et al., 1994). The hypervariability of ISSR markers enables small differences to be detected, even at the subspecies level (Wolfe et al., 1998). ISSRs were used to detect subtle somaclonal differences in asexually propagated Camellia (Devarumath et al., 2002) and Phalaris (Gyulai et al., 2003). In Camellia, ISSRs detected 8.5% more polymorphic loci than did RAPDs (Devarumath et al., 2002).

Amplified fragment length polymorphisms (AFLPs) are random, robust genome-wide markers providing high sensitivity with many polymorphic markers: primer pair (Ridout and Donini, 1999; Vos et al., 1995). Highly repetitive DNA sequences or large genome sizes frequently inhibit the use of AFLPs (Fay et al., 2005). Such has proven to be the case in L. longiflorum with its 77.1 pg/2C nucleus (Lim et al., 2001), a genome ≈550× that of Arabidopsis thaliana (Bennett et al., 1982). Lilium henryii was found to contain ≈13,000 copies of a single retrotransposon (Smyth et al., 1989). As a result, Zlesak et al. (2007) could not use AFLPs to detect clonal differences of L. longiflorum ‘Nellie White’ due to a lack of repeatability and consistency for scorable bands. Thus, AFLPs cannot be used to assess genetic variability and clonal integrity in L. longiflorum.

The objective of this study was to employ ISSRs to determine whether genetic differences exist between clonal ramets of L. longiflorum ‘Nellie White’ [obtained from the easter lily bulb growers and the Easter Lily Foundation (Brookings, OR) in 2002 and 2003] used in morphological (Zlesak and Anderson, 2007) and AFLP studies (Zlesak et al., 2007).

Materials and Methods

DNA samples of most L. longiflorum ‘Nellie White’ genotypes previously analyzed for morphological variation in two forcing cycles (Zlesak and Anderson, 2007) and AFLP analyses (Zlesak et al., 2007) were available for this study (Table 1). A total of 144 (2002) and 165 (2003) ‘Nellie White’ genotypes, and comparison genotypes of L. longiflorum ‘Ace’ clones (two), three seedlings of closely related L. formosanum (section Leucolorion), and one L. ×hybridum was used (Table 1). DNA extraction from the ‘Nellie White’ bulbs and all others was conducted after the completion of forcing cycle 2 in Zlesak and Anderson’s (2007) morphological and greenhouse forcing study. ‘Nellie White’ clonal bulbs had been randomly collected from lily bulb growers in 2002 and 2003 by two independent parties to keep grower identities confidential (Zlesak and Anderson, 2007). Each ‘Nellie White’ genotype was coded with the year of collection (2002 or 2003), grower number, and bulb number (e.g., 5-2-1 designated a bulb collected from grower 5 in 2002 and bulb No. 1). Two ‘Ace’ clonal genotypes were obtained from L. Riddle (Easter Lily Research Foundation) in 2002. Three Lilium formosanum genotypes included 8265–4 (Wayside Gardens, Hodges, SC), 92–224–1 [North American Lily Society Seed Exchange (Eau Claire, WI), sourced in 2002 from Florida, code designation 4–1 (Zlesak, 2006)], and 92–225–1 [North American Lily Society Seed Exchange, sourced in 2002 from Avonsleigh, Australia, code designation 5–1 (Zlesak, 2006)]. Lilium ×hybridum W-2 was an orange-flowered oriental lily (Lilium sp.) × tiger lily (L. tigrinum) hybrid from W. Gray’s...
lily breeding program (University of Minnesota, Morris) obtained in 2001.

**PCR conditions.** All previously extracted DNA samples (Qiagen kit; Qiagen, Valencia, CA) at a concentration of 1 ng·µL⁻¹ (Zlesak et al., 2007) of sufficient volume were diluted 1:100 (v/v) in deionized, distilled water before conducting ISSR analyses. Five UBC (Michael Smith Laboratories, 2008) primers from an initial screening process (Y. Sun, unpublished data) that had a high level of polymorphism and scorability were used for ISSR, PCR amplification (Table 2). Each PCR contained 0.25 units of Flexitaq DNA polymerase (Promega, Madison, WI), 20 µM of a single primer, 10 mM dNTP, 1 mM MgCl₂, 2 µL of diluted DNA solution, and 5× Flexi buffer ™ (Promega), which was supplied with the polymerase, for a total volume of 25 µL in each reaction (Yamagishi et al., 2002). Amplifications were carried out in a thermocycler (PTC-100; MJ Research, Hayward, CA) for 7 min of initial heating at 94 °C followed by 50 cycles of temperatures (denaturing at 94 °C for 30 s, annealing at 43 °C for 30 s, and gel electrophoresis (three replications) for all genotypes in a microwave at high power for ≈2 min until melted. Ethidium bromide was added to the gel (final concentration of 0.5 µg·mL⁻¹) at this point to facilitate DNA visualization. Electrophoresis chambers were filled with 1× Tris Acetate EDTA plus 5 µg·mL⁻¹ ethidium bromide (Sambrook et al., 1989). Sample DNA volumes (12 mL) were then loaded and a current of 75 V was applied for 2 to 2.5 h. Gels were visualized under ultraviolet light and were recorded with a Fluro Chem 500 camera (Alpha Innotech, Santa Clara, CA).

**GEL ELECTROPHORESIS.** A 1.5% agarose (mixed with 100 mL of electrophoresis, 1× Tris Acetate) buffer was made and heated in a microwave at high power for ≈2 min until melted. Ethidium bromide was added to the gel (final concentration of 0.5 µg·mL⁻¹) at this point to facilitate DNA visualization. Electrophoresis chambers were filled with 1× Tris Acetate buffer plus 5 µg·mL⁻¹ ethidium bromide (Sambrook et al., 1989). Sample DNA volumes (12 mL) were then loaded and a current of 75 V was applied for 2 to 2.5 h. Gels were visualized under ultraviolet light and were recorded with a Fluro Chem 500 camera (Alpha Innotech, Santa Clara, CA).

**DATA ANALYSES.** Unequivocally scorable and consistently reproducible amplified DNA fragments across all three replications were transformed into binary character matrices (1 = presence, 0 = absence). Cross Checker (Buntjer, 1999) was used to score and analyze binary interpretation of DNA fingerprints. To minimize variation, gels were equilibrated to comparable background coloration and alignment using the Cross Checker Gel Manager macro. Intra- and intergroup (population) variations were measured using Hedrick’s index [I (Hancock and Bringham, 1979)]. To obtain interpopulation variance estimates, Russell and Rao (1940) coefficients of similarity were calculated. Clustering and dendograms for individual growers or pooling all growers within each year were created by the unweighted pair group method (UPGMA) using the SANN-clustering and TREE programs from NTSys (version 2.02; Exeter Software, Setauket, NY). ISSR primer data for all bulb growers were pooled by years (2002 and 2003) to evaluate the genetic variation among growers. It was not feasible to also pool all growers and years together into one analysis due to software limitations: i.e., Excel® (Microsoft, Redmond, WA) did not have sufficient line numbers to handle the number of data sets. Genetic diversity among and within populations of ‘Nellie White’ clones and comparisons (binary diploid data) were estimated with an analysis of molecular variance (AMOVA) in GenAlEx (Peakall and Smouse, 2006). The Ht tested = there is no genetic difference among the populations (Φpt = 0); i.e., they are

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### Table 1. *Lilium* bulb genotype samples from easter lily bulb growers (Smith River, CA and Brookings, OR) in 2002 and 2003 and comparison species/cultivars used for intersimple sequence repeat analysis.

| Species           | Cultivar         | Bulbs sampled (no.) |
|-------------------|------------------|---------------------|
|                   | Grower | 2002 | 2003 |
| *L. longiflorum*   | Nellie White | 14  | 14  |
|                   | 2       | 14  | 14  |
|                   | 3       | 13  | 15  |
|                   | 4       | 14  | 14  |
|                   | 5       | 14  | 14  |
|                   | 6       | 14  | 14  |
|                   | 7       | 14  | 14  |
|                   | 9       | 0   | 14  |
|                   | 11      | 14  | 14  |
|                   | 12      | 14  | 14  |
|                   | 13      | 14  | 14  |
|                   | 14      | 5   | 10  |
| Total no. of bulbs|       | 144 | 165 |

**Comparisons**

- *L. longiflorum* Ace
- *L. formosanum* 8265–4, 4–1, 5–1
- *L. ×hybridum* W-2

### Table 2. Five University of British Columbia [UBC (Michael Smith Laboratories, 2008)] intersimple sequence repeat primer configurations of 5’-3’ sequence, length in basepairs, melting temperature (Tm), total loci, and total polymorphic loci or amplicons used with *Lilium longiflorum* ‘Nellie White’, ‘Ace’ clones, species, and hybrids in this experiment.

| UBC primer | Sequence | Length (bp) | Tm (°C) | Total loci (no.) | Total polymorphic loci [no. (%)] |
|------------|----------|-------------|---------|-----------------|----------------------------------|
| 808        | 5’-(AG)₆C-3’ | 17          | 48.8    | 14              | 12 (85)                           |
| 810        | 5’-(GA)₆T-3’ | 17          | 45.4    | 17              | 11 (64)                           |
| 811        | 5’-(GA)₆C-3’ | 17          | 46.8    | 16              | 11 (68)                           |
| 814        | 5’-(CT)₆A-3’ | 17          | 44.7    | 15              | 11 (73)                           |
| 818        | 5’-(CA)₆G-3’ | 17          | 51.0    | 15              | 11 (73)                           |
| **Total**  |          | 77          |         | 56              | 56 (72.7)                         |

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**Table 2. Five University of British Columbia [UBC (Michael Smith Laboratories, 2008)] intersimple sequence repeat primer configurations of 5’-3’ sequence, length in basepairs, melting temperature (Tm), total loci, and total polymorphic loci or amplicons used with *Lilium longiflorum* ‘Nellie White’, ‘Ace’ clones, species, and hybrids in this experiment.**
clones that have the opportunity to cross-pollinate due to close proximity of all easter lily bulb growers and, thus, may all be “part of a single large random mating genetic population” (Peakall and Smouse, 2006).

**Results**

A total of 77 well-defined (clear and unequivocal bands) and scorable markers across replications were obtained from five of the selected ISSR primers, 56 (72.7%) of which were polymorphic (Table 2). Replications did not differ significantly in their banding patterns. The size of the scorable amplified fragments ranged from 1018 to 8144 bp. Total numbers of scorable loci/prime ranged from 14 (Primer 808) to 17 (Primer 810) for ‘Nellie White’ clones and comparison genotypes (Table 2), whereas the number of scorable, polymorphic loci/prime ranged from 11 (all primers, except Primer 808) to 12 (Primer 808 only) (Table 2). This set of 56 polymorphic marker loci enhanced the stringency for discernment of ‘Nellie White’ clones.

As would be expected with varying numbers of polymorphic loci for each of the five primers, differing numbers of scorable bands for ‘Nellie White’ populations were found. For example, gel profiles of ‘Nellie White’ ramets obtained from grower 7 for 2002 and 2003, using Primers 810 (Fig. 1A) and 811 (Fig. 1B), show differing numbers of scorable bands. Bulb 7-2-2 has three scorable bands with Primer 810 (Fig. 1A, lane 3), whereas only one band is scorable with Primer 811 (Fig. 1B, lane 3). As would be expected, because the 2002 and 2003 samples were randomly collected from each grower, rarely did any bulb genotype from 2002 have a comparable amplified fragment banding pattern with bulbs harvested in 2003. Because bulb numbers were assigned at random within each grower lot and year, one would not expect any relationship between genotypes with the same bulb number if the ‘Nellie White’ clones are genetically divergent, other than what might occur by chance alone. Comparable, scorable loci for ramet 7-2-2 with only one scorable locus using Primer 811 (Fig. 1B, lane 3) is expressed in the 2003 genotypes 7-3-1, 7-3-5, 7-3-8, 7-3-10, 7-3-11, and 7-3-14, although each of these also have additional scorable loci for this primer and, thus, are not identical to 7-2-2 as would be expected in a clone (Fig. 1B).

Similar analogies commonly surfaced with all clonal ramets from the same bulb grower between years and within primers. Thus, ‘Nellie White’ clonal ramet populations among growers—either between or within sampling years—were not homogeneous for the ISSR polymorphic loci and could not be pooled (Table 3).

**Fig. 1.** Intersimple sequence repeat gel profiles of *Lilium longiflorum* ‘Nellie White’ clones from grower 7 in years 2002 and 2003 using Primers 810 (A) and 811 (B). For bulb genotypic code designators, refer to the text; M = molecular marker (1-kb DNA ladder; Takara, Madison, WI).

Constructed dendrograms for each bulb grower for 2002 and 2003 differed widely, with no two growers being identical. For instance, for bulbs obtained from grower 11 in year 2002, the dendrogram delineates ramets into two principal groups separating at Nei’s (1972) genetic distance of GD = 0.63 (Fig. 2), with only two subsequent monophyletic singletons (a set with exactly one genotype or clonal ramet) found (11-2-13, 11-2-4). Each principal group further separated into 2, 3, or 4 additional overlapping, contiguous clades (contigs) (Fig. 2). Pairs of clones (bulbs) separated at low to higher GDs, e.g., GD = 0.69 for 11-2-6 and 11-2-14; GD = 0.73 for pairs 11-2-7/11-2-3 and 11-2-11 and 11-2-5; GD = 0.75 for pairs 11-2-8/11-2-9 and 11-2-10/11-2-1; and GD = 0.8 for 11-2-2/11-2-12 (Fig. 2). Thus, in this example, the most closely related clones from grower 11 in 2002 are 11-2-2 and 11-2-12 (GD = 0.8).

In contrast with the findings for grower 11 in 2002 (Fig. 2), the principal clades in grower 3 for 2003 bulbs separated at a higher GD of 0.67 (Fig. 3). This separation constituted one singleton, genotype 3-3-13, with all remaining bulbs in a separate clade, which subsequently split into several subclades. The most closely related ‘Nellie White’ clonal genotypes, 3-3-4 and 3-3-5, had a GD = 0.89 (Fig. 3). Thus, these two bulbs are even more related than 11-2-2 and 11-2-12 (GD = 0.8, Fig. 2). A summary of GD values (principal clade separations, most-related
bulbs) from ISSR dendrograms for individual growers for each year (2002 and 2003) are summarized in Table 4. Principal clades ranged from GD = 0.60 (growers 2, 4, and 12) to GD = 0.82 (grower 6) for 2002 bulbs, whereas the 2003 bulbs resulted in GD = 0.51 (grower 4) to GD = 0.78 (grower 14) (Table 4). Across years 2002 and 2003, grower 4 consistently had the lowest principal clade. However, grower 5 had identical principal clade separations of GD = 0.71 in both years (Table 4). The remaining bulb growers had lower or higher principal clade separation GDs between harvest years.

GD values for the most-related bulbs in 2002 ranged from GD = 0.80 (growers 11 and 12) to GD = 0.95 (grower 6) (Table 4). A wider range of values for the most-related 2003 bulbs: GD = 0.70 (grower 4) to GD = 0.91 (grower 6) (Table 4). Again, grower 5 had an identical most-related bulb GD value of 0.89 for 2002 and 2003. While the GD values cannot be statistically analyzed within growers and among years, the trends among years for principal group separations and most-related bulb GDs (Table 4) are intriguing. Six growers had lower GDs between 2002 and 2003 for both categories (growers 1, 2, 4, 6, 7, and 11); one grower had higher GDs between 2002 and 2003 for both categories (grower 12). Two growers had lower principal clade separations and higher mean related bulb GDs (growers 3 and 13); one grower had higher principal group separations and lower mean related bulb GDs (grower 14). One grower had equal values between years (grower 5).

When years were coanalyzed for each ‘Nellie White’ bulb grower population, the principal clade separations occurred for years or the years were scattered between both clades. In the case of grower 7, all 2003 genotypes except one (7-3-10) were in one principal clade while the 2002 bulbs plus 7-3-10 were in the other, separating at GD = 0.53 (Fig. 4). Genotype 7-3-10 (2003) was most similar to 7-2-9 and 7-2-13 (2002), separating at GD = 0.695 (Fig. 4). With other growers, the 2002 and 2003 bulbs were spread between principal clades. In grower 5, ‘Nellie White’ bulbs, the principal clades separated at GD = 0.65 into

Table 3. Analyses of molecular variance (AMOVAs) for Lilium longiflorum ‘Nellie White’ sourced from bulb growers in 2002 (11 populations), 2003 (12 populations), and one comparison population of cultivars (Table 2) on the basis of intersimple sequence repeats (ISSRs).

| Source of variation | 2002 Sum of estimate | Variation | df | Squares of variation (% total) | 2003 Sum of estimate | Variation | df | Squares of variation (% total) |
|---------------------|----------------------|-----------|----|--------------------------------|----------------------|-----------|----|--------------------------------|
| Among populations   | 12                   | 314.3     | 1.69*** | 18.0                           | 13                   | 402.5     | 2.01*** | 18.0                           |
| Within populations  | 138                  | 1075.8    | 7.97   | 82.0                           | 158                  | 1314.7    | 9.01  | 82.0                           |
| Total               | 150                  | 1390.1    | 9.66   | 100.0                          | 171                  | 1717.2    | 11.02 | 100.0                          |

*Genetic distance between populations ($\Phi_{PT}$) = 0.175 ($P \leq 0.01$).

**$\Phi_{PT}$ = 0.183 ($P \leq 0.01$).

Fig. 2. Dendrograms of relatedness for Lilium longiflorum ‘Nellie White’ clones (grower 11, year 2002) derived from polymorphic, scorable bands for all five intersimple sequence repeat primers used (808, 810, 811, 814, and 818) and based on using Nei’s (1972) coefficients.

Fig. 3. Dendrograms of relatedness for Lilium longiflorum ‘Nellie White’ clones (grower 3, year 2003) derived from polymorphic, scorable bands for all five intersimple sequence repeat primers used (808, 810, 811, 814, and 818) and based on using Nei’s (1972) coefficients.
one singleton (5-3-7) and all other 2002 and 2003 bulbs (Fig. 5). These principal clades then subsequently split into two subclades at GD = 0.67 with a) 10 genotypes (2003) and three genotypes (2002), b) 11 genotypes (2002) and three genotypes (2003). Such inconsistencies between years within growers were common for all bulb growers (data not shown). Thus, non-uniform clones exist with all ‘Nellie White’ bulb growers for each year (2002 and 2003), and the level of genetic variation is also inconsistent between years.

When bulbs from all the growers in year 2002 were pooled together, the principal clades separated at GD = 0.54 with the three L. formosanum genotypes (4-1, 5-1, and 8265-4) in one monophyletic clade and all other genotypes (all ‘Nellie White’, ‘Ace’ clones, and L. ×hybridum W-2) in the second clade (data not shown). Within L. formosanum, 8265-4 separated into a subclade at GD = 0.75 with genotypes 4-1 and 5-1 differentiated at GD ≈ 0.76 (data not shown). The other principal clade separated at GD = 0.573 into one subclade with all of grower 7 ‘Nellie White’ genotypes and the other clade containing all of the remaining ‘Nellie White’, ‘Ace’, and W-2 genotypes (data not shown). Clearly, in 2002, there is significant genetic variation between grower 7 and all other ‘Nellie White’ bulb growers. The next subclade within L. longiflorum split at GD = 0.594 with all of grower 11 ‘Nellie White’ genotypes in one subclade, completely separate from the remaining clones (data not shown). Bulbs from grower 11 in 2002 had enough distinctive genetic variation to distinguish them from growers 1-6, 9, and 12-14. Following the first subclade, there were a total of 15 subgroupings splitting the genetic variation between ‘Nellie White’ bulb growers, ‘Ace’, and W-2. No clear patterns emerged with ‘Nellie White’ bulbs from each

Table 4. Genetic distance (GD) coefficients for intersimple sequence repeats using five University of British Columbia (UBC) primers [808, 810, 811, 814, and 818 (Michael Smith Laboratories, 2008)] for the 2002 and 2003 bulb lots from easter lily bulb growers of Lilium longiflorum ‘Nellie White’. GDs of the principal group separations and the most-related bulbs within each bulb lot are derived from polymorphic, scorable bands for all five primers, based on Nei’s (1972) coefficients.

| Grower no. | 2002 GDs | 2003 GDs |
|------------|----------|----------|
| Principal clad | Most-related separation bulbs | Principal clad | Most-related separation bulbs |
| 1 | 0.74 | 0.93 | 0.58 | 0.84 |
| 2 | 0.60 | 0.85 | 0.54 | 0.83 |
| 3 | 0.73 | 0.85 | 0.67 | 0.89 |
| 4 | 0.60 | 0.85 | 0.51 | 0.70 |
| 5 | 0.71 | 0.89 | 0.71 | 0.89 |
| 6 | 0.82 | 0.95 | 0.74 | 0.91 |
| 7 | 0.63 | 0.85 | 0.58 | 0.80 |
| 9 | — | — | 0.59 | 0.89 |
| 11 | 0.63 | 0.80 | 0.61 | 0.75 |
| 12 | 0.60 | 0.80 | 0.62 | 0.83 |
| 13 | 0.61 | 0.83 | 0.53 | 0.84 |
| 14 | 0.76 | 0.89 | 0.78 | 0.85 |

Fig. 5. Dendrograms of relatedness for Lilium longiflorum ‘Nellie White’ clones (grower 7 pooled for years 2002 and 2003) derived from polymorphic, scorable bands for all five intersimple sequence repeat primers used (808, 810, 811, 814, and 818) and based on using Nei’s (1972) coefficients.
grower dispersed narrowly or widely through multiple sub-clades (data not shown).

In contrast with the 2002 dendrogram, when the 2003 bulb lots were pooled for all growers (data not shown), the principal clades consisted of a) six ‘Nellie White’ genotypes (7-3-08, 2-3-06, 4-3-08, 13-3-02, 13-3-04, 11-3-04, and 13-3-05) and b) all other L. longiflorum, L. formosanum, and L. ×hybridum accessions. The latter principal clade subsequently split into two subclades at GD = 0.524 containing a) three genotypes: L. formosanum 5-1, 4-1, and L. longiflorum ‘Nellie White’ 14-3-04; b) all other remaining genotypes in the experiment (data not shown). Subclade b then split at GD = 0.557 into two subgroups (ba) seven bulbs from grower 1 (1-3-03, 1-3-02, 1-3-05, 1-3-06, 1-3-07, 1-3-08, and 1-3-04) and one bulb from grower 5 (5-3-07); bb) all remaining genotypes (data not shown). The latter subgroup bb split again into 18 additional subgroupings with scattered representation of bulbs from each grower spread narrowly or broadly across subgroupings (data not shown). Genetic variation displayed in the 2003 bulb lots differed widely from the variation in 2002 with an even more diverse set of ‘Nellie White’ clones, some of which exceeded the genetic variation found between L. formosanum, L. ×hybridum, and L. longiflorum.

AMOVAs showed higher levels of genetic diversity within (82% of variation), rather than among (18%), ‘Nellie White’ and comparison populations (Table 3). Idetical percentages of variation results were found within and among populations across years (2002 and 2003), although the sums of squares and df differed (Table 3). For ‘Nellie White’ bulbs from 2002, Φπ = 0.175 and the null hypothesis were rejected (P < 0.01). Thus, all ‘Nellie White’ clones are not part of a single, random-mating population (Peakall and Smouse, 2006). Likewise, in 2003, Φπ = 0.183 (P < 0.01).

Discussion

ISSRs distinguished among- and within-bulb grower populations of ‘Nellie White’ clones (ramets) and confirmed the lack of a single clone for this cultivar. The five selected UBC primers, with A, T, C, or G anchors at the 3’ end, provided 56 polymorphic, unequivocally scorable markers of fragment sizes 1018 to 8144 bp (Table 2). A, T, C, and G anchors, C-T repeats, or C-A and G-A repeats at the 3’ end have been useful for ISSR markers in Glycine (Wang et al., 1998), asiatic Lilium (Yamagishi et al., 2002), and interspecific Lilium hybrids (Anderson et al., 2009; Wang et al., 2009). This set of polymorphic marker loci enhanced the stringency for discernment of ‘Nellie White’ clonal ramets. The number of primers:scorable markers used for L. longiflorum ‘Nellie White’ [5:56 (Table 2)] is similar to previous reports involving ISSRs for L. longiflorum ‘Gelria’ [4:38 (Wang et al., 2009)] and within species or closely related clones {2:56 for Linaria vulgaris populations (Ward et al., 2008), 4:14 for the mother ortet (the original hybrid plant from which clones or ramets are derived) and clonal ramets (vegetative propagules derived from the original ortet or subsequent ramets) of Camellia (Devarumath et al., 2002), but slightly smaller than those for more divergent species analyses [12:103 for Prunus (Liu et al., 2007)].

‘Nellie White’ clonal ramets among growers—between or within sampling years—are not homogeneous for the ISSR polymorphic loci and are not part of a single, random-mating population (Table 3; Peakall and Smouse, 2006). Rather, there are several random-mating grower populations. Within growers, as few as two ramets shared similarity with each other, ranging from Nei’s GD = 0.7 [grower 4, 2003 (Table 4)] to GD = 0.95 [grower 6, 2002 (Table 4)]. Within years, the most...
closely related clones among growers for 2002 were four at GD = 0.96: 4-2-13 and 13-2-3; 2-2-2 and 4-2-2 (data not shown) and in 2003 10 clones at GD = 0.98: 7-3-3 and 14-3-3, 5, 6; 14-3-1 and 3-3-14, 15; 14-3-9, 10 and 3-3-10 (data not shown) representing 2.78% [4/144 (2002)] and 6.1% [10/165 (2003)] nuclear genome similarity. Conversely, 97.2% (2002) and 93.9% (2003) of ‘Nellie White’ were clonally variable. These levels of nuclear genome variation are significantly higher than those reported for Coffea [4.38% (Rani et al., 2001)] and Camellia [7.43% (Devarumath et al., 2002)]. The nearly identical ‘Nellie White’ clones involved a majority of growers—three in each year [growers 2, 4, and 13 (27.3% in 2002); growers 3, 7, and 14 (25% in 2003)]—rather than a majority, as would have been expected with a clonal cultivar with absolute genetic fidelity. These low levels of ISSR genetic similarity within ‘Nellie White’ resemble what has been previously reported for clonal Camellia (Devarumath et al., 2002), Achillea (Wallner et al., 1996), Populus (Rani et al., 1995, 2001), and Coffea (Rani et al., 2000).

Because the identical 2002 and 2003 harvested ‘Nellie White’ bulbs used in this ISSR study were also evaluated in two forcing cycles (FC) to record phenotypic (morphological) responses (Zlesak and Anderson, 2007), comparative analyses of the common genotypes is possible herein (even though ISSR markers are usually neutral in expression). The FCs consisted of FC1 (freshly harvested bulbs forced to flower the following spring) and FC2 (reforcing the FC1 bulbs an additional year) for the 2002 and 2003 harvested bulb lots from all growers. All FC1 bulbs, while grown in grower fields of close proximity in Brookings, OR, and Smith River, CA, had differing environmental factors that could compound G × E interactions for phenotypic analyses only (Zlesak and Anderson, 2007). In contrast, the FC2 bulbs had the uniform greenhouse FC1 and subsequent cooling environments that minimized the G × E variation inherent in FC1. Zlesak and Anderson (2007) evaluated critical commercial greenhouse forcing [e.g., quick emergence and flowering, particularly for early Easters, high flower bud number, and shorter plant height (Lange and Heins, 1990; Wilkins and Roh, 1976)], and other traits that make an ideal flowering potted plant. In general for all traits, grower 5 ranked among the top five growers (1, 3, 4, 5, and 6) for all key traits in 2003 FC1 and all traits except plant height in 2002 FC1, and number of days to stem emergence in 2003 FC2 (Zlesak and Anderson, 2007). Grower 3 was also ranked in the top five for all traits, except for days to stem emergence in 2003 FC2 and plant height in 2002 FC1, 2003 FC2. In 2002, growers 3 and 5 ‘Nellie White’ genotypes were clustered across the ISSR subgroup separating at GD = 0.594 that included all top five growers [1, and 3–6 (Zlesak and Anderson, 2007)], plus a couple of outlier genotypes from growers 12 and 13. They were distributed across similarly large ISSR subclades in 2003 (separating at GD = 0.557) that included most of the top five growers (plus a few outliers from grower 4), but others as well.

The number of days to stem emergence in 2002 FC1 followed the trend for the pooled morphological data, with growers 3 and 5 emerging significantly earlier than growers 2, 7, 11 through 14, and all of the top five growers (1, and 3–6) overlapping (Zlesak and Anderson, 2007). Likewise, visible bud and anthesis dates were significantly earlier for the top five growers in 2002 FC1 (Zlesak and Anderson, 2007). These top five growers were in the same ISSR subclade (data not shown), as noted earlier. In 2003 FC1, only grower 11 emerged significantly later than growers 3, 5, and 13, while the number of days to visible bud were earliest for growers 3 and 5 (Zlesak and Anderson, 2007), but the ISSR genetic distances showed a widespread distribution across clades for grower 11 bulbs (Fig. 6B). In contrast, for 2003 FC2, only grower 4 was significantly earlier to emerge than grower 1 while the rest overlapped (Zlesak and Anderson, 2007). Genetic distances of growers 1 and 4 bulbs were intermixed across the dendrogram (data not shown).

Flower number differed slightly from trends noted with days to emergence, visible bud, and anthesis in 2002 FC1 with growers 1, 3, 5, 6, and 7, and grower 13 in 2003 FC1 (rather than growers 1, and 3–6) having the significantly highest bud counts (Zlesak and Anderson, 2007). Flower number does not fit the ISSR GD configurations quite as tightly as emergence and flowering traits. Plant height and leaf measurements (length, width, and length:width ratios) also followed a similar trend. Thus, ISSR genetic distances only matched some (days to emergence, visible bud, and anthesis), but not all, morphological traits for the top five growers in 2002 and 2003. These results are not unusual, as “functional” phenotypic variation is only inferential but “cannot be measured directly” (Ward et al., 2008).

Higher levels of genetic diversity were consistently found within (82%) than among ‘Nellie White’ grower populations for each of the years sampled (Table 3). Thus, only an 18% difference in genetic variation exists between growers—an astounding fact since each grower serves as their own propagator of ‘Nellie White’ (Zlesak and Anderson, 2003, 2007). However, the high within-grower variation is significant and reflective of each grower deliberately selecting from among differing clonal ramet populations for propagules (Zlesak and Anderson, 2003, 2007), causing genetic drift. Low among-grower variation levels may be influenced by growers purchasing bulbs from their competitors to generate additional ramets. Despite the lack of a central propagator to ensure clonal integrity, bulb growers may select a suite of superior ramets with a similar range in phenotypes (82%) to ensure that greenhouse forcers can produce a marketable crop each year. Even though grower field conditions can vary significantly, the close proximity of bulb growers may create enough environmental similarity among growers that favor a similar range of ramet selections. In years when Easter is early, the 18% among grower variation could be enough to offset reaching the target flowering date if a grower were, by chance, to purchase bulbs that performed significantly different from the norm (82%). Commercial greenhouse bulb forcers of easter lily—both currently for ‘Nellie White’ and previously with older clones [‘Ace’, ‘Croft’, ‘Chetco’, ‘Harbor’ (Roberts et al., 1968)]—have been advised to purchase from multiple growers each year to avoid chance variation in bulb lots, which could mean missing the Easter target flowering date, particularly when Easter is early (Zlesak and Anderson, 2003, 2007). Phenotypic variation during the four forcing phases (i.e., shoot elongation, flower initiation, flower development, and senescence) can be environmentally manipulated by the grower to ensure that the crop flowers for Easter (Fanelli and De Hertogh, 2002; Roberts et al., 1983). Erwin and Engelen-Eigles (1998) postulated that phenotypic variation in potted easter lily (days to emergence, leaf and flower number, height, and internode length) was also attributable to bulb maturity, dormancy level differences (bulb lots and years), and environmental factors during shipping and greenhouse forcing. Zlesak and Anderson (2007) forced the same bulbs
tested in our present ISSR analysis over two forcing cycles and found, among bulb lots, the existence of significant morphological variation for all traits except visible bud and anthesis dates. The low (18%) level of genetic variation between grower bulb lots may be a significant underlying genetic/epigenetic factor affecting the significant phenotypic (morphological) differences operating in tandem with other inherent (lily symptomless virus titer) or external (initial bulb size, the previous forcing cycle environment) factors (Roberts et al., 1968; Zlesak and Anderson, 2007).

It is unknown whether there were one or more ‘Nellie White’ bulbs (ortet/s) used to propagate this clone. Regardless of whether there were one or two ortets, both would have been submitted to multiple asexual generations without meiosis, which allows for the continued accumulation of mutations due to the operation of Muller’s ratchet (Anderson and Ascher, 1994; Muller, 1964). Clearly, there were several mutational populations, rather than one or two. Whether this is due to the origination of ‘Nellie White’ from one ortet or several, numerous populations of mutants have arisen during the >60 years of asexual propagation. The most closely related clones of ‘Nellie White’ in 2002 and 2003 had a wide range of GDs, i.e., 0.7 (grower 4, 2003) to 0.95 [grower 6, 2002 (Table 4)]. It would have been expected that all ‘Nellie White’ clones, particularly those from the same grower (within populations) would be closely related (GD ≈ 1.0) if they were derived from the same ortet and remained free of mutation(s). However, because 82% of the variation consistently occurred within populations (growers) in 2002 and 2003, it is more likely that mutations, regardless of the number of original ‘Nellie White’ ortets, are accumulating within growers bulb stock due to the operation of Muller’s ratchet (Anderson and Ascher, 1994; Muller, 1964). Due to a lack of the meiotic sieve of sexual cycles, every clonal (asexual) generation can accumulate deleterious recessive alleles (i.e., every asexual generation is a click of the ratchet). Eventually, such mutations can result in lowered fertility levels (male/female; Anderson and Ascher, 1994) or phenotypic differences that have caused growers to purchase from multiple bulb growers (Zlesak and Anderson, 2003, 2007). This scenario will continue in ‘Nellie White’ until such mutations become severe enough to necessitate clonal replacement. The large genome size of lily may predispose ‘Nellie White’ clones to survive numerous clicks of the ratchet before becoming phenotypically visible. This, coupled with the levels of viruses present (e.g., lily symptomless virus), which are removed only when the viral titer becomes high enough to stunt the plants, may have been the causal factor in the removal of ‘Ace’ and ‘Croft’ from the market after they began developing high rates of leaf scorch (Zlesak and Anderson, 2003).

Comparison cultivars and species also varied in their genetic relatedness to each year’s ‘Nellie White’ ramet populations. For instance, in 2002, all _L. formosanum_ genotypes were genetically distinct from ‘Nellie White’ clones as well as ‘Ace’ and _L. xhybridum_, separating into a separate clade at GD = 0.54 (data not shown). In contrast, _L. formosanum_ in 2003 were not in a distinct clade, but separated out at a higher GD (data not shown), with two genotypes (5-4 and 4-1) in the same subgroup as ‘Nellie White’ 14-3-4. ‘Ace’ clones and _L. xhybridum_ were in the middle of the entire ‘Nellie White’ clonal distribution, changing only slightly between years (data not shown). It would have been expected that _L. xhybridum_ would have been the most genetically distant from _L. longiflorum_ ‘Nellie White’ and/or ‘Ace’ across years, with _L. formosanum_ more closely related, since _L. formosanum_ and _L. longiflorum_ are in the Section Leucoleirion, whereas _L. xhybridum_ is from different sections of the genus (Booy et al., 1998; Yamagishi et al., 2002). Reasons for the genetic distinction of _L. formosanum_ from _L. longiflorum_ in 2002, but the closer similarity in 2003, are unclear. This could have been due to chance mutations accumulating with the operation of Muller’s ratchet (Anderson and Ascher, 1994; Muller, 1964) and/or inadvertent cross-hybridization between _L. longiflorum_ and _L. formosanum_ because the latter species has been grown at the Easter Lily Research Station, which is adjacent to several bulb growers’ fields. Grower 14’s location is unknown to the authors, but one of its ‘Nellie White’ bulbs, 14-3-4, was in the same 2003 subclade as _L. formosanum_ (data not shown). Also equally likely is the chance that 14-3-4 and _L. formosanum_ share common genetic makeup for the tested nuclear DNA sequences because ISSRs are arbitrary multiloci markers (Levinson and Gutman, 1987).

Analyzing populations of clonally propagated plants is difficult because a single genetic individual (ramet) may comprise many morphological units that appear distinct (Parks and Werth, 1993). Clearly, ‘Nellie White’ clones are genetically distinct from each other primarily within growers over years and, to a lesser extent, among growers. The current study is the first to use ISSR markers to characterize genetic variation in _L. longiflorum_ ‘Nellie White’ clones. Without molecular genetic investigations, such clonal drift would not have been detected. For instance, phenotypic variation showed clonal ramets from growers 3 and 5 as the best-performing sources (Zlesak and Anderson, 2007), but the molecular variation (ISSR data herein) show these populations to be intermediate in GD (Table 4). Nonetheless, the clonal variation in ‘Nellie White’ is significant and a cause for greenhouse growers to continue exercising caution to reach critical forcing landmarks for early Easters. There is also a need for increased clonal integrity within this important floricultural crop to ensure reduced genetic variation. Natural mutations (Veilleux and Johnson, 1998), Muller’s ratchet, and intraclonal ‘Nellie White’ selection by bulb growers results in different commercial strains within and among growers (Zlesak and Anderson, 2003). While only 18% of the variation is among growers, this significantly impacts greenhouse forcers, causing them to purchase from multiple growers. The Easter Lily Research Foundation could create an elite stock program, such as that developed for _Pelargonium_ disease control (Wager-Page et al., 2006), that incorporates within- and among-bulb grower genetic variation to ensure crop viability and avoid potential genetic drift. A central propagator to source propagules to the lily bulb growers should also be created.

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