Cybridization of Grapefruit with ‘Dancy’ Mandarin Leads to Improved Fruit Characteristics

Aditi D. Satpute, Chunxian Chen¹, Fredrick G. Gmitter, Jr., Peng Ling, Qibin Yu, Melinda R. Grosser², and Jude W. Grosser³

Citrus Research and Education Center, University of Florida/IFAS, 700 Experiment Station Road, Lake Alfred, FL 33850

Christine D. Chase
Horticultural Sciences Department, University of Florida/IFAS, 2215 Fifield Hall, Gainesville, FL 32611

ABSTRACT. In cybridization, new combinations of nuclear and cytoplasmic genes result in a unique genotype that may bring cellular, physical, physiological, and biochemical changes to the plant. This has been demonstrated in the unexpected cybrids generated from the fusion of citrus (Citrus sp.) protoplasts in two independent experiments. The first experiment was conducted to generate potentially seedless triploids by fusing diploid protoplasts of embryogenic ‘Dancy’ mandarin (Citrus reticulata) suspension culture cells with haploid ‘Ruby Red’ grapefruit (C. paradisi) protoplasts derived from tetrad-stage microspores. After multiple attempts, only one triploid was recovered, but several diploid plants with typical grapefruit morphology were also regenerated. In the second experiment, protoplasts derived from embryogenic ‘Dancy’ mandarin suspension culture were fused with nonembryogenic protoplasts from ‘Duncan’ grapefruit leaves in an effort to produce an allotetraploid somatic hybrid. The fruit from the resulting trees resembled grapefruit in morphology and type, and maintained excellent quality throughout the summer, when commercial grapefruit rapidly loses quality. Fruit on these trees remained firm with exceptional sweetness and good flavor into August, and without seed germination. The regenerants obtained in the protoplast fusion experiments were confirmed as cybrids by genetic marker analyses. The test grapefruit were identical to commercial ‘Ruby Red’ grapefruit at six nuclear simple sequence repeat (SSR) marker loci, but identical to ‘Dancy’ with respect to a mitochondrial intron marker. The plastid genomes of individual trees originated from either fusion partner. In the first experiment, haploid ‘Ruby Red’ protoplast preparations must have also contained contaminant diploid protoplasts. Apart from the value of altered fruit quality attributes in the marketplace, these plants provide an opportunity to understand the contributions of cytoplasmic organelle genetics to important citrus fruit-breeding objectives.

The genetic improvement of citrus is a challenge because of long generation times, polyembryony, and sexual incompatibilities (Grosser and Gmitter, 1990). To overcome these barriers, protoplast fusion and plant regeneration techniques have gained importance in citrus-breeding programs (Grosser and Gmitter, 2011; Grosser et al., 2000; Vardi et al., 1989). The novelty of somatic hybridization via protoplast fusion lies in the creation of new combinations of cytoplasm and nuclear genotypes (Grosser et al., 2000; Guo et al., 2013). Two types of outcomes are recovered from somatic cell fusions in which protoplasts isolated from the embryogenic callus/cell suspension are fused with the protoplasts derived from the nonembryogenic leaf mesophyll cells—somatic hybrids and cybrids (Grosser and Gmitter, 1990). Somatic hybrids contain a hybrid tetraploid nucleus, whereas cybrid cells retain the diploid nucleus from the nonembryogenic parent (Grosser et al., 2000; Guo et al., 2013). In citrus, diploid cybrids are often generated by interspecific fusion events (Cabasson et al., 2001; Moreira et al., 2000a; Moriguchi et al., 1996), whereas tetraploid or amphidiploid somatic hybrids commonly result from intergeneric as well as interspecific combinations (Grosser and Gmitter, 2011; Grosser et al., 1992; Kobayashi et al., 1988, 1991; Louzada et al., 1992; Moriguchi et al., 1996; Ohgawara et al., 1985). These cell fusion outcomes also differ with respect to the cytoplasmic organelle (mitochondria and chloroplast) genotypes. In citrus, plants regenerated from protoplast fusion events almost always possess the mitochondrial DNA (mtDNA) of the embryogenic parent (Cabasson et al., 2001; Moriguchi et al., 1996; Saito et al., 1993; Yamamoto and Kobayashi, 1995), with some exceptions where restriction fragment length polymorphism markers revealed nonparental or recombinant mtDNA genotypes in somatic hybrids and cybrids (Moreira et al., 2000b; Moriguchi et al., 1997; Motomura et al., 1995; Vardi et al., 1989). These findings suggest that mitochondria of the embryogenic lines have a critical role in regeneration of cybrids and somatic hybrids (Grosser et al., 1996; Moriguchi et al., 1996; Saito et al., 1993; Yamamoto and Kobayashi, 1995). With respect to the chloroplast genotype of the regenerants, no specific pattern is seen. Either (Cabasson et al., 2001; Kobayashi et al., 1991; Moriguchi et al., 1996; Vardi et al., 1989) or both (Moreira et al., 2000b) of the fusion partner chloroplast DNAs (cpDNAs) are observed in cybrids as well as in the somatic hybrids. In summary, cybridization gives an opportunity to create novel, alloplasmic combinations of the nuclear and organelle genomes that are present in the fusion
partners, whereas somatic hybridization produces hybrid nuclear genomes in combination with various patterns of organelle genotypes.

In citrus cybrids and somatic hybrids, plant morphology can be variable (Cabasso et al., 2001; Ohgawara et al., 1985). Morishita et al. (1996) reported that cybrids between 'Seminole' tangelo (Citrus reticulata × C. paradisi) + 'Lisbon' lemon (C. limon) and between 'Hazzara' mandarin + rough lemon (C. jambhiri) have nonembryogenic parent 'Lisbon' or rough lemon leaf morphology, whereas somatic hybrids between 'Ohta' ponkan mandarin + rough lemon showed intermediate leaf morphology. Embryogenic parent morphology resembling sour orange (C. aurantium) or rough lemon + Microcitrus sp. (Vardi et al., 1989).

The different combinations of cytoplasmic and nuclear genomes in cybrids and somatic hybrids suggest the potential of protoplast fusion to create a wider range of unique germplasm resources for citrus improvement programs. Over the years, continuously evolving technology and the well-established protocol for cybridization have created interspecific as well as intergeneric cybrids and somatic hybrids in different citrus-breeding programs (Grosser and Gmitter, 2008, 2011; Grosser et al., 1988a, 1988b, 2000; Guo et al., 2013). Cybridization and somatic hybridization have addressed three main economically important objectives of citrus improvement: rootstock improvement (Grosser and Chandler, 2003; Grosser and Gmitter, 1990; Grosser et al., 2000) for disease resistance and soil adaptations, ploidy manipulation in the scion to create triploid seedless fruits or vigorous tetraploids (Grosser and Gmitter, 1990; Grosser et al., 2000). Fusions were conducted in 1 N HCl for 30 s followed by 15 min immersion in 20% commercial bleach containing a few drops of a surfactant (Liquinox; Alconox, White Plains, NY), then washed three times in double-distilled H₂O. Immature anthers were dissected from the buds and gently macerated into 2 mL BHM medium (Grosser and Gmitter, 1990) in 60 × 15-mm petri dishes. To provide sterile explants for leaf protoplast isolation, fruit of 'Duncan' grapefruit were washed thoroughly and surface disinfected in 1.5% sodium hypochlorite for 30 min, followed by a rinse in sterile water. Then fruit was sprayed with 95% ethanol. Fruits were cut with a sharp knife at the equatorial zone, avoiding the core where seeds are embedded. Fruit were twisted with each half in opposite directions until separated to allow for removal of intact seeds. Seeds were germinated and seedlings were maintained in Magenta vessels (GA-7; Magenta Corp., Chicago, IL) containing 50 mL RMA medium (Grosser and Gmitter, 1990; Grosser et al., 2010b) and wrapped with a sealing film (Nescofilm®; Bando Chemical Industries, Kobe, Japan). Tender, expanded leaves from the cultured nucellar seedlings were used for protoplast isolation. Shoots of the seedlings were subcultured as needed to provide a continuous source of sterile leaves.

**Materials and Methods**

**Plant materials**

**Explants for protoplast isolation.** The embryogenic suspension culture of 'Dancy' mandarin was started from unfertilized ovule-derived friable embryogenic callus cultures maintained in the citrus embryogenic callus collection of the University of Florida’s Citrus Research and Education Center, Lake Alfred [UF–CREC (Grosser and Gmitter, 1990)]. Suspension-derived protoplasts were isolated from 1- to 2-year-old suspension cultures maintained in H+H medium (Grosser and Gmitter, 1990) on a 2-week subculture cycle, with protoplasts isolated during days 4–12 (Grosser and Gmitter, 1990; Grosser et al., 2010b). Prepollen haploid protoplasts from 'Ruby Red' grapefruit were isolated as follows (Grosser and Gmitter, 1990): large elongated unopened flower buds were collected and meiotic stages were estimated by squashing anthers in water and examining the squashes using an inverted microscope at ×200 magnification. The properly staged flower buds were decontaminated by immersing in 1 N HCl for 30 s followed by 15 min immersion in 20% commercial bleach containing a few drops of a surfactant (Liquinox; Alconox, White Plains, NY), then washed three times in double-distilled H₂O. Immature anthers were dissected from the buds and gently macerated into 2 mL BHM medium (Grosser and Gmitter, 1990) in 60 × 15-mm petri dishes. To provide sterile explants for leaf protoplast isolation, fruit of 'Duncan' grapefruit were washed thoroughly and surface disinfected in 1.5% sodium hypochlorite for 30 min, followed by a rinse in sterile water. Then fruit was sprayed with 95% ethanol. Fruits were cut with a sharp knife at the equatorial zone, avoiding the core where seeds are embedded. Fruit were twisted with each half in opposite directions until separated to allow for removal of intact seeds. Seeds were germinated and seedlings were maintained in Magenta vessels (GA-7; Magenta Corp., Chicago, IL) containing 50 mL RMA medium (Grosser and Gmitter, 1990; Grosser et al., 2010b) and wrapped with a sealing film (Nescofilm®; Bando Chemical Industries, Kobe, Japan). Tender, expanded leaves from the cultured nucellar seedlings were used for protoplast isolation. Shoots of the seedlings were subcultured as needed to provide a continuous source of sterile leaves.

**Protoplast isolation and fusion.** Protoplasts were isolated from the parental suspension cultures in a 2.5:1.5 (v:v) mixture of 0.7 M BH3 protoplast culture medium and enzyme solution as described previously (Grosser and Gmitter, 1990; Grosser et al., 2010b). Sterile leaves from in vitro cultured nucellar seedlings were feather-cut with a sharp scalpel, and macerated anthers were incubated overnight (including a 25-min vacuum infiltration) in an 8.3 (v:v) mixture of 0.6 M BH3 protoplast culture medium and modified enzyme solution. Protoplasts from both suspension and leaf sources were purified by passing through a 45-μm stainless-steel mesh screen and then by centrifuging on 25% sucrose and 13% mannitol gradient. Haploid protoplasts are not amenable to purification on a sucrose–mannitol gradient, but were washed in 25% sucrose containing citrus protoplast wash (Grosser and Gmitter, 1990) nutrients to facilitate removal of debris before fusion.

In the first group of experiments, embryogenic suspension culture-derived protoplasts of 'Dancy' mandarin were fused with the tetrad-derived haploid protoplasts from 'Ruby Red' grapefruit (Fig. 1). In the second group of experiments, our highly successful model method of fusing embryogenic suspension culture-derived protoplasts of one parent with leaf-derived protoplasts of the second parent was used (Grosser and Gmitter, 1990; Grosser et al., 2000). Fusions were conducted in 60 × 15-mm polystyrene petri dishes using our standard 40% polyethylene glycol volumetric plating method (Grosser and Gmitter, 1990). Following fusion, protoplasts were cultured in a 1:1 (v:v) mixture of 0.6 M BH3 and 0.6 M EME protoplast culture media (Grosser and Gmitter, 1990), and petri dishes sealed with Nescofilm® were maintained in plastic boxes under low light in a tissue culture room.

**Plant regeneration and ploidy analysis.** Regenerating embryogenic calli and embryos were transferred to solid EME medium containing 0.5 g L⁻¹ malt extract (Perez et al., 1998) for somatic embryo induction according to Grosser and Gmitter (1990). After 4 weeks, embryos were cultured for one passage on EME-1500 media containing 1.5 g L⁻¹ malt extract for embryo enlargement, followed by one or two passages on B⁺ embryo germination medium (Grosser and Gmitter, 1990). Large embryos that failed to germinate, often exhibiting abnormal shapes, were sliced horizontally with a sharp scalpel and transferred directly to DBA3 (Deng et al., 1992) medium for adventitious
shoot induction. Recovered adventitious shoots and shoots from germinated embryos with poor root systems were cut and rooted on RMAN (Grosser and Gmitter, 1990) medium in Magenta boxes. Germinated embryos with good roots were also transferred to RMAN for plantlet enlargement. Recovered rooted plantlets were screened for ploidy level using a table top flow cytometer (Partec, Münster, Germany), as previously described (Khan and Grosser, 2004). Numerous fruit-bearing trees each from an independent regeneration event and with typical grapefruit morphology, grafted to ‘Swingle’ citrumelo (C. paradisi × Poncirus trifoliata) rootstock, were recovered from each group of experiments and were planted (1989) in the North-40 grove at UF–CREC.

DNA extraction. Plant leaf materials used for DNA extraction were collected from the experimental scions, control ‘Ruby Red’, ‘Marsh’ and ‘Duncan’ grapefruit, and ‘Dancy’ mandarin trees located at UF–CREC. All experimental scions were 25 years old and grafted onto ‘Swingle’ citrumelo rootstock. ‘Dancy’ mandarin and ‘Marsh’ grapefruit DNAs were used in the initial screening procedure to identify chloroplast primer pairs that revealed polymorphism between mandarin and grapefruit polymerase chain reaction (PCR) amplicons.

Plant genotyping

Regenerants recovered from the ‘Dancy’ + ‘Ruby Red’ experiments (C1, C2, C3, G3, G4, and G7) and ‘Dancy’ + ‘Duncan’ experiments (G1, G2, G5, and G6) were selected for genotyping (Table 1). Total genomic DNA was extracted from the experimental scions, ‘Ruby Red’, ‘Duncan’ and ‘Marsh’ grapefruit, and ‘Dancy’ mandarin using GeneElute™ plant genomic DNA Mini prep kit (Sigma-Aldrich, St. Louis, MO). Extracted DNA was quantified using a spectrophotometer (Smartspec™; Bio-Rad Laboratories, Hercules, CA), and dilutions were made to the concentration of 10 ng·µL⁻¹ in 0.1X NTE buffer [1 mM Trizma base, 0.1 mM ethylenediaminetetraacetic acid (EDTA), 0.001 M NaCl]. For nuclear genotyping, six expressed sequence tag, simple sequence repeat (EST-SSR) primers, CX6F04B, CX6F18G, CX6F29Y, CX0010B, CX0035G, and CX2021R, along with the genotyping procedure, were previously described (Chen et al., 2006, 2008). Fluorescently labeled PCR products produced on DNA templates prepared from regenerants and parental controls were fractionated on a genetic analyzer (3130 ×L; Applied Biosystems, Foster City, CA). The alleles of each EST-SSR were scored from the files with a marker analysis software (GeneMarker™; SoftGenetics, State College, PA).

The PCR primers used for organelle genotyping are described in Table 1. Forward and reverse primers flanking the mitochondrial NADH dehydrogenase subunit 7 intron 1 (nad7i1) (Grosser, 2011) were used for mitochondrial genotyping. For chloroplast genotyping, the 23 primer pairs developed for amplification of monocot cpDNA (Tsuruta et al., 2008) were tested to identify those that amplified citrus DNA and revealed polymorphism between the fusion partners. PCR reactions contained 10 ng·µL⁻¹ of genomic DNA and of the master-mix. For mitochondrial markers, 23 µL of master-mix contained 2 µM forward and reverse primer stock each (Sigma-Aldrich), 2.5 mM dNTP mix, 10X Ex Taq® Hot-Start reaction buffer (TaKaRa Bio, Otsu, Japan), 5 units/µL of TaKaRa Ex Taq® DNA Polymerase Hot-Start Version and sterile distilled water. For chloroplast markers, the PCR mixture contained 2 µM forward and reverse primer each, 2X GoTaq® Hot Start Colorless Master Mix (Promega Corp., Madison, WI) and target DNA template (10 ng·µL⁻¹). Amplification of the target DNA was achieved in a programmable thermocycler (Bio-Rad Laboratories). PCR amplification was 30 cycles of denaturation at 94 °C for 1 min, followed by annealing at 55 °C for 2 min and extension at 72 °C for 3 min.

PCR amplification was first confirmed by electrophoresis through a 1% agarose gel. For mitochondrial markers, 5 µL of PCR product, 5 µL of distilled water, and 3 µL of 5X loading dye (Bioline, Taunton, MA) was loaded into each agarose gel lane. For chloroplast markers, 5 µL of PCR product and 2 µL of loading dye were loaded in each lane. The DNA Hyperladder II™ (Bioline) was used to estimate amplicon size. Gels were run in 1X Tris-Borate-EDTA (TBE) buffer (0.1 mM Trizma base, 0.1 mM boric acid, and 0.0025 M Na₂EDTA at pH 8.2) for 45 min at 90 V for chloroplast and 60 V for mitochondrial PCR products. Gels were stained in ethidium bromide (1 µg·mL⁻¹) solution for 15 min and visualized over an ultraviolet transilluminator (Bio-Rad Laboratories). Successful PCR reactions were also fractionated on 10% precast polyacrylamide gel (Criterion™, Bio-Rad Laboratories) to determine whether polymorphisms were present between fusion partners ‘Dancy’ mandarin and ‘Ruby Red’ or ‘Duncan’ grapefruit. Acrylamide gel samples of 2 µL contained 0.5 µL PCR-amplified DNA, 0.5 µL water, and 1 µL 6X orange-blue loading dye (Promega Corp.). The G210A 100-bp DNA ladder (Promega Corp.) was used to estimate amplicon size. Acrylamide gels were run at 90 V for 4 h in 1X TBE buffer. Gels were stained in ethidium bromide (1 µg·mL⁻¹) for 15 min and visualized over an ultraviolet transilluminator. Confirmation of the mitochondria or chloroplast polymorphisms among the fusion partners and experimental scions was obtained by electrophoresis of mixed PCR products; ‘Dancy’ with experimental scion, ‘Ruby Red’ with experimental scion, and ‘Dancy’ with ‘Ruby Red’. Mixed samples were diluted in distilled water in the proportion of 1:1 to decrease the band intensity and observe the clear separation between the mixed amplicons in the same gel lane.

Fruit quality traits

In each experiment, a total of 25 fruits were harvested from experimental scions at three different time points, pooled together to analyze juice quality and yield. The Florida Department of Agriculture and Consumer Services approved juice extractor (JBT Corp., Chicago IL) and automated brix-acid unit system was used to determine fruit quality parameters—soluble solid content, total acid content, juice weight, and fruit weight.
Table 1. Citrus parents used in the protoplast fusion experiments.

| Cybrid identification no. | Embryogenic parent | Nonembryogenic parent |
|---------------------------|--------------------|------------------------|
| C1                        | Dancy              | Ruby Red               |
| C2                        | Dancy              | Ruby Red               |
| C3                        | Dancy              | Ruby Red               |
| G1                        | Dancy              | Duncan                 |
| G2                        | Dancy              | Duncan                 |
| G3                        | Dancy              | Ruby Red               |
| G4                        | Dancy              | Ruby Red               |
| G5                        | Dancy              | Duncan                 |
| G6                        | Dancy              | Duncan                 |

1Protoplast derived from cell suspension culture of ‘Dancy’ mandarin.
2Prepollen haploid protoplasts isolated from all ‘Ruby Red’ grapefruit fusion parents.
3Leaf mesophyll cells used to obtain protoplast from all ‘Duncan’ grapefruit fusion parents.

Results and Discussion

Genetic characterization of scions recovered from ‘Dancy’ + ‘Ruby Red’ and ‘Dancy’ + ‘Duncan’ fusion experiments. On the basis of six EST-SSR markers used to test the nuclear genomic composition of the experimental scions, all regenerants in both the ‘Dancy’ + ‘Ruby Red’ and ‘Dancy’ + ‘Duncan’ protoplast fusion experiments carried the allelic composition of grapefruit nuclear DNA. The allelic combinations of ‘Dancy’ mandarin were different from those of the experimental scions (Table 2).

Of the 23 monocot chloroplast primer pairs tested for PCR amplification of ‘Dancy’ and ‘Marsh’ DNA, six amplified both ‘Marsh’ and ‘Dancy’ templates, whereas two amplified ‘Marsh’ grapefruit DNA only. The six primers that amplified both citrus templates targeted intergenic and intron regions and the ndhk genomic region of the cpDNA (Table 3). Two of these six markers produced amplicons that revealed polymorphism between ‘Dancy’ and ‘Marsh’ DNA. These primers targeted the trnD-psbM intergenic region and ndhk region of the plastid genome (Table 3). These two markers were also polymorphic between ‘Dancy’ and ‘Ruby Red’ grapefruit when PCR products were fractionated on a 10% acrylamide gel (not shown). The mixture of ndhk PCR products from ‘Dancy’ and ‘Ruby Red’ or ‘Duncan’ showed better separation than the mixture of the trnD-psbM intergenic region PCR products. Therefore, the ndhk marker was used to determine the chloroplast genotype of the cybrids (Fig. 2).

PCR products amplified with the plastid ndhk primer pair revealed no polymorphism between cybrids C1, C2, C3, G3, G4, and G7 and the ‘Dancy’ fusion partner (Fig. 3A and B), but well-separated bands were observed when the PCR amplicon of ‘Ruby Red’ was mixed with that of C1, C2, C3, G3, G4, or G7 (Fig. 3A and C). PCR products amplified from the cybrids G2 and G6 revealed no polymorphism when mixed with the ‘Dancy’ PCR product. However, mixed PCR products from G1 with ‘Dancy’ and from G5 with ‘Dancy’ showed a separation of bands on the acrylamide gel (Fig. 3B). Well-separated bands were observed when the PCR amplicon of ‘Ruby Red’ was mixed with that of G2 or G6, but not when the ‘Ruby Red’ amplicon was mixed with that of the G1 or G5 amplicons (Fig. 3C). Therefore, the grapefruit cybrids G2 and G6 carried the ‘Dancy’ chloroplast genotype, whereas G1 and G5 carried the grapefruit chloroplast genotype.

The mitochondrial nad7i1 primer pair amplified the ‘Marsh’, ‘Ruby Red’, ‘Dancy’, and putative cybrid DNAs as observed on a 1% agarose gel, but did not reveal a clear polymorphism (not shown). Polymorphism was, however, observed in the mixture of PCR products amplified from ‘Dancy’ mandarin and from ‘Ruby Red’, ‘Duncan’, and ‘Marsh’ grapefruit DNA when these products were fractionated on a 10% acrylamide gel (Fig. 2). In the ‘Dancy’ + ‘Ruby Red’ protoplast fusion experiment, polymorphism was observed in the mixture of PCR product from each of the experimental scions C1, C2, C3, G3, G4, and G7 with the ‘Ruby Red’ amplicon (Fig. 4A and C), while band separation was not observed when each scion PCR product was mixed with the ‘Dancy’ amplicon (Fig. 4A and B). Therefore, all of the experimental scions were confirmed to be cybrids carrying the ‘Dancy’ mitotype in combination with the grapefruit nuclear genotype.

The mitochondrial nad7i1 primer pair was also used to identify the mitotype of each regenerant (G1, G2, G5, and G6) created by the protoplast fusion of ‘Dancy’ + ‘Duncan’.

Table 2. Organelle and nuclear genotypes of experimental citrus cybrids.

| Sample   | ndhk (bp) | nad7i1 (bp) | CX0010B | CX0035G | CX2021R | CX6F04B | CX6F18G | CX6F29Y |
|----------|-----------|-------------|---------|---------|---------|---------|---------|---------|
| Ruby Red | 650       | 1300        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| Dancy    | 550       | 1225        | 220/229 | 173/229 | 148/150 | 150/162 | 154/160 | 151/154 |
| C1       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| C2       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| C3       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G1       | 650       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G2       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G3       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G4       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G5       | 650       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |
| G6       | 550       | 1225        | 217/229 | 182/229 | 150/152 | 162/174 | 154/160 | 151/154 |

1Expressed sequenced tags-simple sequence repeats (EST-SSR) primer amplified amplicon size in DNA base pairs (bp) for nuclear genotyping.
2Amplicon size of nad7i1 mitochondrial primer pair amplified PCR product expressed in the bp.
3Amplicon size of ndhk plastid primer pair amplified PCR product expressed in the bp.
Polymorphism was observed in the mixture of PCR-amplified DNA of regenerants and ‘Ruby Red’ amplicon on 10% acrylamide gel (Fig. 4C). However, this separation was not observed when PCR-amplified G1, G2, G5, and G6 DNAs were mixed with ‘Dancy’ PCR-amplified product (Fig. 4B). The regenerants of fusion experiment ‘Dancy’ + ‘Duncan’ were, therefore, also cybrids carrying ‘Dancy’ mitochondrial genotype in combination with the grapefruit nuclear genotype.

Plant phenotypes

Cybrids generated in both fusion experiments had fruit type and leaf morphology similar to their respective nonembryogenic parents (Fig. 5A–C). Fruit obtained in the ‘Dancy’ + ‘Ruby Red’ fusion events contained very few seeds, characteristic of the ‘Ruby Red’ fusion parent. Fruit obtained from ‘Dancy’ + ‘Duncan’ fusion events were seedy, similar to that of the ‘Duncan’ fusion parent. In contrast to the wild-type ‘Duncan’ seeds, those of the cybrids did not germinate as the fruit aged on the tree (Fig. 5A and B). Examination of 10 fruit each from the cybrid ‘Duncan’ and original wild-type Duncan cultivar in late July (2014) revealed that nearly every seed in the original ‘Duncan’ had germinated, whereas it was difficult to find a germinated seed in the cybrid ‘Duncan’. Vivipary was also almost completely absent in the ‘Ruby Red’ cybrids.

Fruit quality and yield traits

Fruit and juice quality data (Table 4) were obtained at three different time points for each of the parental and cybrid selections, and for one additional commercial grapefruit cultivar—Thompson. Juice weight calculated as a percent of fruit weight showed that juice contributed greater than 50% of fruit weight in all but one sample. The soluble solids concentration and soluble solids concentration/acid ratios were always consistently higher in the cybrid fruits. In fact, some individual cybrid fruit exhibited soluble solids concentration levels higher than 13%. This increase was consistent in late summer also (Table 4). The soluble solids concentration/acid ratio is an important juice quality determinant, and fruits obtained in the fusion experiments showed a high ratio at all three time points compared with commercial grapefruit cultivars—Thompson, Duncan, and Ruby Red.

Discussion

The citrus scions regenerated from the fusion of protoplasts in two different experiments (embryogenic ‘Dancy’...
callus + tetrad ‘Ruby Red’ microspore and embryogenic ‘Dancy’ callus + nonembryogenic ‘Duncan’ leaf) resembled grapefruit in leaf morphology and fruit type. Furthermore, nuclear genotyping of all recovered putative cybrids matched that of diploid grapefruit for all six loci tested. These observations were surprising in the case of the ‘Dancy’ + ‘Ruby Red’ experiment. These scions were expected to be triploids carrying the diploid genome of the ‘Dancy’ fusion partner and one haploid genome from the ‘Ruby Red’ microspore. Contamination of the ‘Ruby Red’ haploid microspores with diploid ‘Ruby Red’ microspore mother or surrounding anther cells (Fig. 1B) is the likely explanation for this outcome. A doubled genome contribution from haploid ‘Ruby Red’ microspores is ruled out by the heterozygosity of the experimental scions at six nuclear microsatellite markers (Table 3). Grosser et al. (1996) have discussed the requirements for cybridization and the probable reasons that one of the parental nuclei degenerates to result in a cybrid as opposed to a somatic hybrid. It is also possible that imbalanced or asynchronous conditions during cell division lead to inheritance of only one of the parental nuclei into cybrids (Guo et al., 2013).

PCR assays for the analysis of mitotype showed all 10 of the experimental scions to be cybrids carrying the mitochondrial marker from the embryogenic callus fusion partner ‘Dancy’ combined with the nuclear markers derived from the non-embryogenic grapefruit fusion partner. In the absence of fully sequenced genomes, some nuclear genome contribution from ‘Dancy’ or mitochondrial genome contribution from grapefruit cannot be excluded. Plant mitochondrial genomes are relatively large and exhibit a complex organization due to active recombination (Gualberto et al., 2014; Maréchal and Brisson, 2010). This, along with the capability for mitochondrial fusion (Logan, 2010), generates the opportunity for creation of chimeric (recombinant) or multiple (heteroplasmic) mitochondrial genotypes during somatic cell fusion events. Nevertheless, our findings are consistent with the previous studies yielding citrus cybrids (Moreira et al., 2000a; Moriguchi et al., 1996; Saito et al., 1993; Yamamoto and Kobayashi, 1995). The mechanism for the biased inheritance of the mitochondrial markers from the embryogenic callus parent of the cybridization event is still unknown. There are different hypotheses proposed. Bonnema et al. (1992) interpreted the results of organelle and nuclear inheritance in the cybrids created between tomato (Solanum lycopersicum) and Solanum penellii.
as the inheritance of the organelles influenced by the nuclear background in the cybrids, whereas Earle (1995) proposed that it is spontaneous, culture-related changes rather than the effect of the fusion event. The amount of mtDNA in embryogenic cells culture is about four times more compared with non-embryogenic mesophyll cells (Moreira et al., 2000a), but this difference seems an unlikely basis for consistent transmission of the embryogenic cell mitotype. Biased partitioning of mitochondria following cell fusion might be explained by the status of mitochondria in dedifferentiated as compared with dedifferentiating cells. Massive mitochondrial fusion is observed in dedifferentiating protoplasts of tobacco (Nicotiana tabacum) mesophyll cells, whereas protoplasts of fully dedifferentiated suspension and callus cultures have numerous, small, uniformly distributed mitochondria (Sheahan et al., 2005). Hence, the physical distribution of mitochondria in the dedifferentiated embryogenic callus of ‘Dancy’ might promote the inheritance of ‘Dancy’ mitochondria to the cybrids. A critical role of mitochondria from the embryogenic callus parent in somatic embryogenesis is supported by the observation that unfused mesophyll protoplasts are unable to divide in absence of embryogenic suspension protoplasts (Moriguchi et al., 1996).

Cybrids C1, C2, C3, G2, G3, G4, G6, and G7 carried the chloroplast marker from the embryogenic callus ‘Dancy’, whereas cybrids G1 and G5 carried the marker from the nonembryogenic grapefruit fusion partner. Thus, the ‘Dancy’ + ‘Ruby Red’ fusion resulted in transmission of only the ‘Dancy’ cpDNA, whereas the ‘Dancy’ + ‘Duncan’ fusion led to transmission of the cpDNA from either parent. Additional regenerants should be investigated to determine whether there was a true bias in cpDNA transmission in the former experiment. Such a bias might result from the nature of the plastids in the unknown, diploid ‘Ruby Red’ fusion cell, as compared with the ‘Duncan’ leaf protoplast cell. Inheritance of cpDNA from either fusion partner was expected based upon past studies of cybridization (Grosser et al., 1996; Moreira et al., 2000a). In addition to the effect of alloplasty, heteroplasm present within cytoplasmic organelles is also suspected to bring desirable or undesirable nonparental changes in the regenerants. The expression of nonparental traits because of heteroplasm was observed to be quantitative and documented in both plant and animal kingdom (Picard et al., 2014; Szklarczyk et al., 2014). In this study, the absence of information of mitochondrial and chloroplast genome suggests the possibility of the presence of quantitative distribution of heteroplasm mitochondrial and chloroplast DNA sequences and its role in improving the traits in the cybrids.

New combinations of nuclear and cytoplasmic genes are predicted to bring changes to the cellular, physical, physiological, and biochemical aspects of the tree. Rootstock development, ploidy manipulation, and generation of elite germplasm for the conventional breeding are breeding goals of cybridization (Grosser and Gmitter, 2005, 2011). To date, few cybridization experiments have created economically viable cultivars, but the new cybrids created between ‘Dancy’ and grapefruit-derived protoplasts have a market potential due to attributes of fruit quality lasting well past the current harvest season. The high sugar and low acidity, combined with optimum juice content, demonstrate the potential of these newly created cybrids to stand out in competition with commercial grapefruit in the Florida juice industry and international fruit market. This finding was

Table 4. Comparison between commercial citrus cultivars and cybrid fruit for amount of juice, concentration of soluble solids and the total acid level (data from Florida Department of Agriculture and Consumer Services approved test facility).

|                   | Thompson grapefruit | Ruby Red grapefruit | Duncan grapefruit | Ruby Red cybrid | ‘Duncan’ cybrid |
|-------------------|---------------------|---------------------|-------------------|-----------------|-----------------|
|                   | JC wt (%) | SS (%) | AC (%) | JC wt (%) | SS (%) | AC (%) | JC wt (%) | SS (%) | AC (%) | JC wt (%) | SS (%) | AC (%) | JC wt (%) | SS (%) | AC (%) |
| December          | 58        | 9.93   | 1.21   | 57        | 9.33   | 1.32   | 62        | 8.52   | 1.28   | 56        | 11.64 | 1.44   | 45        | 10.87  | 1.38  |
| April             | 36        | 10.43  | 1.21   | 67        | 9.52   | 1.3    | 53        | 9.12   | 1.37   | 61        | 13.27 | 1.2    | 57        | 12.47  | 1.17  |
| June              | 64        | 10.36  | 1.18   | 67        | 9.51   | 1.08   | 51        | 9.42   | 1.23   | 52        | 11.67 | 1.19   | 54        | 14.05  | 1.03  |

*Percentage juice content present in the fruit weight of each genotype. Sample size of each genotype is between 20 and 30 fruit.

*Measures the percentage of concentration of total soluble solids in the fruit juice of experimental cybrids and commercial grapefruit cultivars.

*Measures percentage of total acids present in the juice.

JC = juice content; SS = soluble solids; AC = acid content.
unexpected based on the results of Bassene et al. (2008), who showed that in ‘Eureka’ lemon cybrids carrying the ‘Willowleaf’ mandarin mitochondria, fruit quality traits, including sugar content, largely matched those of the nuclear genome donor. One clone from this work, cybrid ‘Ruby Red’ C2 has already been released by UF for commercialization under the proposed name ‘Summer Gold Grapefruit’. This clone has the potential to extend the grapefruit packing season, and high concentration of soluble solids grapefruit brings higher prices in several international markets. A cybrid ‘Duncan’ clone will be released subsequently for dooryard use. (The high seed content precludes commercial use.)

Further research is required to understand and predict the contributions of cytoplasmic organelle genetics to important breeding traits. It was recently shown that nuclear gene expression is changed after acquiring foreign mitochondria from ‘Willowleaf’ mandarin under nuclear background of ‘Eureka’ lemon in the cybrids created by fusion between embryogenic ‘Willowleaf’ mandarin and nonembryogenic ‘Eureka’ lemon (Bassene et al., 2011). The transcriptome and proteome analysis conducted to quantify the differential expression between cybrids and its fusion partners also suggests the modulation of mitochondrial genes under the effect of foreign nuclear genotype or vice versa (Wang et al., 2010; Zheng et al., 2012). The cybrid fruit traits observed in the fusion between ‘Dancy’ + ‘Ruby Red’ and ‘Dancy’ + ‘Duncan’ will provide another approach toward understanding nuclear and cytoplasmic interaction in the control of advantageous plant phenotypes.

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