Polyamines and ethylene during in vitro rooting of *Prunus avium* L.

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Introduction

Polyamines are widely occurring organic polycations now recognized as plant growth substances. They appear to be involved in cell division, to delay senescence and usually accompany active growth and metabolism (Bagni et al., 1982). Although a requirement for polyamines has been postulated in some *in vitro* morphogenetic processes, a full understanding of their role in organogenesis has yet to come. The interaction between ethylene and auxin is a well-known phenomenon. Our primary interest in studying the action of ethylene in adventitious root formation stems from auxin usually being the key factor in root induction. Secondly, since ethylene and polyamines share a common precursor (S-adenosylmethionine) in their biosynthetic pathways, there is some evidence for a possible interrelationship between them. The aim of the present work was to begin examining the changes in the endogenous content of the polyamines, putrescine (PUT), spermidine (SPD) and spermine (SPM) and in the ethylene production accompanying the *in vitro* rooting of *Prunus avium* shoots. In addition, the effects of externally supplied polyamines, in particular SPM, of the ethylene precursor 1-aminocyclopropane-1-carboxylic acid (ACC) and of some inhibitors of polyamine and ethylene biosyntheses, namely α-difluoromethylornithine (DFMO), α-difluoromethylarginine (DFMA), dicyclohexylamine (DCHA), methylglyoxal-bis-guanilhydrazone (MGBG) and aminooxyvinylglycine (AVG) and aminooxyacetic acid (AOA) were examined.

Materials and Methods

The experimental material is part of a routine micropropagation program for selected clones of *P. avium* L. The establishment of bud cultures and the shoot multiplication and elongation phases have been described elsewhere (De Paoli and Rocchi, 1984). Rooting experiments were carried out by transferring elongated shoots onto agar-solidified medium composed of half-strength macro- and micro-nutrients of Murashige and Skoog (1962) (MS), MS vitamins, 30 g/l sucrose, pH 5.9–6.0, prior to autoclaving. Indole butyric acid (IBA) or indole acetic acid (IAA) was supplied at various concentrations (2.5, 5, 10, 25, 50 μM) to induce
rooting. Spermine, DFMO, DFMA, MGBG, AVG, AOA and ACC were filter-sterilized and added to the autoclaved medium. Cultures were kept in a growth chamber with a 16 h photoperiod at 22 ± 2°C. Polyamine levels were individually monitored at intervals in the shoot apical and basal 3 mm portions, stem and leaves. Dansylated derivatives of the polyamines were separated by thin-layer chromatography and fluorometrically determined. The incorporation of labeled PUT, the precursor of SPD and SPM, was evaluated by adding 185 kBq to 100 ml of medium (15 shoots). Shoot portions were extracted on d 2, 7 and 15 and polyamines were analyzed as described above. Radioactivity was measured in spots comigrating with SPD and SPM. A system was developed to measure 3,4-[14C]methionine (74 kBq/20 ml of medium) incorporation into ethylene. Ten shoots were placed in small flasks equipped with a side-arm, into which filter paper soaked in KOH was inserted, and a center well containing either 0.1 M mercuric acetate (in methanol) or 0.25 M mercuric perchlorate in order to capture the labeled ethylene formed. The radioactivity in the ethylene traps was measured after a 24 h incubation under normal culture conditions. The time course of methionine uptake was determined by extracting the different shoot portions in 10% trichloroacetic acid.

**Results**

Of the two auxins and different concentrations tested, 5 μM IBA gave the best rooting percentages after 12–15 d and was thus used in all subsequent experiments. Endogenous polyamine content during the rooting phase was characterized by a peak in PUT levels on d 9–11 in all shoot portions. Spermidine levels did not change significantly in leaves but showed maximum accumulation on d 9 or 11 in other shoot portions. Spermine was always absent or present in traces. Although no labeled PUT incorporation into SPM was observed, it is worth noting the sharp peak in SPD synthesis observed in the basal portion of shoots on d 7. Exogenously supplied SPM (10, 50 or 100 μM), either in the presence of optimal or suboptimal IBA levels, had no significant effect on rooting. DCHA plus MGBG (0.5 mM each) markedly reduced rooting but this inhibition was only partially reversed by the simultaneous application of 0.5 mM SPD. At higher concentration, the drugs provoked visible toxicity symptoms. DFMO plus DFMA (1 mM each) drastically reduced rooting as well and 0.2 mM PUT again partially reversed this effect. As expected, treatment with these specific inhibitors of PUT biosynthesis caused a severe decline in endogenous PUT levels. DCHA plus MGBG, however, caused only a minor reduction in SPD content. ACC was supplied only at 1 mM concentration, which proved to be lethal and thus no rooting was observed. Finally, AVG (25 or 50 μM) did not affect rooting percentages but enhanced the number of roots formed per rooted shoot. Results showed that mercuric perchlorate is a more efficient and reliable ethylene trap than mercuric acetate. The latter, being dissolved in methanol, quickly evaporated and, due to the presence of water vapor in the flasks, formed a yellow precipitate which was difficult to collect. The time course of labeled methionine uptake indicated that the compound was rapidly taken up (within 30 min) and reached a plateau around 15 h in all shoot portions. Preliminary data concerning ethylene biosynthesis indicate that, upon transfer to a rooting medium containing 5 μM IBA, ethylene production was of the order of 2.2 pmol/shoot/h. Finally, 0.5 mM AOA was shown to significantly reduce ethylene biosynthesis and, to a lesser extent, so did a 48 h exposure to DCHA plus MGBG.

**Discussion**

The results outlined above suggest that polyamines may be involved in the rooting
process, probably in the stages of active
cell division. In fact, increases in intracel-
lular PUT and SPD levels preceded root
protrusion and may have coincided with
maximum primordium development. Also,
a peak in SPD synthesis was observed on
d 7 in the basal portions, which are the
site of root formation. Finally, although
there does not seem to be a requirement
for SPM, either endogenous or exo-
genous, specific inhibitors of PUT and
SPD biosyntheses had a clear inhibitory
action on rooting. Our preliminary data
seem to indicate that DCHA plus MGBG
do not enhance ethylene production; this
may be due to the fact that these drugs
were ineffective in blocking polyamine
synthesis or may suggest that the 2 bio-
synthetic pathways are not competitive.

Further work on the role of ACC, AVG and
ethylene production in adventitious root
formation is in progress.

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