Breeding programs ideally seek to identify parents that combine and generate a high number of superior progeny. The term combining ability defines this concept (Sprague and Tatum, 1942) and denotes how well a parent performs in producing superior progeny when crossed with many other parents [general combining ability (GCA)] or with specific individuals [specific combining ability (SCA)]. Procedures that enable the identification of desirable parental combinations are likely to facilitate the development of superior cultivars.

Systematically identifying desirable parental combinations requires 1) crossing all possible parents amongst themselves in a diallel cross, and 2) evaluating progeny for fitness for a given trait or in totality for all important attributes. Corn (Zea mays L.) has been served as a theoretical and practical model for determination of GCA- and SCA-based crop improvement. Its ease of crossing, genetically uniform hybrid progeny, and lack of incompatibility barriers are a few of the reasons why combining ability tests are routinely used in corn improvement programs. In contrast, sweetpotato (Ipomoea batatas) is poorly suited for SCA-based improvement. This crop has complex self- and cross-incompatibility barriers (Jones et al., 1986; Martin, 1982) that often prevent the successful hybridization of selected parents. Hence, sweetpotato breeding programs rely on the use of open-pollinated (polycross) nurseries to generate progeny. While the maternal parent in this type of hybridization procedure is easily identified, the paternal parent could potentially be any one of the 12 to 15 heterozygous pollen parents in the nursery. The only way to identify parents exhibiting SCA is to hand-cross-selected parents, generate the subsequent generation from seed, clonally increase the progeny, and evaluate the progeny for desirable characteristics. Few sweetpotato breeding programs, with the exception of the Japanese and Chinese programs (Yamakawa, 1989), can afford to evaluate lines for combining ability by hand-crossing. Yet, targeting specific parental crosses exhibiting SCA would significantly enhance the efficiency of sweetpotato breeding programs. We have found that the majority of the superior progeny retained in the LSU AgCenter sweetpotato breeding program can be traced to two or three of the original maternal parents. We speculate that paternity may be similarly limited to only a few lines. The identification of superior lines a priori, is seen as a means to increase the efficacy of the polycross nursery.

Two basic approaches are typically used to establish paternity; both were developed for human population studies and forensic applications, and are still not universally accepted in their entirety (Chakraborty et al., 1974; Thompson 1975, 1986; Valentin, 1980). The first approach, referred to as paternity exclusion, is based on near conclusive proof of nonpaternity as determined by parent–offspring marker genotype data (Chakraborty et al., 1988). Paternity exclusion compares the progeny genotype with the maternal genotype, subtracts the maternal contribution, and compares the remaining paternal gametic contribution with all putative paternal genotypes. The individuals that cannot provide the observed paternal gametic contribution are excluded, and paternity is assigned to the individual that remains. In large natural mating populations, the true paternal parent cannot be assigned with 100% certainty (Chakraborty et al., 1988). The second...
approach, referred to as the most-likely parent method, calculates paren-
ty likelihood based on segregation (Mendelian) probabilities. Paren-
ty is assigned to the putative parent with the maximum (numerically highest) log-likelihood or LOD score. In essence, progeny alleles (genetic markers) for a given locus are weighted (based on their relative frequency in the parental popula-
tion) as to the likelihood that they could have come from a given paternal parent. The more alike the two genetic marker patterns are, the higher the LOD score for the paternal parent. For example, an allele D that is scarce in a parental population would be uniquely invaluable in matching a lone parent and all progeny possessing this allele. In the case of tied LOD scores, no parent is selected. This approach allows paternity assignment to a higher number of progeny than paternity exclusion (Devlin et al., 1988; Smouse and Meagher, 1994). Caveats and details about these approaches are discussed in Buteler et al. (1997).

The characteristics of a genetic marker that make it suitable for paternity analysis are as follows: 1) the marker should be unambiguously inherited, 2) it should segregate independently in the population, and 3) it should lead to lower levels of ambiguity than the parentage uncertainty to be solved (Smouse and Meagher, 1994). Ideal markers for plant paternity analysis would disclose the parentage uncertainty to be solved (Smouse and Meagher, 1994). Caveats and details about these approaches are discussed in Buteler et al. (1997).

The second group consisted of seven parents (Nacional, Jarret and Bowen, 1994). SSR markers have been successfully applied on animal and plant diploid popula-
tions. Parentage is assigned to the putative parent with the maxi-
mum number of progeny than paternity exclusion (Devlin et al., 1988; Smouse and Meagher, 1994). Caveats and details about these

PLANT MATERIAL
Two groups of plant material were used. The
first group consisted of eight parents (´Resisto´, 86-33, NC-C75, 90-223, 91-153, ´Excel´, ´Beauregard´, and 80-62) from a popula-
tion from Louisiana Agricultural Experiment Station (LAES), North Carolina Agricultural Experiment Station, and USDA, U.S. Vegetable Laboratory breeding lines (LAES population). This group also contained 14 progenies derived from controlled crosses among these parents.

The second group consisted of seven parents (Nacional, Jarret and Bowen, 1994). SSR markers have been successfully applied on animal and plant diploid popula-
tions to determine genealogy structure and kin relationships (Adato et al., 1995; Morin et al., 1994; Saghai Maroof et al., 1994). There are no known data relative to the use of SSR for parentage assignment in plants.

The objective of this study was to assess paternity exclusion and most-likely parent methods for paternity assignment in hexaploid sweetpotato using microsatellites.

Materials and Methods

DIPLOID PATERNITY ANALYSIS

The conditions for PCR were as follows: 2 min denaturation at 95 °C; 5 cycles of 1 min at 94 °C; 1 min at 65 °C for primer IB-316, 1 min at 63 °C for IB-318; and 1 min at 72 °C; 10 cycles of 1 min at 94 °C; 1 min at 64 °C for primer IB-316, 1 min at 62 °C for primer IB-318; and 1 min at 72 °C; and 25 cycles in which the denaturation conditions were 1 min at 90 °C while the annealing and extension (1 min) temperature remained unchanged from the previous cycles. All reactions contained a terminal elongation step of 72 °C for 7 min. These conditions were programmed in a GeneAmp PCR System 9600 (Perkin-Elmer Corp., Foster City, Calif.). The amplified DNA fragments (2-mL samples) were resolved by electrophoresis in 6.5% non-denaturing polyacryla-
mide gels (0.4 mm thick, 38.5 cm long) with 1× TBE buffer (89 mM Tris-borate and 2 mM EDTA, pH 8.0). The gels were run at 65 W for 2 h. Three lanes with X174 HindI markers (Promega Corp., Madison, Wis.) were included. The gels were stained with silver nitrate following the procedure of Bassam et al. (1991) as modi-
fied by He et al. (1994). The parent and progeny genotypic profiles were then scored.

The second group consisted of seven parents (Nacional, Jarret and Bowen, 1994). SSR markers have been successfully applied on animal and plant diploid popula-
tions to determine genealogy structure and kin relationships (Adato et al., 1995; Morin et al., 1994; Saghai Maroof et al., 1994). There are no known data relative to the use of SSR for parentage assignment in plants.

The second group consisted of seven parents (Nacional, Jarret and Bowen, 1994). SSR markers have been successfully applied on animal and plant diploid popula-
tions to determine genealogy structure and kin relationships (Adato et al., 1995; Morin et al., 1994; Saghai Maroof et al., 1994). There are no known data relative to the use of SSR for parentage assignment in plants.

The procedure for PCR was as follows: 2 min denaturation at 95 °C; 5 cycles of 1 min at 94 °C; 1 min at 65 °C for primer IB-316, 1 min at 63 °C for IB-318; and 1 min at 72 °C; 10 cycles of 1 min at 94 °C; 1 min at 64 °C for primer IB-316, 1 min at 62 °C for primer IB-318; and 1 min at 72 °C; and 25 cycles in which the denaturation conditions were 1 min at 90 °C while the annealing and extension (1 min) temperature remained unchanged from the previous cycles. All reactions contained a terminal elongation step of 72 °C for 7 min. These conditions were programmed in a GeneAmp PCR System 9600 (Perkin-Elmer Corp., Foster City, Calif.). The amplified DNA fragments (2-mL samples) were resolved by electrophoresis in 6.5% non-denaturing polyacryla-
mide gels (0.4 mm thick, 38.5 cm long) with 1× TBE buffer (89 mM Tris-borate and 2 mM EDTA, pH 8.0). The gels were run at 65 W for 2 h. Three lanes with X174 HindI markers (Promega Corp., Madison, Wis.) were included. The gels were stained with silver nitrate following the procedure of Bassam et al. (1991) as modified by He et al. (1994). The parent and progeny genotypic profiles were then scored.

PATERNITY ANALYSIS
An algorithm performed the paternity analysis based on LOD scores. The algorithm has two principal routines; the first one excludes all the putative parents that show genetic incompatibility with each offspring. To perform this, all possible gametes from the female (known) and male progenitors are extracted from the respective genotypes; the female contribu-
tion is subtracted from the offspring genotype and the remainder is compared with the putative male parent contribution. The second routine calculates the likelihood of paternity for each nonexcluded putative male parent. The two algorithms were programmed in FORTRAN (Microsoft FORTRAN, Power Station 1993) to be run on a PC.

In a polycross nursery only two genealogical situations need to be considered: 1) relationship A—F is the female parent of O and M is unrelated; and 2) relationship B—both, F and M, are parents of O. The LOD score for a parent pair is (Meagher, 1986; Thompson and Meagher, 1987):

\[
L(B|g_o, g_{f1}, g_{m}) = \sum_{loci} \log_2 \left( \frac{P(g_o|h_{f1}, g_{m})}{P(g_o|h_{f1}g_{m})} \right) = \sum_{loci} \log_2 \left( \frac{M(g_o|h_{f1}, g_{m})}{M(g_o|h_{f1}g_{m})} \right)
\]

where \(P(g_o)\) is the probability of genotype \(g_o\) in a random mating population and \(M\) the Mendelian probability. The symbols, \(g_o, g_{f1}\) and \(g_{m}\), represent the offspring (progeny), female parent and male parent, respectively. This algorithm calculates: 1) the probability that the genotype of a given locus \[e.g., the probability based on Mendelian frequencies that \(A_1A_1\) in a 3 allele \((A_1A_2A_3)\) diploid locus model\] occurs in progeny given the set of parental genotypes mating at random \(P(g_{o|A_1A_2A_3})\); and 2) the probability that the known maternal parent \((e.g., A_1A_2)\) passes on its essential allele found in the progeny \((i.e., A_1)\) \(P(g_{o|A_2A_3})\). In a breeding
nursery the hypothesized relationship is B because A is known, so it can be used as the base-point alternative. To compare the likelihoods of distinct relationships (A and B) the difference in the natural logarithm of the likelihoods is considered (LOD scores). Because offspring’s loci are independent conditionally on the parental genotypes, LOD scores are summed across all loci.

Even though this approach allows paternity assignment to a higher number of progeny, it presents two limitations. First, categorical assignments are not possible for all the progeny for ambiguous progeny genotypic profiles. Second, a statistical bias in favor of homozygotes for a homoyzygous putative parent will always result in a higher LOD score for a given locus than a heterozygous individual (Devlin et al., 1988; Smouse and Meagher, 1994). Both limitations can be overcome by increasing the number of genetic markers.

Allele frequencies and gene diversity or average heterozygosity \( (H) \) \( (H = 1 - \sum_{i=1}^{n} p_i^2) \) for each population were calculated according to Nei (1987). \( H \), as a measure of degree of genetic variability, is the probability that two randomly chosen alleles with frequencies \( p_i \) the population frequency for the \( i \)th allele, are different. A higher \( H \) value represents a higher expected frequency of heterozygous individuals (Ott, 1992).

**Results**

**Allele frequencies.** In the LAES parental population, five microsatellite alleles were detected in parents for locus IB-318 (Table 1). Allele frequencies in the parent population were skewed towards the 120 bp allele. This allele, at a 75% frequency, and the 114 bp allele accounted for 90% of the allelic population. This poor allelic diversity in progeny is reflected in a gene diversity (even distribution) that was high and comparable to locus IB-316. Gene diversity was high and comparable to locus IB-316 for the LAES progeny population. All alleles fit within the expected class size for the two loci (Buteler et al., 1999).

**Paternity allocation.** In the LAES population, two out of 13 progeny were correctly allocated to the paternal parent by paternity exclusion (progeny 8 and 9, Table 2). All potential parents, except for the true paternal parent showed allelic incompatibility. Most-likely parent analysis (Table 2) correctly allocated one additional progeny (10) to the true paternal parent. Ambiguous assignments, i.e., two parents with identical LOD scores, occurred for two additional progeny (3 and 13). Misassignments were made for the remaining eight progeny. Of these, the true paternal parent had the second highest LOD score for the progeny. Four progeny were similarly narrowed to within three potential paternal parents. The parent 91-153 had six misassignments associated with it; five were traceable to parents with similar ancestry to 91-153.

In the CIP population, two out of eight progeny were correctly allocated to the paternal parent by paternity exclusion (progeny 3 and 4, Table 3). Most-likely parent analysis correctly allocated five additional progeny to the true male parent. Only one progeny was narrowly misassigned.

**Discussion**

Paternal allocation by the most-likely parent method was more successful when the CIP population was used as compared to the LAES population, i.e., seven out of eight progeny were allocated to the correct paternal parent vs. three out of 12 for the LAES population. The main difference between the data sets is the low level of allelic diversity at the LAES IB-318 locus. In contrast, allelic diversity (even distribution) was high for both loci in the CIP progeny population. These results are consistent with our expectations that the LAES population has a narrower genetic base. He et al. (1995) estimated that most cultivars within the United States could be traced to several common ancestors; by implication, we assume greater homogeneity among LAES parents. The CIP population, however, consists of parents from the United States and South America. In this more genetically diverse population, the informativeness of the two microsatellite loci was sufficient to determine the paternity. However, we would expect the CIP population to behave more like the LAES population as homogeneity increases under intense selection. These data sup-

<table>
<thead>
<tr>
<th>Locus designation</th>
<th>Primer pairs</th>
<th>Expected Size</th>
<th>CIP population</th>
<th>LAES population</th>
</tr>
</thead>
<tbody>
<tr>
<td>IB-316</td>
<td>CAAAACGACACGCTTAC (GA)(_4) (GA)(_4)</td>
<td>150</td>
<td>146 0.68 0.2500 0.68 0.4038</td>
<td>125 0.0192</td>
</tr>
<tr>
<td></td>
<td>CAAAACGACACGCTTAC (GA)(_4) (GA)(_4)</td>
<td>142 0.1429 0.0385</td>
<td>138 0.1429 0.2692</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAAAACGACACGCTTAC (GA)(_4) (GA)(_4)</td>
<td>133 0.4643 0.2885</td>
<td>123 0.2143 0.0000</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAAAACGACACGCTTAC (GA)(_4) (GA)(_4)</td>
<td>119 0.0714 0.0385</td>
<td>125 0.0192</td>
<td></td>
</tr>
</tbody>
</table>

\(^1\) Band size in base pairs.

\(^2\) Measure of degree of polymorphism, expected heterozygosity.
port the evidence obtained by simulation studies that skewed allele frequencies (i.e., two alleles among five account for more than 90% of the alleles present at the IB-318 locus in the LAES population) increase the number of misassignments and the variability of the estimates (Buteler et al., 1997). A second difference in these data sets is parent population size. The CIP parental population was smaller than the LAES parental population, i.e., five parents vs. eight parents, respectively. Buteler et al. (1999) showed via computer simulation that misassignments increased as a function of increasingly larger parental populations.

The algorithm used in the present study allocated parental assignments based on paternity exclusion and LOD scores. Paternity exclusion correctly allocated two progeny in each population and represented a 100% confidence level, given that no other paternal parents were a part of the experimental population. Yet, the most-likely parent method correctly allocated additional progeny in the populations. The confidence that one places on accurate parentage assignment is based on the numbers of loci and alleles used in the data set. Buteler et al. (1997), using computer simulation studies with a haploid, showed that the number of loci scored for a 10-parent population should not be less than 20 in the case of three alleles per locus, and no more than 10 loci for a five allele per locus model for parental discrimination with negligible errors or misassignments. A three loci, five allele (even allele frequency) model is accurate to within 15%. An incremental increase in the number of alleles enhances discriminatory power to a greater extent than an increase in the number of loci. Albeit, our present work was limited to two loci; other microsatellite loci were tested but these failed to segregate in a Mendelian fashion (Buteler et al., 1999). Microsatellite loci in sweetpotato show insertions, deletions, and base substitutions in addition to point mutations within the repeat and adjacent regions (Buteler et al., 1999). These aberrations may impact Mendelian segregation patterns and complicate discovery of new loci. Nevertheless, additional microsatellites are being developed for multiple uses in sweetpotato (Zhang et al., 2001).

In conclusion, paternity can be allocated with few loci in an experimental population. The most-likely parent method showed greater discriminatory power when compared with paternity exclusion for paternity determination in both populations. These

Table 2. LOD score matrix for paternity allocation in the CIP population for loci IB-316 and IB-318.

<table>
<thead>
<tr>
<th>Parents</th>
<th>Resisto</th>
<th>86-33</th>
<th>NC-C75</th>
<th>90-233</th>
<th>91-153</th>
<th>Excel</th>
<th>Beauregard</th>
<th>80-62</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>–0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>2</td>
<td>0.000000</td>
<td>0.08661434</td>
<td>0.000000</td>
<td>0.000000</td>
<td>3.063637</td>
<td>1.2716085</td>
<td>1.2716085</td>
<td>0.8661434</td>
</tr>
<tr>
<td>3</td>
<td>1.721809</td>
<td>0.000000</td>
<td>1.721809</td>
<td>0.000000</td>
<td>0.000000</td>
<td>0.000000</td>
<td>0.000000</td>
<td>1.0286623</td>
</tr>
<tr>
<td>4</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>–0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>5</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>6</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>7</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>8</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>9</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>10</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>11</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>12</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
<tr>
<td>13</td>
<td>0.000000</td>
<td>–</td>
<td>0.355317</td>
<td>0.000000</td>
<td>1.453930</td>
<td>0.337829</td>
<td>–0.337829</td>
<td>–1.436441</td>
</tr>
</tbody>
</table>

2The higher LOD score determines parental allocation. Female parents are as follows for offspring: 1 = Resisto, 2 = Resisto, 3 = 86-33, 4 = Resisto, 5 = NC-C75, 6 = 90-223, 7 = NC-C75, 8 = NC-C75, 9 = Resisto, 10 = NC-C75, 11 = NC-C75, 12 = 91-153, 13 = Excel.
3Excluded male parent (LOD = 0.0).
4Misassigned male parent (underlined).
5True male parents (bold).
6Ambiguous parental allocation.

Table 3. LOD score matrix for paternity allocation in the LAES population for loci IB-316 and IB-318.

<table>
<thead>
<tr>
<th>Parents</th>
<th>Nacional</th>
<th>Huarmeyano</th>
<th>ST87.006</th>
<th>LM87.045</th>
<th>Jewel</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>–0.595754879255</td>
<td>1.483686624258</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>2</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>0.224414006944</td>
<td>2.526729099988</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>3</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>30.024131703372</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>4</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>5</td>
<td>–0.354592823244</td>
<td>1.2716085</td>
<td>1.2716085</td>
<td>0.8661434</td>
<td>2.65790285</td>
</tr>
<tr>
<td>6</td>
<td>0.369326016789</td>
<td>2.161085486017</td>
<td>0.0000000000000000</td>
<td>3.35105085</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>7</td>
<td>0.0000000000000000</td>
<td>0.908322517521</td>
<td>0.0000000000000000</td>
<td>2.161085486017</td>
<td>0.0000000000000000</td>
</tr>
<tr>
<td>8</td>
<td>1.679431239551</td>
<td>1.456287688237</td>
<td>0.06993327117</td>
<td>1.719252733738</td>
<td>1.574070723893</td>
</tr>
</tbody>
</table>

2The higher LOD score determines parental allocation. Female parents are as follows for offspring: 1 = Nacional, 2 = Nacional, 3 = Huarmeyano, 4 = Huarmeyano, 5 = ST87.006, 6 = ST87.006, 7 = Nacional, 8 = LM87.045.
3Excluded male parent (LOD = 0.0).
4Misassigned male parent (underlined).
5True male parents (bold).

data sets underscore the utility of the most-likely parent method, i.e., no LOD based paternal allocation is made without first eliminating paternal parents based on paternity exclusion. Our results demonstrate the feasibility of microsatellite-based paternity analysis in polyploid species and statistical approaches to paternity determination in plant breeding programs.

Literature Cited


