Rapid Testing of Salinity Effects on Pistachio Seedling Rootstock

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Abstract. A laboratory procedure was used to evaluate saline tolerance of pistachio rootstock species. Results were compared to those from a 2-year, outdoor lysimeter study to test reliability of the method. Excised root tips from seedlings of Pistacia atlantica Desf., P. terebinthus L. (two selections), and P. integerrima Stewart × atlantica (Pioneer Gold II, or PG II), were exposed to laboratory solutions that simulated soil solution electrical conductivity (EC) and Na : Ca ratios in the lysimeters. Following 24 hours of incubation, the efflux of ultraviolet (UV)-absorbing solutes was measured, providing an indication of cell membrane permeability. Leakage occurred with saline solutions comparable to lysimeter soil water salinity that increased leaf Na concentrations and decreased average root growth (175 mm NaCl with 12.5 mm Ca, or EC of 18.1 dS·m⁻¹). Cell injury increased linearly with salinity ($R^2 = 0.81$) and was highest in root tips of a P. terebinthus selection having least Na exclusion capability in the lysimeters. On average, these excised roots lost 38% more solutes than roots of a stronger Na-excluding genotype. There were no differences in leakage responses of the other species and selections. Leakage intensity was independent of various stress media, including isosmotic CaCl₂, mannitol, and the simulated Na/Ca mixtures in molar ratios of 10:1 to 20:1. With no Ca, however, damage caused by isosmotic NaCl was 76% to 87% higher, indicating that for these species, the Na : Ca ratio can alter root cell membrane permeability. Correlation between long-term observations in the lysimeters and leakage occurrence in the laboratory indicates that solute leakage tests with roots may aid in characterizing Pistacia spp. rootstocks for saline condition.

Empirical screening of rootstock for saline resistance can involve lengthy and extensive evaluations, but a simpler method could aid in the characterization process. The reliability of a rapid procedure could be determined by comparing results to responses obtained in a more natural environment, such as outdoor lysimeters. For example, Greenway (1970) proposed that short-term studies of excised roots, which reveal immediate consequences of salinity, may provide useful comparisons to long-term responses measured in natural conditions. Based on results with various crop species, Nassery (1979) suggested that measurement of K loss from excised root segments may accurately and rapidly estimate salt resistance of plants. Milbocker (1988) achieved a close correlation between the induction of cell plasmolysis in root tips and a conventional, long-term screening procedure for the ranking of salt resistance of seven azalea cultivars.

Passive release of UV-absorbing solutes from plant tissues also measures short-term salinity effects. Rauser and Hanson (1966) showed Na and K salts (50 mm concentrations and above) induced leakage of UV-absorbing materials from excised root tips of soybean. Several workers have applied this criterion to the study of salt-affected leaf disks (Leopold and Willing, 1984; Redmann et al., 1986). Leakage of UV-absorbing solutes upon injury is interpreted as direct injury to cell membranes (Leopold and Willing, 1984; Willing and Leopold, 1983).

We could find no reports incorporating long-term observations of growth and ion uptake of salt-treated rootstock with short-term laboratory responses using the UV-absorbing solute technique. Cell membranes of roots must endure immediate effects of salinity due to direct exposure to the soil solution (Maas and Nieman, 1978), a condition that could be simulated in the laboratory and observed by UV-absorption measurement.

The objectives of this study were to 1) assess salinity responses of root tips from two pistachio rootstock species and one hybrid using the UV-absorbing solute technique, and 2) compare these results to those obtained from a previous, long-term study involving the same rootstock.

Materials and Methods

Seedling culture and test material. Seedlings of P. atlantica (PI 246336), P. terebinthus (PI 246341 and 246342, referred to as A and B, respectively), and P. integerrima × atlantica (PG II) were maintained in an open shadehouse for 3 years in 10-liter plastic pots containing Hueco sandy loam soil (calcareous, coarse silty, mixed, thermic, Petrocalcic Paleargid). The seedlings were from the same experimental population of a previous salinity study made in outdoor lysimeters (Picchioni et al., 1990) but had not been used previously. On 22 Mar. 1989, dormant plants were moved to a greenhouse and tops were pruned back by ~25% (to 0.3 to 0.6 m high). Roots were washed free of soil, and individual seedlings were transplanted into sealed, 16-liter plastic pots containing complete, full-strength Hoagland nutrient solution #1 (Hoagland and Arnon, 1950). Aeration was provided continuously by “Micro II” tubing (Entek Corp., Grapevine, Texas), glued to the base of each pot, and connected to a pressure pump providing a flow rate of 3.2 × 10⁻³ m/sec per pot. The greenhouse was maintained between a minimum

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of 21°C at night and maximum of 27°C during the day. Throughout the 7-week study, nutrient solution pH varied between 5.5 and 6.5, and was renewed and adjusted to pH 5.5 (with 1 N NaOH or HCl) every 14 days. Midway between changes, solution was added to replace transpirational losses (ranging from 150 to 300 ml·day-1) and maintain upper structural roots below the surface of the solution.

Within 4 weeks, seedlings had vigorous tops with fully expanded leaves, three to five elongating shoots, and new roots. Such roots are described for other woody species as having rapid extension growth and subsequent secondary thickening; e.g., white "extension" roots of grapevine (Richards, 1983) and white "axial" roots of various rootstock (Rem, 1987). They are likely an important zone for water and nutrient absorption. In our study, visible lateral roots (≥5 cm proximal to the tip) were formed no sooner than 3 to 4 weeks after appearance of the primary structures. In soil environments, new roots of P. inglefieldiana have similar characteristics (e.g., white tips with rapid elongation) when observed through rhizotrons (Goldhammer et al., 1989; D. Goldhammer, personal communication). With the exception of diameter (0.5 to 1.5 mm), selected roots were morphologically similar.

Testing procedure. Four solute leakage trials (1 to 4) were made between 21 Apr. and 6 June 1989. All laboratory procedures were developed in preliminary experiments (Picchioni, 1989). One root per plant per treatment was removed in the greenhouse, then shortened to 2 cm from the tip using a razor blade. Preliminary observation indicated that the distal 1 to 2 cm was the most salt-sensitive zone of new roots. Root tips were taken to a laboratory in 5 ml of 0.5 mm CaCl₂, shaken for 24 h, and transferred to saline solutions for an additional 24 h. To accelerate leakage rates (Leopold and Willing, 1984), roots were next place in distilled water for 4 h. This released 95% of the solutes resulting from saline injury. Absorbance at 260 nm (A_260) of the 4 incubation solutions (3-ml samples) was measured using a Spectronic 21 UVD Spectrophotometer (Bausch and Lomb Instruments, Inc., Rochester, N.Y.). Solutions were returned to their original containers, sealed, and frozen at −25°C for 24 h. Preliminary results showed this temperature, which was the most practical during the experimental period, provided 96% ± 1% of solutes released by −5°C. The A_260 was again recorded after thawing and 4-h incubation, which liberated 75% of the total possible solutes. The sampling of incubation solutions (before and after freezing) was made at 4 h ± 5 sec. Solutions were centrifuged at 1260 × g before A_260 measurements. All incubations (5-ml volumes) were made in 125-ml flasks (slightly angled) at pH 6.1 using a reciprocating shaker (240 rpm) at the lowest setting. The laboratory was at 23.5 ± 0.5°C, with lighting (cool-white fluorescent) providing <10 µmol·s⁻¹·m⁻².

Relative leakage ratio (RLR) was calculated as the fraction of A_260 after salt treatment, but before freezing, to that after freezing. The background leakage from control roots, incubated in distilled water, was corrected to calculate an index of treatment injury (I):

\[ I(\%) = \frac{RLR_c - RLR_s}{1 - RLR_s} \times 100 \]  

where RLR_c and RLR_s refer to salt and control treatment samples, respectively. This equation uses the RLR of each treated replicate and the RLR of the control treatment mean (Flint et al., 1967).

Species comparison (trial 1). Six salt concentrations that approximated EC and Na : Ca molar ratios at the mean soil water content in the lysimeter soils (Picchioni et al., 1990) were applied to P. atlantica, P. terebinthus A, and PG II roots. The EC values for the control, 125:12.5, 175:17.5, and 250:25.0 (mM Na : Ca) treatment solutions were 2.3 × 10⁻¹, 1.4 × 10⁻¹, and 2.9 × 10⁻¹, respectively. To more closely simulate lysimeter soils, which had a CaSO₄ volubility limit in solution (Picchioni et al., 1990), Ca concentrations were held constant in two additional solutions at the highest Na concentrations, providing increased Na : Ca ratios (14:1 and 20:1). Salinity of the additional 175:12.5 and 250:12.5 solutions was 17.4 and 22.3 dS·m⁻¹, respectively.

Responses of P. terebinthus genotypes (trial 2). The leakage from roots of P. terebinthus A and B selections was compared in duplicate trials, and the results were combined. P. terebinthus B seedlings have been shown to maintain the lowest root and basal stem but highest leaf Na concentrations at high salinity compared to other rootstock selections (Picchioni et al., 1990). Only the control, 175:12.5, and 250:12.5 treatments or those of equivalent osmolality were applied in this and in subsequent trials. The EC values were 2.5 × 10⁻¹, 17.6, and 23.2 dS·m⁻¹, respectively.

Isosmotic solutions (trials 3 and 4). To evaluate solute specificity, Na/Ca, CaCl₂, mannitol, and NaCl without Ca (Na/0 Ca) were applied to PG II (trial 3) and P. atlantica (trial 4). The treatments were isosmotic to the 175:12.5 and 250:12.5 Na/Ca solutions (359 and 497 mOsm·kg⁻¹, respectively). Osmolality, measured with a cryoscopic osmometer (Osmette S, Precision Instruments, Sudbury, Mass.), varied by <0.3% among the isosmotic solutions.

Statistical analyses. Twelve seedlings (three per species or selection) were maintained in a randomized complete block in the greenhouse. For laboratory incubations, three blocks were arranged in split plots. The main plots (salinity level in trials 1 and 2, osmolality in trials 3 and 4), were randomized completely within each block. Each seedling provided one root per incubation solution and was assigned to a specific incubation block. Subplots were rootstock species or selection (trials 1 and 2) and solute type (trials 3 and 4). Incubation and solution sampling began with the first block and ended with the third. Analyses of variance were made for the split-plot according to Little and Hills (1978). For linear effects of salinity (orthogonal contrast in trial 1), rootstock responses were evaluated by slope and intercept comparisons (Snedecor and Cochran, 1989). Main plot and subplot effects were verified by F test, and subplots were distinguished by least significant difference (LSD), calculated using the formulas of Little and Hills for a split plot (1978). Multiple subplot comparisons were made using Duncan’s multiple range test. For interactions in trials 3 and 4, orthogonal contrasts were made between Na/O Ca and the remaining solutions.

Results

Rootstock species. The I values in Eq. [1] represent leakage from root tips resulting only from saline treatment, even though control roots lost appreciable amounts of UV-absorbing solutes. In trial 1, however, RLR did not increase until NaCl was raised to 175 mM (Fig. 1). Within 175- and 250- mM NaCl levels, Ca application rate (Na : Ca ratios of 10:1 to 20:1) did not alter root injury. For all rootstocks combined, I averages were 31.5% vs. 32.3% for 175:17.5 (10:1) and 175:12.5 (14:1), and 72.3% vs. 68.7% for 250:25.0 (10:1) and 250:12.5 (20:1) solutions, respectively (LSD = 14.6%). Data from both Ca treatments were then combined for further analyses.
Fig. 1. Comparison of relative leakage ratio (RLR) in control (0 NaCl) and saline media with Na : Ca molar ratios of 10:1, 14:1, and 20:1 (trial 1). Data represent the main plot averages for each treatment solution, for \( P. \) atlantica, \( P. \) terebinthus A, and PG II combined. Bars are the standard deviation of the mean of nine observations (LSD = 0.11, \( P = 0.05 \)). Subplots (rootstock species) were the average of three replicates each.

Fig. 2. Leakage from root tips (I\(_t\)) of rootstock in trial 1 (I\(_t\) calculated as in Table 1). The Na : Ca ratios are next to points representing the mean of three replicates, with data from both Na : Ca ratios averaged at 175 and 250 mM NaCl. Combined regression for line shown: \( I_t = 4.6 + 0.53 (mM \ NaCl - 125) \), \( R^2 = 0.81 \), \( n = 27 \).

Salinity effects resulted from the linear increase in \( I_t \) with salt concentration (the linear contrast accounted for 100% of the salinity variation). On average, 175 mM NaCl raised \( I_t \) by \( \approx 8 \) times, with high salinity (250 mM) resulting in a further doubling (Fig. 2). There were no \( I_t \) differences between rootstock species (nonsignificance for F test and regression comparisons); thus, all data were combined for an overall regression. Within this range of salinity and Na : Ca ratios, \( I_t \) increased 5.370 with each 10-mM increase in NaCl concentration.

\( P. \) terebinthus genotypes. Large increases in leakage occurred with \( P. \) terebinthus A and B roots exposed to high salinity (Table 1), the magnitude (average of \( \approx 2.8 \) times) being similar to that of the same treatment interval of trial 1. \( P. \) terebinthus B roots lost an average of 38% more solutes than \( P. \) terebinthus A, with the greater difference at low salinity. With 175:12.5 solution, there was high variability in leakage response of \( P. \) terebinthus A (\( I_t \) averages of 20.3% and 7.9%, compared with 36.0% and 56.1% for \( P. \) terebinthus B in duplicate trials).

\( P. \) atlantica responses in trials 1 and 4 were combined to show the influence of increasing salinity and Na : Ca ratio (Fig. 3). The Na/0 Ca treatments, 194.3 and 269.3 mM NaCl, were isosmotic to 14:1 and 20:1 solutions, respectively, whereas 10:1 solutions had 5% to 7% higher osmolality. All \( I_t \) values are grouped within the 175 and 250 mM concentrations for simplicity.

Table 1. Leakage of UV-absorbing solutes (I\(_t\)) from root tips of \( P. \) terebinthus seedling genotypes exposed to two Na : Ca ratios (trial 2).

| Genotype          | Na : Ca concn (mm) | \( I_t \) (%) |
|-------------------|-------------------|--------------|
|                   | 175:12.5          | 250:12.5     |
| \( P. \) terebinthus A | 14.1 b            | 82.4 a       |
| \( P. \) terebinthus B | 46.1 a            | 87.1 a       |

\( I_t \) expressed as \( \frac{(RLR_s - RLR_c)}{(1 - RLR_c)} \times 100 \), where RLR\(_s\) = salt treatment for each replicate, and RLR\(_c\) = control treatment mean.

\( \gamma \)Mean separation in columns by F test (\( P = 0.05 \)). Each value is the mean of two trials of three replicates each.

Table 2. Leakage of UV-absorbing solutes (I\(_t\)) from root tips of PG II (trial 3) and \( P. \) atlantica (trial 4) in various isosmotic solutions.

| Solution* | Osmolality (mOsm·kg\(^{-1}\))* |
|-----------|--------------------------------|
| Na/Ca     | 14.3 b                         | 42.3 b       |
| CaCl\(_2\) | 15.2 b                         | 52.1 ab      |
| Mannitol  | 26.8 b                         | 37.0 b       |
| Na/0 Ca   | 45.9 a                         | 64.4 a       |
| PG II     | 359                            | 497          |
| Na/Ca     | 23.3 b                         | 77.6 a       |
| CaCl\(_2\) | 9.8 b                          | 75.3 a       |
| Mannitol  | 17.4 b                         | 50.8 b       |
| Na/0 Ca   | 68.8 a                         | 90.0 a       |

\( I_t \) calculated as in Table 1.

Low and high osmolality correspond to Na : Ca mixtures of 175:12.5 and 250:12.5 mM, respectively.

Discussion

Salinity damaged root tips of \( P. \) atlantica, \( P. \) terebinthus, and PG II at 175 mM NaCl with 12.5 to 17.5 mM Ca (average EC of 18.1 dS·m\(^{-1}\) in trial 1). This is similar to the soil solution salinity (17.9 dS·m\(^{-1}\)) and Na : Ca ratio (174 : 13.5) that decreased root growth an average of \( \approx 60\% \) (Picchioni et al., 1990). We did not test leakage between 125 and 175 mM Na. Additional study within this salinity range would determine how
to the presence of a significant interaction in trial 4 (osmolality concentrations of Na and Cl in root tips of a related species, Poovaiah and Leopold, 1976; Hanson, 1966). About 10 to 25 mg fresh weight of tissue was available for each 24-h incubation. This raised the possibility that excised roots of P. terebinthus B are less tolerant to saline incubation than those of P. terebinthus A is of interest. In the 174:13.5 lysimeter solution, P. terebinthus B contained ≈140% more Na in leaf tissue and ≈33% to 66% less Na in whole roots and basal stems, respectively, than four other pistachio rootstock (Picchioni et al., 1990). This is of practical importance for rootstock selection, but leaves unanswered the question of whether the Na storage capacity was merely limited (Jacoby, 1964), or whether the storage was directly involved in root injury (Bernstein, 1975). For example, Levitt (1980) stated that saline tolerance may be controlled by the composition of cell membranes (lipid and bound protein constituents).

We experienced leakage variability between trials with all species and selections, which was highest for P. terebinthus A (=60% in trial 2). A possible explanation is the time elapsed during the duplicate trials (=30 days), even though we could not detect macroscopic root differences between sampling periods. Redmann et al. (1986) found leakage of UV-absorbing solutes from 24-h salt-treated leaves of aspen (Populus tremuloides Michx.) declined with leaf age during a 30-day sampling period, but there was no consistent relationship between root salt tolerance and culture time in this study. Differences in root growth rates, as well as background leakage (Flint et al., 1967), may also have caused between-trial variation.

Coefficients of variation averaged 35% for solutions causing root injury. This indicates that precision may be improved by increasing the number of observations or the number of roots per incubation, even though these were limitations in our study. Other studies have used shorter treatment durations (2 to 6 h) and greater amounts of root tissue (Nassery, 1975; Rauscher and Hanson, 1966). About 10 to 25 mg fresh weight of tissue was available for each 24-h incubation. This raised the possibility that irreversible damage by CaCl2 and mannitol eliminated specific effects of Na/Ca simulations. However, the effects of all three osmotica were similar even with a 75% reduction in exposure time (6 h), except Amax measurements were ≈30% lower before freezing (data not shown). Thus, much of the cell injury occurred within 6 h, but longer duration increased test sensitivity without altering the results.

The preparation of seedling rootstock material need not be as prolonged as in the above conditions. For example, root tips from 2-month-old P. atlantica responded much like those from

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**Fig. 3. Solute leakage (L) from root tips of P. atlantica as influenced by Na : Ca ratio and NaCl concentration (Lcalculated as in Table 1).** Center columns with Na : Ca ratios of 14:1 (175:12.5) and 20:1 (250:12.5) represent combined averages of trials 1 and 4 (n = 3 for each trial). End columns are the averages of three observations in trial 1 (10:1) and trial 4 (0 Ca). Mean separation within concentrations (main plots) by Duncan’s multiple range test, P = 0.05.
seedlings aged 3 years in the leakage trials. We also found that young lateral roots from pecan seedlings [Carya illinoensis (Wangen.) C. Koch], which are less saline-resistant than Pistacia spp. in lysimeter conditions (Miyamoto et al., 1985), are also less tolerant to saline incubation (e.g., leakage at 100 mM NaCl). Thus, this approach also may have potential for other tree crops.

The correlation between lysimeter data and that obtained in the present study indicates that solute leakage tests with roots might provide a simple and rapid means of characterizing salinity responses of rootstock. Such aspects could include saline resistance and differential avoidance of adverse ion relations.

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