A Tomato Triploid Hybrid Whose Double Genome Parent Is the Male

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Abstract. A spontaneous tomato (Lycopersicon esculentum Mill.) triploid hybrid was analyzed by isozyme and restriction fragment length polymorphism profiles. The double chromosome complement donor was shown to be the male parent, contrary to the prevailing hypothesis.

Obtaining triploid plants from crosses between tetraploids and diploids does work for some crops such as watermelon (Citrus lanatus Thunb.) (Kihara, 1951) and lettuce (Lactuca sativa L.) (Eenink, 1980). However, this result is prevented by a strong block in other crops, such as alfalfa, (Medicago sativa L.) (Veronesi et al., 1986), clover, (Trifolium pratense L.) (Parrot et al., 1985), potato, (Solanum tuberosum L.) (Hanneman and Peloquin, 1968), and tomato (Nilsson, 1950). This problem is due not to lack of fertilization or nonviability of the embryo, but rather to endosperm failure (Cooper and Brink, 1945) stemming from chromosome imbalance (Johnston et al., 1980). Thus, any cross that fails to produce the right ratio of 2 maternal : 1 paternal chromosome complement will result in unbalanced endosperm, and, ultimately, in abortive seeds.

Apparantly, viable 2n ovules and pollen grains are produced in 2n plants like potato (den Nijs, 1977), maize (Zea mays L.) (Rhoads and Dempsey, 1966), alfalfa (Veronesi et al., 1986), and clover (Parrot et al., 1985). A theory of direct tetraploidization through 2n gametes has been published (Parrot et al., 1985). However, triploid formation through such 2n gametes resembles the difficult cross between tetraploids and diploids in that it produces unbalanced endosperm (Veronesi et al., 1986). However, spontaneous triploids may be obtained for many plants such as cannot be obtained by 4x x 2x crosses, including soybean [Glycine max (L.) Merr.] (Chen et al., 1985), apple (Malus pumila Mill.) (Singh and Wafai, 1984), tomato (Rick, 1945), and cabbage (Brassica oleracea L.) (Mackay and Low, 1975). The prevailing hypothesis is that spontaneous triploids probably arise through the female parent as the donor of the double chromosome complement (Einset, 1948; Chen and Palmer, 1985). This path has been shown by Chyi and Weeden (1984) for certain triploid apple (Malus domestica Borkh.) cultivars. A possible mechanism may be a fertilized endosperm cell overturning a degenerated fertilized egg (Munyamma, 1977).

Our study resulted from research aimed at estimating the commercial potential of triploid tomato fruits. An organized search for spontaneous triploids in commercial greenhouses was undertaken in the Negev (southern) area of Israel. Several hybrid triploids were discovered, and one, arisen in a greenhouse of the ‘FC121’ tomato hybrid, was studied. With appropriate hormonal treatment its fruits are juicy, seedless, good tasting, and 50% larger than those of their diploid hybrid counterpart (Kagan-Zur et al., 1991). Its commercial potential is currently being assessed.

Since the triploid was a hybrid of two distinct lines, it enabled us to attempt to identify the double genome donor parent. Here we present evidence that spontaneous triploidy can be effected through the male parent and may be a common occurrence in tomato.

Materials and Methods

A spontaneous triploid ‘FC121’ tomato hybrid spotted in a commercial greenhouse in the Negev area and the diploid ‘FC121’ were propagated through cuttings. Parent lines of the ‘FC121’ were grown from seeds provided by N. Kedar. Cuttings were placed in half-strength Hoagland’s solution (Hoagland and Arnon, 1950) and root primordia formed about a week later.

Root tips of freshly rooted cuttings were fixed by placing them in a saturated water solution of α-bromo-naphthalene for 165 min and then into 1:3 ethanol acetic acid for 12 to 24 hr. The tips were then placed for 5 min in 1 ethanol : 1 HCl (concentrated), followed by 10 min of soaking in Carnoy solution. Orcein at 1% in 1 glacial acetic-acid : 1 water (aceto-orcein) was used as a chromosome stain as modified from Darlington and La Cour (1962).

Isozyme study. Leaf disks (1 cm²) were macerated on ice in a mortar with a mixture of 0.1 M Tris-HCl (pH 7.5) buffer containing 1.5% reduced glutathione. This crude extract was used without further treatment. Samples were electrophoresed for 6 hr at 4C in 12.5% starch gels containing 0.005 M citric acid at pH 7.0.

The gels were stained for aconitase (EC 4.2.1.3) according to Tansley (1984) with the following modifications: NADP 10 mg, MTT (thiosulphate blue) 10 mg, PMS (phenasine methasulfate) 2 mg, isocitrate dehydrogenase 30 units, cis-aconit acid 50 mg, all per 50 ml of 0.1 M Tris-HCl buffer at pH 7.5. Gels stained for aconitase activity were scanned for estimation of relative stain density using an LKB 2202 Ultrascan laser densitometer (LKB, Bromma, Sweden). A units are arbitrary.

Restriction fragment length polymorphism analysis. Total DNA was extracted from leaves of the triploid ‘FC121’, the diploid ‘FC121’, and their male and female parents. DNA isolation, restriction digests, agarose gel electrophoresis, Southern blots, hybridizations, and autoradiography were as described by Ben-Natzky and Tansley (1986). Genomic DNA at 15 µg was digested with Eco RV and hybridized to the radiolabelled DNA clone TG101 (Young et al., 1988).
Results and Discussion

Triploid tomato plants should have 2n = 3x = 36 chromosomes. One triploid (Fig. 1) was verified among plants of the tomato hybrid line 'FC121'. This line, a hybrid of two distinctive parental cultivars, made a suitable subject for testing the prevailing hypothesis that a spontaneous triploid contains two maternal genome complements and one paternal genome (Chen and Palmer, 1985; Chyi and Weeden, 1984).

In some cases the number of gene copies of a particular isozyme can be established by examining the stain intensity of an electrophoresed sample (Tsaftaris et al., 1981). We tested (10) enzymes known to have more than one electrophoretic form. Only two differed between the two parents. In the esterase (EC 3.1.1.-) system the zymograms of both the 2n and the 3n samples were identical to the male parent, thus showing complete dominance and making esterase unsuitable for our purposes. In the aconitase system, only the monomeric ace-l was variable (Fig. 2). The female parent was homozygous for a fast-migrating allozyme and the male parent for a slower-migrating one. The diploid ‘FC121’ exhibited both allozymes at an intensity ratio of 3 male (slow-migrating) : 1 female (fast-migrating) (Table 1). This ratio probably reflects the differential expression of the two alleles in the heterozygote. The known phenomenon of allozyme imbalance in the heterozygote has been shown in maize to be due to different initial rates of translation (Freeling, 1983). The intensity ratio between male and female allozymes in the triploid was 6:1. If we assume that the relative expression of the alleles is the same as in the diploid hybrid, these results indicate that the triploid has two copies of the male allozyme and that it is the male parent that had contributed the double chromosome complement. Thus, our triploid plant may be the product of a haploid egg and a diploid pollen nucleus.

As the above result was somewhat surprising, a second, more conclusive approach was adopted. The male parent of the ‘FC121’ hybrid was homozygous for the Tm-2 gene that confers resistance to the tobacco mosaic virus (TMV), while the female parent was susceptible to TMV. Young et al. (1988) demonstrated that the DNA clone TG101 was tightly linked to Tm-2 and that restriction fragment length polymorphism exists be-

Table 1. Scan of an aconitase isozyme gel (using a Laser densitometer) from 3n and 2n tomato cv. FC121 hybrids and their parents.

<table>
<thead>
<tr>
<th>Migration from origin (mm)</th>
<th>Triploid (3n hybrid)</th>
<th>Diploid (2n hybrid)</th>
<th>Male parent</th>
<th>Female parent</th>
</tr>
</thead>
<tbody>
<tr>
<td>19</td>
<td>1466</td>
<td>1491</td>
<td>1460</td>
<td>1416</td>
</tr>
<tr>
<td>24(a)</td>
<td>425</td>
<td>289</td>
<td>397</td>
<td>0</td>
</tr>
<tr>
<td>26(b)</td>
<td>69</td>
<td>106</td>
<td>20</td>
<td>438</td>
</tr>
<tr>
<td>a/b</td>
<td>≈6/1</td>
<td>≈3/1</td>
<td>---</td>
<td>---</td>
</tr>
</tbody>
</table>

Fig. 1. A representative triploid tomato root tip cell containing 36 chromosomes (magnification: 800 × 4). (Left) Photograph, (right) scheme.

Fig. 2. Gel electrophoresis representation and photograph of aconitase isozymes of the triploid hybrid, the diploid hybrid, and parents of the 'FC121' cultivar. Migration in millimeters from origin on gel. (Experiment repeated four times, yielding similar results.) (Left) Photograph, (right) scheme.

between TMV-resistant and -susceptible lines. *TG101* was hybridized to a filter containing DNA of the two parents and the diploid and triploid hybrids digested with *Eco* RV. As expected (Young et al., 1988), the male parent proved to be homozygous to an *Eco* RV fragment of 8.5-kb size, while the female parent was homozygous to a 4.5 kb-size fragment. The diploid hybrid showed heterozygosity for the two allelic forms with similar intensities in the two bands. In the triploid hybrid, the hybridization signal of the 8.5-kb band (male parent) was stronger than that of the 4.5-kb band (female parent) (Fig. 3). When the lanes of the diploid and triploid hybrids were compared, it became evident that the higher molecular weight band was more intense in the triploid hybrid (Fig. 3B), indicating again that the double complement donor parent was the male.

Dempsey (1961) reached a similar conclusion regarding the identity of the double complement donor parent through a statistical analysis of trisomic offspring of a cross between a triploid hybrid and carefully chosen diploid plants. Preliminary results for two more triploid hybrids currently under investigation in our laboratory point in the same direction. This combination may prove to be a general phenomenon in spontaneously arising triploid tomato plants.

The question arises how a viable seed could have resulted from such a cross [if the pollen grain is a true diploid, one should expect the endosperm not to develop properly (Cooper and Brink 1945; Johnston et al., 1980)]. Of all conceivable and rare schemes, the most likely to have occurred is either that both generative nuclei fertilized the egg while the vegetative (usually disintegrating) nucleus merged with the pre-endosperm nucleus to form a normally triploidic endosperm, or that when the haploid generative nucleus was about to fertilize the egg it underwent mitosis before merging of the two.

**Literature Cited**


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