Regeneration of *Brassica oleracea* from Peduncle Explants

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**Abstract.** Peduncle explants from 12 *Brassica oleracea* L. lines representing five varieties [broccoli (*italica*), cabbage (*capitata*), cauliflower (*botrytis*), Chinese broccoli (*alboglabra*), and rapid-cycling *B. oleracea*] readily regenerated shoots in vitro. Average regeneration rates of more than 75% were obtained for most lines, with up to 35 shoots per explant. Shoots were visible within 7 to 10 days. Initial regeneration was polarized, occurring mainly from the basal end of explants. Linsmaier-Skoog-based medium containing 1 mg BA/liter was suitable for shoot regeneration from all 12 lines tested. Plants were rooted on hormone-free medium and transferred to soil. Chemical name used: benzyladenine (BA).

*Brassica oleracea* is an economically important vegetable species, containing 14 vegetable varieties, including broccoli, cabbage, cauliflower, Chinese broccoli. Practically every part of the plant can be used, including the leaves (cabbage), axillary buds (brussels sprouts), stems (kohlrabi), floral primordia (cauliflower), and flower buds (broccoli). These vegetables are amenable to in vitro manipulation; shoot regeneration has been demonstrated for almost all varieties using explants from various vegetative organs, and also from floral organs (reviewed in Christey, 1989, and in Zee and Johnson, 1984).

The use of peduncle explants for the regeneration of *Brassica* was first reported in *B. napus* by Stringam (1977). Regeneration frequencies of more than 80% were obtained with multiple buds per explant. Peduncle explants have since been used for regeneration and transformation of *B. napus* (Fry et al., 1987), but not for *B. oleracea*. Although previous studies have demonstrated shoot regeneration from many varieties of *B. oleracea*, rates of shoot regeneration and number of shoots per explant were generally low. We aimed to determine whether peduncle explants could be used to obtain high levels of shoot regeneration with *B. oleracea* and, if so, whether the procedure was suitable for several *B. oleracea* varieties. Shoot regeneration from peduncle explants of 12 *B. oleracea* lines representing five botanical varieties is reported here. The use of this regeneration system for transformation and in vitro propagation of desirable genotypes is discussed.

**Plant material.** The plant materials used in this study are described in Table 1. The four vegetables (Chinese broccoli, cauliflower, cabbage, and broccoli) were chosen to represent a range of *B. oleracea* varieties grown in the northeastern United States. Lines were also included to permit comparison of cytoplasm effects and to test regeneration from rapid-cycling (RC) *B. oleracea* (Williams, 1985).

Peduncle explants from broccoli and Chinese

Received for publication 12 July 1990. Support for this work was provided by a National Research Advisory Council fellowship from the New Zealand government to M.C.C. and from U.S. Dept. of Agriculture grant 85-CRCR-1-1608 to E.D.E. The cost of publishing this paper was defrayed in part by the payment of page charges. Under postal regulations, this paper therefore must be hereby marked advertisement solely to indicate this fact.

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broccoli were obtained from plants that were bolting. The peduncles chosen had closed flower buds and few, if any, open buds. Only the top 5 to 7 cm was used to avoid the harder, woody lower tissue. Flower buds and pedicels were discarded.

Peduncles were surface sterilized by immersion in 70% ethanol for 3 to 5 min and then 0.5% sodium hypochlorite for 10 to 15 min, followed by three 5-min rinses with sterile water. Tissue damaged by the bleach was removed before cutting each peduncle into five to ten 5-mm explants. Explants were placed horizontally onto Linsmaier-Skoog (LS) medium (Linsmaier and Skoog, 1965) containing 3% sucrose and 1 mg BA/liter and solidified with 0.25% Gelrite (Scott Laboratories, Carson, Calif.). The order and orientation (basal vs. apical) of each explant was noted. About 20 to 25 explants were cultured per 9-cm petri plate.

After 3 to 4 weeks, the entire regenerating ends of some explants were excised and transferred to hormone-free LS medium (LS-N). Individual shoots (5-10 mm) were excised 10 to 14 days later and placed on LS-N for rooting. After transfer to soil, high humidity was maintained either by placing plants in a mist unit or by covering them with a plastic bag. After 1 to 2 weeks, the plants were either removed from the mist unit or holes were cut in the bag before its removal 1 week later.

RC Brassica oleracea cultures were maintained at 25C under a 16-h photoperiod provided by cool-white fluorescent lights, 80 µmol·m⁻²·s⁻¹. All other cultures were maintained at 25C under a 16-h photoperiod provided by an equal number of cool-white and Gro and Sho (General Electric, Schenectady, N.Y.) fluorescent lights, 45-70 µmol·m⁻²·s⁻¹. Plants in soil were grown at 25C under a 16-h photoperiod with lighting from cool-white fluorescent lights, 35 µmol·m⁻²·s⁻¹. After 3 to 4 weeks, some plants were transferred to the greenhouse as described above.

Shoot regeneration was readily obtained from peduncle explants of all lines tested (Table 2). Average regeneration frequencies were high, ranging from 41% to 98% but usually >75%. Buds per explant were also high, ranging from one to 35 and generally

<table>
<thead>
<tr>
<th>Variety</th>
<th>Common name</th>
<th>Cultivar or line</th>
<th>Cytoplasm*</th>
<th>Description</th>
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<td>F</td>
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<td>Jourdan et al., 1989</td>
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<td></td>
<td>Broccoli</td>
<td>Green Comet (GC)</td>
<td>F</td>
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<td>CMS, oogre</td>
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<td>Rapid cycling</td>
<td>Broccoli</td>
<td>CN16#3</td>
<td>F</td>
<td>Rapid cycling</td>
<td>Williams, 1985</td>
</tr>
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</table>

*ATR = atrazine-resistant chloroplasts; F = male fertile; CMS = cytoplasmic male sterile.
**Developed by M. Dickson, New York State Agricultural Experiment Station, Geneva.
***BN#4 and BN#5 materials included two original plants as well as plants regenerated from leaf protoplasts from them. The two sources of plants were not distinguished for this work.

**Table 1. Brassica oleracea lines used as sources of peduncle explants.**

Fig. 1. Shoot regeneration from peduncle explants. (a) Broccoli peduncle explant after 8 days in culture. Note the swollen basal end (B) with shoot buds (S) (x5). (b) Shoot buds on a Chinese broccoli peduncle explant after 10 days in culture (x 20). (c) Cauliflower peduncle explants after 14 days in culture (x 0.5).

Shoot regeneration was readily obtained from peduncle explants of all lines tested (Table 2). Average regeneration frequencies were high, ranging from 41% to 98% but usually >75%. Buds per explant were also high, ranging from one to 35 and generally
more than 10 per explant. Shoots were usually visible after 7 to 10 days. The same medium could be used for all lines tested.

Effect of culture conditions. In preliminary experiments, 0.8% Bacto Agar (Difco, Detroit, Mich.) was used to solidify the medium, but browning of the explants in the region in contact with the medium was noted. Such browning did not occur with Gelrite, which was routinely used in further work.

Initially, explants were cultured vertically with the basal end in contact with the medium, as done by Stringam (1977) and Fry et al. (1987). The shoots that formed often appeared abnormal, so explants were subsequently cultured horizontally. Use of horizontal explants gave more normal shoots and also increased the number of shoots per explant because shoots often developed from both ends.

General response to culture. Peduncle explants from all 12 lines showed a similar response to culture. The first change noted was a swelling of the entire explant, followed by further swelling of the basal end (Fig. 1a). After 7 days, both ends were covered by a small amount of white or pale-green callus. The surface area of the basal end was about twice that of the apical end. After 28 days, the surface area of the basal end (6-8 mm in diameter) was about five times that of the apical end (2-4 mm). Fresh weight of the explants increased dramatically during the experiment, from 40 to 50 mg to 400 to 600 mg after 28 days.

The first signs of regeneration were slightly raised areas on the lower edge of the end of the explants. These areas preceded the appearance of buds which then pushed through from under the surface callus. Shoot regeneration was first noted 7 to 10 days after culture of explants (Fig. 1a and b). After 2 weeks, numerous vegetative buds were present. Shoot formation was initially polarized: buds were concentrated on the basal end of explants in the region closest to the medium (Fig. 1a), except on ‘Guy Len’ explants, which showed initial regeneration from the apical end. Often more than 80% of the explants regenerated shoots (Table 2, Fig. 1c). Root formation was only rarely noted. By the end of the experiment (day 28), there was no further increase in the number of regenerating explants, but the number of buds per explant had increased and shoots had elongated. Shoots often developed from both ends, although bud number was still usually higher on the basal end. While polarity in the shoot regeneration pattern per explant was noted, there was little effect of original explant position within a peduncle.

Shoot buds were usually pale green with slight reddening of the tips for some lines. Buds from the cauliflower lines were deep red for the first 2 to 3 weeks of culture but turned green as the shoot tips elongated.

Transfer to soil. After 4 weeks, when shoots had elongated, at least 10 shoots from most lines were excised and transferred to LS-N medium. Within 7 days, several short roots had developed from the cut end of more than half the shoots. After 3 to 4 weeks, more than 80% of the cuttings showed sufficient root and shoot growth for transfer out of culture (Fig. 2a). More than 95% of these were successfully transferred to soil, but most were not kept for further growth. Plants from BN#4 and BN#5 peduncle explants were grown to maturity and showed no obvious morphological changes.

From the RC line, 166 plants were transferred to soil, with 98% surviving to maturity (Fig. 2b). These plants retained the RC phenotype, with flower buds visible 14 days after transfer to soil. Plants were fertile and set seed. All appeared morphologically normal, except for two plants (1.2%) that had cream-colored flowers instead of yellow ones. Sectoring of color within individual petals and flowers was apparent on these two plants.

Effect of cultivar on regeneration. The effect of cultivar was studied for broccoli where shoot regeneration percentages for ‘Packman’ were lower than those for ‘Green Comet’ (Table 2). In one experiment 7 with ‘Packman’ peduncle explants, raising the BA level from 1 to 2 or 5 mg liter$^{-1}$ increased the percentage of explants that formed shoots from 68%
to 83% or 94%, respectively. Although the number of explants per treatment was small (16-23), it appears that an increase in BA can increase regeneration to levels comparable to those in other lines. In *B. napus*, raising the BA level to 10 mg·liter⁻¹ also increased the percentage of peduncle explants regenerating shoots (data not shown).

**Effect of cytoplasm on regeneration.** The effect of cytoplasm on regeneration was tested by comparing regeneration from two fertile and three cytoplasmic male-sterile (CMS) broccoli lines, all with 'Green Comet' nuclear backgrounds. These lines contained either the *B. oleracea* cytoplasm ('Green Comet' and 'Green Comet' regenerant), the *ogura* cytoplasm from *Raphanus sativus* (84-3185), or the *B. nigra* cytoplasm (BN#4 and BN#5); the latter two cytoplasts confer male sterility. The fertile lines had higher percentages of regeneration than either type of CMS line (Table 2). Material with the *B. nigra* cytoplasm showed more shoot regeneration than the *ogura* cytoplasm, and experiments were more reproducible. Raising the BA level did not increase the percentage of CMS explants with shoot regeneration.

Our results indicate that peduncle explants from the five tested *B. oleracea* varieties readily regenerate shoots. Regeneration was rapid with multiple shoots per explant. Average regeneration percentages were high; only two lines, 'Packman' and 84-3185, had average regeneration percentages below 75%. The medium used for regeneration from peduncle explants of *B. napus* (Fry et al., 1987; Stringam, 1977) could also be used for all *B. oleracea* lines tested. Optimum regeneration from leaf discs of *B. campestris*, *B. oleracea*, and *B. napus* required a different hormone combination for each species (Dunwell, 1981).

The regeneration percentages and shoot production per explant and growth patterns obtained here for *B. oleracea* lines are similar to those reported for *B. napus* by Stringam (1977) and Fry et al. (1987). In contrast, only rare shoot regeneration was obtained from peduncle explants from three RC *B. campestris* lines cultured on this medium and more than 50 other combinations of growth regulators (mainly BA and NAA) (Christie, 1989). These observations of differential regeneration from peduncle explants of *B. oleracea*, *B. campestris*, and their amphidiploid derivative *B. napus* confirm previous studies in which regeneration of *B. oleracea* and *B. napus* was also higher than that of *B. campestris* (Dietert et al., 1982; Dunwell, 1981; Narasimhulu et al., 1988).

The culture of intact peduncle explants in this study with *B. oleracea* and previously with *B. napus* (Fry et al., 1987; Stringam, 1977; M.C.C., unpublished observations) produced only vegetative shoots. The plants regenerated required the normal growth period to induce flowering even though the explants were obtained from the flowering zone. In these experiments, regeneration was first seen on one end of the explants, usually the basal end. Such polarity was also noted with *B. napus* var. *oleifera* cv. Oturu peduncle explants cultured in a similar manner (data not shown). There was no corresponding polarity of root regeneration from the opposite end. In contrast, Lazzeri and Dunwell (1986) observed strong polarity of shoot and root regeneration from root and hypocotyl explants of 'Green Comet' broccoli.

A disadvantage of using peduncle explants for shoot regeneration is the time required to obtain them. The use of the RC *B. oleracea* line, from which peduncle explants were available 6 weeks after planting seed, overcame this disadvantage, allowing evaluation of regenerants for morphology and fertility 10 to 12 weeks after culture of explants. Peduncle explants from RC *B. oleracea* could thus be used as a model system for the production and study of transgenic plants.

There are several advantages to using peduncle explants for in vitro culture and regeneration. The culture procedure is simple, as one medium was suitable for all lines tested. Numerous explants can be obtained from a plant nondestructively, allowing continued growth of the plant for collection of additional explants or for further evaluation and seed set. In *B. oleracea*, the only other explants showing comparable rates of regeneration are derived from pedicels (Anderson and Carstens, 1977). The use of both pedicel and peduncle explants could enable a further increase in the shoots recovered from a single peduncle.

Peduncle explants could be useful for the in vitro propagation and rapid multiplication of important or novel genotypes. To be successful such a system has the following requirements: 1) multiple buds per explant and high regeneration, 2) short time from culture to potted plant, 3) reproducibility, and 4) high survival after transplanting from culture to soil.

These features have been met in the peduncle regeneration system reported here for *B. oleracea*. From a single plant, it is possible to obtain >150 explants; assuming a regeneration percentage of 75% with 10 buds per explant, >1125 shoots could be regenerated from one plant. A further increase in the propagation rate per plant could be obtained by including pedicel explants. The regeneration time is short-only 10 to 12 weeks from initial explant to potted plants. Survival on transfer to soil is high (>95%) provided care is taken to ensure a gradual decrease in humidity. In addition, little callus is formed before shoot formation, so it is likely that cytological aberrations and somaclonal variation will be slight.

Peduncle explants could also be used for transformation of *B. oleracea* via cocultivation of explants with *Agrobacterium tumefaciens* or by use of particle acceleration methods. The high percentages of regeneration, with quick production of numerous shoots, mean that transgenic plants might be rapidly obtained, as has been done in *B. napus* (Fry et al., 1987).

**Literature Cited**


Christie, 1989. Cell and tissue culture studies of *Brassica oleracea* and *B. campestris*. PhD Diss., Cornell Univ., Ithaca, N.Y.


