The Influence of Stock Plant Fertilization on Tissue Concentrations of N, P and Carbohydrates and the Rooting of Prosopis alba Cuttings

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ABSTRACT


Clonal propagation techniques are required for Prosopis due to great variability in biomass productivity, pod productivity, and nitrogen fixation from seed propagated stock. Previous studies identified a decline in the rooting percentage of cuttings taken from stock plants that were grown in the growth chamber and harvested every four weeks. To determine the influence of the mineral nutrient status of the stock plants on the rooting of cuttings, fertilized and nonfertilized Prosopis alba clone B2V50 stock plants were established under 1000 W metal halide lamps in the greenhouse. A complete macro- and micronutrient solution was used. The regrowth was harvested after 4, 8, 12, 16, and 20 weeks, and alternate nodes from each stem were taken for the rooting assay and for N, P, K, Ca, Mg, Fe, Zn and total available carbohydrate evaluation (N=79). The mean rooting percentage for the five harvest cycles was 58% for the fertilized stock plants, and 44% for the nonfertilized stock plants (P=0.087). There was no significant correlation between rooting percentage and leaf carbohydrate (P=0.964), stem carbohydrate (P=0.876) and leaf plus stem phosphorus (P=0.319). The nonsignificant correlation between percent rooting and carbohydrates was negative (r= -0.018). However, there was a significant correlation with stem nitrogen and percent rooting (P=0.009) and an inverse correlation between stem nitrogen and leaf carbohydrate content (r=-0.353, P=0.001). This suggests that unbalanced fertilizers, rich in nitrogen and low in P, K, and other nutrients, might stimulate stem N and be successful in maintaining high rooting percentages from stock plants.

INTRODUCTION

Nitrogen fixing trees of the genus Prosopis are drought resistant and well adapted to poor soils of dry regions (National Academy of Sciences, 1979). In addition to producing biomass for use as a fuel, the trees provide pods which can be used for human or livestock food and they are useful for soil and environmental amelioration in general.
Previous research encountered large variability in *Prosopis* in mean biomass productivity per tree between species (0.2–29 kg per tree) and within species (1.9–23.9 kg per tree) (Felker et al., 1983). Considerable variation was also exhibited in tolerance to cold, salinity, and in pod production (Felker et al., 1981, 1982, 1984). This variability can be attributed to the fact that several *Prosopis* species are self-incompatible (Simpson, 1977) and thus the plants are obligate outcrossers that do not propagate true to type from seed.

Rooting of *Prosopis* cuttings has been previously reported by Leakey and Last (1980), Felker and Clark (1981), and De Souza and Nascimento (1984). The latter two groups reported strong seasonal influences on the rooting of cuttings. The influences of environmental parameters on rooting of cuttings were examined in growth chamber studies by Klass (1984) and Klass et al. (1985). These workers found that a temperature of 35°C, a light intensity of 520 μE m⁻² s⁻¹, and a 12 h photoperiod were necessary for optimum rooting of *P. alba* clone B₂V₅₀.

An unexpected result of the experiment to determine the optimum environmental parameters for rooting was a significant decline in all rooting parameters as the one year old, previously uncut stock plants were harvested every 3–4 weeks for rooting material (Klass et al., 1985). Suggested causes for this decline included: decreased juvenility of the stock plants, inadequate mineral nutrient status of the stock plants due to overharvest, and decreased supply of carbohydrate in the stem and root system of the stock plant.

This study was conducted to determine (1) if fertilization could overcome the decline in rooting percentages associated with successive harvests of stock plants for rooting material and (2) the correlations between rooting percentages and macronutrients, micronutrients, and carbohydrates in the leaves and stems of the stock plants.

**MATERIALS AND METHODS**

*Prosopis alba* clone B₂V₅₀ (referred to as V₅₀B₂ in Felker et al., 1983) used in this experiment was selected in a progeny trial to screen for biomass productivity under heat/drought stress in the California Imperial Valley (Felker et al., 1983). The parent accession of this clone was *P. alba* 0194 that was collected from ornamental trees growing along the Colorado River in Big River, California. These South American trees were introduced into southern California or southern Arizona by unknown means in the early 1950's.

To determine the influence of the mineral nutrient status of the stock plants on the rooting of cuttings, fertilized and nonfertilized stock plants were established in the greenhouse. The fertilized plants received one liter of a complete macro- and micronutrient solution three times a week. The nonfertilized stock plants received no fertilizer addition. A commercial fertilizer formulation (ValGro) was diluted to the following composition: nitrogen (800 mg/l), phospho-
rus (500 mg/l), potassium (300 mg/l), zinc (2 mg/l), iron (0.2 mg/l), magnesium (0.15 mg/l), copper and manganese (5 mg/l). The plants and vegetative regrowth were sprayed once a week with the fungicide benomyl [methyl-1-(butyl-carbamoyl)-2-benzimidazole carbamate].

Previous studies established the need for a light intensity of at least 200 μE m$^{-2}$ s$^{-1}$ for the stock plants (Klass, 1984; Klass et al., 1985). Since these experiments were partially conducted in the winter when light intensities in the greenhouse were as low as 100 μE m$^{-2}$ s$^{-1}$, 1000 W metal halide lamps were used to supplement the light intensity to a minimum of 300 μE m$^{-2}$ s$^{-1}$. Two nonfertilized and two fertilized stock plants were established under each of four metal halide lamps. Thus a randomized complete block design was used with each light source as a block.

The 2.5 m tall stock plants, in 20 l pots, were about one year old and had a basal diameter of about 3 cm. Before the experiment, the stock plants were cut at about 1 m in height leaving a severed stem about 2 cm in diameter.

The stock plant regrowth was successively harvested after 4, 8, 12, 16, and 20 weeks. The fertilized stock plants typically produced 5 to 10 new shoots about 60-80 cm long in the 3–4 week period, while the nonfertilized plants produced 2–3 shoots 30–40 cm long. At each harvest alternating nodes from each stem were taken for chemical analysis and the rooting assay. The leaves and stems were analyzed separately for N, K, Ca, Mg, Fe, Zn, and total available carbohydrate, but were pooled for the phosphorus analyses. From each of the eight fertilized and eight nonfertilized stock plants, 16 cuttings were taken for the rooting assay and two samples (leaf and stem) for chemical analyses.

The cuttings had two nodes with the leaves removed from the lower node. Each cutting was dipped in 0.3% IBA in talcum, then placed in a round plastic pot (13×13 cm) filled with fine vermiculite previously soaked for two days. Each pot plus vermiculite received 80 ml of a 0.5 g/l drench of the fungicide Banrot [5-ethoxy-3-trichloromethyl-1,2,4-thiadiazole and dimethyl-4,4-O,O-phenylenebis(thioallophanate)]. Eight cuttings were planted per pot, then placed in a polyethylene bag (25×41 cm), sprayed with benomyl + captan (0.5 g l$^{-1}$) and the top sealed with rubber band. The cuttings were placed in a light bank with a 12 h photoperiod and 300 μE m$^{-2}$ s$^{-1}$ of fluorescent light. Three weeks after placement under the light bank, the rooting percentage, the number of roots per cutting, and the length of the longest root were measured.

Samples for N and total available carbohydrate evaluation were dried for 36 h at 70°C, while samples for P, K, Ca, Mg, Fe, and Zn were dried for 24 h at 70°C. The dried samples were ground to pass through a 40-mesh screen using a Wiley mill equipped with stainless steel hardware. Tissue N was measured using a micro-Kjeldahl digestion, followed by colorimetric determination of the ammonium as described elsewhere in this volume (Cline et al., 1986). A concentrated nitric acid digestion was used for the remainder of the elements, also as described by Cline et al. (1986). Phosphorus was measured by the col-
orimetric procedure of Murphy and Riley (1962) and the remaining elements were measured using atomic absorption spectrophotometry. To evaluate the effect of K, Ca, Mg, Fe and Zn on rooting, 24 samples (leaf plus stem) were selected that represented the full range in rooting percentage.

The total available carbohydrates were extracted according to Smith et al. (1964), and measured spectrophotometrically using the anthrone reagent (Yemm and Willis, 1954).

RESULTS

The effect of stock plant fertilization on the number of roots per cutting, maximum root length, and rooting percentage is presented in Fig. 1. Each data point is the mean of eight plants, two pots per plant with eight cuttings per pot. The vertical lines represent the 95% confidence interval for the means (T-test). A decline in rooting percentage was observed on the third harvest. The rooting percentage started off higher (73%) for the fertilized than for the nonfertilized stock plants (60%), but they declined to 33% and 35% respectively for the third harvest. These mean rooting percentages were not statistically different except in the fourth and fifth harvest, where the rooting percentage for the fertilized stock plants was higher than for nonfertilized plants.

The maximum root length of the cuttings from the fertilized stock plants decreased from 12.1 cm (first harvest) to 7.2 cm (fourth harvest), and increased...
again to 13.3 cm at the last harvest. A similar response was obtained from the nonfertilized stock plants. The number of roots in both treatments also decreased at the third harvest, but further remained constant until the last observation. As can be seen in Fig. 1, the 95% confidence intervals for the root number and root length overlapped for all harvest periods.

There was a wide variation in rooting between clonal stock plants of the same treatment. For example, the rooting percentage of the eight fertilized stock plants ranged from 38% to 100%, 25% to 75%, 0% to 63%, 13% to 88% and 38% to 88% for the first, second, third, fourth, and fifth harvests, respectively.

A comparison of the rooting percentage with the nitrogen, carbohydrate, and phosphorus contents of the leaves and stems is presented in Fig. 2. The nitrogen contents of the fertilized leaves and stems are high, i.e. equivalent to 31% and 12% crude protein, respectively. The nitrogen and phosphorus contents of the stems and leaves from the fertilized stock plants are greater than the nonfertilized stock plants. In contrast, the leaf and stem carbohydrate contents are higher for the nonfertilized stock plants. The strong dip in the rooting percentage at the third harvest is paralleled by a dip in phosphorus percentage and an increase in stem carbohydrate. The changes in the stem nitrogen content are not as large as the other parameters, but the stem N appears to track the rooting percentage for both the fertilized and nonfertilized stock plants better than any of the other parameters.

The statistical analysis for the influence of stock plant fertilization on rooting parameters and chemical composition of \textit{P. alba} cuttings is presented in Table 1. When averaged over all five harvests, the rooting percentage for the
TABLE 1

Effect of stock plant fertilization on percent rooting and chemical composition of Prosopis alba B7V50 cuttings for all five harvest periods combined

<table>
<thead>
<tr>
<th>Plant characteristics</th>
<th>Fertilized</th>
<th>Non-fertilized</th>
<th>Level of significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Root number</td>
<td>7.90</td>
<td>7.60</td>
<td></td>
</tr>
<tr>
<td>Root length (cm)</td>
<td>10.30</td>
<td>8.48</td>
<td></td>
</tr>
<tr>
<td>Rooting (%)</td>
<td>58.00</td>
<td>44.00</td>
<td>0.0866</td>
</tr>
<tr>
<td>Leaf N (%)</td>
<td>4.85</td>
<td>3.79</td>
<td>0.0031</td>
</tr>
<tr>
<td>Stem N (%)</td>
<td>1.88</td>
<td>1.40</td>
<td>0.0033</td>
</tr>
<tr>
<td>Leaf carbohydrate (%)</td>
<td>3.15</td>
<td>3.77</td>
<td>0.0222</td>
</tr>
<tr>
<td>Stem carbohydrate (%)</td>
<td>3.21</td>
<td>3.26</td>
<td>0.8595</td>
</tr>
<tr>
<td>Leaf + stem P (%)</td>
<td>0.30</td>
<td>0.23</td>
<td>0.0940</td>
</tr>
</tbody>
</table>

*Each entry is the mean of 5 replications in time, consisting of 40 observations for fertilized and 39 for nonfertilized.

cuttings from the fertilized stock plants was greater \((P=0.087)\) than for the nonfertilized stock plants. The greater leaf and stem N in the fertilized stock plants were very highly significant \((P=0.003\) in both cases). Leaf carbohydrate was significantly greater for the nonfertilized stock plants \((P=0.022\) ), but stem carbohydrate \((P=0.859)\) and leaf plus stem P \((P=0.94)\) were not significantly different between treatments.

The correlation between rooting percentage and leaf and tissue concentrations of N, carbohydrate and P presented in Table 2 support the hypothesis that stem N tracks the percent rooting better than the other parameters. For example, no significant correlation was detected between rooting percentage

TABLE 2

Correlation coefficients of rooting percent with N, carbohydrates, and P content of P. alba B7V50 cuttings*

<table>
<thead>
<tr>
<th>Plant characteristics</th>
<th>Rooting percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Correlation coefficient</td>
</tr>
<tr>
<td>Leaf nitrogen</td>
<td>0.205</td>
</tr>
<tr>
<td>Stem nitrogen</td>
<td>0.368</td>
</tr>
<tr>
<td>Leaf carbohydrates</td>
<td>-0.005</td>
</tr>
<tr>
<td>Stem carbohydrates</td>
<td>-0.018</td>
</tr>
<tr>
<td>Leaf + stem phosphorus</td>
<td>0.114</td>
</tr>
</tbody>
</table>

*For 79 pairs of observation, chemical analyses versus rooting \((\%)\).
TABLE 3
Correlation coefficients of nitrogen with carbohydrate and phosphorus contents of *P. alba* B₂V₅₀ cuttings*

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Leaf nitrogen</td>
<td>-</td>
<td>-</td>
<td>0.508</td>
<td>0.0001</td>
</tr>
<tr>
<td>Stem nitrogen</td>
<td>0.508</td>
<td>0.0001</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Leaf carbohydrates</td>
<td>-0.344</td>
<td>0.0019</td>
<td>-0.353</td>
<td>0.0014</td>
</tr>
<tr>
<td>Stem carbohydrates</td>
<td>-0.010</td>
<td>0.9312</td>
<td>-0.167</td>
<td>0.1416</td>
</tr>
<tr>
<td>Leaf + stem phosphorus</td>
<td>0.356</td>
<td>0.0013</td>
<td>0.355</td>
<td>0.0013</td>
</tr>
</tbody>
</table>

*For 79 pairs of observation.

and leaf or stem carbohydrate or phosphorus concentrations. Leaf N was slightly correlated \( r=0.205, P=0.070 \) with rooting percentage while stem N was highly correlated \( r=0.368, P=0.0009 \). Since stem N was so significantly correlated with rooting percentage, correlations between stem N and the other leaf and stem parameters were examined (Table 3). Stem nitrogen was significantly negatively correlated \( r=-0.353, P=0.001 \) with leaf carbohydrate, but not with stem carbohydrate. In contrast, leaf plus stem phosphorus was significantly and positively correlated \( r=0.355, P=0.001 \) with stem nitrogen.

The correlations between K, Ca, Mg, Fe, Zn and rooting percentage and N are presented in Table 4. No significant correlations were detected between K, Ca, Mg, Fe, or Zn and rooting percentage. However, Leaf N was correlated with

TABLE 4
Correlation coefficients between percent rooting, tissue N, and K, Ca, Mg, Fe, Zn concentrations of *P. alba* B₂V₅₀ cuttings*

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>Potassium</td>
<td>0.071</td>
<td>ns</td>
<td>0.004</td>
<td>ns</td>
<td>0.117</td>
<td>ns</td>
</tr>
<tr>
<td>Calcium</td>
<td>-0.053</td>
<td>ns</td>
<td>0.220</td>
<td>ns</td>
<td>0.090</td>
<td>ns</td>
</tr>
<tr>
<td>Magnesium</td>
<td>0.085</td>
<td>ns</td>
<td>0.525</td>
<td>0.05</td>
<td>0.520</td>
<td>0.05</td>
</tr>
<tr>
<td>Iron</td>
<td>-0.360</td>
<td>ns</td>
<td>0.286</td>
<td>ns</td>
<td>-0.182</td>
<td>ns</td>
</tr>
<tr>
<td>Zinc</td>
<td>0.084</td>
<td>ns</td>
<td>0.460</td>
<td>0.05</td>
<td>0.218</td>
<td>ns</td>
</tr>
</tbody>
</table>

*For 24 pairs of observation, chemical analyses versus rooting (%). ns: P-value below 0.20.
Mg ($r = 0.525, P = 0.05$) and Zn ($r = 0.460, P = 0.05$). Stem N was only correlated with Mg ($r = 0.520, P = 0.05$).

**DISCUSSION**

Through use of fertilization, 70% rooting was achieved at fifth harvest, while 59% was obtained when no fertilizer was used. In spite of an unexplained decrease in rooting observed at the third harvest, the mean for all five harvests was 58% rooting for the fertilized stock plants and 44% for the nonfertilized ones.

The 58% mean rooting for all five periods is considerably less than the 83% mean for the 12 h stock plant/12 h cutting photoperiod observed by Klass (1984) and Klass et al. (1985) in growth chamber studies. The lower percentage rooting in the present study is, no doubt, partially due to less control of environmental parameters in the greenhouse as compared to the growth chambers. Nevertheless, the 58% rooting in the greenhouse in the winter is a considerable improvement over the 15% rooting previously found by Felker and Clark (1981) in the greenhouse during the winter.

Despite the fact that the fertilization did not greatly enhance the rooting percentage, the nonfertilized stock plants grew very slowly compared to the fertilized stock plants, making it difficult to obtain sufficient cutting material for the experiments. Thus, frequent fertilization will be required to maintain the growth rates required for commercial production.

Carbohydrate reserves in the leaves and stems of cuttings has long been considered an important determinant for rooting success (Hess, 1969; Eliasson, 1978; Hartmann and Kester, 1983), and the fact that we observed no correlation between leaf and stem carbohydrate with rooting percentage was surprising. It is curious that this clone has such an obligate requirement for high light intensities (Klass et al., 1985) and yet has such a poor correlation between total available carbohydrate and rooting percentage. De Souza and Nascimento (1984) have demonstrated that pruning *Prosopis* leaves prior to incubating them for rooting greatly decreases the rooting percentage.

These *Prosopis* leaves and stems were considerably higher in crude protein (31% and 12%, respectively) than other nonlegume hardwood species and it is possible that legume cuttings have a greater demand for nitrogen than for carbohydrate. Production of new leaves during rooting may represent a significant N sink. Alternatively, these high N contents may serve as a protein source during root production. The fact the De Souza and Nascimento (1984) have demonstrated an obligate requirement for leaves, coupled with the high leaf crude protein, suggests a strong demand for nitrogen in the cuttings.

The high leaf carbohydrate observed in cuttings obtained from the nonfertilized stock plants may be due to the reduced growth associated with limiting nutrients. On the other hand, the fertilized stock plants are less nutrient lim-
ited and are capable of more rapid growth, thus diluting the carbohydrate concentration in the leaves and stems.

Since N was positively correlated with rooting percentage and negatively correlated with leaf carbohydrate, this indicates that unbalanced fertilizers, rich in N and low in other nutrients, might increase stem N, decrease stem carbohydrate, and be successful in maintaining high rooting percentages from stock plants.

Although P, Mg and Zn were not correlated with rooting percentage, they were correlated with N content, suggesting a need to understand the relationship between these elements with nitrogen and possible effects on rooting.

The wide variation in rooting percentage from replicated clonal stock plants in the same treatment, at the same time period i.e. from 38% to 100% at the first harvest and from 0% to 63% at the depressed third harvest, implies lack of control of an additional rooting factor. Furthermore, the third harvest period had a low rooting percentage for both the fertilized and nonfertilized stock plants for no apparent reason. There were no mechanical failures and the environmental records indicated no unusual events. Thus, there still are major unknown factors besides exogenously supplied auxin concentrations, air temperature, light regimes, and nutrient requirements that dramatically influence rooting percentages and routinely preclude obtaining near 100% rooting success.

ACKNOWLEDGEMENTS

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