Promotive Effects of DCPTA on Seedling Development and Growth of Radish

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Abstract. A radish (Raphanus sativus L. cv. Scarlet turnip white tipped) seedling growth test was developed to examine promotive effects of 2-(3,4-dichlorophenoxy) triethylamine (DCPTA) on seedling vigor and plant development. Compared with controls, seed treatment using 30 µM DCPTA significantly (P = 0.05) enhanced the rates of root and hypocotyl elongation and seedling dry weight. Enhanced hypocotyl development by DCPTA showed a significant linear correlation (r = 0.83) with the increased taproot yield of mature plants grown from DCPTA-treated seeds. The harvestable taproot yield and harvest index of plants grown from seeds treated with 30 µM DCPTA were increased 109% and 38%, respectively, as compared with controls. Incubation of radish seeds in 30 µM DCPTA with actinomycin-D, alpha-amanitin, anisomycin, or cordycepin significantly reduced DCTPA-mediated seedling growth. These results indicate that nuclear gene expression and translation of mRNA on 80S ribosomes are required for the acceleration of seedling development by DCPTA.

The tertiary amine bioregulator DCPTA has been shown to increase the vegetative growth and harvestable yield of crop plants (Keithly et al., 1990, 1991a, 1991b). In addition, DCPTA promotes seed germination and increases seedling vigor. The rate of guayule (Parthenium argentatum Gray) seed germination was increased significantly by DCPTA treatment to dry seeds (Hayman and Yokoyama, 1990). Application of DCPTA to ungerminated cotton (Gossypium hirsutum L.) seeds significantly increased the root : shoot ratio of seedling plants (Gausman et al., 1988). In tomato (Lycopersicon esculentum Mill.), DCPTA treatment significantly increased seedling root and hypocotyl elongation growth and increased the relative growth rate of treated plants when compared with controls (Keithly et al., 1991a). In this study, a radish seedling growth test was used to further investigate the correlation of DCPTA-enhanced seedling vigor with improved harvestable yield.

Materials and Methods

Chemicals. DCPTA was synthesized and purified by the methods of Poling et al. (1977) and Echols et al. (1981), respectively. All other chemicals were purchased from Sigma Chemical Co. (St. Louis).

Seedling growth studies. Radish seeds (cv. Scarlet turnip white tipped) were provided by the Ferry Morse Seed Co., Modesto, Calif. Solutions of DCPTA that ranged in concentration from 3 µM to 50 mM (pH 4.3) were prepared using distilled water. All DCPTA solutions contained 0.1% Tween 80 (v/v). Radish seeds were hydrated for 6 h at 22C in 0.1% Tween 80. Hydrated seeds were then incubated in either 0.1% Tween 80 (control) or DCPTA solutions at 22C for 6 h. After treatment, seeds were germinated (95% germination frequency) in darkness at 27± 1C for 48 h using a ragdoll seed germination method (Gausman et al., 1988). Each ragdoll contained 60 to 65 seeds, and four replicates were prepared for each DCPTA treatment. After 48 h, primary root and hypocotyl lengths of 50 randomly selected germinates were determined for each replicate. Root tissue was distinguished from hypocotyl tissue by the emergence of root hairs along the primary root axis. Seedling growth experiments were performed five times.

Inhibitor studies. Radish seeds were hydrated as described for seedling growth studies. Seeds were then co-incubated for 6 hat 22C in 30 µM DCPTA and one of the following metabolic inhibitors: actinomycin-D, 4 µM; alpha-amanitin (AM), 5 µM; anisomycin, 19 µM; and cordycepin, 40 µM. Control seed lots were co-incubated for 6 h in 0.1% Tween 80 and inhibitor. For AM pulse studies, seeds were incubated for 6 h in 0.1% Tween 80 and then transferred to a solution of 30 µM DCPTA. Alpha-amanitin (5 µM) was added to the DCPTA solution as a pulse treatment; from 0.5 to 1, 1 to 2, 2 to 3, 3 to 4, 4 to 5, or 5 to 6 h after the DCPTA treatment. After treatment with AM, seeds were rinsed five times in distilled water and reincubated in fresh 30 µM DCPTA for a total of a 6-h DCPTA seed treatment. After the AM pulse treatment, seeds were germinated in ragdolls as described for seedling growth studies.

Greenhouse studies. Treated and control seeds were planted in 3.5-liter pots filled with a mixture of 2 fir bark : 1 sand :1 sphagnum peat (by volume). Seedlings were thinned to one plant per pot at the two true-leaf stage of development. Greenhouse-grown plants received a photosynthetic photon flux (PPF) of 1200 to 1400 µmol·m⁻²·s⁻¹ photosynthetically active radiation (PAR). Photoperiod was adjusted to 12 h with metal halide lamps (PPF at plant height, 450 µmol·m⁻²·s⁻¹ PAR). The greenhouse was maintained at a 24 ± 4C/16 ± 2C day/night cycle. Plants were fertilized weekly with 3 g 20N–27P–16K/liter. Each treatment group contained 20 replicates and each pot was considered to be an experimental unit. All pots were arranged as a completely randomized block. Foliage treatments of 0.1% Tween 80 (v/v) that contained 30 µM DCPTA (3 ml solution per plant) were performed at 6, 16, and 23 days after seed planting. Treatments were applied using a trigger-action hand sprayer.
Plants were separated into root and shoot samples 30 days after seed planting. Taproot diameters were recorded. Total leaf areas were determined using a Delta T video leaf area meter (Decagon Devices, Pullman, Wash.). Harvest indices were determined from fresh weight and dry weight values, after the plants were dried to constant weights at 75°C. Plant growth and yield studies were performed five times.

Lycopene analysis. Cotyledon pairs (1 g fresh weight per sample) were homogenized in acetone using a Polytron. Total carotenoids were extracted into acetone and were saponified (Benedict et al., 1985). Lycopene was quantified by absorption spectrophotometry (Benedict et al., 1985).

Statistical analyses. All growth data were subjected to analysis of variance with linear and quadratic regression analyses (Zar, 1974). Separations among treatment means were also determined using Duncan’s multiple range test at $P = 0.05$ (Zar, 1974).

Results

Application of 30 µM (10 ppm) DCPTA as a soak treatment to hydrated radish seeds increased the rate of primary root and hypocotyl elongation during seedling development (Fig. 1) when compared with the growth of controls. Radish seeds were fully hydrated after 6 h of imbibition, and all chemical treatments were performed before radicle emergence. Promotive and inhibitory effects of DCPTA on the development of etiolated radish seedlings were observed (Fig. 2). The largest numerical increases in root and hypocotyl elongation were observed in the 30 µM DCPTA-treatment group when compared with controls. Treatments of seed with >150 µM DCPTA inhibited seedling growth. Lycopene accumulation in cotyledons was induced by DCPTA treatments that were inhibitory to seedling growth. Treatments using >3 mM DCPTA (1000 ppm) inhibited seedling growth and lycopene accumulation in cotyledons and resulted in seedling tissue necrosis. The largest numerical improvements in seedling dry weight and root : shoot ratio were observed with 30 µM DCPTA when compared with controls (Table 1). However, quadratic responses indicated inhibitory effects of 300 and 900 µM DCPTA-treatments on seedling development.

Metabolic inhibitors of transcription and translation effectively prevented seedling growth enhancement by DCPTA (Table 2). Coincubation of radish seeds in 30 µM DCPTA with 5 µM AM, 4 µM actinomycin-D, 19 µM anisomycin, or 40 µM cordycepin significantly ($P = 0.05$) inhibited DCPTA-mediated seedling growth without inhibition of controls. Seedling growth enhancement by DCPTA was effectively blocked by 1-h pulses with AM that were applied simultaneously with DCPTA (Fig. 3a). However, significant growth inhibition by AM was not observed when DCPTA was applied 4 h before ungerminated seeds were hydrated.

![Fig. 1. Enhanced elongation growth of etiolated radish seedlings by DCPTA.](image)

![Fig. 2. Concentration dependence of DCPTA-mediated seedling growth and lycopene accumulation by cotyledons. Data represent means ($n = 50$) ± se. Root length, ■; hypocotyl length, ▼; cotyledon lycopene, □.](image)

Table 1. Enlarged dry weight of etiolated radish seedlings by DCPTA. Shoot values represent the combined weights of hypocotyl and cotyledon tissues. Root : shoot ratios ($R : S$) were determined from dry-weight values. Data represent mean ($n = 50$) values from five independent experiments.

| DCPTA (µM) | Dry wt (mg x 100) | Root | Shoot | Total | R : S |
|------------|-------------------|------|-------|-------|-------|
| 0          | 94                | 668  | 758   | 0.141 |
| 3          | 98                | 684  | 782   | 0.143 |
| 30         | 106               | 732  | 838   | 0.145 |
| 300        | 94                | 658  | 752   | 0.143 |
| 900        | 72                | 660  | 732   | 0.109 |

Significance: Q* = Significant at $P = 0.05$; Q = quadratic.

Table 2. Inhibition of DCPTA-mediated root and hypocotyl elongation.

| DCPTA (µM) | Inhibitor      | µM | Length (mm) |
|------------|----------------|-----|--------------|
| 0          | ---            | --- | 48.3 b       |
| 30         | ---            | --- | 77.6 a       |
| 30         | Actinomycin-D  | 4 (5) | 50.9 b       |
| 30         | Alpha-amanitin | 5 (5) | 51.3 b       |
| 30         | Anisomycin     | 5   | 48.4 b       |
| 0          | Anisomycin     | 19 (5) | 53.2 b       |
| 30         | Cordycepin     | 40 (10) | 52.3 b       |
| 30         | Cordycepin     | 40  | 59.3 b       |

Mean separation within columns ($n = 50$) by Duncan’s multiple range test, $P = 0.05$.

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seeds were pulsed with AM (Fig. 3a). Control seeds were insensitive to AM that was applied as a short-term pulse treatment (Fig. 3b).

Leaf and taproot development of mature radish plants were increased significantly by the application of DCPTA to hydrated seeds (Table 3). Significant 'quadratic responses were observed for all growth variables that were measured. Compared with controls, an enhanced dry-matter maintenance was observed in DCPTA-treated seedlings (Table 1), which suggests an effect of DCPTA on the respiratory efficiency (Bewley and Black, 1983) of germinating seeds. Mature radish plants grown from DCPTA-treated seeds showed significant improvements in total dry weight and harvest indices as compared with controls (Table 3). These results suggest that DCPTA may increase net CO₂ fixation per unit leaf area and significantly increase photosynthetic partitioning to developing root sinks during plant growth. Previous studies have shown DCPTA treatments to enhance net photosynthesis of mature sugarbeet (Keithly et al., 1990) and spinach (Keithly et al., 1991b) leaves due to improvements in chloroplast compartment size. In addition, competition for photosynthate by root and leaf sinks may increase net CO₂ fixation in mature, photosynthate-exporting leaves of DCPTA-treated plants (Keithly et al., 1990, 1991b).

Derepression of DNA-dependent RNA synthesis has been implicated as the mode of action of tertiary amine bioregulators on carotenoid biosynthesis (Benedict et al., 1985; Greenblatt et al., 1986). Our study has shown that inhibitors of transcription and translation effectively inhibit DCPTA-enhanced seedling development (Table 2). Alpha-amanitin and actinomycin-D inhibit nuclear RNA polymerase II activity and mRNA chain elongation, respectively, whereas cordycepin induces premature mRNA chain termination. Anisomycin is an inhibitor of protein synthesis on 80S ribosomes. Our study indicates that enhanced radish seedling development by DCPTA depends on de novo nuclear gene expression and translation of mRNA on 80S ribosomes. Growth inhibition of DCPTA-treated seedlings by means of pulsing with AM (Fig. 3) suggested that a rapid alteration of nuclear RNA polymerase II activity was induced by DCPTA. The conservation of mRNA and ribosomal RNA during seed maturation (Aspart et al., 1984; Bewley and Black, 1983) would explain the observed insensitivity of control seeds to inhibitor treatments (Table 2; Fig. 3). Our results (Table 2; Figs. 2 and 3) and the results of previous studies (Benedict et al., 1985; Greenblatt et al., 1986) suggest that a common mode of action may regulate vegetative plant growth and carotenoid biosynthesis in tertiary amine-treated crops. Application of DCPTA as a seed-treatment appeared to be the most effective dosage method to increase radish taproot yield (Table 4). These results suggest that plant growth regulation by DCPTA may involve the tem-

Table 3. Enhancement of leaf growth and taproot yield of radish by DCPTA. Plants were harvested 30 days after seed planting (n = 20).

| DCPTA (µM) | Leaf | Root |
|------------|------|------|
|            | Dry wt (g) | Area (dm²) | Dry wt (g) | Diam (mm) | Harvest index* |
| 0          | 1.1   | 3.7   | 0.5       | 18.2      | 0.32       |
| 3          | 1.1   | 3.7   | 1.0       | 28.4      | 0.47       |
| 15         | 1.6   | 5.2   | 1.1       | 30.1      | 0.41       |
| 30         | 1.4   | 4.8   | 1.2       | 32.6      | 0.46       |
| 150        | 1.1   | 3.6   | 0.6       | 22.1      | 0.37       |
| Significance | Q*   | Q*   | Q**       | Q**       | Q*         |

* Harvest index = g dry weight taproot/g total plant dry weight.
** Significant at P = 0.05 or 0.01, respectively. Q = quadratic.

Table 4. Effect of the time of application of DCPTA on the taproot yield of mature radish plants. Plants were harvested 30 days after seed planting. Letters within columns indicate mean (n = 20) separations according to Duncan's multiple range test, P = 0.05. Harvest indices (HI) based on root and shoot fresh-weight values.

| DCPTA (µM) | Type of application | Time of application | Fresh wt (g) | Diam (mm) | HI* |
|------------|---------------------|---------------------|---------------|-----------|-----|
| 0          | Seed                | ---                 | 18.4          | 27.5      | 0.54 |
| 30         | Seed                | ---                 | 27.7 a        | 36.8 a    | 0.67 a |
| 30         | Foliage             | 6                   | 23.8 ab       | 34.5 a    | 0.61 ab |
| 30         | Foliage             | 16                  | 21.3 ab       | 31.8 ab   | 0.59 b |
| 30         | Foliage             | 23                  | 17.8 b        | 27.1 b    | 0.62 ab |

* Days after seed planting.
* HI = g fresh weight taproot/g total plant fresh weight.
poral control of nuclear gene expression during radish seedling development.

Separate experimental methodologies are generally used to describe the biological activities of tertiary amines on carotenoid biosynthesis (Benedict et al., 1985; Greenblatt et al., 1986) and crop growth (Gausman et al., 1988). However, the results of this study indicate that the radish seedling growth method may be used to examine vegetative growth and carotenoid induction by DCPTA. In addition, the seedling growth test would provide a useful bioassay of the growth-inducing activities of newly synthesized tertiary amine bioregulators.

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