Fruit Quality in Induced Polyploids of *Actinidia chinensis*

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Abstract. Fruit of colchicine-induced tetraploids of *Actinidia chinensis* were 50% to 60% larger than those of their diploid progenitors. In addition to fruit size, fruit quality is a key issue in any commercialization of these autotetraploids. We have made the first detailed study of the effects of chromosome doubling on fruit quality parameters other than size: these attributes include flesh firmness, color, soluble solid content (SSC), dry matter content (DM), vitamin C content, fruit skin thickness, and red pigmentation distribution in red-fleshed kiwifruit. Four selections from colchicine-induced tetraploids from the yellow-fleshed kiwifruit cultivar Hort16A were trialed for the stability of their fruit quality. Analysis of fruit at harvest over 3 years showed that fruit of the induced autotetraploids were significantly softer (lower flesh firmness), had lower DM, and had a less intense golden flesh color than fruit of their diploid progenitor. During development, SSC of fruit of the autotetraploid plants started to increase earlier than in the diploid Hort16A. This has been confirmed by replicated trials. No difference was found in vitamin C content between fruit of autotetraploids and diploids. Autotetraploids had significantly thicker skins than diploid Hort16A. Induced autotetraploids from three female genotypes of red-fleshed *A. chinensis* showed similar trends to autotetraploids of Hort16A in fruit flesh firmness and outer pericarp flesh color, DM, SSC, and vitamin C. All the traits analyzed indicated that fruit of the autotetraploid plants matured earlier than those of their diploid progenitors. Furthermore, red pigmentation, one of the most important traits for red-fleshed kiwifruit breeding, showed a reduction in both intensity and distribution in the autotetraploids compared with their diploid progenitors. There was considerable variation among fruit of autotetraploid plants regenerated from each diploid progenitor. Therefore, selection among the regenerants may be required to achieve the best outcome after ploidy manipulation in kiwifruit breeding.

We thank Nihal de Silva for statistical advice and Ian Hallett for guidance on measurements of skin thickness; Robert Campin, Paul Datson, Meng Meng, Eric Wu, and Hao Wu for assistance with fruit analysis and collection of some data; Tony Corbett for figure design; Jem Burdon, Jinquan Feng, and Anne Gunson for critical comments on the manuscript; and Mike Currie, Pauline Mooney, and Alan Seal for useful discussions.

Received for publication 23 Jan. 2013. Accepted for publication 11 Apr. 2013.

Materials and Methods

Plants. The plants used comprised 11 clonally propagated vines of the diploid *A. chinensis* cultivar Hort16A, and 77 independently derived colchicine-induced tetraploid regenerants of this cultivar. There were 11 clonally propagated plants of three genotypes of *A. chinensis* with red flesh, ‘Hort22D’, Selection 1 and Selection 2 (Wu et al., 2012), and 88 independently derived colchicine-induced autotetraploid plants produced from these three genotypes. Autotetraploids of ‘Hort16A’ were grouped into Type A (55 regenerants with large fruit of the same general shape as those of their diploid progenitor, ‘Hort16A’) and Type B (22 regenerants with smaller “fasciated” fruit); no such variation was observed in the autotetraploid plants regenerated from the three diploid red-fleshed genotypes of *A. chinensis* (Wu et al., 2012). All the regenerated autotetraploids were grown in vitro and were on their own roots.

The plants of ‘Hort16A’ and the 77 tetra-
males from the Plant & Food Research germplasm collection and a diploid *A. chinensis* male cultivar (Meteor) (1:2) were planted at a ratio of one male:eight female plants.

The three “red-fleshed” genotypes and their autotetraploids were planted in the Plant & Food Research Orchard at Te Puke. The diploid male (‘Meteor’) was planted at a ratio of one male:six females along the same row.

The tetraploid plants with their clonally propagated progenitor plants as controls, and males as pollenizers, were planted randomly in the rows of the two orchards as described by Wu et al. (2012). All vines were managed following standard orchard practices and grown on a modified T-bar structure.

**Fruit quality attributes.** Fruit quality was evaluated on a random sample of 10 fruit/year (2006–08) that had been used to measure individual fruit weights and dimensions as described previously by Wu et al. (2012). Measurements began in the third year after planting in the orchard. Fruit were harvested on 23 May in 2006, 5 Apr. in 2007, and 11 May in 2008. The method of sampling was described previously (Wu et al., 2012). Fruit quality was evaluated over three seasons (2006–08) for diploid ‘Hort16A’ and its autotetraploid regenerants and two or three seasons (2007, 2008, and 2009) for the three diploid red-fleshed genotypes and their autotetraploid regenerants. Fruit attributes other than fruit weight and dimensions were evaluated as follows:

- **Flesh firmness** was measured, using a handheld penetrometer (EFFEGI, Alphonsine, Italy; 8-mm probe), at opposite sides at the equator of the fruit after removal of a 1-mm layer of skin and outer pericarp. The two readings were averaged (Feng et al., 2003).
- **Flesh color** in the outer pericarp was measured at opposite sides at the equator of each fruit using a Minolta CR-300 Chroma meter (Osaka, Japan) after removal of a 2-mm layer of skin and outer pericarp. The two readings were averaged and color was expressed as hue angle (h°), chroma (C*), and lightness (L*) (Feng et al., 2003).
- **Soluble solids content** was measured using a handheld refractometer (Atago®, Tokyo, Japan) with one drop of juice from each end of the fruit combined.
- **Dry matter content** was measured by weighing a 4-mm thick equatorial slice of fruit cross-section (including skin, flesh, seeds, and core tissue) before and after drying at 60 °C for 24 h.
- **Vitamin C** (ascorbic acid) was measured by the method described by Ferguson and MacRae (1992) using three batches of 15 fruit from each plant. A composite sample from each batch of ~300 g was prepared by taking a sector, approximately one-sixth, from each fruit. The reduced ascorbic acid of the samples was measured by titration against 2,6-dichlorophenolindophenol (Deutsch, 1996).

Skin thickness was measured by taking a section of the skin using a Vibratome sectioning system (Series 1000, USA), photographing with a digital camera attached to an Olympus microscope (Vanox-AHBT3), and determining skin thickness using an ImageJ system (image process and analysis in Java).

**Pigmentation in red-fleshed Actinidia.** The intensity of red pigmentation was scored on a 0 to 5 scale with no red scored as 0 and scores 1 to 5 (right to left, Fig. 1) representing a slight tinge of red, pale pink or red color, distinct red, dark red, and intense red, respectively, in equatorial cross-sections (Cheng et al., 2007). Pigment distribution was scored separately in fruit cross and longitudinal sections. The transverse distribution was scored by taking an equatorial cross-section of the fruit and recording the extent to which red pigments were distributed radially from the outside of the core to the skin using a score of 0 to 10 based on 10% increments of the radial length. The longitudinal distribution was scored by recording the extent to which red pigmentation extended from the stalk end to the stylar end based on 10% increments of the locule length (Fig. 2).

**Fruit flesh firmness**

‘Hort16A’. Fruit of induced autotetraploid regenerants of ‘Hort16A’ varied in flesh firmness. Data from three consecutive harvests showed that firmness varied by year (because of different harvest dates), vine, regenerant, and fruit type (Fig. 3). Fruit of induced autotetraploid Type B plants were significantly softer than fruit of Type A plants in 2007 and 2008, but no difference was detected in 2006; fruit of both Type A and Type B autotetraploids were softer than those of their progenitor, diploid ‘Hort16A’, at each harvest date (Table 1). This trend was consistent over years.

Fruit from the replicated trial of four different autotetraploid Type A regenerants were also softer than those of their progenitor, diploid ‘Hort16A’, at all sampling times. Firmness measured at different sampling dates indicated that the fruit from autotetraploids
began to mature earlier than those of ‘Hort16A’ (Fig. 4A).

**Red-fleshed genotypes.** Fruit of induced autotetraploids were much softer; flesh firmness was \( \text{C}25 \) \( \text{50 to 60 N} \) lower than that of fruit of their diploid progenitors in all three genotypes analyzed from the 2009 evaluation (Table 2). However, these differences, although dramatic, are for only one season.

**Flesh color**

‘Hort16A’. The differences in outer pericarp flesh color were similar to those found in fruit firmness: ‘Hort16A’ had significantly higher fruit flesh color reading values than the autotetraploids induced from it (\( P < 0.01 \)) (Table 3).

Significant differences in fruit flesh color of the outer pericarp between years were detected, but the relative differences between the diploids and tetraploids were maintained. Variation in outer pericarp flesh color was also observed among induced autotetraploid regenerants and vines of diploid ‘Hort16A’ (data not shown).

Flesh color reading values, e.g., hue angle (\( h^* \)), from fruit of the replicated trial of four selected autotetraploid Type A regenerants of ‘Hort16A’ also showed similar trends in that they were lower than in their progenitor ‘Hort16A’ during the study (Fig. 4B–D).

**Red-fleshed genotypes.** Similar results were found for the genotypes that had red pigmentation in the inner pericarp. Each year, significantly higher values for lightness and chroma for outer pericarp flesh (yellow) were recorded for all three diploid genotypes of red-fleshed *A. chinensis* than for the corresponding induced autotetraploids. The one exception was that in 2009, the value for lightness of Selection 2 was not significantly higher. Hue angle varied with year and with genotype (Table 4).

In all 3 years, the intensity of the fruit red pigmentation in the inner pericarp was significantly greater in diploids of ‘Hort22D’ than in autotetraploids induced from it (\( P < 0.05 \)) (Table 4). Mixed results were obtained for the other genotypes. There was considerable variation in fruit pigmentation among fruit of autotetraploid regenerants derived from the same diploid progenitor. Some autotetraploids had deeper red pigmentation in the cross-section than the diploid progenitor.

**Soluble solids content**

‘Hort16A’. Generally, Type B autotetraploid fruit had significantly higher SSC at harvest than fruit of Type A or of the diploid progenitor (Table 1). SSC varied with vine

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**Table 1.** Fruit flesh firmness, soluble solids content, and dry matter content of diploid *Actinidia chinensis* ‘Hort16A’ and regenerated plants of colchicine-induced autotetraploids derived from it.

<table>
<thead>
<tr>
<th>Group</th>
<th>Ploidy</th>
<th>Fruit flesh firmness (N)</th>
<th>Soluble solids content (°Brix)</th>
<th>Dry matter content (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hort16A</td>
<td>2x</td>
<td>14.7 ± 1.0 a(^*)</td>
<td>69.6 ± 0.2 a</td>
<td>41.2 ± 6.9 a</td>
</tr>
<tr>
<td>Type A</td>
<td>4x</td>
<td>7.8 ± 0.4 b</td>
<td>53.9 ± 1.0 b</td>
<td>32.4 ± 2.0 a</td>
</tr>
<tr>
<td>Type B</td>
<td>4x</td>
<td>8.8 ± 1.0 b</td>
<td>48.0 ± 2.0 c</td>
<td>17.7 ± 2.9 b</td>
</tr>
</tbody>
</table>

\(^*\)Ten-fruit samples taken from each vine during 2006–08. All data are averages ± SEs.

\( \text{Values followed by different letters within the same column for the same item in the same year are significantly different (} P < 0.01) \).
and regenerants among the three genotypes and also with year over the 3 years among the three groups (Fig. 3).

Fruit from the replicated trial of four autotetraploid Type A regenerants showed the same trends in SSC as fruit of their diploid progenitor, ‘Hort16A’, except that all fruit from autotetraploids initially had a higher SSC and a greater rate of increase in SSC than fruit of their diploid progenitor, ‘Hort16A’ (before 4 May). Eventually the SSC values of fruit of the diploid were as high (or even higher) as those of fruit of the autotetraploids (Fig. 4E).

Table 2. Fruit flesh firmness, soluble solids content, and dry matter content of three diploid genotypes of red-fleshed Actinidia chinensis and of autotetraploids induced from them by colchicine.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Ploidy</th>
<th>Fruit flesh firmness (N)</th>
<th>2009</th>
<th>2007</th>
<th>2008</th>
<th>2009</th>
<th>Soluble solids content (°Brix)</th>
<th>2009</th>
<th>2007</th>
<th>2008</th>
<th>2009</th>
<th>Dry matter content (%)</th>
<th>2007</th>
<th>2009</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hort22D</td>
<td>2x</td>
<td>93.2 ± 11.8 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>12.4 ± 1.4 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>16.7 ± 0.7 a</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4x</td>
<td>27.5 ± 3.9 b</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>11.7 ± 1.1 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>15.9 ± 0.3 a</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selection 1</td>
<td>2x</td>
<td>71.6 ± 11.8 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>15.1 ± 0.4 a</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selection 1</td>
<td>4x</td>
<td>16.7 ± 3.9 b</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>—</td>
<td>16.7 ± 0.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>14.2 ± 0.4 b</td>
<td></td>
</tr>
<tr>
<td>Selection 2</td>
<td>2x</td>
<td>67.7 ± 0 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>14.4 ± 2.1 b</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>14.7 ± 0.2 a</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selection 2</td>
<td>4x</td>
<td>14.7 ± 2.9 b</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>16.7 ± 3.2 a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>14.7 ± 0.3 b</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

^All data are averages ± SEs.

^Values followed by different letters within the same column for the same item of the same genotype in the same year are significantly different; for fresh firmness, the significant difference at P < 0.01; for soluble solid content and dry matter content, the significant difference at P < 0.05.

^Insufficient fruit.

Table 3. Outer pericarp flesh color: hue angle (h°), chroma (C*), and lightness (L*) of fruit from vines of diploid Actinidia chinensis ‘Hort16A’ and regenerated plants of colchicine-induced autotetraploids derived from it.

<table>
<thead>
<tr>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Hort16A</td>
<td>2x</td>
<td>99.3 ± 0.1 a</td>
<td>105.0 ± 0.6 a</td>
<td>102.6 ± 0.3 ab</td>
<td>31.6 ± 0.3 a</td>
<td>37.3 ± 0.5 a</td>
<td>32.0 ± 1.0 a</td>
<td>67.1 ± 0.1 a</td>
<td>75.6 ± 0.6 a</td>
<td>69.3 ± 1.0 a</td>
</tr>
<tr>
<td>Type A</td>
<td>4x</td>
<td>98.1 ± 0.2 b</td>
<td>102.8 ± 0.2 b</td>
<td>102.7 ± 0.3 a</td>
<td>27.0 ± 0.4 b</td>
<td>34.2 ± 0.2 b</td>
<td>28.7 ± 0.5 b</td>
<td>62.4 ± 0.4 b</td>
<td>72.9 ± 0.2 b</td>
<td>65.0 ± 0.4 b</td>
</tr>
<tr>
<td>Type B</td>
<td>4x</td>
<td>98.2 ± 0.4 ab</td>
<td>102.2 ± 0.4 b</td>
<td>101.3 ± 0.4 b</td>
<td>27.5 ± 0.7 b</td>
<td>35.7 ± 0.5 b</td>
<td>27.7 ± 0.6 b</td>
<td>58.0 ± 0.6 c</td>
<td>70.8 ± 0.4 c</td>
<td>60.7 ± 0.8 c</td>
</tr>
</tbody>
</table>

^Ten-fruit samples were taken from each vine during 2006–08. All data are averages ± SEs.

^Values followed by different letters within the same column for the same year are significantly different (P < 0.01).
Dry matter content

‘Hort16A’. Fruit of diploid ‘Hort16A’ had the highest DM, autotetraploid Type A fruit had the lowest DM in all 4 years of the study, and Type B fruit had intermediate values (Table 1). Fruit in the replicated trial of the four autotetraploid Type A regenerants gave similar results; at all sampling times, they all had lower DM than did fruit of the diploid progenitor ‘Hort16A’ (Fig. 4F).

Variation in fruit DM between years from 2006 to 2008 was observed among the three groups, the variation of fruit DM from all three groups of fruit having the same trend among the replicate trials (Table 2). Only in 2006 was DM significantly higher than in the other two seasons ($P < 0.01$); no significant differences were found between the other two seasons.

Variation was also observed between vines or regenerants within the same group in any one year (Fig. 3). In 3 years’ observation with subsamples from all vines or regenerants, DM of ‘Hort16A’ fruit was 16.0% to 19.5%, of Type B fruit 13.3% to 19.3%, and of Type A fruit 12.4% to 18.3%.

Red-fleshed genotypes. In both years, fruit of the induced autotetraploids had lower DM than did fruit of the respective diploid progenitors. The differences between the two ploidy levels varied with the genotype (Table 2). DM also varied between years.

Vitamin C content

‘Hort16A’. Fruit of the original diploid ‘Hort16A’ and of the Type A and Type B autotetraploids all contained 100 to 120 mg vitamin C/100 g fresh weight irrespective of ploidy. The differences observed in vitamin C content (although higher in autotetraploids) were not statistically significant.

Red-fleshed genotypes. Differences in fruit vitamin C content between the original diploids and induced autotetraploids were not statistically significant. Differences in fruit vitamin C content of the different diploid genotypes were paralleled in the autotetraploids induced from those diploids: the diploid with the highest vitamin C concentrations produced autotetraploids with the highest concentrations. Similarly, the diploid progenitor with the lowest vitamin C produced autotetraploids with the lowest concentrations (Table 5).

Skin thickness of ‘Hort16A’

All fruit of induced autotetraploids from ‘Hort16A’ had brown skins at harvest, whereas fruit of the diploid progenitor vines often had greenish skins. Such differences in color may be attributable, at least in part, to differences in canopy development and hence exposure of the fruit to light. However, fruit skins of the autotetraploids were also significantly thicker than those of the fruit of the diploid progenitor (Table 6). Type B autotetraploid fruit had the thickest skins.

Discussion

Inducing a doubling of the ploidy of the diploid kiwifruit selections we studied significantly increased fruit size: fruit of the colchicine-induced autotetraploids were, on average, 50% to 60% bigger than those of their diploid kiwifruit progenitors (Wu et al., 2012). Some spontaneously occurring budsports of ‘Hort16A’ also have fruit twice the average size of ‘Hort16A’ fruit (Martin, 2005). A number of these budsports have since been shown by flow cytometry to be tetraploid or mixoploid (diploid + tetraploid) (Ferguson et al., 2009). Increasing the ploidy, either spontaneously or by manipulation, can therefore increase fruit size in some kiwifruit selections. However, fruit size is only one attribute of fruit quality, and many other quality attributes will be important in determining whether the colchicine-induced autotetraploids can be used commercially.

An increase in fruit size was often accompanied by changes in fruit shape (Wu et al., 2012). The colchicine-induced autotetraploid regenerants from ‘Hort16A’ could be divided into two classes, Type A and Type B, on fruit shape. Type B fruit would not be acceptable commercially because of their shape.

Fruit of the colchicine-induced autotetraploids had lower flesh firmness and lower viscosity, but this was not significantly different among the three ploidy levels of the original diploid ‘Hort16A’. This means that fruit firmness and viscosity data were not significantly different for the diploid progenitors of the autotetraploids (Table 2).

In both years, fruit vitamin C content of ‘Hort16A’ was similar to that of the Type A and Type B autotetraploids, indicating that vitamin C content was not significantly different. However, this does not mean that vitamin C content was the same; vitamin C content of ‘Hort16A’ was significantly different from that of the Type A and Type B autotetraploids (Table 5).

Table 5. Vitamin C content in 2008 of fruit of three diploid genotypes of red-fleshed Actinidia chinensis and the autotetraploids induced from them by colchicine.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Ploidy</th>
<th>Number of vines or regenerants</th>
<th>Vitamin C (mg/100 g fresh weight)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hort22D</td>
<td>2x</td>
<td>4</td>
<td>147.9 ± 8.2</td>
</tr>
<tr>
<td>Hort16A</td>
<td>2x</td>
<td>4</td>
<td>156.3 ± 9.4</td>
</tr>
<tr>
<td>Selection 1</td>
<td>2x</td>
<td>4</td>
<td>121.6 ± 5.2</td>
</tr>
<tr>
<td>Selection 2</td>
<td>2x</td>
<td>1</td>
<td>112.9</td>
</tr>
<tr>
<td>Selection 3</td>
<td>2x</td>
<td>4</td>
<td>117.8 ± 6.7</td>
</tr>
</tbody>
</table>

All data are averages ± ses.

Table 6. Fruit skin thickness measured in 2007 of diploid Actinidia chinensis ‘Hort16A’ and the autotetraploids induced from it by colchicine.

<table>
<thead>
<tr>
<th>Group</th>
<th>Ploidy</th>
<th>Vines sampled</th>
<th>Skin thickness (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hort16A</td>
<td>2x</td>
<td>2</td>
<td>0.025 ± 0.001 c</td>
</tr>
<tr>
<td>Type A</td>
<td>2x</td>
<td>2</td>
<td>0.049 ± 0.001 b</td>
</tr>
<tr>
<td>Type B</td>
<td>4x</td>
<td>2</td>
<td>0.046 ± 0.002 a</td>
</tr>
</tbody>
</table>

All data are averages ± ses.

Values followed by different letters are significantly different ($P < 0.01$).
DM than fruit of their diploid progenitors. There was some variation in these characteristics from year to year because fruit were harvested at different maturity in different years but the trends in variation were the same for both autotetraploids and their progenitors. Fruit of Type B autotetraploid had the highest SSC, but the difference in SSC between Type A and ‘Hort16A’ fruit varied with year during the 3-year analysis. These variations with year make it difficult to draw firm conclusions. The variation between years in fruit firmness and SSC was largely the result of the fruit being harvested at different maturities each year. The differences and variation in DM, SSC, and flesh color between ‘Hort16A’ (diploid) and their induced tetraploids were confirmed by a trial of selected four large-fruited regenerants (with Type A fruit). In that trial, the trends were consistent and revealed a change in fruit texture. Unfortunately, data for only 1 year are available, because many of the plants in the trial have since been lost to bacterial canker of kiwifruit (caused by Pseudomonas syringae pv. actinidiae). Likewise, many of the original induced autotetraploid plants of the red-fleshed genotypes have since died.

A preliminary report described how fruit of spontaneous large-fruited budsports of ‘Hort16A’ likewise had lower flesh firmness, lower DM, and lower SSC than fruit of the normal diploid ‘Hort16A’ (Martin, 2005). Increasing ploidy had no effect on the fruit vitamin C concentrations in the kiwifruit selections we studied. Although tetraploids contained slightly higher vitamin C concentrations than their diploid progenitors, the differences were not significant. Differences among the three red-fleshed autotetraploid genotypes in vitamin C concentrations therefore reflected the differences among the diploid progenitors. In some other crops, increases in ploidy have resulted in higher concentrations of vitamin C. In watermelons, doubling the ploidy of nine different cultivars consistently resulted in an increase in vitamin C (Cheng et al., 2008) and similar results were obtained with eggplant (Li et al., 2002), and Ullucus (Viehmannová et al., 2012). Chromosome doubling might therefore be a useful tool for improving the antioxidant properties of some crops, but apparently not in kiwifruit.

The reduction in fruit DM on doubling the ploidy is particularly important. This is not simply an effect of fruit size: when large numbers of ‘Hayward’ kiwifruit were graded in size before export, there was no correlation between fruit size and fruit DM (Woodward and Clearwater, 2008).

Lower DM will mean fruit flavor is likely to be poor after storage (Harker et al., 2009). Fruit that are low in DM will inevitably be poor after storage (Burdon et al., 2004): because fruit of the autotetraploids were lower in DM than their diploid progenitors, they can be expected to be lower in SSC when ripe. DM has been proposed as an indicator of maturity (and harvest time) and is used to determine returns to kiwifruit orchardists in New Zealand (Burdon et al., 2004; Jaeger et al., 2011). ZESPRI Group Limited (the main exporter of kiwifruit from New Zealand) believes that to maintain its international competitive advantage, and hence higher returns to growers, it needs the fruit it exports to be consistently of higher DM content to ensure that customers receive the fruit they expect with enhanced sweetness, flavor, and storage life (Belrose, Inc., 2009; Jaeger et al., 2011). Thus, although the increased size of the fruit of autotetraploids could provide a commercial advantage, the lower DM would be a real disadvantage.

Commercial priorities will determine the relative importance placed on size and flavor in the market acceptance of new cultivars.

Fruit of Type C autotetraploid increased faster than that of ‘Hort16A’ early in the season, although later on the values were the same or even lower than that of ‘Hort16A’. The initial faster increase could be the result of the fruit of the autotetraploid maturing earlier than fruit of the corresponding diploids. The reduced firmness and the earlier changes in SSC imply that the fruit of autotetraploids do mature earlier than fruit of the diploid progenitors. This conclusion is consistent with studies in other crops: in the grape cultivar Muscat Hamburg, fruit of the autotetraploids were larger and ripened 10 d earlier than those of diploid plants (Luo et al., 1997); autotetraploid eggplants showed a similar trend in ripening earlier (Li et al., 2002).

The reduction in fruit firmness and DM could be the result of the fruit of autotetraploids having larger cells with larger intercellular spaces; perceived flesh texture could likewise be affected. Polyploids generally have larger cells than their diploid progenitors, as shown by the increased size of their stomata (Van Laere et al., 2011). In ryegrass, such increase in cell size has been shown to be the result of greater elongation rates (Sugiyama, 2005); this fast growth of autotetraploids results in production of the less biomass and lower DM. Ploidy manipulation in other plants often has the same result. In Ullucus, starch content and DM in the microtubers decreased with increased ploidy (Viehmannová et al., 2012), and in Spathiphyllum wallisii, biomass production of tetraploids was less than that of their diploid progenitors (Van Laere et al., 2011). In other cases, however, doubling the chromosome number tended to improve fruit quality, because autotetraploids had higher SSC, sugar content, or other indicators of quality than the diploids, e.g., autotetraploid ‘Royal Gala’ apple (Liu et al., 2006), autotetraploid ‘Meiwa’ kumquat (Nukaya et al., 2009), and autotetraploid eggplant (Li et al., 2002). These apparently contrasting effects are not really contradictory; different attributes of fruit quality were assessed, e.g., texture (flesh firmness), DM, SSC, and/or sugars; furthermore, the various fruits studied are also very different in structure to kiwifruit and generalizations should be avoided. It is also possible that some of the effects we observed are the result of our comparing plants that had arisen from micropropagation with those resulting from vegetative propagation. Fruit from ‘Hayward’ kiwifruit plants raised by micropropagation had lower DM than fruit from plants vegetatively propagated by cuttings, but no differences were found in firmness and SSC (Monasta and Testoni, 2009). Further work on fruit structure, physiology, and biochemistry could help us to explain these results and further to explain the effects of ploidy levels on flesh texture and composition.

The color of the fruit flesh of ‘Hort16A’ and other yellow- or green-fleshed kiwifruit is determined by the relative amounts of chlorophyll and carotenoids (McGillie and Ainge, 2002). As fruit of ‘Hort16A’ and many other A. chinensis selections mature, chlorophyll is lost and the pericarp changes from an initial light green to light yellow (Montefiori et al., 2009) as shown by a reduction in hue angle. A hue angle of 103° is one of the main indicators that ‘Hort16A’ fruit in commercial orchards have reached harvest maturity (Minchin et al., 2003). The fruit used in our studies were harvested later in the season than is usual in commercial orchards, but outer pericarp color measurements, especially of hue angle, indicated that fruit of the autotetraploids derived from ‘Hort16A’ mature earlier than that of those of their progenitor, diploid ‘Hort16A’. This conclusion is consistent with measurements of their fruit flesh firmness and SSC.

In the red-fleshed selections studied, red pigmentation is largely restricted to the inner pericarp and the decrease in hue angle because the outer pericarp changes from green to yellow can likewise be used as an indicator of harvest maturity. Again it seems that fruit of the autotetraploids might be maturing earlier than those of their diploid progenitors.

If autotetraploid kiwifruit consistently mature earlier, increasing the ploidy could be an interesting way of inducing earlier maturity and fruit harvest, possibly extending the harvest and marketing periods and avoiding the risk of fall frosts. However, to be commercially useful, there would need to be at least several weeks’ advance in maturity dates and in our experiments, this seems unlikely with these induced tetraploid kiwifruit.

Fruit of autotetraploid ‘Hort16A’ have significantly thicker skins than do fruit of diploid ‘Hort16A’’. This could be a real advantage, because many otherwise promising diploid selections of A. chinensis, especially red-fleshed selections, have very thin skins, which make them very susceptible to environmental stress, mechanical damage while still on the vine such as wind rub, or damage during harvest. Thicker skins could also be an advantage during storage because they might reduce water loss. However, we have not yet studied skin anatomy in detail or determined whether thicker skins do in fact result in higher pack-out rates or less damage during storage. Thicker skins did not seem to retard fruit maturation.

Red-fleshed kiwifruit, especially selections of A. chinensis, are currently stimulating great interest among fruit breeders and the kiwifruit industry in general (Ferguson
and Seal, 2008; Jiang et al., 2009). One of the most serious defects affecting commercial development of most red-fleshed selections available at present is their small fruit size (Cui, 1993). This might be overcome by chromosome doubling with colchicine (Wu et al., 2012). However, fruit of autotetraploid regenerants generally were less intensely colored and the distribution of pigments was more limited than in the diploid progenitors. The increase in fruit size after chromosome doubling is attributable mainly to the increase in fruit transverse diameters and this might affect both the concentration and distribution of anthocyanins. However, the distribution and intensity of red pigmentation varied in individual regenerants, suggesting that further selection could result in acceptably colored fruit.

Our results have showed that chromosome doubling significantly affected both fruit quality and fruit maturation in addition to fruit size and morphology. However, we have studied only some of the characters that determine fruit quality and whether a genotype produces fruit that meet commercial requirements. Core size and texture are also important as is the presence of any core cavities. Long storage life is another obvious requirement. However, the variation already observed in some important fruit quality characters among each group of autotetraploids indicates that there is a good chance of selecting desirable regenerants to be used as parents or directly as new commercial cultivars. Manipulation of ploidy certainly shows promise as a way of overcoming ploidy barriers in a genus such as Actinidia with species at different ploidy levels.

Literature Cited


