Tissue proliferation (TP) is a disorder in which abnormal callus-like growths, or tumors, are produced at or near the crown or base of a woody plant (LaMondia et al., 1992). A proliferation of short shoots or buds may form on tumors of certain cultivars (Brand and Kiyomoto, 1992). In addition to tumors and tumor-associated shoot proliferation, Brand and Kiyomoto (1997) observed specific shoot morphological differences between Rhododendron ‘Montego’ plants with and without TP tumors. Genera in which TP has been observed include Rhododendron, Kalmia, and Pieris (Linderman, 1993), but TP has been most prevalent in Rhododendron. Within the genus Rhododendron, genotype has a significant effect on TP incidence and severity (McCulloch and Britt, 1997). At least 40 Rhododendron cultivars with TP have been observed (Zimmerman, 1997).

Although TP appears to be similar to crown gall, previous experimental evidence suggests that TP does not result from Agrobacterium tumefaciens infection (Zimmerman, 1997). The morphology of TP also appears to be similar to that of lignotubers, which are woody outgrowths at the stem base that contain numerous dormant buds (Del Tredici, 1992). However, TP tumors do not generally exhibit nor-mal lignotuber characteristics (Brand and Kiyomoto, 1994; Del Tredici, 1992; Mercure et al., 1998). The most widely accepted theory is that TP results from epigenetic changes that affect the incidence and severity of TP symptoms (Brand and Kiyomoto, 1997). Linderman (1993). Specifically, Brand and Kiyomoto (1997) found that TP-like symptoms in Rhododendron ‘Montego’ were associated with adventitious shoot formation in vitro.

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Almost all instances of TP can be traced back to propagation by tissue culture (Linderman, 1993; Zimmerman, 1997). From nursery surveys, Mudge et al. (1997) determined that TP symptoms developed only on plants that had been propagated by stem cuttings collected from previously micropropagated plants. Growers generally use their existing container stock as the source of cuttings for next year’s Rhododendron crop. Of particular importance to this practice is whether normal-appearing cuttings taken from the tops of plants with TP would be large if TP were transmitted to a new crop by cutting propagation. The objectives of this study were to determine 1) if TP is trans-mitted to plants through cutting propagation, and 2) if stock plant age from micropropagation affects the incidence and severity of TP symptoms in plants propagated from cuttings.

**Materials and Methods**

Propagation and plant maintenance. Terminal cuttings were collected and stuck to root in late November. Rooting was performed when present and cuttings were trimmed at the base to a maximum stem length of 10 cm. Leaves were trimmed to remove the distal one-third of the blade. All cuttings were wounded (1 cm basal wound on two sides of the stem) and were dipped in Hormex, rooting powder No. 16 (1.6% indole-3-butyric acid) (Brooker Chemical, North Hollywood, Calif.). Cuttings were stuck in 35 × 50 × 9-cm black plastic flats filled with 1 sphagnum peat : 1 coarse perlite (v/v). Each flat contained 30 cuttings spaced evenly in six rows of five. Sixty cuttings were stuck for each cultivar × TP status combination. Cuttings were rooted using bottom heat and intermittent mist (electronic leaf-controlled) in the greenhouse for 120 d at 21 °C day/17 °C night. Following rooting, cuttings were acclimated for 45 d in a greenhouse with 30% shade provided by black polypropylene cloth. Microcuttings were also harvested in late November and were rooted in 1 sphagnum peat : 1 fine perlite (v/v) in plastic, clear-lidded humidity chambers. Microcuttings received no wounding or auxin treatment.

Rooted cuttings were potted in 3 pine bark : 2 sphagnum peat : 1 sand (by volume) amended with ground dolomitic limestone and gypsum, both at 3.6 kg·m⁻³. During the first growing season, cuttings were grown in 2.6-L containers (Classic 300S; Nursery Supplies, Fairless Hills, Pa.); in mid-May of the second season they were repotted into 6.1-L containers (Classic 600; Nursery Supplies). All plants were topdressed with Sierrablen 17N–2.6P–8.3K 8- to 9-month formulation (The Scotts Co., Marietta, Ga.) using 10 g per pot following transplanting the first season, and 50 g per pot the second year. During the period from 15 May to 31 Oct. (growing season), 2.6-L containers received ≈1.5 L of water every other day and 6.1-L containers received 4 L of water every other day. Irrigation was provided through trickle emitters using an automated
controller. During the growing season, plants were placed on a gravel-covered growing area in full sunlight. Between growing seasons (1 Nov. through 14 May), plants were overwintered in an unheated white polyethylene-covered hoop house and hand watered as needed. At the conclusion of the overwintering period, plants were pruned according to standard nursery practices by removing much of the last growth flush and further shaping the plants as needed. Plants received no pesticide or herbicide treatments to avoid any effect these products might have on the induction of TP symptoms. Hand weeding was done as needed.

**Plant growth, shoot, and foliar measurements.** Plant size was calculated as the product of two plant widths, taken at right angles to each other, and plant height. A vigorous shoot on each selected plant was chosen for measurement, avoiding shoots with a late-season growth flush. Basal shoot length was measured from the bud scale scars to the first leaf, and total shoot length from the bud scale scars to the base of the terminal bud. The number of leaves per shoot (growth flush) was also counted. Leaf measurements were taken on each selected shoot. Leaf length was measured on 30 mL of medium in 200-mL glass jars sealed with B-caps (Magenta Corp., Chicago). Cultures were incubated under 40 μmol·m–2·s–1 cool-white fluorescent light for 16 h/day at 24 °C ± 2 °C with 6-week subcultures. Prior to use in experiments, cultures were subcultured a minimum of six times.

Once rooted, all four cutting types were grown in containers for 2 years and then examined for TP tumors. Tumors were categorized as: 1) absent; 2) small and individual; 3) encompassing one-quarter of the stem; 4) encompassing one-half of the stem; or 5) surrounding the stem. Morphological and growth data were collected as well.

**Experimental design and statistical analysis.** All experiments were arranged in completely random designs. Analysis of variance was performed on plant growth, shoot, and foliar measurements using the general linear model (GLM) procedure (SAS Institute, 1990). For these variables, within cultivar mean separations between TP(–) and TP(+) plants were performed using multiple Scheffe–Marascuilo comparisons among binomial samples (Marascuilo, 1966).

**Results and Discussion**

**TP symptom development in plants grown from TP(+) and TP(–) cuttings.** The main effects of cultivar and TP status were highly significant at the P ≤ 0.01 level for all foliar and plant growth characteristics measured. Cultivar × TP status interactions were significant (P ≤ 0.05) for all measured characteristics except the number of leaves per shoot (Table 1).

At least one growth or morphological difference between plants from TP(–) and TP(+) cuttings was evident in all cultivars. The nature and number of characteristics affected and the degree of difference between plants from TP(–) and TP(+) cuttings were highly influenced by genotype (Table 1). For example, plants of ‘Besse Howells’ differed only in the number of leaves per shoot, whereas those of ‘Montego’ differed in all characteristics measured. In addition, for any given growth or morphological feature, differences were always of the same nature for each cultivar; i.e., if narrower leaves occurred in plants from TP(+) cuttings of one cultivar, all other cultivars in which this feature was affected had narrower, not wider, leaves. Leaf size and shape differed between plants from TP(–) and TP(+) cuttings in three of the seven cultivars in the study (Table 1). Leaf length and leaf width of plants grown from TP(+) cuttings were either the same as, or less than, those of plants grown from TP(–) cuttings. ‘Montego’ exhibited the most pronounced effect, with differences in length and width of 1.6 and 1.3 cm, respectively. As a result of these differences, plants from TP(+) cuttings of ‘Holden’, ‘Montego’, and ‘Scintillation’ had greater leaf length:width ratios than did plants from TP(–) cuttings. For ‘Montego’, TP(+) leaves had a length:width ratio of 2.8, while TP(+) leaves had a ratio of 3.6, giving them a very narrow, strap-like appearance. Differences in the size and shape of ‘Holden’ and ‘Scintillation’ leaves from plants of TP(+) and TP(–) cuttings were not easily quantified by casual observation. Areas of individual leaves of ‘Montego’ and ‘Scintillation’ plants from TP(+) cuttings were only 58% and 78% as large, respectively, as those leaves produced from TP(–) cuttings.

Shoot growth was affected in three of seven cultivars (Table 1). For ‘Boule de Neige’, ‘Catawbiense Album’, and ‘Montego’, both basal and total shoot length were shorter for plants from TP(+) cuttings, averaging 62% and 76%, respectively, of the basal and total shoot length on TP(–) cuttings. All cultivars differed in the number of leaves per shoot—plants from TP(+) cuttings producing an average of about one (1.1) more leaf per shoot than plants from TP(–) cuttings.

Reduced canopy size was evident in plants from TP(+) cuttings of four of the seven cultivars (Table 1). This reduction ranged from 4% for ‘Scintillation’ to 43% for ‘Montego’. Tumors were not observed on any plants in this study, regardless of cultivar or TP status of the stock plant. These data indicate that plants grown from cuttings collected from TP(+) plants may exhibit shorter, narrower, and smaller leaves than are typical for the cultivar. In addition, they may have shorter shoots, more leaves per shoot, and reduced canopy sizes relative to their TP(–) counterparts. In an earlier study comparing TP(+) and TP(–) plant morphology for *Rhododendron* ‘Montego’, Brand and Kiyomoto (1997) found that TP(+) plants with tumors had shorter, narrower, smaller leaves and shorter shoots than TP(–) plants. However, Brand and Kiyomoto (1997) did not observe a difference in number of leaves per shoot between TP(+) and TP(–) plants, as was found for all cultivars in this study, including ‘Montego’.

Whether plants grown from TP(+) cuttings...
Table 1. Comparison of shoot and leaf characteristics of seven Rhododendron cultivars, two growing seasons after cutting propagation. Stem cuttings were taken from plants with tissue proliferation tumors [TP(+)] or without [TP(−)] tumors or history of micropropagation.

| Cultivar | TP status | Length (cm) | Width (cm) | Area (cm²) | Length/width | Total | Basal | Shoot length (cm) | No. leaves | Plant size (dm³) |
|----------|-----------|-------------|------------|------------|--------------|-------|-------|------------------|-----------|-----------------|
| BH       | TP(+)     | 8.5 a       | 4.0        | 25.9 a     | 2.1 a        | 9.3 a | 6.5 a | 81.1 b           | 78.0 a    |
|          | mean      | 8.7         | 4.2 a      | 28.0 a     | 2.1 a        | 9.9 a | 6.3 a | 88.8 a           | 78.3 a    |
| BN       | TP(−)     | 9.5 a       | 3.9 a      | 27.8 a     | 2.5 a        | 12.6 a| 9.4 a | 73.0 b           | 47.5 a    |
|          | mean      | 9.5         | 4.0        | 28.6 a     | 2.5          | 11.1  | 7.8  | 78.0             | 42.4      |
| CS       | TP(−)     | 9.8 a       | 3.2        | 23.0 a     | 3.1 a        | 11.7 a| 2.7 a | 12.3 b           | 81.2 a    |
|          | mean      | 9.8         | 3.2        | 22.2 a     | 3.1          | 12.0  | 2.6  | 13.1             | 74.3      |
| CA       | TP(−)     | 8.8 a       | 3.8 a      | 25.2 a     | 2.3 a        | 11.8 a| 7.2 a | 9.3 b            | 58.1 a    |
|          | mean      | 8.7         | 3.7        | 25.2       | 2.4          | 10.4  | 6.3  | 10.1             | 57.9      |
| HO       | TP(−)     | 8.7 a       | 3.4 a      | 23.5 a     | 2.6 b        | 11.1 a| 6.7 a | 10.3 b           | 58.1 a    |
|          | mean      | 8.6         | 3.1 b      | 21.1 a     | 2.8 a        | 9.6 a | 5.7 a | 11.5 a           | 57.6 a    |
| MO       | TP(−)     | 10.6 a      | 3.8 a      | 28.4 a     | 2.8 b        | 12.1 a| 4.7 a | 14.2 b           | 79.4 a    |
|          | mean      | 9.8         | 3.2        | 22.4       | 3.2          | 10.7  | 3.5  | 14.7             | 62.2      |
| SC       | TP(−)     | 10.3 a      | 4.6 a      | 39.1 a     | 2.2 b        | 14.1 a| 9.2 a | 10.2 b           | 44.1 a    |
|          | mean      | 9.9         | 4.3 a      | 34.8       | 2.7          | 13.7  | 9.0  | 10.4             | 43.2      |
|          | mean:     | 9.5         | 3.8        | 27.6       | 2.5          | 11.8  | 6.6  | 10.2             | 55.5      |
|          | mean:     | 9.1         | 3.5        | 24.6       | 2.7          | 10.4  | 5.3  | 11.3             | 46.8      |

Significance of main effects and interaction

| Cultivar | TP status | Microcuttings | 3-year-old TP | 6-year-old TP | TP(−) |
|----------|-----------|---------------|---------------|---------------|-------|
|          | **       | **            | **            | **            | **    |
|          | *        | NS            | NS            | NS            | NS    |

Table 2. Incidence and severity of tissue proliferation (TP) tumors on Rhododendron ‘Montego’ plants grown from stem cuttings collected from plants of varying ages postmicropropagation.

| Degree of tumorization | Microcuttings | 3-year-old TP | 6-year-old TP | TP(−) |
|------------------------|---------------|---------------|---------------|-------|
| No tumor               | 20 (17)b      | 10 (26)b      | 42 (100)a     | 59 (100)a |
| Small individual tumors| 13 (11)a     | 5 (13)b       | 0 (0)b        | 0 (0)b |
| One-quarter of stem surrounded | 23 (20)a | 9 (23)a | 0 (0)b | 0 (0)b |
| One-half of stem surrounded | 26 (22)a | 11(28)a | 0 (0)b | 0 (0)b |
| Stem surrounded         | 34 (29)a     | 4 (10)b       | 0 (0)b        | 0 (0)b |
| Tumor of any size       | 97 (83)a     | 29 (74)a      | 0 (0)b        | 0 (0)b |

*Harvested from in vitro cultures known to produce plants exhibiting TP tumors.

*Cultures were harvested from plants with TP tumors that were 3 and 6 years old, postmicropropagation.

*Mean separation of percent values within each degree of tumorization using multiple Scheffe–Marascuilo comparisons among binomial samples, P ≤ 0.05.

Develop overt and obvious TP symptoms appears to be highly cultivar-dependent; growers who understand this will be better able to manage or control TP in their production. Clearly, to avoid producing noticeably off-type plants with cultivars such as ‘Montego’, cuttings should not be collected from TP(+) stock plants. The absence of tumor development in plants propagated from such plants is encouraging, since tumors are potentially the most problematic TP symptom from both an ornamental and physiological viewpoint. Unfortunately, the absence of tumor development in this study does not prove that plants grown from TP(+) stock plants will not form tumors under certain conditions. Cultural triggers may exist that stimulate tumor formation (Linderman, 1993; Maynard, 1995), and our growing conditions may not have been conducive to tumor development. However, the lack of tumor formation in plants from TP(+) cuttings is not surprising, given that cuttings were collected from the distal parts of the plant. The spatial separation of the apical cuttings from the basal region, where TP symptoms are concentrated, could minimize the influence the basal region has on the apical shoots.

Influence of TP(+) plant age from micropropagation on TP symptom development in rooted cuttings. All plants from microcuttings, 3-year-old TP(+) cuttings, and 6-year-old TP(+) cuttings exhibited small, narrow leaves, shortened shoots, reduced plant canopy size, and an increased number of leaves per shoot when compared with TP(−) plants. Age from micropropagation had no significant effects on plant growth or shoot morphology. However, plants grown from cuttings were more likely to develop tumors when taken from TP(+) plants that were recently micropropagated (Table 2). When plants were grown from microcuttings, 83% of them developed tumors, whereas 74% of those grown from cuttings of 3-year-old TP(+) plants developed tumors. Additionally, plants from microcuttings exhibited larger tumors than other groups in the study, 51% developed tumors that surrounded one-half or more of the stem vs. 38% of plants grown from 3-year-old TP(+) cuttings and 0% of plants grown from TP(−) or 6-year-old TP(+) cuttings. The incidence and severity of TP tumor development in Rhododendron ‘Montego’ decreased with time after removal from tissue culture, but TP-related shoot growth and morphological differences remained unchanged during the first 6 years. Tumor-forming capacity appeared to be lost in the apical portions of plants sometime between 3 and 6 years after tissue culture. The loss of tumor-forming ca-

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Capacity and concurrent retention of altered foliar morphology in *Rhododendron* ‘Montego’ in response to increasing time may be analogous to phase change in woody plants such as *Hedera helix* L. Although tumor-forming capacity in *Rhododendron* ‘Montego’ is rapidly lost with time, alterations in shoot morphology may revert to normal appearance more slowly. Both phase change and TP are believed to be epigenetic in nature, providing additional support for a comparison between the two phenomena (Brand and Kiyomoto, 1997; Hartmann et al., 1997).

Although our data with *Rhododendron* ‘Montego’ indicate that cuttings collected from sufficiently old micropropagated stock plants will not produce plants with tumors, alterations in foliar morphology will persist in propagated plants. Even though ‘Montego’ is at the extreme end of the spectrum with regard to TP symptoms, growers should use caution in collecting cuttings from even relatively old micropropagated stock plants, of any cultivar, if the plants have exhibited TP symptoms.

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