Intraspecific Genome Size Variation in Pumpkin (Cucurbita pepo subsp. pepo)

A. Lane Rayburn
Department of Crop Sciences, University of Illinois at Urbana-Champaign, 1201 West Gregory Drive, Urbana, IL 61801

Mosbah M. Kushad and Wanasari Wannarat
Department of Natural Resources and Environmental Sciences, University of Illinois at Urbana-Champaign, 1201 West Gregory Drive, Urbana, IL 61801

Additional index words: pie pumpkin, flow cytometry, genome size

Abstract. Genome size has recently been reported to vary 16% in pumpkins (Cucurbita spp.). The majority of this variation can be attributed to genome size differences in pumpkins of various taxonomical classes. The purpose of this study was to determine if intraspecific genome size variability could be detected by flow cytometry in Cucurbita pepo subsp. pepo pumpkin cultivars with similar fruit morphology. The pie pumpkins group was chosen for this study because of their similar fruit size, shape, and color. Genome sizes ranged from 1.109 pg in Spooktacular to 1.064 pg in Small Sugar. Spooktacular had a genome size larger than Small Sugar in all three experiments. Therefore, intraspecific genome size variation does exist in C. pepo subsp. pepo among pumpkin cultivars of similar fruit morphology.

Changes in cellular parameters that are brought about by variations in genome size are referred to as nucleotypic effects (Bennett, 1972; Rayburn, 1993a). Nucleotypic effects have been observed in various plant species (Bennett, 1976; Chung et al., 1998; Rayburn et al., 1985). A recent study by Tatum et al. (2006) suggested that nucleotypic selection was occurring in pumpkins. In particular, the hypothesis was that genome size (the amount of DNA within the nucleus of a cell) fluctuated and that this fluctuation influenced cell size in pumpkins and thus ultimately influenced fruit size. One caveat to nucleotypic studies is to ensure that true genome size variation exists within the species in question.

Over the past several years, concerns have been raised over the methods used to determine intraspecific genome size variation (Greilhuber, 2005). These concerns have led to doubts of the existence of intraspecific genome size variation and the amount of such variation. With the exception of maize (Zea mays; Rayburn et al., 1985; Rayburn et al., 1989), most reports of high genome size variation have not stood up under closer scrutiny (Greilhuber, 2005). For instance, soybean (Glycine max) has a reported genome size variation range between 40% and 0%, with two independent studies now agreeing on about a 4% variation (Chung et al., 1998; Rayburn et al., 2004). Several studies have documented a small genome size variation within plant species, with one report documenting about a 1% variation between different wheat cultivars (Wetzel et al., 1999). This small variation was supported by the cytological data of Wetzel and Rayburn (2000).

Pumpkin is a somewhat arbitrary term usually referring to “any squash used for pies, jack-o’-lanterns, or stock feed” (Robinson and Decker-Walters, 1997). Pumpkin encompasses various Cucurbita L. species such as Cucurbita pepo L., C. maxima Duchesne, C. moschata Duchesne, and C. argyrosperma C. Huber.

The pumpkins studied by Tatum et al. (2006) were of two species, C. pepo and C. maxima. The reported fruit weight of these pumpkins ranged from 0.11 to 273 kg (Anonymous, 2005). Tatum et al. (2006) observed up to a 16% variation in genome size in pumpkins, with the smallest genome sizes found in two pumpkin fruit types: miniature and jumbo. These two fruit types represent the extreme diversity found in pumpkin with regard to fruit size. The jumbo pumpkin cultivars are of the species C. maxima and produce fruits ranging from 22.7 to 273 kg (University of Illinois Extension, 2005a). Although it is interesting that the C. maxima cultivars have a smaller genome size than most of the C. pepo cultivars, this fact sheds no light on existence of intraspecific variation because any differences observed would represent interspecific variation. The miniatures, i.e., Jack-B-Little, that Tatum et al. (2006) also found to have small genome sizes have the smallest fruit type of the pumpkins. Plant and fruit size have been hypothesized to correlate with altered genome sizes in plant species (Rayburn et al., 1985; Tatum et al., 2006). Therefore, although the 13% to 16% variation observed by Tatum et al. (2006) may indeed be intraspecific variation, it could be from the extreme fruit size difference among these pumpkins. An additional problem with the inclusion of Jack-B-Little types in intraspecific genome size variation studies in pumpkins is the possibility that these miniature pumpkin types may actually represent another taxonomic class in C. pepo.

Goldman (2004) indicates that Jack-B-Little is not actually a pumpkin but is instead an acorn squash. If Jack-B-Little types are acorn squash, they would be of the subspecies “texana” not “pepo” (Paris et al., 2003). If one truly wants to focus on intraspecific variation, only plants of C. pepo subsp. pepo should be compared. When the miniature pumpkins were removed from the analysis of Tatum et al. (2006), the amount of intraspecific genome size variation, although reduced to ≤4%, was still statistically significant. However, even this small variation included pumpkins of vastly different fruit size. Gasmanova et al. (2007) also noted genome size variation in widely variable C. pepo accessions. The question becomes: Can genome size variation occur in pumpkins of similar morphology?

The purpose of this study was to document genome size variation in C. pepo subsp. pepo pumpkins of similar fruit morphology. The experiment was replicated over time, and statistical analyses were performed to ensure only true genome size variation was reported.

Materials and Methods

Pumpkins tested were of the subspecies C. pepo subsp. pepo with the exception of Jack-B-Quik, which is C. pepo subsp. ovifera (L.) D.S. Decker var texana (Scheele) Filov. The seeds were obtained from Rupp Seed, Inc. (Wauseon, OH). The first 10 cultivars listed in Table 1 of are of similar fruit morphology and are referred to as pie pumpkins based on their sweet, less watery flesh (University of Illinois Extension, 2005b). Little October and Jack-B-Quik, classified as miniature pumpkins (Rupp Seed, Inc. catalog), were selected for their high and low genome sizes, respectively (Tatum et al., 2006).

Plants were grown in the laboratory in Hummer Growing Mix No. 2 Lightweight Mix. One Sylvania Gro-Lux 40w light bulb and one Cool White Deluxe 40w light bulb were used for the light source with an irradiance level of 83 mmole-m⁻²·sec⁻¹ photosynthetically active radiation (400–800 nm). The lights were set on a 16-h light/8-h dark cycle.

The nuclei were isolated according to Rayburn et al. (1989). Briefly, a 2-cm segment of pumpkin leaf tissue from each plant and a rolled leaf portion above the growing point of the maize seedling were cochopped and placed in a beaker containing 10 mL of extraction buffer and 200 μL of 25% Triton X. The material was then homogenized with a tissue grinder for 10 s, and then the homogenate was filtered through a 250-μm and a 53-μm diameter nylon mesh filter. The samples were then stored at −20°C until used.
were centrifuged at 200 g for 15 min at 4 °C, the supernatant was aspirated, and the pellet was resuspended in 300 μL of propidium iodide stain (Bashir et al., 1993). Samples were then placed in a 37 °C water bath for 20 min. Afterward, 300 μL of propidium iodide salt (Bashir et al., 1993) was added to each sample and samples were vortexed for ~2 s. The samples were placed on ice, covered, and stored at 4 °C for at least 1 h.

Samples were analyzed using a Flow Cytometer-Cell Sorter Epics XL-MCL (Coulter Electronics, Hialeah, FL) set at a 488-nm laser wavelength. A histogram plotting the frequency of nuclei versus the fluorescence channel was produced. The relative fluorescence between the pumpkin G1 peak and the maize G1 peak was used to convert fluorescence intensity to picograms of DNA.

The experiment was repeated three times over a 7-month period. A total of five, nine, and four replicates per cultivar were analyzed in the first, second, and third experiments, respectively. Each replication consisted of a single plant, which was analyzed on a separate day. For each replicate, one small leaf of pumpkin and a small leaf segment of the maize line W22 were chopped together. Co-chopping with the maize line W22 provided an internal standard. The reason this maize line was selected is that it has consistently been reported to have 5.35 pg per 2C nucleus (Rayburn, 1993b). The maize line was selected because all of the pie pumpkins relatively small fruit size and is of a different subspecies. Because all of the pie pumpkins were C. pepo var. pepo and had fruit of similar morphology, drastic differences in fruit type did not complicate the present study.

Baby Pam, Touch of Autumn, and Trickster were observed to rank consistently in the top five genome sizes in all three experiments. Although statistical significance did vary among the three experiments, the rankings of these cultivars were reproducible.

The genome size of Little October was observed to have an amount of variation may be exacerbated due to the amount of variation observed by Tatum et al. (2006). Two studies have now confirmed that genome size variation exists in pumpkin cultivars. This genome size variation is found in one specific group of pumpkins, a complex relationship between genome size and fruit size seems to be indicated. Further studies are underway to confirm that the observed genome size variation is associated with nucleotypic characteristics such as fruit size.

Table 1. The 2C genome size of the pumpkin cultivars.

| Cultivar            | Average of three experiments | Expt. 1 | Expt. 2 | Expt. 3 |
|---------------------|------------------------------|---------|---------|---------|
|                     | N Mean                       |         |         |         |
| Spooktacular        | 1.109 pg                     | 5 1.096 | 9 1.095 | 4 1.135 |
| Peek A Boo          | 1.107 pg                     | 5 1.096 | 9 1.098 | 4 1.129 |
| Baby Pam            | 1.090 pg                     | 5 1.089 | 9 1.086 | 4 1.095 |
| Touch of Autumn     | 1.089 pg                     | 5 1.083 | 9 1.088 | 4 1.094 |
| Trickster           | 1.088 pg                     | 5 1.081 | 9 1.088 | 4 1.094 |
| Snackjack           | 1.074 pg                     | 5 1.067 | 9 1.086 | 4 1.062 |
| Hybrid Pam          | 1.073 pg                     | 5 1.064 | 9 1.086 | 4 1.069 |
| Orange Smoothie     | 1.070 pg                     | 5 1.070 | 9 1.068 | 4 1.073 |
| Pik-A-Pie           | 1.065 pg                     | 5 1.072 | 9 1.064 | 4 1.058 |
| Small Sugar         | 1.064 pg                     | 5 1.061 | 9 1.074 | 4 1.056 |
| Little October      | 1.064 pg                     | 5 1.053 | 9 1.052 | 4 1.086 |
| Jack-B-Quik         | 0.967 pg                     | 5 0.960 | 9 0.975 | 4 0.965 |

LSD (0.05) = 0.028

1. Pie pumpkins, C. pepo var. pepo.
2. Miniature pumpkin, C. pepo var. pepo.
3. Miniature pumpkin, C. pepo var. texana.

The similarity of genome sizes between Spooktacular and Little October was selected based on our previous observation that this cultivar had a medium to large genome size compared with other pumpkins (Tatum et al., 2006). The fact that none of the pie pumpkins in this study had a genome size smaller than Little October was unexpected. The genome sizes of the pie pumpkins were larger than expected and exceeded the DNA range of the controls.

With respect to the three experiments, the observed genome sizes were similar in eight of the 12 cultivars examined (Table 1). Three of the cultivars (Small Sugar, Hybrid Pam, and Snackjack) had aberrantly larger genome sizes in Exp. 2, whereas Little October had a larger genome size in Exp. 3 (Table 1). Final comparisons among the cultivars were made based on the average of the three experiments.

In all three experiments, Spooktacular and Peek-a-Boo were observed to have the largest genome sizes and had significantly larger genome sizes than Jack-B-Quik and Small Sugar (Table 1). Overall, when comparing the means, the variation in genome size between Spooktacular and Jack-B-Quik was approximately 15%. This genome size variation is similar to the amount of variation observed by Tatum et al. (2006). Two studies have now confirmed that genome size variation exists in pumpkins and that the variation is approximately 15%. However, this variation includes the miniature pumpkin cultivar Jack-B-Quik. As such, the amount of variation may be exacerbated by the inclusion of this cultivar that has relatively small fruit size and is of a different subspecies. Because all of the pie pumpkins were C. pepo subsp. pepo and had fruit of similar morphology, drastic differences in fruit type did not complicate the present study.

Results and Discussion

Little October was observed to have an average genome size of 1.064 pg, whereas Jack-B-Quik was observed to have 0.967 pg (Table 1). These results confirm the results from our previous study (Tatum et al., 2006) in which we reported the genome size of Little October to be 1.06 pg and the genome size of

| Cultivar            | Average of three experiments | Expt. 1 | Expt. 2 | Expt. 3 |
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| Touch of Autumn     | 1.089 pg                     | 5 1.083 | 9 1.088 | 4 1.094 |
| Trickster           | 1.088 pg                     | 5 1.081 | 9 1.088 | 4 1.094 |
| Snackjack           | 1.074 pg                     | 5 1.067 | 9 1.086 | 4 1.062 |
| Hybrid Pam          | 1.073 pg                     | 5 1.064 | 9 1.086 | 4 1.069 |
| Orange Smoothie     | 1.070 pg                     | 5 1.070 | 9 1.068 | 4 1.073 |
| Pik-A-Pie           | 1.065 pg                     | 5 1.072 | 9 1.064 | 4 1.058 |
| Small Sugar         | 1.064 pg                     | 5 1.061 | 9 1.074 | 4 1.056 |
| Little October      | 1.064 pg                     | 5 1.053 | 9 1.052 | 4 1.086 |
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1. Pie pumpkins, C. pepo var. pepo.
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3. Miniature pumpkin, C. pepo var. texana.

When comparing the mean genome sizes of Peek-A-Boo and Small Sugar (the pie pumpkin cultivar with the smallest genome size), overall, a 4% variation was observed. Although smaller than the 15% variation observed between Jack-B-Quik and Spooktacular, the genome sizes of Peek-A-Boo and Small Sugar were consistently statistically significant at α = 0.05. Given the reproducibility of the results, this variation in genome size appears real. The amount of variation observed here in pumpkins is similar to amount of variability seen in soybean and much less than that observed in maize (Rayburn et al., 2004).

Intraspecific genome size variation has now been confirmed in C. pepo subsp. pepo pumpkin cultivars. This genome size variation is found in one specific group of pumpkins, indicating that genome size variation can occur among pumpkin cultivars of similar morphology. Upon comparing genome size with reported fruit size in these pie pumpkins, a complex relationship between genome size and fruit size seems to be indicated. Further studies are underway to confirm that the observed genome size variation is associated with nucleotypic characteristics such as fruit size.

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