Low Genetic Diversity Indicates the Need to Broaden the Genetic Base of Cultivated Watermelon

Amnon Levi¹ and Claude E. Thomas

U.S. Department of Agriculture, Agricultural Research Station, U.S. Vegetable Laboratory, 2875 Savannah Highway, Charleston, SC 29414

Todd C. Wehner

Department of Horticultural Science, North Carolina State University, Raleigh, NC 27695-7609

Xingping Zhang

Syngenta Seeds, 21345 Road 98, Woodland, CA 95695

Additional index words. Citrullus lanatus, Citrullus colocynthis, RAPD markers, germplasm evaluation, vegetable breeding

Abstract. Genetic diversity and relatedness were assessed among 46 American cultivars of watermelon (*Citrullus lanatus* var. *lanatus*), and 12 U.S. Plant Introduction accessions (PIs) of *Citrullus* sp. using 25 randomly amplified polymorphic DNA (RAPD) primers. These primers produced 288 distinct reproducible bands that could be scored with high confidence among cultivars and PIs. Based on the RAPD data, genetic similarity coefficients were calculated and a dendrogram was constructed using the unweighted pairgroup method with arithmetic average (UPGMA). The cultivars and *C. lanatus* var. *lanatus* PIs differentiated at the level of 92% to 99.6% and 88% to 95% genetic similarity, respectively. In contrast, the *C. lanatus* var. *citroides*, and *C. colocynthis* PIs were more divergent and differentiated at the level of 65% to 82.5% and 70.5% genetic similarity, respectively. The low genetic diversity among watermelon cultivars in this study emphasizes the need to expand the genetic base of cultivated watermelon.

The xerophytic genus Citrullus Schrad. ex Eckl. & Zeyh. belongs to the Cucurbitaceae family. It comprises four known diploid (n = 11) species found in the temperate regions of Africa, Central Asia, and the Mediterranean (Jeffrey, 1975; Whitaker and Davis, 1962). Among these species is C. lanatus (Thunb.) Matsum & Nakai, from which the cultivated watermelon (C. lanatus var. lanatus) originated (Whitaker and Bemis, 1976; Whitaker and Davis, 1962). Watermelon has been cultivated in Central Africa for at least 5000 years, and in Egypt and in the Middle East for over 4000 years. By the 10th century it was introduced to China, which is the world's greatest producer and consumer of watermelon. By the 13th century, watermelon was grown in Europe, and the crop was introduced into North America during the 17th century (Jeffrey, 1975; Whitaker and Davis, 1962). Major U.S. production areas are in Florida, California, Texas, Georgia, and Arizona. U.S. watermelon production has increased from 1.2 million tons in 1980 to 3.9 million tons in 1999 with a farm value of \$270 million [U.S. Dept. of Agriculture (USDA), Agricultural Statistics, 2000]. Over 500 diploid cultivars were developed in the United States during the last two centuries, but there is an ongoing need to improve watermelon, particularly with respect to development of disease and pest resistant cultivars.

There is little information regarding the ancestries of many American watermelon cultivars developed during the 19th and early 20th centuries (G.W. Elmstrom, personal communication). Identification of watermelon cultivars and determination of their genetic purity and relatedness relies mainly on fruit characteristics. Molecular markers can be an effective means to determine genetic relatedness among cultivars and among selections used in watermelon breeding programs. In previous studies designed to examine genetic diversity and phylogenetic relationships among watermelon cultivars using isozymes, most isozymes tested produced monomorphic patterns (Biles et al., 1989; Zamir et al., 1984). In contrast, RAPD procedure provided a sufficient number of informative markers that could distinguish among watermelon cultivars (Hashizume et al., 1993; Zhang et al., 1994). In a recent study (Levi et al., 2000), genetic

diversity and relatedness were examined among U.S. Plant Introduction accessions (PIs) of *C. lanatus* var. *lanatus*, *C. lanatus var. citroides, and C. colocynthis* using RAPD analysis. RAPD markers were also used in the construction of an initial genetic linkage map for watermelon (Hashizume et al., 1996), and to determine genetic relatedness among Asian watermelon cultivars and breeding lines (Lee et al., 1996).

Three hundred and fourteen American cultivars are stored at the USDA, ARS, National Seed Storage Laboratory (NSSL, Fort Collins, Colo.), and are considered an essential germplasm resource for watermelon breeding programs. Among them are 'Allsweet', 'Au-Producer', 'Charleston Gray', 'Crimson Sweet', 'Jubilee', and 'Peacock', that are grown throughout the world and are widely used as parents for many hybrids. Currently, there are no molecular data assessing the genetic relatedness and diversity among these cultivars. The objectives of this study were to: 1) estimate genetic relatedness and diversity among American cultivars; and 2) compare their genetic diversity with that in PI accessions of Citrullus.

Materials and Methods

Plant material. Forty-six watermelon cultivars (Table 1) and twelve PIs representing the three major *Citrullus* groups (*C. lanatus* var. *lanatus*, *C. lanatus* var. *citroides*, and *C. colocynthis*) (Table 2) were evaluated. Seeds of cultivars were obtained from commercial seeds companies, and the USDA NSSL in Fort Collins, Colo. (Table 1). Seeds of all PIs were obtained from the USDA Plant Genetic Resources Conservation Unit in Griffin, Ga. All plants were grown in the greenhouse. Young leaves were collected from four to five, 3week-old-plants of each watermelon accession and stored at –80 °C.

Marker data collection. DNA was isolated from young watermelon leaves as described by Levi and Thomas (1999). One hundred and thirty eight primers (60% to 80% GC content) were screened for polymorphism using the cultivars Blackstone and Stone Mountain #5. These two cultivars were chosen for the initial screening of primers because of their difference in parentage (Table 1). RAPD-PCR reactions were performed as described by Levi et al. (1993). Amplification products were separated by electrophoresis in 1.4% agarose gels in $\times 0.5$ Tris borate buffer (Sambrook et al., 1989). The gels were stained with 0.5 $mg \cdot mL^{-1}$ ethidium bromide solution for 30 min and destained for 15 min in distilled water. DNA fragments were visualized under UV light and photographed using a still video system (Gel Doc 2000; Bio-Rad, Hercules, Calif.). The molecular weights of amplification products were calculated using the "1-Kb Plus DNA Ladder" standards (Gibco BRL, Gaithersburg, Md.).

Data analysis. A pairwise similarity matrix was generated using the Nei-Li similarity index (Nei and Li, 1979) according to the equation: Similarity = $2 N_{ab} / (N_a + N_b)$, where

Received for publication 27 Nov. 2000. Accepted for publication 5 May 2001. We thank Susan Fox and Laura Pence for their technical assistance. We also thank G. Elmstrom, R. Fery, J. Thies, and J. Staub for reviewing this manuscript. The use of trade names in this publication does not imply endorsement by the USDA or the NCARS of the products named, or criticism of similar ones not mentioned. The cost of publishing this paper was defrayed in part by the payment of page charges. Under postal regulations, this paper therefore must be hereby marked *advertisement* solely to indicate this fact.

¹To whom requests for reprints should be addressed. E-mail address: alevi@awod.com

Table 1. Watermelon cultivars evaluated in this study, including source of seeds, parental background, year of introduction, and fruit characteristics.

	Source									
	of	Breeding	Year	Fruit	Fruit	Flesh	Flesh	Rind	Rind	
Cultivar	seeds	parentage	introduced	shape	wt	color	firmness	color	firmness	Maturity
Allsweet	Sunseeds ^z	(Miles x Peacock) x	1972	Long	25	Pink	Firm	Green w/deep	Firm	Late
		Charleston Gray						green wide stripes		
Astrakanski	Seed Saversy	^x								
AU-Golden Producer	Hollar ^w	Selection from AU-Producer	1993	Globe	20	Light red	Firm	Light green w/green	Firm	Mid-early
								wide stripes		
AU-Jubilant	Hollar	Jubilee x PI 271778	1985	Long	25	Light red	Firm	Light green w/green	Firm	Midseason
				-		-		narrow stripes		
AU-Producer	Hollar	Crimson Sweet x PI 189225	1985	Globe	20	Light red	Firm	Light green w/green	Firm	Mid-early
						U		wide stripes		2
Black Diamond	Sunseeds			Globe	15	Light red	Soft	Dark green	Medium	Early
Blackstone	Hollar	Florida Giant, Fairfax		Globe	15	Light red	Soft	Dark green	Medium	Mid-early
Calhoun Gray	Sunseeds			Long	20	Light red	Firm	Light green/gray	Firm	Late
CalSweet	Sunseeds			Oblong	20	Deep red	Firm	Green w/deep	Firm	Late
				0		1		green wide stripes		
Charleston Grav	NSL-5267 ^v	Africa 8. Iowa Belle.Garrison.	1954	Long	20	Light red	Firm	Light green/grav	Firm	Late
	and NKL&G	Hawkesbury, Leesburg		0		0		8 8 8 8 9		
Coles Early	NSL-5270		1892							
Congo	Syngentat	(African x Iow Belle) x Garrison	1949	Long	25	Light red	Firm	Deep green w/dark	Firm	Late
	~)8	(8				green narrow stripes		
Crimson Sweet	Hollar	(Miles x Peacock) x	1963	Globe	20	Light red	Firm	Light green w/green	Firm	Mid-early
	monta	Charleston Gray	1700	01000	20	Light rea		wide stripes		inita carry
Dixielee	Hollar	Texas W5 Wilt resistant Peacock	1979	Globe	20	Deep red	Firm	Light green w/green	Firm	Late
Differen	montu	Fairfax Summit	., .,,,	01000	20	Deepied		narrow stripes		Bute
Dixie Queen	Sunseeds		1890	Globe	20	Deep red	Firm	Light green w/green	Firm	Late
Divite Queen	Buildeeds		1070	01000	20	Deepied	1 1111	narrow stripes	1	Luie
Dunbarton	NSL-6637	(African x Iowa Belle) x Garrison	n 1953							
Fairfax	Sunseeds	Garrison African Iowa Belle	1952	Long	20	Light red	Firm	Light green w/green	Firm	Late
Tuntux	Buildeeds	Leesburg Hawkesbury	1752	Long	20	Eight ied	1 1111	narrow stripes	1	Luie
Family Fun	Syngenta		1973							
Garrison	NSI -2053		1775	Long	20	Light red	Firm	Light green w/green	Firm	Late
Garrison	102 2000			Long	20	Light icu	1 1111	narrow stripes	1 1111	Luie
Garrisonian	Willhites	Africa & Jowa Belle Garrison	1057	Long	20	Light red	Firm	Light green w/green	Firm	Late
Garrisonian	vv innite	Hawkesbury Leesburg	1)57	Long	20	Light red	1 1111	narrow stripes	1 1111	Luie
Georgia Rattlesnake	Seed Savers							narrow surpes		
Golden Honey	Hollar		105/	Globa	12		Soft		Explosive	Forly
Golden Midget	NSI -5288	Mixed strain from Japan Seed Co	1050	Globe	12	Vellow	Soft		Tender	Early
Hawkashury	Syncente	New Hampshire Midget V	1036	GIUDE	12	TCHOW	Son		Tender	Larry
11awACSOULY	Syngenia	Pumpkin Rind	1950							
Ioprida	Synconto	т ишркш кши		Globa	20	Light rad	Firm	Deen green w/derk	Firm	Mideascon
Topride	Syngenta			Giobe	20	Light ieu	rum	groop porrow strings	гшп	whuseason
								green narrow surpes		

 N_{ab} is the number of RAPD fragments shared by two genotypes (a and b) and N_a and N_b are the total number of RAPD fragments analyzed in each genotype. A dendrogram was constructed based on the similarity matrix data by applying unweighted pair-group method with arithmetic average (UPGMA) cluster analysis using the Numerical Taxonomic and Multi-Variant Analysis System for PC (NTSYS-PC version 2) (Rohlf, 1993).

Results and Discussion

Of the 138 primers that were initially screened, only 35 produced polymorphic RAPD patterns. Of these, 25 primers that produced distinct polymorphic bands were used for further analysis with all cultivars and PIs. The 25 primers produced 288 reproducible RAPD bands that ranged in molecular size from 100 to 3000 base pairs (bp) (Fig. 1). Of these bands, 26 were monomorphic for all cultivars and PIs, 9 were polymorphic among cultivars, but were monomorphic for all PIs. One hundred and sixty eight bands were polymorphic among PIs, but were monomorphic for all cultivars, while 85 bands were polymorphic among all cultivars and PIs (Table 3).

The RAPD marker data were used to construct a genetic similarity matrix among cultivars and PIs (data not shown) based on the Nei-Li estimate of similarity (Nei and Li, 1979). The similarity matrix was used in a UPGMA cluster analysis to produce a genetic similarity dendrogram (Fig. 2). High genetic similarity values (92% to 99.6%) were detected among watermelon cultivars and among PIs of C. lanatus var. lanatus (88% to 95%) which is considered the progenitor of the cultivated watermelon. Lower similarity values were found among the C. lanatus var. citroides and among the wild species C. colocynthis PIs (65% to 82.5%, and 70.5%, respectively) (Fig. 2). In an additional study designed to elucidate genetic diversity in Citrullus sp. (Levi et al., 2000), genetic diversity and relatedness were estimated among 17 C. lanatus var. lanatus PIs. 12 C. lanatus var. citroides PIs. 13 C. colocynthis PIs, and five cultivars ('Allsweet', 'Charleston Gray', 'Ironsides', 'Mickylee', and 'New Hampshire Midget') using RAPD analysis. In that study, the five watermelon cultivars grouped in a distinct cluster, indicating that they were derived from common parents. However, the low genetic diversity among C. lanatus var. lanatus PIs in the previous Table 1 continued on next page

study (Levi et al., 2000), and in the present study (Fig. 2) indicates that the lack of genetic diversity among cultivars is due to a narrow genetic base in C. lanatus var. lanatus. Navot and Zamir (1987) found little isozyme variation among watermelon accessions, and suggest that this is due to the domestication of watermelon outside of its center of origin. They base this assumption on the probability that only a small fraction of plants of the progenitor species (a few plants that have desired qualities) were selected and used at the early stages of domestication (Ladizinsky, 1985). Katzir et al. (1996) could not detect any polymorphism among the watermelon cultivars Sugar Baby and Malali (diploids), and Tri-X-313 (triploid) using seven simple sequence repeat (SSR) markers. However, Jarret et al. (1997) could detect genetic diversity among Citrullus PIs using the same SSR markers. In that study, PIs of C. lanatus var. citroides were slightly more divergent than PIs of C. lanatus var. lanatus. Navot and Zamir (1987) considered C. lanatus var. citroides as the wild progenitor of C. lanatus var. lanatus.

In contrast with the low DNA polymorphism, extensive variation in morphological characteristics existed among watermelon cul-

	Source									
	of	Breeding	Year	Fruit	Fruit	t Flesh	Flesh	Rind	Rind	
Cultivar	seeds	parentage	introduced	shape	wt	color	firmness	color	firmness	Maturity
Ironsides	NSL-7369	(Leesburg x Hawkesbury) x Garrison	1950							
Jubilee	Hollar	Africa 8, Iowa Belle, Garrison, Hawkesbury, Leesburg	1963	Long	25	Light red	Firm	Light green w/green narrow stripes	Firm	Mid-season
King and Queen	Hollar			Globe	12	Light red	Soft	Light green	Medium	Early
Kleckely's Sweet	Seed Savers			Globe	12	Light red	Soft	Deep green	Medium	Early
Klondike	Syngenta		1959	Long	20	Light red	Firm	Dark green	Firm	Late
Klondike Striped RS 57	Sunseeds			Long	20	Light red	Firm	Light green w/green narrow stripes	Firm	Late
Klondike Striped- Blue Ribbon	Hollar		1939	Long	20	Light red	Firm	Light green w/green narrow stripes	Firm	Late
Leesburg	NSL-7368	Selection from Kleckley Sweet	1936							
Melitoplisky	Seed Savers									
Mickylee	Hollar	Texas W5, Fairfax, Summit, Graybelle	1986	Globe	15	Deep red	Firm	Light green/gray	Firm	Midseason
Miles	NSL-6688	Dixie Queen x Klondike R-7	1948							
Minilee	Hollar	Texas W5, Fairfax, Summit, Graybelle	1986	Globe	8	Deep red	Firm	Light green/gray	Firm	Early
New Hampshire Midget	Syngenta	Favorite Honey x Dakota Swee	t 1951	Globe	6	Red	Soft	Light green/gray	Explosive	Early
Northern Sweet	Syngenta		1932	Globe	12	Red	Medium	Green w/lines	Medium	Early
Parker	Willhite	\mathbf{F}_{1}								
Peacock	Hollar		1939	Oblong	20	Red	Firm	Dark green	Firm	Late
Prince Charles	Syngenta	F1	1978	Long	20	Red	Firm	Light green/gray	Firm	Midseason
Sangria	Syngenta	\mathbf{F}_{1}	1985	Long	25	Deep red	Firm	Green w/deep green wide stripes	Firm	Late
Sugar Baby	Sunseeds	Selection from Tough Sweets	1955	Globe	15	Orange red	d Soft	Dark green w/dark stripes	Medium	Early
Stone Mountain	Hollar		1924	Oval	20	Light red	Firm	Light green w/green narrow stripes	Firm	Midseason
Stone Mountain #5	Syngenta	Stone Mountain x Iowa Belle	1936							

^zSunseeds Co. (Acampo, Calif.).

^ySeed Savers Exchange (Decorah, Iowa).

^xInformation unavailable.

"Hollar Seeds (Rocky Ford, Colo.).

vAccession number provided for each of the heirloom cultivars kept at the USDA National Seed Storage Laboratory (Fort Collins, Colo.).

^uNK Lawn & Garden (Chattanooga, Tenn.).

^tSyngenta Seeds (Naples, Fla.).

^sWillhite Seed (Poolville, Texas).

Table 2. Twelve U.S. plant introduction accessions (PIs) examined in the pr	resent study, the (Citrullus group to whic	h they belong, the c	ountry from
which they were collected, and their fruit characteristics as described by	y the Germplasm	Resources Information	n Network (www.ar	s-grin.gov).

	Citrullus	Country of	Rind color	Rind color	Flesh	Fruit	Fruit	Frit
PI #	type	origin	background	pattern	color	diameter	shape	maturity
162667	lanatus	Argentina	Medium green	Solid	Red	15/30	Oblong	Midseason
165451	lanatus	Mexico	Medium green	Solid	Pink	18/25	Oblong	Midseason
169289	lanatus	Turkey	Dark green	Solid	Red	36/36	Round	Early
185636	lanatus	Ghana	Medium green	Solid	White	10/10	Round	Late
189316	lanatus	Nigeria	Dark green	Striped	White	12/12	Round	Late
203551	lanatus	U.S.A.	Light green	Striped	White	12/12	Round	Midseason
271778	lanatus	S. Africa	Medium green	Striped	Yellow	22/27	Oblong	Midseason
270564	citroides	S. Africa	Light green	Solid	Yellow	20/25	Oblong	Early
482251	citroides	Zimbabwe	Medium green	Striped	Yellow	12/12	Round	Midseason
271779	citroides	S. Africa	yellow	Striped	Yellow	30/45	Oblong	Midseason
386014	colocynthis	Iran	Light green	Striped	White	9/9	Round	Early
388770	colocynthis	Morocco	Light green	Striped	White	12/12	Round	Early

tivars used in this study (Table 1). These characteristics included rind color and thickness, fruit shape and size, flesh texture and color, sugar content, seed shape and color, days to fruit maturity, and disease resistance. Most of these characteristics are qualitative traits affected by a single or a few gene mutations (Rhodes and Dane, 1999; Rhodes and Zhang, 1995) that could not be readily detected by the RAPD markers in this study. However, some of these mutations may be detected through a bulked segregant analysis procedure (Michelmore et al., 1991) using a large number of RAPD primers.

The watermelon cultivars in the present study are open-pollinated or F_1 hybrid diploid types (n = 11). The parentage records for most of these cultivars are incomplete (Table 1). However, records available for some of the cultivars are consistent with the RAPD-based results (Fig. 2). For instance, the gray-green rind and round fruit type cultivars Mickylee and Minilee were developed from sister plants that had the cultivars Fairfax, Summit, and Texas-W5 in their genetic background (Table 1) (Crall, 1986). 'Mickylee' and 'Minilee' appeared closely related in the present analysis (Fig. 2). The cultivar Garrison contributed to the genetic background of the oblong fruited cultivars Congo, Charleston Gray, and Garrisonian (Table 1). In the present analysis, 'Congo', 'Garrison', and 'Garrisonian' are in the same group, while 'Charleston Gray' is in a closely related group (Fig. 2). The cultivar Allsweet has a green-striped rind and oblong fruit (Table 1), and according to the records it



Fig. 1. PCR-RAPD patterns (on 1.4% agarose-gel) of watermelon cultivars produced by primer UBC-222. Lanes are: 0) 1-Kb Plus DNA ladder markers (Gibco BRL, Rockville, Md.), 1) 'Allsweet', 2) 'Astrakanski', 3) 'AU-Golden Producer', 4) 'AU-Jubilant', 5) 'AU-Producer', 6) 'Blackstone', 8) 'Calhoun Gray', 9) 'Calsweet', 10) 'Charleston Gray' (NSL-5267), 11) 'Charleston Gray' (NK Lawn & Garden Co), 12) 'Coles Early', 13) 'Congo', 14) 'Crimson Sweet', 15) 'Dixielee', 16) 'Dixie Queen', 17) 'Dunbarton', 18) 'Fairfax', 19) 'Family Fun', 20) 'Garrison', 21) 'Garrisonian', 22) 'Georgia Rattlesnake', 23) 'Golden Honey', 24) 'Golden Midget'.

is comprised of 'Charleston Gray' (50%), 'Miles' (25%), and 'Peacock' (25%) (Table 1). The present analysis shows that 'Allsweet' is closer to 'Charleston Gray' than to the two other cultivars (Fig. 2). The cultivar AU-Golden Producer (globular fruit with green, wide-striped rind) is reported to be an orange fleshed mutation of 'AU-Producer' (Table 1). Predictably, these two cultivars are closely related (Fig. 2). 'AU-Producer' is derived from a cross of 'Crimson Sweet' (globular fruit with green-striped rind) with PI 189225 that is reported to have resistance to gummy stem blight (Sowell and Pointer, 1962). In the present study, 'Crimson Sweet' is indeed closely related to 'AU-Producer'. Like 'Allsweet', 'Crimson Sweet' also is comprised of 'Charleston-Gray' (50%), 'Miles' (25%), and 'Peacock' (25%) (Table 1). 'Allsweet' is indeed in the same group with 'Crimson-Sweet' and its descendents 'AU-Producer' and 'AU-Golden Producer' (Fig. 2). The cultivar Blackstone (globular fruit with dark green rind) contains 'Black-Diamond' (also known as Florida-Giant) and 'Fairfax' in its genome (Table 1). Indeed, 'Black-Diamond' and 'Blackstone' have similar characteristics (Table 1) and are in the same group (Fig. 2). 'Dixilee' is in the same group with 'Charleston-Gray' (Fig. 2). A detailed review (data not shown) reveals that these two cultivars are derived from common parents ('Peacock', 'Garrison', 'Leesburg', and 'Hawkesbury', Table 1).

Not all the results are fully consistent with parental records or with morphological characteristics. For instance, the cultivar Leesburg is a selection from 'Kleckely-Sweet' (Table 1). The present analysis shows that these two cultivars are in closely related groups, but are not clustered together (Fig. 2). Also, 'Stone-Mountain #5' is relatively distant from 'Stone-Mountain' (Fig. 2), which is considered one of its original parents, the other being 'Iowa Bell' (Table 1). The cultivar Jubilee is distant from 'Charleston Gray' and 'Garrisonian' (Fig. 2), which supposedly have similar parental backgrounds (Table 1). 'Jubilee' is also distant from its descendent 'Au-Jubilant' (Fig. 2).

Using a contingency test, we could not detect any significant association between RAPD markers and fruit shape. Therefore, it is likely that the RAPD markers here do not provide the resolution required to elucidate gene loci that control fruit characteristics, but elucidate random parts of the watermelon genome. Another indication for this assumption is that cultivars with different morphological characteristics appear closely related. For instance, the cultivar Calhoun-Gray, which has an elongated fruit with light green-gray rind, is in the same group with 'Blackstone' and 'Black Diamond' (Fig. 2), which have globular fruit and dark rind (Table 1).

Most cultivars in the present study have not been examined in previous studies of genetic relatedness using molecular markers. However, three cultivars ('Family Fun','Sugar Baby', and 'New Hampshire Midget') were examined together with watermelon cultivars and breeding lines developed in Asia (Lee et al., 1996). The genetic distance between 'Sugar Baby' and 'New Hampshire Midget' in the present study is comparable with that in the previous study. However, 'Family-Fun' appeared to be distant from most watermelon cultivars (Fig. 2), and not as closely related to 'Sugar Baby' as reported by Lee et al. (1996). This difference may be due to the use of different RAPD primers and RAPD procedures, or it may be due to use of seeds of cultivars that were produced by different sources. Due to the open-pollinated nature of watermelon, different genotypes may occur in a cultivar during seed increase, resulting in

Table 3. The nucleotide sequences of RAPD primers used in the present study, and the number of polymorphic (PM), and monomorphic (MM) band markers produced by each primer. Primer names are according to manufacturer's identification system (Operon Technologies; OP, and University of British Columbia; UBC).

	Nucleotide		
Primer	Sequence (5'—>3')	PM	MM
OPB-05	TGCGCCCTTC	11	2
OPB-11	GTAGACCCGT	24	
OPB-14	TCCGCTCTGG	14	1
OPC-05	GATGACCGCC	18	2
OPC-20	ACTTCGCCAC	15	1
OPD-02	GGACCCAACC	16	
OPD-07	TTGGCACGGG	17	
OPD-20	ACCCGGTCAC	25	
OPE-04	GTGACATGCC	19	
OPJ-06	TCGTTCCGCA	26	
OPJ-13	CCACACTACC	19	
OPK-14	CCCGCTACAC	21	
OPK-20	GTGTCGCGAG	17	
OPT-01	GGGCCACTCA	26	
OPT-05	GGGTTTGGCA	29	
UBC106	CGTCTGCCCG	32	3
UBC115	TTCCGCGGGC	25	1
UBC137	GGTCTCTCCC	21	
UBC147	GTGCGTCCTC	8	
UBC149	AGCAGCGTGG	25	
UBC152	CGCACCGCAC	22	4
UBC155	CTGGCGGCTG	16	1
UBC157	CGTGGGCAGG	24	1
UBC159	GAGCCCGTAG	10	
UBC186	GTGCGTCGCT	28	2
UBC199	GCTCCCCAC	27	4
UBC212	GCTGCGTGAC	21	1
UBC218	CTCAGCCCAG	20	2
UBC222	AAGCCTCCCC	29	1
UBC228	GCTGGGCCGA	31	

different RAPD patterns. To test this possibility, we examined 'Charleston Gray' plants from the original seed stock (NSSL) vs. 'Charleston Gray' plants from seeds provided by NK Lawn & Garden, Co. (Chattanooga, Tenn.). The RAPD patterns (as shown in Fig. 1) confirmed that these two types are alike, but not identical (Fig. 2). Two distinct RAPD markers (470 and 790 bp) produced by primer OPD-07 and UBC-155, respectively, were present in the original Charleston Grav cultivar (NSSL), but not in the type obtained commercially. The 790 bp marker was unique to the 'Charleston Gray' provided by NSSL and was not found in any other cultivar or PI in this study. These differences between two sources of the same cultivar indicate that changes in cultivar genotype may occur during seed increase. Thus, the source of the seeds for each cultivar tested should be taken into account in genetic studies and in DNA fingerprinting of watermelon cultivars.

The high genetic similarities among *C. lanatus* var. *lanatus* PIs, and among watermelon cultivars indicate that many cultivars developed in the United States over the last two centuries have a narrow genetic background. Therefore, it is essential to broaden the genetic base of the cultivated watermelon to reduce its vulnerability to diseases and insect pests. The recent study of Levi et al. (2000) indicated that the wild species *C. colocynthis*, which has the widest geographic



Fig. 2. Dendogram of watermelon cultivars and U.S. Plant Introduction accessions (PIs) produced by UPGMA cluster analysis of similarity matrix. 'AU-Golden Producer' = AU-G. Producer, 'Charleston Gray' (NSL-5267) = Charleston Gray, 'Charleston Gray' (NK Lawn & Garden Co.) = Chrles.-Gray-N, 'Georgia Rattlesnake' = Georgia-RS, 'Klondike Striped Blue Ribbon' = Klondike-SBR, 'New Hampshire Midget' = New-Hampshire-M., 'Klondike Striped RS 57' = RS57.

distribution, also has the highest genetic diversity among *Citrullus* species. That study also indicated higher genetic diversity within the wild subspecies *C. lanatus* var. *citroides*, than in *C. lanatus* var. *lanatus*.

Accessions of C. lanatus var. lanatus are preferred in watermelon breeding programs because of their horticultural qualities and their close genetic proximity to the cultivated watermelon. Previous studies indicate that resistance to anthracnose (Boyhan et al., 1994; Sowell et al., 1980), or watermelon mosaic virus (Gillaspie and Wright, 1993) exists among accessions of C. lanatus var. lantus. Although the two former *Citrullus* types are wild and do not have desirable fruit qualities. they might be an essential source of genes that would confer resistance to major diseases and pests. Resistance to gummy stem blight and Fusarium wilt, which are major diseases of watermelon, may exist among accessions of C. lanatus var. citroides (Dane

et al., 1998; Martyn, and Netzer, 1991; Sowell, and Pointer, 1962). In a recent study (Simmons and Levi, 2000), C. colocynthis PIs had high resistance, while all C. lanatus PIs were highly susceptible to whitefly (Bemisia tabaci). The Citrullus germplasm collection at the USDA, Agricultural Research Service, Plant Genetic Resources Conservation Unit (Griffin, Ga.), contains 1480 C. lanatus var. lanatus PIs, but only 102 C. lanatus var. citroides PIs, and 21 C. colocynthis PIs (Germplasm Resources Information Network, www.ars-grin.gov). Thus, in order to broaden the genetic base of watermelon cultivars, further efforts are required to expand the collection of C. lanatus var. citroides and C. colocynthis PIs, and to evaluate additional accessions for resistance to important diseases and pests of watermelon.

Literature Cited

Biles, C.L., R.D Martyn, and H.D. Wilson. 1989. Isozymes and general proteins from various watermelon cultivars and tissue types. HortScience 24:810–812.

- Boyhan, G.E., J.D. Norton, B.R. Abrahams, and N.H. Wen, 1994. A new source of resistance to anthracnose (Race 2) in watermelon. HortScience 29:111–112.
- Crall, J.M., 1986. Minilee and Mickylee two icebox-type watermelon cultivars with excellent fruit quality and resistance to *Anthracnose* and *Fusarium* wilt. Florida Agr. Expt. Sta. Circ. S-336.
- Dane, F., L.K. Hawkins, and J.D. Norton, 1998. New resistance to race 2 of *Fusarium oxysporum* f. sp. *niveum* in watermelon. Cucurbit Genetics Coop. Rpt. 21:37–39.
- Germplasm Resources Information Network (GRIN, Online Database, National Germplasm Resources Laboratory, USDA, ARS, Beltsville, MD; www.ars-grin.gov).
- Gillaspie, A.G., Jr., and J.M. Wright, 1993. Evaluation of *Citrullus* sp. germ plasm for resistance to watermelon Mosaic Virus 2. Plant Dis. 77:352– 354.
- Hashizume, T., T. Sato, and M. Hirai. 1993. Determination of genetic purity of hybrid seed in watermelon (*Citrullus lanatus*) and tomato (*Lycopersicon esculantun*) using random amplified polymorphic DNA (RAPD). Jpn. J. Breed. 43:367–375.
- Hashizume, T., I. Skimamoto, Y. Harushima, M. Yui, T. Sato, T. Imai, and M. Hirai. 1996. Construction of a linkage map for watermelon [*Citrullus lanatus* (Thunb.) Matsum & Nakai] using random amplified polymorphic DNA (RAPD). Euphytica 90:265–273.
- Jarret, R.L., L.C. Merrick, T. Holms, J. Evans, and M.K. Aradhya. 1997. Simple sequence repeats in watermelon [*Citrullus lanatus* (Thunb.) Matsum. & Nakai]. Genome 40:433–441.
- Jeffrey, C. 1975. Further notes on Cucurbitaceae: III. Some African taxa. Kew Bul. 30:475–493.
- Katzir, N., Y. Danin-Poleg, G. Tzuri, Z. Karchi, U. Lavi, and P.B. Cregan. 1996. Length polymorphism and homologies of microsatellites in several Cucurbitaceae species. Theor. Appl. Genet. 93:1282–1290.
- Ladizinsky, G., 1985. Founder effect in crop plant evolution. Econ. Bot. 39:191–199.
- Lee, S.J., J.S. Shin, K.W. Park, and Y.P. Hong. 1996. Detection of genetic diversity using RAPD-PCR and sugar analysis in watermelon [*Citrullus lanatus* (Thunb.) Mansf.] germplasm. Theor. Appl. Genet. 92:719–725.
- Levi, A., L.J. Rowland, and J.S. Hartung. 1993. Production of reliable randomly amplified polymorphic DNA (RAPD) markers from DNA of woody plants. HortScience 28:1188–1190.
- Levi, A. and C.E. Thomas. 1999. An improved procedure for isolation of high quality DNA from watermelon and melon leaves. Cucurbit Genetics Coop. Rpt. 22:41–42.
- Levi, A., C.E. Thomas, A.P. Keinath, and T.C. Wehner. 2000. Estimation of genetic diversity among *Citrullus* accessions using RAPD markers. Acta Hort. 510:385–390.
- Martyn, R.D. and D. Netzer, 1991. Resistance to races 0, 1, and 2 of Fusarium wilt of watermelon in *Citrullus* sp. PI-296341-FR. HortScience 26:429–432.
- Michelmore, R.W., I. Paran, and R.V. Kesseli. 1991. Identification of markers linked to disease resistance genes by bulked segregant analysis: A rapid method to detect markers in specific genomic regions by using segregating populations. Proc. Natl. Acad. Sci. U.S.A. 88:9828– 9832.
- Navot, N. and D. Zamir. 1987. Isozyme and seed protein phylogeny of the genus *Citrullus* (Cucurbitaceae). Plant Syst. Evol. 156:61–67.

- Nei, M. and W. Li. 1979. Mathematical model for studying genetic variation in terms of restriction endonucleases. Proc. Natl. Acad. Sci. USA 76:5269–5273.
- Rhodes, B. and X. Zhang. 1995. Gene list for watermelon (*Citrullus lanatus*). Cucurbit Genetics Coop. Rpt. 18:69–84.
- Rhodes, B. and F. Dane. 1999. Gene list for watermelon (*Citrullus lanatus*). Cucurbit Genetics Coop. Rpt. 22:61–77.
- Rohlf, F.J. 1993. NTSYS-PC numerical taxonomy and multivariate analysis system, Version 2.00 Exeter Publishing, Ltd., Setauket, N.Y.

Sambrook, J., E.F. Fritsch, and T. Maniatis. 1989.

Molecular cloning. A laboratory manual. Second ed. Cold Spring Harbor Lab. Press. Plainview, N.Y.

- Simmons, A.M. and A. Levi. 2000. Evaluation of watermelon germplasm for resistance to *Bemisia*. Entom. Soc. of Amer. Annu. Meeting. Abstr. No. 15611.
- Sowell, G. and G.R. Pointer. 1962. Gummy stem blight resistance of introduced watermelons. Plant Dis. Rptr. 46:883–885.
- Sowell, G., Jr., B.B. Rhodes, and J.D. Norton. 1980. New Sources of resistance to watermelon anthracnose. J. Amer. Soc. Hort. Sci. 105:197–199.
- U.S. Dept. of Agriculture. 2000. Agricultural statis-

tics. U.S. Govt. Printing Office, Wash., D.C.

- Whitaker, T.W. and G.N. Davis. 1962. Cucurbits botany, cultivation and utilization. Interscience Publishers, New York.
- Whitaker, T.W. and W.B. Bemis. 1976. Cucurbits, p. 64–69. In: N.W. Simmonds (ed.). Evolution of crop plants. Longman, London.
- Zamir, D., N. Navot, and J. Rudich. 1984. Enzyme polymorphism in *Citrullus lanatus* and *C. colocynthis* in Israel and Sinai. Plant Syst. Evol. 146:163–137.
- Zhang, X.P., B.B. Rhodes, and H.S. Skorupska. 1994. RAPD molecular markers in watermelon. Cucurbit Genetics Coop. Rpt. 17:116–119.