**Influence of in vitro Micropropagation Growth Conditions on Stomatal and Morphological Characteristics of Mature *Pistacia vera* L.**

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**ABSTRACT:** This research was conducted to reveal the stomatal anatomy, stomatal index and water loss (%) of mature pistachio leaves as well as the leaves of different phases (multiplication, rooting, hardening and regenerated plant) of micropropagation of mature pistachio trees obtained from the *in vitro*. Microscopic observations on surfaces of these leaves showed variety from elliptical to ovate stomata with length of 0.81-2.02 µm and width of 1.58-3.80 µm. An increase in stomatal index (SI) in the leaves of plants grown in vitro was observed most specifically in the hardening phase. (17.49±0.04). The stomatal index declined in the leaves of plantlets transferred to *in vivo* conditions subsequent to the hardening phase. In order to measure water loss, leaves obtained from all types of samples were dried in the oven between 30 minutes and 2 hours and weighed. The percent water loss of *in vitro* leaves of multiplication phase was greater than the other phases. The stomatal differentiation was found to be influenced by the different hardening regimes applied. Hardening by covering the pots with polyethylene bags improved the survival rate. This study indicates that optimization of *in vitro* micropropagation stages is necessary to avoid transplantation stress.

**Keywords:** Hardening, *in vitro* culture, stomatal characteristics, water loss, stomatal index, *P. vera* L.

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**Olgun Antepfıstığı’nın (*Pistacia vera L.*) Stomatal ve Morfolojik Özellikleri Üzerine In vitro Mikropropagasyon Büyüme Koşullarını Etkisi**

Özet: Bu çalışma, olgun antepfıstığı ağaçlarının *in vitro* olarak mikroçoğaltılmının farklı evrelerinden (çoğaltma, köklendirme, alıştırma ve rejenere bitki) elde edilen yaprakların stomal anatomisini, stoma indeksini ve su kaybını (%) ortaya çıkarmak için yapılmıştır. Yaprak yüzeyinde yapılan mikroskobik gözlem, 0.81-2.02 µm uzunlukunda ve 1.58-3.80 µm genişliğinde eliptiktan ovat stomaya kadar çeşitlilik göstermiştir. *In vitro* yetiştirilen bitkilerin özellikle alıştırma aşamasında yapraklarında stoma indeksinde (SI) bir artış göstermiştir (17.49±0.04). Alıştırma aşamasından sonra *in vivo* koşullara transfer edilen bitki yapraklarının stoma indeksi azalmıştır. Su kaybını ölçmek için, her çeşit numunenin elde edilen yapraklar, 30 dakika ile 2 saat arasında fırında kurutularak tartışmıştır. Çoğaltma aşamasındaki su kaybı yüzdesinin diğer aşamalara göre daha büyük olduğu tespit edilmiştir. Stoma farklılaşmasının, uygulanan doğal şartlara aktarım metodu tarafından da etkileşimi tespit edilmiştir. Saksıların polietilen poşetlerle kapatılması yoluya gerçekleştilen aklimatizasyon yönteminin yaşam oranını arttırdığı tespit edilmiştir. Bu çalışma, transplantasyon stresinden kaçınmak için *in vitro* mikro-coğaltma aşamlarının optimizasyonunun gerekli olduğunu göstermektedir.

**Anahtar Kelimeler:** Alıştırma, *in vitro* kültür, stoma özellikleri, su kaybı, stoma indeksi, *P. vera* L.
INTRODUCTION

Micropropagation that cause rapid growth and multiplication of shoots under *in vitro* conditions often lead to the formation of plantlets with abnormal morphology, anatomy and physiology (Mohamed and Alsadon, 2010). Therefore, a significant amount of micropropagated pistachio plantlets could be lost or harmed especially during the period of acclimatization to *ex vitro* conditions (Zobayed et al., 2001). The main parameters of these controlled culture conditions include constant temperature, high relative humidity (RH), low photosynthetic photon flux (PPF), optimized concentrations of salts, carbohydrates and plant growth regulators (Kubato et al., 1997). Among them incubation of plantlets at high RH could be considered to be the most important parameter as it could cause high mortality rate during *ex vitro* transfer mainly due to the excessive water loss attributed to reduced leaf epicuticular wax (Sutter and Langhans, 1979) high stomatal densities (Desjardins, 1988) and poor stomatal functioning (Blanke and Belcher, 1989). The reason for the poor functioning of stomata could be due to the guard cell wall characteristics (Ziv et al., 1987), deformation of stomata (Blanke and Belcher, 1989) or K⁺ overflow through the guard cells (Assmann, 1993).

Although several improvements were made for the initiation, multiplication and rooting of pistachio, up to now, only one study was reported on stoma morphology of micropropagated pistachio (Barghchi, 1982; Bustamante-Garcia, 1984; Al Ramadhani, 1985; Martinelli, 1988; Abousalim, 1990; Gonzales and Frutos, 1990; Yang and Ludders, 1993; Onay, 1996; Onay et al., 1997; Onay, 2000; Ozden-Tokatlı et al., 2005; Tilkat et al., 2008; Tilkat and Onay, 2009; Tilkat et al., 2009). However, a detailed study on stomatal anatomy, stomatal index and water loss (%) should be carried out in micropropagated pistachio (Namlı and Ayaz, 2007) plantlets to reveal out the status and function of stoma during different phases of micropropagation. Besides, the transfer of plantlets from *in vitro* culture containers to air conditions is still considered to be the one of the most challenging steps of some woody species together with pistachio (Chandra et al., 2001; Joshi et al., 2006; Mišalová et al., 2009; Balakrishnan et al., 2009; Moyo et al., 2012) as well as pistachio. Thus, the purpose of this study was to investigate stomatal and morphological characteristics of *P. vera* L. leaves during different stages of micropropagation and to develop a method for successful hardening of pistachio plantlets during their rapid propagation.

MATERIALS AND METHODS

Establishment of *In vitro* Culture Conditions

Vigorous new shoots were collected from adult *Pistacia vera* L. trees (25-year-old) at Pistachio Research Institute in Gaziantep, South-east of Turkey, during the month of April, for the experiment. Shoot cultures, were started from aseptically grown adult trees of *P. vera* L. according to the method defined by (Tilkat, 2006). Adventitious buds from the initiation medium were repeatedly subcultured every 3 weeks on a MS (Murashige and Skoog, 1962) media supplemented with 1.0 mg L⁻¹ BA (6-benzyladenine). This culture was preserved for more than three years. Long shoots (> 3 cm long) collected after the third subculture were used for *in vitro* rooting. Shoots longer than 3 cm were transported to a rooting media consisting of MS media with 5.5 g L⁻¹ agar, 30 g L⁻¹ sucrose and 2.0 mg L⁻¹ IBA (indole-3-butyric acid). Cultures were maintained at 25 ± 2 °C, with 40 μmol m⁻² s⁻¹ for 16 h photoperiod and 60 - 70 % relative humidity.

Applied Hardening Conditions

*In vitro* regenerated plantlets were subjected to the following mode of hardening: 1) Rooted microplants were directly transported to *ex vitro* conditions. 2) The plants in pots were covered with
polyethylene bags for 20 d followed by gradual opening and complete removal after the next 20 d 3) Rooted microplants were hardened in vitro in a culture coated with pyrex glasses containing autoclaved peat and perlite moistened with ½ MS nutrients after the 14th day, followed by gradual opening and complete removal after the next 40 days. All those operations were carried out in a growth chamber where the temperature was 25 ± 2°C and the relative humidity was decreased from 90% to 50% by using an evaporative cooling system.

**Stomatal Analyses**

When the plants were completely weaned (4 to 6 weeks), they were exposed to greenhouse conditions. For peeling preparation, leaves were collected and used from the top node at different stages of micropropagation (multiplication, rooting, hardening, regenerated plant). Controls included leaves obtained from field-grown trees of *P. vera* L. (25 years old). In order to determine the stomata characteristics, together with control, five plantlets from each phase, twenty leaves from each plantlets and ten peels from each leaf were examined. In the case of the stomatal index (the number of stomata per unit area), the length of the stomata, the width of the stomata, the epidermis of the ventral parts were all peeled and placed on a glass slide, mounted with distilled water and covered with a cover slip. Prepared slides were observed under an inverted microscope. The stomatal index (SI) was determined on the leaf lamina zones, using the following formula of Salisbury, 1927:

\[
SI = \frac{S}{(E+S)} \times 100,
\]

where S is the number of stomata per unit leaf area and E is the number of epidermal cells per unit leaf area.

The size of the stomata was measured with the aid of an ocular micrometer, pre-calibrated with a stage micrometer. The shape and the size of the leaves (width and length) were measured and compared. The morphological characteristics of the leaves obtained during different phases of micropropagation were also reported. Moreover, leaf colours were identified as either light green (translucence) or dark green (opaque).

**Water Loss Analyses**

Almost, in order to determine the amount of water loss, fresh weight of leaves with similar size (two out of five plants) were weighed to determine water loss (Table 4). Leaves were allowed to transpire by keeping their abaxial edges, on clean bench at room temperature of 23-27 °C and relative humidity of 35-40 %. Each leaf was weighed at 30 min. interval for 2 h. Lastly, all tested leaves were dried at 60 °C to obtain dry mass. Water loss was calculated as defined by Brainerd and Fuchigami, 1981:

Water loss (%) = \([(\text{Initial Fresh Mass–Mass After Holding})/(\text{Fresh Mass–Dry Mass})] \times 100\]

**Statistical Analyses**

For all treatments were conducted using a completely randomized block design was used. Significance was determined by an analysis of variance, and the least significant (P≤0.05) differences among mean values were estimated using Duncan’s New Multiple Range test. Data presented in percentages were subjected to chi-square (χ²) analysis. Analysis of variance, was used to find differences between treatments in comparing stomatal density, diameter and length. Duncan Multiple Range Test, was used to differentiate mean values and significant effects were accepted at p≤0.05. Data calculations were performed using SPSS for Windows (version 15.0, SPSS®; Chicago, USA).
RESULTS AND DISCUSSION

Morphological Characteristics

Morphology of the leaves changed in shape (width x length) and colour as shown in Figure 1. A-E and Table 1. The leaves developed in the multiplication phase were a light green and elliptic shapes, whereas leaves developed rooting phase had greenish yellow colour and ovate or elliptic shapes. Moreover the leaves of *in vitro* hardening phase with elliptic-ovate shapes with dark green and the leaves of regenerated plantlets had round-ovate shapes with dark green colour. In case of the control (field-grown-tree) leaves broad-ovate shapes these were thin and with light green or green colour and elliptic or ovate shape. The shape measured as the width and the length of the leaves was also altered in the micropropagation phases tested; these had greater length than width. The width and length of mature tree leaves were significantly higher than the leaves obtained from different stages of micropropagation. However, leaves taken from all tested phases had greater length than width, similar to the mature tree.

As a result of this study, it was determined that *in vitro* differentiation of leaf morphology and stoma were affected by micropropagation phases and the type of hardening. This confirms the previous data on stomatal morphology reported on different plant species (Marin, 1988; Zacchini, 1997). Thus, epidermal peels of different phases of micropropagation appear to be different with decreasing stomatal index. This trend was particularly marked with the hardening type tested. Leaf shape and color deviations was apparent at different micropropagation stages. The reversibility, differentiated between the different phases induced by the above-mentioned treatment (Table 1), was different in the leaves investigated *in vitro* at different stages and appears to exclude/include the possibility that the stomatal reaction, which was obscured at such high percentages, could be due to completely physical and chemical factors. Hyperhydricity caused translucent thin and brittle leaves including also malformation of stoma. This phenomenon is also a serious problem during *in vitro* culture of mangosteen (Te chato et al., 2005).

Table 1: Morphological characteristics of leaves obtained in the course of different phases of micropropagation.

| Source of Leaf | Shape of Leaf | Colour           | Width (cm) (±SD)* | Length (cm) (±SD)* |
|----------------|---------------|------------------|-------------------|-------------------|
| Multiplication Phase | Elliptic     | Light Green      | 0.81±0.53        | 1.58±0.12        |
| Rooting Phase      | Elliptic-Ovate | Greenish Yellow  | 0.83±0.36        | 1.58±0.10        |
| *In vitro* Hardening | Elliptic-Ovate       | Dark Green       | 0.85±0.08        | 1.65±0.30        |
| Regenerated Plantlets (1 year old) | Round Ovate | Dark Green       | 1.56±0.13        | 2.75±0.20        |
| Mature Tree (30 years-old) | Ovate or Broad Lanceolate | Dark Green | 2.02±0.36        | 3.80±0.30        |

Stomatal Characteristics

According to our analysis, the epidermal cells on the leaves obtained from the cultures at the end of the replication phase were polygonal, having no curved fluctuations in the anticline walls. (Fig. 1F). The stomata of the leaves obtained from the cultures of this phase were either wide open, or with equal guard cells. These stoma had a rounded morphology with more porous area. Such irregularities, were also common during the rooting and hardening phase (Fig. 1G -1H). Irregularly shaped epidermal cells (with or without fluctuations) were observed in the micropropagation phase while curved fluctuations were detected during the rooting phase in pistachio.(Fig. G). The fact remains that, stomata in leaves collected from the regenerated plantlet (Fig. 1I) and mature tree of pistachio (Fig. 1K) were anomocytic, elliptical (different from kidney formed guard cells) and isodiametric epidermal cells. The significantly highest stomatal index (17.49) was obtained in leaves collected from *in vitro* hardening phase while the lowest index was found in leaves of regenerated plantlets and mature tree (10.85 and 10.96, 802
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respectively). It was observed that the stomata in the leaves obtained from all stages of the micropropagation were similar in length and width, but the multiplication phase was different from them (Table 2). When the stomata in the leaves obtained from the multiplication phase were compared with the mature pistachio tree, the stomata in the multiplication phase were almost round and spherical porous (20.27 μm length and 18.19 μm width), while the stomata of the mature pistachio tree were elliptical (22.64 μm length and 17.40 μm width). The stomatal density was highest in leaves from hardening cultures (17.49). The values obtained in other micro-propagation stages were determined in multiplication phase (13.31), rooting phase (11.35), regenerated plants (10.85) and mature tree (10.96), respectively.

The stomatal index during the \textit{in vitro} hardening phase was almost twice of regenerated plantlets and the mature tree. This differential behaviour could be correlated with the percentage of malformed stomata. Stomatal differentiation was found to be influenced by the different hardening regimes applied (Table 3). The highest value of stomatal index was induced by transferring the plantlet to pots covered with polyethylene bags. Relatively lower stomatal index was obtained in leaves collected from pots covered with pyrex glasses while data could not be taken from leaves of direct transfer, those plantlets could not survive beyond a few hours. Although the length of the stomata was higher in leaves collected from pots covered with polyethylene bags, the width of the stomata was relatively lower in this treatment than stomata on leaves of pots covered with pyrex glasses. Significant differences have emerged in the successful acclimatisation of plantlets between the different types of hardening (Table 3). The highest density of acclimatised plantlets was 80 % in pots covered with polyethylene bags and 90 % of these plants survived when transferred to soil. Pots covered with pyrex glasses also resulted in high frequency
of acclimatized plantlets (70 %). 90 % of viable regenerants was also obtained after 6 months transfer to growth room conditions.

Table 2. Stomatal characteristics of *Pistacia vera* L. leaves in the course of different phases of micropropagation (Means ± SE, n = 5)

| Source of Leaf                     | Stomatal Index (%)* | Length of Stomata [μm]* | Width of Stomata [μm]* |
|-----------------------------------|---------------------|--------------------------|------------------------|
| Multiplication Phase              | 13.31 ± 0.02b       | 20.27 ± 0.08d            | 18.19 ± 0.06a          |
| Rooting Phase                     | 11.35 ± 0.04c       | 23.52 ± 0.08b            | 15.42 ± 0.06d          |
| In vitro Hardened                 | 17.49 ± 0.04a       | 24.70 ± 0.05e            | 15.28 ± 0.06d          |
| Regenerated Plantlets             | 10.96 ± 0.06d       | 24.18 ± 0.07e            | 16.92 ± 0.04c          |
| Mature Tree                       | 10.85 ± 0.04d       | 22.64 ± 0.04a            | 17.40 ± 0.06b          |

* Different lower-case letters in each column indicate that these values are statistically different at P ≤ 0.05 according to Duncan’s Multiple Range Test.

Table 3. Effect of hardening type on stomatal characteristics and viable regenerants (%) of *P. vera* L. leaves.

| Type of Hardening                        | Stomatal Index* | Length of Stomata [μm]* | Width of stoma [μm]* | % of Acclimatized Regenerant** | Viable Regenerants (%) After Six Months Transfer to Growth Room*** |
|------------------------------------------|----------------|--------------------------|----------------------|-----------------------------|------------------------------------------------------------------|
| Pots covered with Polyethylene Bags     | 23.32 ± 0.07a  | 23.16 ± 0.08a            | 14.42 ± 0.04b        | 80                          | 90                                                               |
| Pots Coated with Pyrex Glasses          | 21.78 ± 0.08b  | 20.25 ± 0.02c            | 17.24 ± 0.04a        | 70                          | 90                                                               |
| Direct Transfer                          | 0.00 ± 0.00c   | 0.00 ± 0.00c             | 0.00 ± 0.00c         | 0                           | 0                                                                |
|χ² (df)                                   |                |                          |                      | P ≤ 0.05                    | P ≤ 0.05                                                         |

* Different lower-case letters in each column indicate that these values are statistically different at P ≤ 0.05 according to Duncan’s Multiple Range Test.

**Data were recorded on day- 30 of acclimatization and represent the means of 10 explants per treatment with 2 repetition of the experiment.

*** Data were recorded 6 months after transfer to a growth room.

There may be many causes of decreased stoma function over time *in vitro*. Probably as a result of high relative humidity, *in vitro* plantlets cellulose deposits seem to be reduced, compared to *in vivo* conditions (Marin, 1988).

According to this evidence, the earliest leaves, in our experiments also could have a lower stomata index since they were submitted for a longer period of time to high relative humidity. Similar results have also been reported by (Zacchini et al., 1997). When the *Wrightia tomentosa* cultures were transferred to a low relative humidity level, no deformation of the stoma morphology was recorded in the leaves of *in vivo* hardened plants.

Our results strongly suggest that the stomatal index is affected by the different phases of micropropagation; index, width and length of the stomata are also influenced by the hardening type used. (Table 3.) Therefore, the results of this study indicate that enhancement of the micropropagation phases *in vitro* is necessary to avoid transplantation stress. This would then lead to a higher survival rate and a greater yield. A relationship was observed between the stoma index and different stages of micropropagation. However, the gradual change in stomata shape from elliptical in the early phases (multiplication and rooting) to circular in mature leaves is noteworthy.

**Water Loss**

The maximum water loss (69.70 %) was observed in the leaves from multiplication phase within the first 30 min. (Table 4). In the same period, the water losses was 65.62 % for leaves of rooted
microplants, 61.80 % for in vitro hardened plants, 34.50 % for regenerated plants, 13.95 % for the leaves of the mature tree, respectively.

Except the mature tree in other all cases, the percentage of water loss decreased with time. Within 2 hours, maximum water loss was observed in multiplication phase at the rate of 97.23 %. When the seedlings were gradually exposed to in vivo conditions, water loss control have increased with time. In vitro study of Actinidia deliciosa plantlets found that the water loss value gradually decreased (Moncalean et al., 2009). This result is similar to our study.

Table 4. Water loss (%) from decomposed leaves of plantlets in the course of different phases of micropropagation in Pistacia vera L. (Means ± SE of 5 plants).

| Time (min) | Multiplication* | Rooting* | In vitro Hardened Plants* | Regenerated Plantlets* | Mature Tree* |
|-----------|-----------------|----------|--------------------------|------------------------|-------------|
| 0- 30     | 69.70 ± 1.85a   | 65.62 ± 0.28a | 61.80 ± 0.67a          | 34.50 ± 0.25a          | 13.95 ± 0.25bc |
| 30- 60    | 23.77 ± 0.71b   | 17.94 ± 0.35b | 15.18 ± 0.27b          | 15.17 ± 0.91b          | 22.63 ± 0.36a |
| 60- 90    | 2.75 ± 0.13c    | 7.49 ± 0.02c  | 12.12 ± 0.02c          | 11.66 ± 0.71c          | 13.16 ± 0.38c |
| 90- 120   | 1.01 ± 0.19d    | 1.99 ± 0.05d  | 7.44 ± 0.18d           | 4.74 ± 0.04d           | 14.75 ± 0.06b |

* Different lower-case letters in each column indicate that these values are statistically different at P ≤ 0.05 according to Duncan’s Multiple Range Test.

CONCLUSION

This study was performed to reveal the stomatal anatomy, stomatal index and water loss (%) of mature pistachio leaves as well as the leaves of different phases (multiplication, rooting, hardening and regenerated plant) of micropropagation of mature pistachio trees obtained from the in vitro. The results of our study were corroborate the results obtained by Zacchini et al., (1997) who suggest that the circular shape of in vitro stomata is not a characteristic of the early differentation stages, but rather occurs as a gradual process over time. It can be expressed that it is necessary to optimization micropropagation steps in order to obtain high survival rates of in vitro propagated pistachio seedlings after adaptation to in vivo conditions.

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