Optimization of the micropropagation of elite adult trees of *Sequoia sempervirens*: forest species of interest in the Basque Country, Spain

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**Abstract.** Forest trees are renewable sources of timber and other valuable non-timber products. Nowadays, the increase in population and demand for forestry products results in overexploitation of forestry. Therefore, there is an urgent need to produce elite plants with higher productivity under stress derived from climate change to have available to afforestation. For this reason, propagation methods should be improved to be more efficient in terms of quality and productivity. The main species planted in the Basque Country is *Pinus radiata*; during the last three years, *Pinus radiata* plantations have suffered a fungus attack affecting mainly needlings until the tree’s death. This crisis is caused by the combined action of two fungi of the genus *Dothistroma* and *Lecanosticta acicola*, whose expansion seems to have been enhanced by weather-related factors, such as humid and hot summers. Although we have evidence of this disease’s presence in our mountains since 1942, the disease has had a speedy expansion with an aggressive effect for reasons that are not scientifically known today. For the above, Basque Country forestry sector is looking for alternative species to be used in its plantations. Part of the forestry sector considers that *Sequoia sempervirens* could be a good choice for plantations. Besides, its high-quality wood and its tolerance to the attack of several pathogens and other diseases derived from climate change are characteristics that could confer some advantage over other forest species. The main goal of this study was to optimize the micropropagation of adult elite trees of *Sequoia sempervirens*. The effect of 6-benzylaminopurine, meta-Topolin and Kinetin, and 4 types of explant in the multiplication stage were analyzed to carry out this objective. Furthermore, the effect of two types of auxins: 1-naphthalene acetic acid, indole-3-butyric acid, and a mixture of both, were evaluated on the induction of roots and their subsequent effect on the acclimatization process. The best multiplication index was obtained when 4.4 µM 6-benzylaminopurine and apical explants longer than 1.5 cm of length were used. The root induction percentage was 75% in the most responsive genotype analyzed when 4.4 µM 6-benzylaminopurine was used on the induction stage, and 50 µM of 1-naphthalene acetic acid was used for rooting. Finally, after 3 months in the greenhouse, the explants cultured with Kinetin and rooted in a culture medium with indole-3-butyric acid showed the highest acclimatization success (94%).

**Key words:** Acclimatization, auxins, cytokinins, multiplication index, organogenesis, rooting, shoot induction.

**Introduction**

The Coast redwood or California redwood *Sequoia sempervirens* (D. Don.) Endl. (Taxodiaceae) is a valuable forest species and occurs naturally in Western North America, especially in California1,2. This species is the tallest tree on earth with a high volume of standing biomass, in some stands exceeding 3500 metric tons/hectare3.

This conifer has been introduced and domesticated in European countries such as Romania, Spain, France, Great Britain, Russia, and Turkey, and it can be used for reforesting due to the quality of its wood and for ornamental purposes. In Spain, this species has been used by foresters for its productivity, its tolerance to the attack of several pathogens and other diseases. Moreover, its reforestation is recommended on valley bottoms and fotthills4.

Nowadays, redwood shows seed reproduction difficulties, displaying low germination and rooting rates, dormancy of the shoot, and low seedling viability5,6. Therefore, biotechnological in vitro techniques of plant tissue culture as micropropagation, organogenesis, adventitious or auxiliary shoot/bud regeneration, shoot tip culture, micrografting, and somatic embryogenesis emerged as useful tools for the propagation and conservation of germplasm7,8,9.

There are few investigations on micropropagation in the specific case of *Sequoia sempervirens*, either through organogenesis or somatic embryogenesis. Arnaud et al.10 developed a protocol for micropropagation and rejuvenation of the species using direct organogenesis and somatic embryogenesis. Years later, Mihaljević et al.1 investigated the root formation in micropropagated shoots of *Sequoia sempervirens* using *Agrobacterium*. In the 21st century, Korban and Soul11 developed a procedure for the micropropagation of juvenile and adult material, and Lui et al.12 worked on the shoot regeneration and somatic embryogenesis from needles of redwood. The medium-term conservation of *Sequoia sempervirens* was investigated by Ozudogru et al.13, and Meneguzzi et al.14 evaluated shoot multiplication of two *Sequoia sempervirens* genotypes with the addition of Kinetin.

Considering all the information mentioned above, our study’s objective was to optimize the micropropagation process of elite selected adult trees of *Sequoia sempervirens*. We focused on improving 1) the multiplication phase using different types of explants and cytokinins, 2) the rooting using different auxins, and the subsequent acclimatization phase under ex vitro conditions.

**Materials and methods**

**Plant material**

Stem sections were collected from the basal parts of three different mother trees (7, 11 and 12) of *Sequoia sempervirens*...
located in Ataun, Gipuzkoa, Basque Country (Spain), at the geographic coordinates: 42° 58′ 41″ N and 2° 10′ 53″ O.

Shoot induction

In vitro micro shoots growing in half-strength ARN medium (Arnaud et al.) were selected to carry out the experiments. Four types of explants were used: A) apical explants of 1.5 cm in length (AG), B) apical explants of 1.0 cm length (AP), C) basal explants of 1.5 cm length (BG), and D) basal explants of 1.0 cm in length (BP). Explants were cultured in glass jars (Merck) with 25 mL of ARN multiplication medium supplemented with 3% sucrose and 8 gL⁻¹ Difco Agar®. The effect of different cytokinins for the induction of shoots was evaluated: A) 6-benzylaminopurine (BAP), B) meta-Topolin (m-T), and D) kinetin (K) at a concentration of 4.4 µM. The pH of the medium was adjusted to 5.8 before autoclaving (121°C, 20 min). The shoots were induced in the different induction media for four weeks in the growth chamber at a temperature of 21 ± 1°C, under 16-h photoperiod with 120 µmol m⁻² s⁻¹ of light intensity cool white fluorescent tubes (TLD 58 W/33; Philips, France).

Shoot elongation

The explants induced in the multiplication phase were cultivated for six weeks in shoot elongation medium. This medium was half strength ARN (Arnaud et al.) supplemented with 2 gL⁻¹ activated charcoal and solidified with 9 gL⁻¹ Difco Agar®. The pH of the medium was adjusted to 5.8 before autoclaving (121°C, 20 min).

Root induction and acclimatization of plants

After the elongation phase, stems with at least 2.5 cm in length were used for root induction. Traceability of the explant was maintained according to its origin (genotype and cytokinin treatment used along the shoot induction stage). Subsequently, the stems were grown in root inducing medium (RIM), which consisted of 1/3 strength ARN basal medium supplemented with three different auxin treatments: A) 50 µM 1-naphthalene acetic acid (NAA), B) 50 µM indole-3-butyric acid (IBA), and C) a mixture of 40 µM NAA µM + 10 µM IBA. All the different media were solidified with 9 gL⁻¹ Difco Agar®. The culture medium’s pH was adjusted to 5.8 before autoclaving (121°C, 20 min).

The stems were placed in the dark at 21 ± 1°C for 8 days. After this period, the stems were cultured in the root expression medium (REM), which consisted of 1/3 strength ARN basal medium (Arnaud et al.) without plant growth regulators and solidified with 9 gL⁻¹ Difco Agar®. The pH of the medium was adjusted to 5.8 before autoclaving (121°C, 20 min). The cultures were placed in the growth chamber at a temperature of 21 ± 1°C, under 16-h photoperiod at 120 µmol m⁻² s⁻¹. After six weeks of culture in REM medium, explants with visible roots were transferred to wet peat: vermiculite mixture (2:1, v/v) and acclimatized in the greenhouse under controlled conditions at 21 ± 1°C and decreasing humidity progressively along one month from 95 to 80%.

Data collection and statistical analysis

To assess the genotype’s effect on each of the variables of this study, an analysis of variance was performed (ANOVA), followed by a Tukey’s post hoc test (α=0.05). The genotype had a significant effect on all variables studied. For this reason and to obtain robust conclusions, the genotype factor was introduced into all the models as a block variable to reduce variability and analyze the effect of the other variable factors (the type of explant, cytokinin treatment, and auxin treatment) more accurately.

The shoot induction and the number of shoots per explant were recorded after six weeks. The experiment used a completely randomized design with 25 explants per treatment. Logistic regression was performed to assess the effect of the type of explant and the cytokinin treatment on the shoot induction percentage, followed by a Tukey’s post hoc test (α=0.05) for multiple comparisons. The number of shoots per explant variable did not fulfill homoscedasticity and normality assumptions for ANOVA; then, a Kruskal-Wallis was performed.

The root induction percentage, the number of roots per plant, and the root length were recorded after six weeks in REM. A completely randomized design was carried out using ten stems per cytokinin treatment. As an exception, in genotype 7, four stems were employed when K was used in the induction stage.

A logistic regression model was used to analyze the auxin and cytokinin treatment’s effect on the root induction percentage. Tukey’s post hoc test (α=0.05) was used for multiple comparisons. The number of roots per explant variable did not fulfill homoscedasticity and normality assumptions for ANOVA; then, a Kruskal-Wallis was performed.

Data on the length of the longest root were square-root transformed, and an ANOVA was performed; multiple comparisons were made using Tukey’s post hoc test (α=0.05). After twelve weeks under ex vitro conditions, the survival percentage was calculated. The survival percentage variable did not fulfill homoscedasticity and normality assumptions for ANOVA; then a Kruskal-Wallis was performed.

Results

Shoot induction

When the genotype effect on the shoot induction percentage was evaluated, genotypes 12 and 11 showed significantly higher percentages (98.98%) than genotype 7 (79%, Table 1). As shown in Table 1, the number of shoots obtained in genotype 12 (5.04) was significantly higher than those obtained in genotypes 11 and 7 (3.21 and 3.28, respectively).

The deviance analysis of the factors studied showed a significant effect on both the shoot induction and the number of shoots per explant developed from each explant and cytokinin type.

There was a significant interaction between the explant type and the cytokinin type (Table 2).

<table>
<thead>
<tr>
<th>Variable</th>
<th>Genotype</th>
<th>Shoot induction (%)</th>
<th>N^3 shoots /explant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7</td>
<td>11</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>79.00±0.03 b</td>
<td>98.98±0.01 a</td>
<td>98.98±0.01 a</td>
</tr>
<tr>
<td></td>
<td>3.28±0.12 b</td>
<td>3.21±0.07 b</td>
<td>5.04±0.16 a</td>
</tr>
</tbody>
</table>

Table 1. Shoot induction (%) and number shoots per explant from Sequoia sempervirens elite adult trees cultured in ARN medium (Arnaud et al.)

Data are presented as mean values ± S.E. Different letters within a row indicate significant differences (p<0.05).
Significantly higher shoot induction percentages were obtained in AP, BG and BP explants (between 91% and 94%) than in AG explants (84%) (Table 3).

Since AG explants showed the best appearance, a comparative figure was done to describe the aspect of regenerated shoots in cytokinins’ presence (Figure 3). The most vigorous shoots were observed in the presence of BAP and K (Figure 3 (A-C, G-H)).

Root induction

A significantly higher root induction percentage was obtained in genotype 12 (93%) (Table 5). The lowest response was genotype 7 (31%), and intermediate values were recorded for genotype 11 (55%). Also, a significantly higher number of roots per explant was observed in genotype 12 (4.73) than genotype 7 and 11 (Table 5).

On the contrary, significantly longer primary roots were recorded for genotype 11 (3.73 cm, Figure 4A) and genotype 7 (2.95 cm, Figure 4B) than for genotype 12 (2.18 cm, Figure 4C).

The auxin type and cytokinin type used in the multiplication stage showed a significant effect on the root induction percentage. Further, the cytokinin type showed a significant effect on the longest root’s length (Table 6). Regarding the number of roots per explant, no significant differences were found for the variables tested.

As shown in Table 7, a significantly higher root induction percentage was obtained in the treatment with NAA (70%) than in treatments with IBA or IBA/NAA mixture. As mentioned before, there was no effect of the auxin treatment on the number of roots per explant (ranging from 2.98 to 3.94) or on the length of the primary root (ranging from 2.23 to 3.09 cm) (Table 7).

When evaluating the effect of the cytokinin type over root induction percentage, significantly higher values were obtained from shoots induced in BAP treatment when compared with those induced in m-T and K treatment (Table 8).

As aforementioned, no significant differences were found for the number of roots per explant, independently of the shoot induction treatment applied (ranging from 3.11 to 3.88) (Table 8).

The longest primary roots (3.32 cm) showed significantly higher values when shoots were induced in BAP treatment (Table 8).

Acclimatization of rooted microplants

The analysis of deviance for survival (%) of rooted shoots propagated in vitro showed a significant effect of the auxin type applied in the root induction stage and of the cytokinin type applied in the shoot induction phase (Table 9).

When the effect of auxin type on the survival percentage of rooted shoots was analyzed, the presence of IBA in the rooting medium provoked a significantly higher survival percentage (94%) than the mixture IBA/NAA and NAA treatments (Table 10).
Figure 1. Number of shoots per explant in different explant types from *Sequoia sempervirens* elite adult trees cultured in ARN medium (Arnaud et al. 2010) supplemented with different cytokinins (BAP, m-T, K), (apical explants of 1.5 cm in length (AG), apical explants of 1.0 cm length (AP), basal explants of 1.5 cm length (BG) and basal explants of 1.0 cm in length (BP)). Data are presented as mean values ± S.E. Different letters indicate significant differences (p<0.05).

Figure 2. Shoot induction from *Sequoia sempervirens* elite adult trees cultured in ARN multiplication medium (Arnaud et al. 2010) cultured for 6 weeks: (A) apical explant > 1.5 cm of length (AG) of genotype 7 on ARN medium + 4.4 µM BAP, bar = 3 mm; (B) basal explant > 1.5 cm of length (BG) of genotype 7 on ARN medium + 4.4 µM m-T, bar = 5 mm; (C) apical explant < 1.5 cm of length (AP) explant of genotype 7 on ARN medium + 4.4 µM m-T, bar = 3 mm.

Table 5. Root induction (%), number roots per explant, and the longest root from *Sequoia sempervirens* elite adult trees cultured in ARN medium (Arnaud et al. 2010). Data are presented as mean values ± S.E. Different letters within a row indicate significant differences (p<0.05).
Figure 3. Shoot induction from *Sequoia sempervirens* elite adult trees cultured in ARN multiplication medium (Arnaud et al. 10) cultured for 6 weeks: (A) apical explant > 1.5 cm of length (AG) of genotype 7 on ARN medium + 4.4 µM BAP, bar = 3 mm; (B) AG of genotype 11 on ARN medium + 4.4 µM BAP, bar = 3 mm; (C) AG explant of genotype 12 on ARN medium + 4.4 µM BAP, bar = 3 mm; (D) AG explant of genotype 7 on ARN medium + 4.4 µM m-T, bar = 5 mm; (E) AG explant > 1.5 of genotype 11 on ARN + 4.4 µM m-T, bar = 3 mm; (F) AG explant of genotype 12 on ARN medium + 4.4 µM m-T, bar = 5 mm; (G) AG of genotype 7 on ARN medium + 4.4 µM K, bar = 3 mm; (H) AG explant of genotype 11 on ARN medium + 4.4 µM K, bar = 5 mm; (I) AG of genotype 12 on ARN medium + 4.4 µM KIN, bar = 3 mm.

Table 6. Analysis of deviance for root induction (%), number roots per explant, and length of longest root from *Sequoia sempervirens* elite adult trees cultured in ARN medium (Arnaud et al. 10).

*Significant differences at p<0.05, n.s Non-significant at p<0.05, df Degrees of freedom.

Table 7. Root induction (%), number of roots per explant and length of the longest root in *Sequoia sempervirens* elite adult trees cultured in ARN medium (Arnaud et al. 10) supplemented with different auxin types (IBA, IBA/NAA or NAA). Data are presented as mean values ± S.E. Different letters within a row indicate significant differences (p<0.05). *ns Non-significant.
Figure 4. Root induction from *Sequoia sempervirens* elite adult trees micro shoots cultivated *in vitro* for 6 weeks on root expression medium (REM) (Arnaud et al.10): (A) explant of genotype 7 induced in root inducing medium (RIM) (Arnaud et al.10) + 50 µM NAA (BAP applied in shoot induction stage), bar = 2 cm (B) explant of genotype 11 induced in RIM medium + 50 µM NAA (BAP applied in shoot induction stage), bar = 2 cm; (C) explant of genotype 12 induced in RIM medium + 50 µM NAA (BAP applied in shoot induction stage), bar = 2 cm.

Table 8. Root induction (%), number roots per explant, and length of longest root in different cytokinin types from *Sequoia sempervirens* elite adult trees cultured in ARN medium (Arnaud et al.10). Data are presented as mean values ± S.E. Different letters within a row indicate significant differences (p<0.05). n.s Non-significant.

<table>
<thead>
<tr>
<th>Cytokinin type applied in the root induction stage</th>
<th>BAP</th>
<th>m-T</th>
<th>K</th>
</tr>
</thead>
<tbody>
<tr>
<td>Root induction (%)</td>
<td>66.00±0.04&lt;sup&gt;a&lt;/sup&gt;</td>
<td>58.00±0.04&lt;sup&gt;b&lt;/sup&gt;</td>
<td>62.00±0.04&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>N° roots/explant</td>
<td>3.11±0.24&lt;sup&gt;n.s&lt;/sup&gt;</td>
<td>3.89±0.33&lt;sup&gt;n.s&lt;/sup&gt;</td>
<td>3.16±0.29&lt;sup&gt;n.s&lt;/sup&gt;</td>
</tr>
<tr>
<td>Lenght of longest root (cm)</td>
<td>3.32±0.22&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.39±0.21&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.48±0.24&lt;sup&gt;b&lt;/sup&gt;</td>
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</table>

Table 9. Analysis of deviance for survival (%) of rooted shoots propagated *in vitro* coming from *Sequoia sempervirens* elite adult trees, after twelve weeks under *ex vitro* conditions.
*Significant differences at p<0.05, n.s Non-significant at p>0.05, df Degrees of freedom.

<table>
<thead>
<tr>
<th>Source</th>
<th>df</th>
<th>Survival (%)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>X² Test</td>
</tr>
<tr>
<td>Auxin type (A)</td>
<td>2</td>
<td>244.40</td>
</tr>
<tr>
<td>Cytokinin type (C)</td>
<td>2</td>
<td>223.62</td>
</tr>
<tr>
<td>A x C</td>
<td>4</td>
<td>216.66</td>
</tr>
</tbody>
</table>

Table 10. *Ex Vitro* survival (%) of in vitro rooted shoots from *Sequoia sempervirens* elite adult trees, shoots were rooted in ARN medium (Arnaud et al.10) supplemented with different auxins (IBA, IBA/NAA, or NAA). Data are presented as mean values ± S.E. Different letters indicate significant differences (p<0.05).

<table>
<thead>
<tr>
<th>Auxin type</th>
<th>IBA</th>
<th>IBA/NAA</th>
<th>NAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Survival %</td>
<td>94.00±0.03&lt;sup&gt;a&lt;/sup&gt;</td>
<td>79.00±0.06&lt;sup&gt;b&lt;/sup&gt;</td>
<td>68.00±0.05&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>
Regarding the effect of the cytokinin applied in the shoot induction stage as shown in Table 11, the shoots induced in K treatment showed a significantly higher ex vitro survival percentage than those induced in BAP or m-T treatments.

At the end of the experiment, the micro propagated plants from the three elite adult trees evaluated showed normal development, profitable growth, and uniformity after 5 months under ex vitro conditions in the greenhouse (Figure 5).

**Discussion**

The genotype is an endogenous factor that has a significant role in regenerative potential in the overall repeatability and reliability of tissue culture protocol\(^{13}\). In this study, the shoot induction, the number of shoots per explant, the root induction, and the number of roots per explant were affected by the explant’s genotype. Our results are in concordance with the results reported by Meneguzzi \(et\ al\).\(^2\), who observed different genotypes’ responses in shoot multiplication of \(S.\) sempervirens. Similarly, Sul and Korban\(^{14}\) found differences in shoot proliferation and shoot elongation of \(S.\) sempervirens.

The organogenesis processes are influenced by endogenous and exogenous factors\(^{14,15,16}\). There are different exogenous factors as the components in the tissue culture medium or the culture’s environmental conditions. In this work, we observed that the type of explant and chemical conditions in the culture medium (cytokinin and auxin type) significantly affected shoot induction and rooting response.

During the evaluation of the shoot induction rate, the explants with the most extended length (AG, AP and BG) showed the best induction rate, and the highest number of shoots per explant were obtained. Similar results were found by Clapa \(et\ al\).\(^{17}\) and Meneguzzi\(^{18}\), who developed an in vitro protocol for \(S.\) sempervirens using shoot explants bigger than 1.5 cm in length. George\(^{16}\) reported that larger explants coming from more extensive parts of shoot apex or stem segments bearing one or more lateral buds could show advantages over smaller size explants.

Several studies about tissue culture protocols have reported positive effects on shoot proliferation, shoot multiplication rate, alleviating physiological disorders, better rooting, and acclimatization when using toplins for shoot induction\(^{15,19}\). In our study, the explants induced in the presence of m-T exhibited the best results. In this sense, De Diego \(et\ al\).\(^{20}\), obtained a high rate of organogenic response in adult buds of \(P.\) sylvestris using m-T and suggested that it could be used as an alternative cytokinin BAP in micropropagation. Likewise, Bairu \(et\ al\).\(^{21}\) obtained outstanding multiplication rates in \(A.\) polyphylla when used the same cytokinin. However, in the present study, these explants from m-T treatments were more

**Table 11.** Ex vitro survival (%) of in vitro rooted shoots from \(S.\) sempervirens elite adult trees; shoots were propagated in ARN medium (Arnaud \(et\ al\).\(^{10}\)) supplemented with different cytokinins (BAP, m-T or K). Data are presented as mean values ± S.E. Different letters indicate significant differences by Tukey’s post hoc test (\(p<0.05\)).

<table>
<thead>
<tr>
<th>Cytokinin type</th>
<th>BAP</th>
<th>m-T</th>
<th>K</th>
</tr>
</thead>
<tbody>
<tr>
<td>Survival (%)</td>
<td>80.00±0.05(^b)</td>
<td>65.00±0.05(^b)</td>
<td>94.00±0.03(^a)</td>
</tr>
</tbody>
</table>

**Figure 5.** Plants of \(S.\) sempervirens cultured in vitro after 5 months growing in ex vitro conditions in the greenhouse, bar = 4 cm.
yellowish and less vigorous. In contrast, in Prunus rootstocks, *Pterocarpus marsupium*, and *Corylus colurna* a positive effect of m-T on the growth and quality of micro propagated shoots was found [22,23,24].

According to Sul and Korban [14], Meneguzzi et al. [15], Valverde et al. [16] and Montalbán et al. [17] in forestry species, BAP stimulated the axillary bud breakage and shoot elongation. In our study, a similar result was obtained in the BAP treatment. It is essential to mention that in the treatment with BAP the quality of the shoots formed (robustness and color) was superior to those shoots generated in the presence of m-T. In conclusion, in our study, BAP was the best cytokinin in shoot induction and the number of shoots per explant. This is in accordance with the results reported by Moncaleán et al. [18], Reau et al. [19], Aremu et al. [20], and Bairu et al. [21], confirming that BAP is the most used cytokinin in micropropagation due to its effectiveness and affordability. Regarding K treatment, in this work, we obtained the worst results in agreement with those found in *Barleria greenii* and *Eucalyptus globulus* [22,23]. It is essential to mention, that the rooting percentages obtained in the present studies were higher than those recorded by Huang et al. [24], who obtained about a 30% of rooting competence using adult stem sections of *Sequoia sempervirens* in the presence of IAA/K. The number of roots per explant obtained in the IBA/NAA treatment presented the highest values. In species such as *Eucalyptus sideroxylon*, *Rosa hybrida* and *Citrus aurantium* the best root induction was obtained when using a mixture of IBA/NAA. Nonetheless, this treatment did not lead to the highest values in the number of roots per explant [25,26,27]. The highest value in the longest root length was also obtained in NAA treatment, being under the results obtained in *Citrus aurantium* [28].

In our research, the induction stage’s cytokinin type had a significant effect on root induction. Bairu et al. [21], Werbroock et al. [22], and Aremu et al. [23] found that m-T stimulated in vitro rooting activity, but the m-T treatment presented the worst in our work results. In this sense, Bairu et al. [21] and Escalona et al. [24] obtained negative carryover effects on rooting at too high m-T levels, so the m-T concentration applied in our cultures might have had a detrimental effect in the subsequent stages of development.

Regarding the acclimatization stage, the highest survival percentage was observed in plantlets rooted in IBA and developed in the presence of K. This result is in agreement with those found in *Eucalyptus globulus*, where the low concentration of IBA showed the best survival rates. In *Arachis paraguariensis* cultured in polyethylene terephthalate glycol vessel where IBA treatment was the best in survival [25,26].

Aremu et al. [27] explained that cytokinins generally have inhibitory effects on rooting, resulting in low acclimatization rates afterward. In this sense, the plants from m-T treatment showed the lowest survival. In contrast, several studies have shown that plantlets coming from m-T induction treatments have been successfully acclimatized [28,29].

In conclusion, the role of auxins and cytokinins in the micropropagation of different types of explants and their relationship with the survival and acclimatization of seedlings in ex vitro conditions was analyzed. Our study results demonstrated that the apical explant 1.5 cm length, and BAP showed the best results in the shoot induction stage. Moreover, in vitro shoots rooted with IBA, led to a higher ex vitro survival. Finally, the results shown allow the development of forthcoming studies for large-scale propagation of this species in semi-solid systems and bioreactors.
Optimization of the micropropagation of elite adult trees of Sequoia sempervirens: forest species of interest in the Basque Country, Spain


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