A Genetic Map of Squash (Cucurbita sp.) with Randomly Amplified Polymorphic DNA Markers and Morphological Markers

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ABSTRACT. A molecular and morphological marker map would improve our knowledge of Cucurbita genetics, and would facilitate efforts to breed improved summer and winter squash cultivars. Random amplified polymorphic DNA (RAPD) markers were used to construct a partial map of the Cucurbita genome. The mapping population was the BC1 progeny of the Cucurbita pepo L. yellow straightneck inbred A0449 and the tropical Cucurbita moschata Duchesne ex Lam. landrace ‘Nigerian Local’. A0449 was the recurrent parent. This cross was chosen because of the relatively greater economic importance of summer squash, traits of value to be introgressed from the C. moschata parent, and maximized genetic variation from the interspecific cross. The map contains 148 RAPD markers in 28 linkage groups. Loci controlling five morphological traits were placed on the map. The map covers 1,954 cM, which is estimated to be 75% of the Cucurbita genome. The qualitative traits placed on the map include the B gene for fruit which turn yellow before anthesis, the M gene for silver mottling of leaves, and a locus controlling the intensity of rind color on mature fruit. Quantitative trait loci (QTL) associated with fruit shape and the depth of the indentations between primary leaf veins were identified.

The Cucurbita pumpkins, squash and gourds are represented by five species that were domesticated in the Americas. Most widely grown in North America, C. pepo includes summer squash (yellow straightneck and zucchini types), Jack O’ Lantern pumpkins, acorn and delicata winter squash, and small ornamental (yellow straightneck and zucchini types), Jack O’ Lantern pumpkins, acorn and delicata winter squash, and small ornamental vegetables are important in a healthy diet (USDA National Agricultural Statistics Service, 1997). The Cucurbita can supply carbohydrates, soluble fiber and provitamin A from the flesh, and protein, vegetable oils and vitamin E from the edible seeds. To meet increased demand for squash and pumpkin cultivars, additional research is needed. A genetic map for Cucurbita would improve our knowledge of genetics, and would facilitate breeding efforts.

The development of molecular tools for use in breeding Cucurbita species is still in the early stages and lags behind the major vegetable crops as well as other Cucurbitaceae. All Cucurbita species have twenty pairs of relatively short chromosomes. This compares to seven pairs in cucumber (Cucumis sativus), eleven in watermelon (Citrullus lanatus), and twelve in melon (Cucumis melo). Most morphological traits in Cucurbita appear to be unlinked, and many markers are required to adequately map the genome.

The genus Cucumis contains the best-mapped species of the Cucurbitaceae. There are currently five published maps of melon, and three maps of cucumber, with multiple versions of each map (Baudracco-Arnas and Pitrat, 1996; Brotman et al., 2000; Danin-Poleg et al., 1998; Kennard et al., 1994; Liou et al., 1998; Oliver et al., 2000; Park et al., 2000; Perin et al., 2000; Serquen at al., 1997; Wang et al., 1997). Efforts are currently underway to merge the cucumber maps (Staub and Serquen, 2000) and the melon maps, and to do comparative mapping between the two species (Danin-Poleg et al., 2000). Two of the melon maps are essentially complete, with twelve linkage groups and an average distance of <5 cM between markers. The maps contain stable, sequence based markers such as RFLPs, microsatellites, and markers based on expressed sequences of known function, in addition to RAPD makers. AFLPs and morphological markers (Oliver et al., 2000; Perin et al., 2000). The most extensive cucumber map is that of Park et al. (2000). It is less complete than the melon maps, having twelve linkage groups instead of the seven expected.

No linkage map of any Cucurbita species existed prior to the development of molecular mapping. Weeden and Robinson (1986) used isozyme markers to build the first map of Cucurbita. It was based on the F2 of the cross C. maxima × C. ecuadorensis, and contained 11 isozyme loci in five linkage groups. Lee et al. (1995)
developed a RAPD map of an F₂ population of a *C. pepo* × *C. moschata* interspecific hybrid. The map of Lee et al. (1995) was small, consisting of 28 markers in five linkage groups. No morphological traits or other types of markers were included on the map. The markers on their map were labeled arbitrarily, rather than following the standard practice of identifying the markers by the primer and the band size in base pairs. This prevents comparisons between their map and other *Cucurbita* maps. The map of Lee et al. (1995) was the only published molecular map of *Cucurbita* prior to this study.

The *Cucurbita* map presented here is intended as a framework to be built upon in further research as we seek to locate markers useful to breeders, determine the genetics of complex traits, and study evolution and species divergence in *Cucurbita*, and higher-level divergence among the Cucurbitaceae.

**Materials and Methods**

**Plant materials.** The mapping population consisted of 162 BC₁ individuals from a cross between *C. pepo* A0449 and *C. moschata* ‘Nigerian Local’. A0449 is a summer squash, an elite inbred line of the yellow straightneck type, developed by Sunseeds (Morgan Hill, Calif.). ‘Nigerian Local’ is a tropical pumpkin landrace that has been widely used as a source of virus resistance. The initial cross A0449 × ‘Nigerian Local’ was made in the greenhouse at Sunseeds in Brooks, Ore., in 1997. A single F₁ plant was obtained. The backcross population used in this experiment had A0449 as the female parent, and the F₁ as the male.

The BC₁ was seeded into 7.5-cm pots in growth chambers on 19 Sept. 1999. When plants had one to two true leaves, the leaves were harvested for DNA extraction and the seedlings transplanted to 20-L pots in the greenhouse. Plants were grown to maturity in the greenhouse. They were arranged in three rows running the length of each bench, with =10 cm between pots. All plants were staked to keep them upright, except for the ‘Nigerian Local’ plants, which were trellised. Temperatures ranged from 24 to 32 °C; supplemental light (16 h day, 8 h night) was provided by high intensity metal halide and sodium vapor lights hung 1.3 m above the benchtop. The BC₁ plants were hand-pollinated to produce fruit for data on fruit traits.

**Morphological traits.** Leaf mottle was scored as the presence or absence of distinctive silver-white areas in the axils of the leaf veins. The extent of mottling varied both between plants and among the leaves of a given plant; the degree of variation was not examined.

Precocious yellow fruit is controlled by two genes, *B* and *Ep*. *B* regulates the expression of the phenotype, while *Ep* controls the extent of expression. Both genes are incompletely dominant. Phenotypes for *B* and *Ep* were scored on immature fruit at or before anthesis. Fruit pattern was scored as bicolor or solid yellow; the extent of yellow or green on the peduncle was rated using a four-point scale from solid green to solid yellow. The data for the two traits were combined and related to known genotypes using information in the literature (Shifriss, 1981; Shifriss and Paris, 1981). Two *Ep* loci have been identified in *C. pepo*, *Ep-1* and *Ep-2* (Shifriss and Paris, 1981). The loci are phenotypically indistinguishable; we do not know which locus is present in A0449.

Mature fruit were examined visually to determine the rind color. Hue (orange, yellow, green, or tan) and intensity (light or dark) were recorded separately, as they have been shown to be controlled by different loci (Paris et al., 1985; Paris and Nerson, 1986). Hue and intensity were determined relative to the fruit of the parent lines and the other BC₁ plants. No color chart was used.

The basic fruit shape in squash ranges from disc-shaped through round to elongate. The shape was quantified by measuring the length and width of mature fruits from closeup photographs. Length was measured from the point of peduncle attachment to the blossom scar, and width was measured across the widest point, perpendicular to the length axis. The measurement used in the genetic analysis was the length: width ratio, which was calculated by dividing the length by the width. This value was independent of actual fruit size or the scale of the photograph.

For the mapping population, the degree of indentation between the primary veins of the leaves was scored on a 0 to 6 scale, where 0 = an entire leaf margin, and 6 = deep indents between all lobes similar to the cut leaf trait in zucchini. The scale is illustrated in Fig. 1. The genetic control of leaf margin indentation is unknown in *C. pepo* or *C. moschata*, although it is controlled by a single recessive gene in *C. maxima*, which normally has an entire leaf margin (Dyutin, 1980).

**DNA extraction.** DNA was extracted using a modified CTAB (hexadecyl trimethyl ammonium boride) technique as described in Brown et al. (1998). Concentration was determined using a DNA fluorometer (Hoefer DyNA Quant 200; Pharmacia Biotech Inc., San Francisco, Calif.) according to the manufacturer’s directions. The DNA was further purified by running 20 μg of...
DNA from each sample through GeneMate PCRpure spin columns (ISC BioExpress, Kaysville, Utah) and reeling with 50 μL of Tris buffer. The DNA stocks were diluted to 5 ng·μL⁻¹ in sterile ultrapure water to create working stocks for polymerase chain reaction (PCR).

Polymerase chain reaction. DNA from the five plants representing each parental line was bulked, and these bulks were used to screen primers. Primers that amplified poorly on the first screening were screened a second time. Operon primer sets A through R (Operon Technologies, Inc., Alameda, Calif.) and University of British Columbia (UBC) primers 405, 411, 417, 419, 421, 423, 425, 430, 432, 447, 489, 533, 546, 550, 577, 589, 592, and 597 (Nucleic Acid-Protein Service Unit, Vancouver, B.C.) were used to screen the parents; those primers which had strong bands present in ‘Nigerian Local’ but not in A0449 were used to amplify the mapping population. Because the mapping population was a BC1, only bands present in that parent did not segregate in the progeny. Reaction mixtures contained 30 ng of plant genomic DNA, 1.5 mM MgCl₂, 0.1 mM dNTPs, 3 pmole of primer (Operon or UBC), 0.5 units of Taq DNA polymerase (Promega, Madison, Wis.), and 1× Taq buffer (Promega) in a total volume of 15 μL. The amplification program was 2 min at 94 °C followed by 5 cycles of 5 s at 94 °C, 1 min at 37 °C, 30 s at 54 °C, a 7 min ramp to 72 °C, and 2 min at 72 °C, and then 30 cycles of 5 s at 94 °C, 1 min at 37 °C, 30 s at 54 °C, and 2 min at 72 °C, ending with 15 min at 72 °C. Because of the size of the population, it was necessary to amplify the population in two batches for each primer. Parental and H₂O controls were included in each batch. In the interest of speed, four thermocyclers [one MJR-100 (MJ Research, Cambridge, Mass.), one PE 9600, and two PE 9700s (PerkinElmer, Wellesley, Mass.)] were used. To minimize the effects of differences between machines, both batches for each primer were run on the same machine. In addition, the amplification profiles were adjusted to be as identical as possible on the four machines. The PCR products were separated on 1.5% agarose (Seakem LE; FMC, Rockland, Maine) or Ultrapure (Gibco, Grand Island, N.Y.) in 0.5× TBE buffer. PCR product molecular weight was determined with a 100-bp ladder (Promega). The gels were stained with ethidium bromide and separated on 1.5% agarose (Seakem LE; FMC, Rockland, Maine) or Ultrapure (Gibco, Grand Island, N.Y.) in 0.5× TBE buffer. PCR product molecular weight was determined with a 100-bp ladder (Promega). The gels were stained with ethidium bromide and recorded on Polaroid 667 film. Only strong, repeatable bands were scored.

Data analysis. RAPD bands were scored as present or absent. Because the mapping population was the BC₁ to inbred A0449, bands present in that parent did not segregate in the progeny. Reaction mixtures contained 30 ng of plant genomic DNA, 1.5 mM MgCl₂, 0.1 mM dNTPs, 3 pmole of primer (Operon or UBC), 0.5 units of Taq DNA polymerase (Promega, Madison, Wis.), and 1× Taq buffer (Promega) in a total volume of 15 μL. The amplification program was 2 min at 94 °C followed by 5 cycles of 5 s at 94 °C, 1 min at 37 °C, 30 s at 54 °C, a 7 min ramp to 72 °C, and 2 min at 72 °C, and then 30 cycles of 5 s at 94 °C, 1 min at 37 °C, 30 s at 54 °C, and 2 min at 72 °C, ending with 15 min at 72 °C. Because of the size of the population, it was necessary to amplify the population in two batches for each primer. Parental and H₂O controls were included in each batch. In the interest of speed, four thermocyclers [one MJR-100 (MJ Research, Cambridge, Mass.), one PE 9600, and two PE 9700s (PerkinElmer, Wellesley, Mass.)] were used. To minimize the effects of differences between machines, both batches for each primer were run on the same machine. In addition, the amplification profiles were adjusted to be as identical as possible on the four machines. The PCR products were separated on 1.5% agarose (Seakem LE; FMC, Rockland, Maine) or Ultrapure (Gibco, Grand Island, N.Y.) in 0.5× TBE buffer. PCR product molecular weight was determined with a 100-bp ladder (Promega). The gels were stained with ethidium bromide and recorded on Polaroid 667 film. Only strong, repeatable bands were scored.

Morphological traits were scored in a variety of ways, depending on expression. For those traits where segregation appeared to fit a 1:1 segregation ratio, the morphological data were recoded by setting the phenotype of the yellow squash parent equal to 0, and the ‘Nigerian Local’ or nonparental phenotype equal to 1. Leaf mottle, precocious yellow fruit, mature fruit color intensity and peduncle color were analyzed this way using Mapmaker. The relationship of quantitative traits to the mapped loci was analyzed using QTL Cartographer (Basten et al., 1999). The LOD threshold for each trait was calculated using the Permutations function with 150 iterations.

Results and Discussion

Morphological traits

Leaf mottle. Leaves of A0449 are uniformly green, while some plants of ‘Nigerian Local’ have silvery-white mottling in the axes of the leaf veins. The silverying is caused by air spaces within the palisade cell layer and between that layer and the epidermis (Scarchuk and Lent, 1965), and is distinct from the leaf silverying caused by insect feeding. This mottling is controlled by a single dominant gene (M) with an unknown number of modifiers in both C. moschata and C. pepo (Coyne, 1970; Scarchuk, 1954). The BC₁ progeny contained some plants with silver-
mottled leaves, indicating that the ‘Nigerian Local’ parent carried the dominant allele at M. The degree of mottling in the BC₁ progeny ranged from one plant with completely silvered leaves to plants with only tiny silver patches in the axes of the primary veins. Mottling was scored as either present or absent, because the variation in degree of mottling was continuous, and changed from leaf to leaf within a plant. The ratio of mottled to plain leaves was 67:95, which differs from the expected single gene ratio at P > 0.05, but does not differ at P > 0.01 (Table 1). Shifriss (1984) showed that the mottled phenotype is more fully expressed under conditions of high light intensity. The excess of plain-leafed plants may have resulted from low light intensity in the greenhouse. The mottled leaf trait was mapped despite its distorted segregation, and was placed on linkage group 6 (Fig. 2).

**Leaf Indentation.** Plants of *C. moschata* normally have leaves with little to no indentation between the veins, while the leaves of *C. pepo* plants can be deeply indented (Whitaker and Davis, 1962). Sixty of the BC₁ progeny had leaves that were more deeply indented than either parent, as shown in Fig. 3. A0449 has shallow indents on either side of the apical vein, which fits class 2 on the scale (Fig. 1). Most ‘Nigerian Local’ plants had very shallow indentations, fitting class 1 on the scale. This trait was somewhat difficult to classify, as the degree of indentation varied among leaves of a given plant. The distribution is typical of a quantitative trait. A significant QTL was identified on linkage group 5 between the marker loci F10_400 and G2_400, with the point of maximum likelihood at the locus K11_950 (Fig. 2). The LOD threshold was 12.59. The transgressive segregation seen in the mapping population suggests that two or more genes are involved, some of which may come from ‘Nigerian Local’.

**Fruit Shape.** ‘Nigerian Local’ has round fruit, while A0449 has long, slightly flared fruit. The length: width ratios are 0.9 and 3.1, respectively (Fig. 4). The F₁ had elongated pear-shaped fruit, and fruit shape in the BC₁ ranged from the long shape of the recurrent parent through oval shapes to fruit nearly as round as that of ‘Nigerian Local’ (Fig. 4). Inheritance of fruit shape is known to be complex and multigenic, with elongated shapes sometimes being dominant and sometimes recessive (Sinnott and Hammond, 1930). Narrowing of the fruit at the peduncle end, resulting in a pear shape, appears to be inherited separately from the fruit length or width, in that pear shapes of all lengths occurred in the BC₁, from short, nearly round pear shapes to long crookneck shapes. Round and round-pear shapes usually represent intermediate (heterozygous) conditions in *C. pepo* (Sinnott, 1922; Whitaker, 1932), but may not in *C. moschata*.

A significant QTL for the length: width ratio was identified on linkage group 10 between the RAPD markers L19_800 and P19_400, with the point of maximum LOD at B8_900 (Fig. 2). The LOD threshold was 10.09. Extensive studies of fruit shape in *C. pepo* have shown that separate physiological factors are responsible for ovary lengthening and ovary wall thickening, resulting in long or wide fruit shapes, respectively (Sinnott and Durham, 1929). However, it is not clear how many genes control these factors.

**Fruit Rind Color.** Fruit rind color is probably the best-studied trait in *C. pepo*. At least eleven genes, two of them having more than two alleles, are involved. A0449 has yellow fruit from the time the female flower bud develops, indicating that it is of genotype *B/B* (precocious yellow). Being a yellow squash, it is also of genotype *Y/Y* (Sinnott and Durham, 1922). The peduncle and the base of the corolla are also yellow, indicating that A0449 carries at least two dominant *Ep* alleles. The *Ep* alleles are extenders of the precocious yellow phenotype. Two *Ep* loci are known; they are additive and incompletely dominant. The degree of extension of the precocious yellow phenotype depends on the number of dominant *Ep* alleles present, and on the genotype at *B*.
(B/B or b/b). The Ep loci have no known effect on genotype b/b (Shifriss and Paris, 1981). The ‘Nigerian Local’ parent had green immature fruit which ripened black-green with a green peduncle.

Immature fruit of plants in the BC1 generation segregated for bicolor vs. precocious yellow fruit. Peduncles segregated for green, bicolor yellow and green, and yellow. No fruits were completely green, indicating that B was dominant over the color genes from ‘Nigerian Local’. This agrees with the results of attempts to move B from C. pepo to C. moschata (Paris et al., 1986). About 25% of the fruit were bicolored, indicating that they were heterozygous for B and contained no more than one Ep allele (Table 2). Another 25% of the plants had yellow peduncles as well as uniformly yellow fruits like A0449, suggesting that they were homozygous for B and for the Ep gene (Table 2). Of the remaining 50%, half had a thin band of yellowing where the peduncle joined the fruit, indicating that they were homozygous for B but had only one Ep allele (Shifriss and Paris, 1981). The others had uniformly green peduncles and were presumably heterozygous for B but carried two Ep alleles (Table 2). The segregation fit the 1:1:1:1 ratio expected for a two-gene model in the backcross to the parent carrying the dominant alleles. If contrasting phenotype classes were compared, both B and Ep fit the single gene model (Table 1). The Ep gene present in A0449 remains unlinked (Fig. 2).

Mature fruit colors ranged from orange through the golden yellow of A0449 to plain yellow and dull yellow-tans to greenish yellows. Fruit color intensity was independent of B in that bicolor fruit occurred in both light and intense colors. If plants were separated into those bearing intensely colored fruit (orange or golden yellow) like A0449, and those bearing lighter-colored fruit (yellow, yellow-tan, and greenish yellow) the data fit both a one-gene model and a three-gene model (Table 1). If the one-gene model is accepted, then the segregating gene is most likely an unidentified dominant factor from ‘Nigerian Local’ that decreases the intensity of rind color associated with B, but has no effect in a bb background. Such a factor was observed in the black-fruited C. moschata PI 165561, but was not explored further (H.S. Paris, personal communication). Segregation of L-I from ‘Nigerian Local’ would also result in a ratio of 1 intense : 1 light for rind color, but in that case A0449 should have had light-colored fruit. Intensely-colored fruit versus light-colored fruit was mapped to linkage group 8a, but the identity of the locus is unclear.

**RAPD markers**

A total of 378 primers were screened on the parental bulks; 72 failed to amplify clear bands in one or both parents, even when repeated. Of the remaining 306 primers, 162 (52.9%) amplified bands that were polymorphic in ‘Nigerian Local’ and distinct from bands in A0449. In total, 212 bands were scored, 175 of which were included in the data set for map construction. The remaining 37 bands were discarded because of nonbiological segregation distortion or questionable scoring. In total, 148 RAPD markers were placed on the map. All of the RAPD markers amplify DNA sequences from ‘Nigerian Local’, so are linked in repulsion to alleles controlling traits in A0449.

**The map**

The map contains 151 loci: 148 RAPD markers and 3 morphological marker loci. There are 28 linkage groups (Fig. 2); the haploid genome of *Cucurbita* contains 20 chromosomes. Two pairs of linkage groups, 3a and 3b and 8a and 8b, are considered to be single linkage groups by Mapmaker at LOD 5 and a maximum of 30% recombination, but when the loci are arranged in the best possible order they are more
than 36 cM apart. The map covers a total of 1,954 cM with an average of 12.9 cM between loci. However, the loci are not evenly distributed, so many gaps are more than 20 cM. Two QTL have been identified, one influencing fruit shape and the other influencing leaf indentation. In addition to the 28 linkage groups included here, nine RAPD markers and one qualitative locus remain unlinked.

This Cucurbita map is much more complete than that of Lee et al. (1995), which contained only 28 RAPD markers in 5 linkage groups. The Cucumis melo map of Perin et al. (2000), the most comprehensive map in the Cucurbitaceae, covers 1590 cM and is believed to cover the 12 chromosomes of melon in their entirety. The 1,981 cM of this map cover approximately 75% of the Cucurbita genome. However, significant gaps remain to be filled on this map.

The segregation at 21 loci placed on the map differed significantly from the expected 1:1 ratio. One of these loci was the mottled leaf trait; the excess of plain-leaved plants may be due to the low light intensity under which the plants were grown. The other distorted loci were RAPD markers. Five linkage groups contained more than one distorted marker. In all cases the distorted markers were grouped together.

Distorted segregation ratios in segregating populations can arise from competition among gametes during fertilization, which is a result of gametophyte genes expressed in the haploid genotype. Distorted segregation can also be a result of hybrid sterility genes causing the abortion of certain genotypes (Faris et al., 1998). Lethal and sublethal genes can also result in segregation distortion if they result in embryos that are too weak to germinate properly, or plants that die before maturity. Distortions can also be caused by unequal crossing over and chromosome mispairing during meiosis. This is frequently a problem in interspecific crosses, and is one reason for their generally lower fertility. The synteny of the genomes of C. pepo and C. moschata is not known.

Table 2. Immature fruit color in the BC1 of the cross Cucurbita pepo A0449 × (A0449 × C. moschata ‘Nigerian Local’). The scale used to rate the leaves is shown in Fig. 1.

| Genotype | Phenotype | Plants (no.) |
|----------|-----------|--------------|
| BBEpEp   | Solid yellow with a completely yellow peduncle | 32 |
| BBEpEp   | Solid yellow with a mostly green peduncle; a thin yellow band at fruit attachment | 50 |
| BhEpEp   | Solid yellow with a completely green peduncle | 40 |
| BhEpEp   | Bicolor green and yellow with a solid green peduncle | 33 |

χ² = 5.60, P = 0.13

²Fit to a two-gene model with ratio of 1:1:1:1.
serve as a reference point for identifying markers closely linked to the \( B \) locus.

A dominant allele from \( C. moschata \) was mapped for the \( M \) locus. Mottled leaves are found in many species of \( Cucurbita \), and the same gene symbol has been assigned for \( C. pepo \), \( C. moschata \), and \( C. maxima \). However, it is not entirely certain that the genes are allelic in all three species. Wessel-Beaver (personal communication) has indicated to R.N. Brown that further research failed to substantiate the linkage.

The mapping population segregated for many other traits in addition to the ones discussed above. Methods and data for these traits are discussed in detail in Brown (2001). Some of the traits could be traced to the action of known single genes; others were genetically more complex or have not been studied. Traits which appeared to be controlled by a single dominant gene included orange vs. pale fruit flesh, warty vs. smooth fruit, soft vs. stiff leaf hairs, rounded vs. pointed leaf lobes, and the development of foliar mosaic after inoculation with zucchini yellow mosaic virus. None of these traits were linked to any of the RAPD markers identified. Additional quantitative traits for which data were collected included the number of days from planting to first flower, the node at which the first flower was located, average internode length, response to powdery mildew, and virus titer in young tissue 28 d after inoculation with zucchini yellow mosaic virus. \( A0449 \) is resistant to powdery mildew; under severe disease pressure some sporulation occurs but no chlorosis or necrosis develops. \( A0449 \) is fully susceptible to zucchini yellow mosaic virus. ‘Nigerian Local’ is highly susceptible to powdery mildew, with extensive sporulation, chlorosis, and necrosis even under light disease pressure. It is entirely resistant to zucchini yellow mosaic virus, showing no symptoms and no detectable virus even after repeated inoculations. ‘Nigerian Local’ is a vine squash, with long internodes; it flowered 10 to 12 weeks after planting. \( A0449 \) is a compact bush, with extremely short internodes. It flowered 5 weeks after planting (Brown, 2001). QTLS were identified for response to powdery mildew and for the number of days from planting to first flower. However, they were of borderline significance. No QTLS were identified for the other traits examined.

The map presented here represents a starting point for the construction of high-density maps in \( Cucurbita \). More markers are needed to increase saturation, to complete the coverage of the genome and to permit the construction of a true framework map with markers evenly distributed throughout the genome. The map needs to be anchored with markers that are not population-specific, such as simple sequence repeats (SSRs) or RFLPs. No SSRs or RFLPs are available specifically for \( Cucurbita \). SSRs have been identified and primers developed for melon and cucumber (Danin-Poleg et al., 2001) and watermelon (Jarret et al., 1998). Small scale analysis of \( Cucurbita pepo \) genotypes using primers for 50 \( Cucumis \) SSRs found that seven (14%) were functional and polymorphic (Katizr et al., 2000). However, it will probably be necessary to develop SSRs specific to \( Cucurbita \) to identify sufficient markers to fully anchor the map. More morphological traits are needed, particularly traits specific to \( C. pepo \). Markers need to be more tightly linked to the morphological traits, and longer, more specific primers need to be developed for best results in marker-aided selection. Better characterization of the parental genotypes and the development of additional populations would also assist in adding morphological traits to the map. In particular, a map that could be used in breeding zucchini-type \( C. pepo \) cultivars would be of interest to the vegetable seed industry. The population used to construct this map was well suited to mapping in that the parents were morphologically quite disparate. However, ‘Nigerian Local’ is of interest only as a source of virus resistance, and yellow squash are economically important only in the southeastern United States.

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