Hybridization Assays in Strawberry Tree toward the Identification of Plants Displaying Increased Drought Tolerance

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Abstract: *Arbutus unedo* L. is a small Ericaceae tree with a circum-Mediterranean distribution. It has a huge ecological impact on southern Europe forests and a great economic importance as a source of phytochemicals with bioactive properties and for fruit production. On the foreseen climate change context, breeding toward drought tolerance is necessary in order to ameliorate plant performance. Therefore, the aim of this work was to study the reproduction mechanisms of the strawberry tree, obtain new genetic combinations by hybridization, and select genotypes more tolerant to drought stress. A morphological analysis of flowers and pollen was carried out, and controlled pollinations were performed both in vitro and ex vitro. The very first approach on strawberry tree breeding by means of hybridization is also presented. Several physiological parameters were evaluated on 26 genotypes submitted to a water-deficit regime. Plant behavior under drought greatly varied among genotypes, which showed high phenotype plasticity. Three genotypes that were able to cope with water restriction without compromising net CO₂ assimilation were identified as highly tolerant to drought stress. The results obtained elucidate the reproduction mechanisms of the strawberry tree and open the way for a long-term breeding program based on the selection of drought-tolerant plants.

Keywords: *Arbutus unedo* L. artificial pollination; breeding; drought stress; microscopy; pollen; physiological performance

1. Introduction

Strawberry tree (*Arbutus unedo* L.) is an Ericaceae widely distributed mainly around the Mediterranean basin, but also in the Atlantic coasts of Portugal, France, and Southern Ireland [1]. Members of this species grow spontaneously on poor rocky and well-drained acidic soils and can stand a wide range of temperatures, easily thriving in marginal soils where other species of trees or bushes can hardly survive. Moreover, its ability to rapidly regenerate after forest fires prevents the spread of common invasive species in Mediterranean ecosystems such as *Acacia* spp. or *Ailanthus altissima*, avoids soil degradation, and helps retain water, which are aspects that demonstrate the relevant ecological role of this species. Strawberry tree has several uses, mainly on pharmaceutical, cosmetics, and food industries, due to the variability and amount of phytochemicals present in their tissues and organs [2]. However, the edible berries used to produce an alcoholic distillate that reaches high prices in the market still remain the principal income source for farmers and other stakeholders [3].

Strawberry tree orchards are usually established using seed-derived plantlets on marginal dry areas where water is usually a scarce resource. Considering the expected increase on the frequency and severity of drought events in southern Europe in the near future [4], it is urgent to obtain new genotypes more tolerant to drought stress, in order to ameliorate their performance and increase productivity. Several studies have been...
carried out to study the effects of drought stress on water relations, growth rate, and photosynthesis in *A. unedo* under field conditions [5]. Although crucial, data focusing on drought performance at early stages of plant development are missing. Thus, early selection decisions are currently only based on productivity/fruit quality traits. Considering the increasing demand of high-quality plant stocks of *A. unedo*, in vitro propagation protocols were developed to clone selected genotypes [6–8], and studies to evaluate how these in vitro propagation systems change drought tolerance of regenerated plants have been carried out [9].

Some studies have been conducted on strawberry tree pollen, and a morphological description has been provided [10,11]. However, as far as is known, no work has been done in order to improve strawberry tree throughout conventional breeding, although some extensive experiments have been carried out on other Ericaceae, such as *Rhododendron* [12,13] and *Vaccinium* [14,15] species. Although conventional breeding is a lengthy process, particularly in tree species with long life-cycles, improved varieties of several tree species such as *Populus* spp., *Platanus* spp., and *Malus x domestica* have been produced through classical breeding [16,17]. The first step to initiate the development of new cultivars based on conventional breeding is a deep knowledge of the mechanisms of sexual plant reproduction, in particular the compatibility between the male and female reproductive structures, as well as the time of their maturation and phenology [18].

In order to set up the basis for a long-term breeding program on strawberry tree, the aim of this work was to study the reproduction system of *A. unedo*, from pollen morphology to pollen–stigma interactions, and analyze the tolerance of the F1 plants toward drought. For this purpose, a morphological analysis of flowers and pollen was carried out, and controlled pollinations were made in vitro and in situ to obtain hybrid plants. Moreover, the plants obtained by the artificial crossings were tentatively selected based on its drought tolerance. For this purpose, they were submitted to a water deficit regime in order to identify individuals able to maintain higher photosynthetic levels under water deficit conditions, which might be used on future micropropagation and/or breeding programs.

2. Materials and Methods

2.1. Plant Material

Flowers from three different populations were used in this study: CH (N 41°42′31.868″ W 7°26′32.506″, altitude 579 m), from Chaves (North Portugal) and populations C1 (N 40°12′17.472″ W 8°23′40.929″, altitude 103 m) and C2 (N 40°11′33.604″ W 8°23′37.163″, altitude 123 m) from Coimbra (Central Portugal). Plants were selected based on their fruit quality and production. A tree from population CH was used as a pollen donor for morpho-histological analysis (Section 2.2), germination studies (Section 2.3), and in vitro and in situ pollination assays (Section 2.4). For pollen release and gathering, anthers were removed from the flowers and placed on a Petri dish coated with aluminum foil for 1–2 days at room temperature. In vitro and in vivo pollinations (Section 2.4) were carried out using emasculated flowers from ten trees from C1 and C2 populations (5 from each population) and the collected pollen as described before (Figure 1).
2.2. Reproductive Phenology and Anatomy

Trees from populations C1 and C2 were monthly monitored throughout the year in order to characterize A. unedo reproductive phenology. Flowers and fruits were gathered and characterized, including anther position and fruit maturation stages. To analyze its morphology, pollen from CH was treated by the standard method of acetolysis [19]. Briefly, after being washed in water and acetic glacial acid (100%, v/v), pollen grains were treated with the classic acetolysis mixture (9:1, acetic anhydride:sulphuric acid) and heated in a water bath at 70 °C for 5 min. After being treated with acetone (100%, v/v), acetylated pollen material was mounted in glycerin jelly (Sigma-Aldrich, St. Louis, MO, USA). Measurements (D: diameter of the tetrad; d: single grain diameter; and the ratio D/d) were taken under light microscopy (Nikon EclipseCi, Nikon Instruments Europe BV, Amsterdam, Netherlands) with an ocular micrometer, from 30 randomly chosen pollen tetrads from 3 different slides (10 per slide). Terminology based on that of Punt et al. [20] was used for pollen morphology characterization. For scanning electron microscopy, pollen was placed on stubs and coated with gold on a JEOL JFC 1100 apparatus (JEOL, Musashino, Japan). Pollen observations were performed on a JEOL JSM 5400 microscope (JEOL, Murashino, Japan). For another anatomy studies, whole anthers were fixed for 3 h at room temperature in glutaraldehyde (1.5%, v/v, Sigma), prepared with phosphate buffer (0.1 M), and postfixed in osmium tetroxide (1%, w/v, Sigma) prepared with the same buffer. Samples were further dehydrated with ethanol and embedded with Spurr resin [21]. After the polymerization, ultrathin sections (1.5 μm) were obtained on an LKB Ultratome III and the cross-sections were stained with toluidine blue (1%, w/v) [22] and observed on a light microscope (Nikon EclipseCi, Nikon Instruments Europe BV), and photographs were collected with a Nikon DS-Fi3 camera (Nikon Instruments Europe BV) and processed with the software NIS-Elements D (version 4.60, Nikon Instruments Europe BV).

2.3. Pollen Germination

Previously to the pollination assays, the viability of the collected pollen was checked, and the effect of sucrose concentration was evaluated. Mature pollen was cultured on Petri dishes containing a basal germination medium [23] composed of H3BO3 (5 mg L−1), CaCl2 (15 mg L−1), KNO3 (10 mg L−1), agar (8%, w/v, Duchefa Biochemie BV, Haarlem, The Netherlands), and different concentrations of sucrose (0, 3, 6, 9, 12, 15, and 18%, w/v, Duchefa), for 6 and 24 h, at room temperature. Then, pollen grains were stained with acetocarmine and observed under a light microscope. As A. unedo pollen grains are dispersed as

![Methodology scheme applied in this work to study the phenology and reproductive anatomy of strawberry tree as well as selection of drought tolerant plants.](image-url)
tetrad units [11], germination rates were determined by scoring 100 pollen tetrads from 5 replicates (a total of 500 tetrads and 2000 pollen grains). A pollen tetrad was considered germinated when the pollen tube length of at least a pollen grain surpassed the diameter of a pollen grain. In a second experiment, different carbon sources as well as the effect of some plant growth regulators on pollen germination were also tested. For this purpose, pollen was cultivated for 6 h on the medium described before, with sucrose, glucose, and fructose at three different concentrations (3%, 9%, and 15%). The effect of NAA (1-naphthaleneacetic acid, Sigma), IBA (indole-3-butyric acid, Sigma), and GA$_3$ (gibberellic acid, Sigma) was also tested in three concentrations (10, 100, and 500 mg L$^{-1}$) on the same basal germination medium containing 15% sucrose.

2.4. In Vitro and In Situ Pollination Assays

For in vitro pollination, flowers immediately before anthesis from C1 and C2 populations were used. After emasculation, a total of 120 pistils from each population was placed on baby food jars (5 pistils per container) with a jellified medium for support (with water and 8 g L$^{-1}$ agar). Then, pollen from population CH collected as described in Section 2.2. was carefully placed at the stigma using a spatula. Open and closed non-pollinated flowers were used as controls and all treatments were done in triplicate. From each population, a total of 75 pistils were crosspollinated (15 pistils/tree from 5 different trees), 15 autopollinated (from a single tree), and 30 used as negative and positive controls (15 each from a single tree). Following artificial pollination, the pistils were kept in the dark at 25 °C, for 24 h, and the efficiency of the pollination was evaluated. For this purpose, pistils were fixed in FAA (formalin:acetic acid:ethanol, 5:5:90, v/v/v) at room temperature for 24 h, washed in water, softened on a NaOH solution (8 N), and mounted with aniline blue (0.1%, w/v, Sigma) as described by Martin [24]. The observations were carried out in a fluorescence microscope (ex: 370 nm, Leica DM4000 B, Leica Microsystems GmbH, Wetzlar, Germany), and pollination was considered efficient when pollen germination was observed on the stigma, and pollen tubes were grown along the style and reached the ovaries.

For in situ pollination assays, pollen with viability over 80% from a single tree (population CH) was used to hand-pollinate flowers from the trees used for in vitro pollination. After the emasculation with forceps, the pollen was carefully placed on the stigma, and the pollinated flowers were covered with polypropylene pollination bags for 7 days in order to avoid pollen contamination. Then, 10 flowers from three different inflorescences (a total of 30 per tree) were pollinated on each of the 10 trees. All the immature and old flowers from the pollinated inflorescences were removed. During the assays, the minimum absolute temperature was 7.1 °C, and the maximum absolute temperature was 26.9 °C, while the total precipitation recorded was 145.5 mm, according to the data provided by the meteorological station of Coimbra/Cernache (www.ipma.pt).

2.5. Seed Germination and Plant Development

Mature fruits resulting from hand-pollination were gathered and washed with tap water. Isolated seeds were washed with distilled water for 10 min. Following a 30 s surface sterilization with ethanol (70%, v/v, Merck), the seeds were sterilized in a calcium hypochlorite solution (5%, w/v, Sigma) and 2–3 drops of Tween 20 for 10 min, washed 3 times with distilled sterilized water, and sowed on sterilized Petri dishes (9 cm) with cotton wool imbibed with sterile distilled water and covered with filter paper. The seeds were kept at 4 °C for 30 days and then transferred to a culture chamber (25 °C) for another 60 days. After this period, the germination rate was recorded, and Relative Germinability (RG) was calculated: $RG = (\text{number of seeds produced} \times 100)/\text{number of viable seeds germinated}$ [25]. Then, seedlings were transferred to acclimatization containers with sterilized perlite and kept in a growth chamber at 25 °C and 70% relative humidity, under a 16 h daily illumination regime of 15–20 µmol m$^{-2}$ s$^{-1}$ photosynthetically active radiation (PAR, cool-white fluorescent lamps). After 15–30 days, the plants were transferred to
individual containers (5 dm$^3$) with a substrate composed of peat and perlite (3:1, $v/v$, Siro, Mira, Portugal) for further growth on a greenhouse.

### 2.6. Drought Stress Assays

Then, three-year-old plants resulting from cross-pollination (a tree from each population) were submitted to water stress. A total of 26 plants (#1–13 from population C1 and #14–26 from population C2) were watered to full field capacity, and plant performance was evaluated after 24 h (t0). After that period, watering was interrupted, and plants were submitted to 3 weeks (t3) of water deficit. Leaf gas exchange was evaluated on t0 and t3, while due to the destructive nature of leaf water potential ($\Psi_w$) and leaf relative water content (RWC) measurements, sampling was performed at the end of the experiment (t3). The experiment was conducted during July, and the temperature ranged from 13 °C (15.8 ± 1.4) to 32 °C (24.1 ± 2.7). The average temperature at each sampling point was 20.5 ± 4.5 °C (t0) and 17.5 ± 6.4 °C (t3).

In situ leaf gas exchange measurements (net CO$_2$ assimilation rate: A, transpiration rate: E, stomatal conductance: gs and intercellular CO$_2$ concentration: ci) were measured on a young and fully expanded leaf (normally the fifth leaf from the top) using a portable infrared gas analyzer coupled to a broad leaf chamber (LCpro+, ADC, Hoddesdon, UK), operating in open mode and under the following conditions: photosynthetic photon flux density—650 µmol m$^{-2}$ s$^{-1}$ (based on a light curve: 0–1750 µmol m$^{-2}$ s$^{-1}$); air flux—200 mol s$^{-1}$; block temperature—25 °C; and atmospheric CO$_2$ and H$_2$O concentration. Data were recorded when the measured parameters were stable (2–6 min). Water potential was measured with a Scholander-type pressure chamber (PMS Instrument Co., OR, USA). Relative water content (RWC) was calculated as: $\text{RWC} (%) = (\text{FW} - \text{DW})/(\text{TW} - \text{DW}) \times 100$, where FW is the fresh weight of the leaf, TW is the turgid weight (after 24 h on distilled water at 4 °C), and DW is the dry weight (after drying at 70 °C for 48 h).

### 2.7. Statistical Analysis

Pollen germination and physiological data were analyzed using ANOVA (GraphPad Prism for Windows v. 6.01) followed by Tukey’s multiple comparison test ($p < 0.05$). Data expressed as percentages were first submitted to arcsine transformation. A heatmap with a dendrogram and principal component analysis (PCA) were carried out using R software (version 4.0.3, R Foundation for Statistical Computing, Vienna, Austria) [26] to evaluate the interaction and significance of all the physiological parameters measured on the analyzed trees. A heatmap with physiological data from all the samples was constructed using the Heatmap function and the package ComplexHeatmap [27]. The dendrogram within the heatmap was calculated with Euclidean distance as a dissimilarity measure. Finally, data were classified with a PCA, using the prcomp function and the package ggbiplot [28].

### 3. Results

#### 3.1. Reproductive Phenology and Anatomy

The reproductive cycle of strawberry tree is long, and it lasts for almost two years (Figure 2A). During this period, three distinct stages can be identified: flower buds, flowers at anthesis, and fruit development. During June, the inflorescences (panicles) start to appear from terminal meristems of young stems (Figure 2B). Flower development proceeds through summer months, and flower anthesis usually begins in October (Figure 2C). The flowering period can be long, from early October to late January depending on the trees and location. The flower is complete, bell-shaped, sympetalous, and white to slightly pink (Figure 2C). Each pistil is formed by a pentalocular ovary, a style, and a stigma that becomes receptive to pollen just before flower anthesis. Each stamen possesses a hairy filament and an anther with two pores located at the top. During flower development and just before flower anthesis, anthers suffer an inversion process from an extrorse to an introrse position and develop two appendages on the apical end (Figure 2D). After pollination,
the slow fruit development process begins. Each infructescence will usually bear between 1 and 20 fruits (Figure 2E) that will develop along the year until fully ripened (Figure 2F). Consequently, fruit ripening occurs simultaneously with the next flowering period, during autumn (Figure 2A). Fruits at different developmental and ripening stages can be found at the same time on a tree. When fully ripped, fruits present a variable size and shape and a bright red color (Figure 2G).

Anthers of *A. unedo* have four microsporangia or pollen sacs arranged in pairs (Figure 3A). Pollen were dispersed in groups of four, and each anther contains an average 500 pollen units. At the earlier developmental stages, some pollen grains were found to be aborted on the pollen tetrad (Figure 3B). Pollen tetrads became mature and are released just after flower anthesis. Single pollen grains are 3-zonocolporate and have a circular or slightly elliptic outline on optical slice (Figure 3C,D). The ectoapertures are long colpus with a granulate membrane, and the endoapertures are pores with a regular outline (Figure 3E,F). The exine (≈ 1.5 µm) has a psilate surface and is tectate and slightly columel-
late. The size of the pollen tetrads (D) ranged from 42 to 67 μm (53.8 μm ± 3.6), whereas the size of single pollen grains (d) varied from 22 to 36 μm (29.5 μm ± 0.8). The relation D/d was between 1.6 and 2.2 (1.8 ± 0.1).

![Figure 3. Anther and pollen morphology of strawberry tree: (a) anther cross-section stained with toluidine blue; (b) pollen sac with aborted pollen grains; (c) pollen tetrad section stained with toluidine blue showing a bi-nucleated pollen grain; (d) non-acetolized pollen tetrad; (e) pollen tetrad on SEM; (f) aperture and ornamentation detail of the pollen tetrad on SEM.](image)

3.2. Pollen Germination

Pollen germination was higher on media with higher sucrose concentrations for both periods analyzed (6 and 24 h). After 24 h, the best germination rates were obtained on 15%, 18%, and 21% sucrose, without statistical differences. However, a decrease on germination was observed on the medium with 18% sucrose after 6 h (Figure 4A,B). The highest germination rates were obtained on a medium with 21% sucrose, after 24 h (83.29% ± 10.85), and 15% sucrose after 6 h (80.52% ± 12.55; Figure 4A,B). In most of the pollen tetrads scored as germinated, only one of the pollen grains developed a pollen tube (70.98% ± 1.81), whereas the germination of more than two pollen grains was only observed occasionally (Figure 5A). When different carbon sources were tested, sucrose gave the best results on the three concentrations tested, with statistical differences when compared to glucose and fructose. The maximum germination rate was obtained with 15% sucrose (70.33% ± 1.89). Although the glucose was not as efficient as sucrose, a germination rate of 57.00% ± 4.55 was obtained with the maximum concentration tested (Figure 4C). No pollen germination was observed when fructose was used as a carbon source. Likewise, NAA and IBA had an inhibitory effect on pollen germination, even on the lowest concentration tested (10 mg L⁻¹). When these two auxins were applied at higher concentrations, pollen germination was completely inhibited with statistical difference between concentrations for both NAA and IBA (Figure 4D). On the other hand, GA3 highly promoted pollen germination, and germination rates of 93.33 ± 3.09 and 94.67 ± 1.25 were obtained with 100 mg L⁻¹ and 500 mg L⁻¹ of GA3, respectively (Figure 4D). These values are higher than those obtained when a concentration of 10 mg L⁻¹ was applied, with statistic significant differences. When the concentration of CaCl₂ was highly incremented on the germination medium (10× increase), similar results were obtained.
3.5. Plant Water Status and Gas Exchange

Figure 4. (a) Effect of different sucrose concentrations (0, 3%, 6%, 9%, 12%, 15%, and 18%) on pollen germination rates after 6 h and (b) 24 h; (c) effect of different carbon sources (sucrose, glucose, and fructose) at three different concentrations (3%, 9%, and 15%) on pollen germination after 24 h; (d) effect of plant growth regulators (indole-3-butyric acid (IBA), 1-naphthaleneacetic acid (NAA), and gibberellic acid (GA3)) on germination rates at different concentrations (10, 100, and 500 mg L−1) after 24 h. Values (%) are means ± standard deviations, n = 5, different letters indicate significant differences between treatments at p ≤ 0.05.

Figure 5. Strawberry tree pollen germination: (a) germinated pollen tetrad stained with aceto-carmine; (b) germination of pollen grains on stigma; (c) pollen tube entering the ovule micropyle; (d) pollen germination on the stigma, and pollen tube growth along the pistil.
3.3. In Vitro and In Situ Pollination Assays

From the 12 combinations of crosses carried out in vitro, including two auto-pollinations, the average success rate obtained was 78.9% ± 22.7. Pollen germination was observed on stigma 1–2 h after the pollination (Figure 5B). The pollen tubes grow along the stile, reaching the ovary in 24 h (Figure 5D), and the tips of pollen tubes enter the micropyle (Figure 5C). The effectiveness of the cross-pollinations was 82.51% ± 19.81, while the effectiveness of the self-pollinations was 71.65 ± 29.50. We observed the accumulation of callose along the pollen tubes as well as on the tips. In some cases, pollen showed no signs of germination, both on self- and cross-pollinations. The growing pattern of pollen tubes seemed to be very similar on all the crosses made. In most of the flowers from the positive control (open flowers), pollen germination and pollen tube growth were observed, while all the flowers from the negative control (closed flowers) showed no signs of pollen in the stigma. Most of the pollinated flowers in situ were lost along the fruit developmental process. From the total of 300 pollinated flowers, after one year under development, only three fruits reached the mature stage, which represents a very low success rate of only 1%.

3.4. Seed Germination and Plant Development

From the three fruits retrieved from the field, a relative germination rate of 85.0% was obtained for group C1 and 86.7% was obtained for C2. After in vitro germination, seedling development proceeded rapidly, and after the acclimatization period, the root system was well developed (Figure 6A). The hybrid plants were morphological diverse (Figure 6B) in terms of height and leaf morphology. A total of 35 plants were obtained, 17 from group C1 and 13 from C2. After 3 years under development, 13 plants from each group were submitted to drought stress.

3.5. Plant Water Status and Gas Exchange

Before the imposed water stress deficit regime (t0), a considerable variance was found among genotypes on all the physiological parameters measured. While some of the tested genotypes presented higher net CO₂ assimilation rates (e.g., 3, 5, 13, 21, and 25), others had considerably lower values (e.g., 2 and 17) (Table 1, Figure 7A). After 3 weeks under water...
deficit, stomatal conductance, net CO\textsubscript{2} assimilation, and transpiration rates decreased (Table 1, Figure 7B). On the other hand, intercellular CO\textsubscript{2} concentration increased on most of the plants throughout the imposed water stress. Although a great reduction on stomatal conductance, transpiration and net CO\textsubscript{2} assimilation rates was observed on genotype 13 after drought stress, these parameters are still considerably higher than on most of the evaluated genotypes. A similar behavior was observed on genotypes 12 and 17, but with a less marked decreased on net CO\textsubscript{2} assimilation rate. Meanwhile, these parameters remained unchanged or slightly increased on genotypes 14, 15, and 18. In general, relative water content and water potential were higher on plants with a higher net CO\textsubscript{2} assimilation rate.

| Genotype | ci \textsubscript{t0} | E \textsubscript{t0} | gs \textsubscript{t0} | A \textsubscript{t0} | RWC \textsubscript{t0} | WP \textsubscript{t0} |
|-----------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1         | 181             | 166             | 2.76            | 0.86            | 0.32            | 0.06            | 25.87           | 7.67 | 63.4 | 50 * |
| 2         | 243             | 220             | 1.98            | 0.34            | 0.18            | 0.02            | 11.81           | 2.31 | 72.8 | 50 * |
| 3         | 181             | 159             | 3.5             | 0.64            | 0.51            | 0.04            | 32.36           | 5.49 | 70.5 | 50 * |
| 4         | 215             | 302             | 2.88            | 0.14            | 0.32            | 0.01            | 21.2            | 0.45 | 47.5 | 50 * |
| 5         | 176             | 223             | 3.34            | 0.55            | 0.42            | 0.03            | 29.84           | 3.07 | 70.9 | 50 * |
| 6         | 193             | 258             | 1.98            | 0.53            | 0.16            | 0.03            | 14.7            | 2.21 | 59.4 | 50 * |
| 7         | 186             | 204             | 2.48            | 0.84            | 0.23            | 0.05            | 19.59           | 5.18 | 63.9 | 50 * |
| 8         | 172             | 230             | 2.91            | 0.45            | 0.31            | 0.02            | 25.23           | 2.32 | 53.0 | 50 * |
| 9         | 209             | 364             | 2.82            | 0.19            | 0.28            | 0.01            | 19.85           | 0.15 | 48.6 | 50 * |
| 10        | 205             | 296             | 3.52            | 0.29            | 0.43            | 0.02            | 26.22           | 0.82 | 50.0 | 50 * |
| 11        | 192             | 435             | 2.96            | 0.27            | 0.3             | 0.01            | 22.51           | 0.46 | 45.7 | 50 * |
| 12        | 196             | 220             | 2.96            | 2.52            | 0.3             | 0.22            | 22.1            | 15.61 | 84.6 | 21   |
| 13 +      | 165             | 165             | 3.5             | 1.63            | 0.43            | 0.11            | 31.25           | 12.51 | 81.2 | 32   |
| 14 ++     | 160             | 171             | 2.11            | 2.7             | 0.16            | 0.23            | 16.92           | 20.27 | 84.7 | 24   |
| 15 ++     | 163             | 201             | 2.8             | 3.27            | 0.24            | 0.33            | 22.5            | 22.19 | 79.4 | 14   |
| 16        | 179             | 232             | 2.89            | 0.63            | 0.24            | 0.03            | 20.86           | 2.89 | 59.6 | 50 * |
| 17 +      | 199             | 190             | 1.87            | 1.59            | 0.12            | 0.1             | 11.24           | 10.47 | 78.8 | 26   |
| 18 ++     | 139             | 208             | 2.58            | 3.41            | 0.2             | 0.34            | 20.96           | 21.7  | 71.7 | 10.5 |
| 19        | 157             | 305             | 2.85            | 0.91            | 0.22            | 0.05            | 21.26           | 2.01  | 73.3 | 18.5 |
| 20        | 163             | 260             | 2.82            | 0.68            | 0.22            | 0.03            | 20.68           | 2.45  | 44.0 | 50 * |
| 21        | 105             | 302             | 3.28            | 0.63            | 0.26            | 0.03            | 29.13           | 1.24  | 74.0 | 31.25|
| 22        | 125             | 260             | 2.93            | 0.6             | 0.23            | 0.03            | 24.77           | 2.16  | 42.3 | 50 * |
| 23        | 76              | 188             | 2.64            | 0.64            | 0.18            | 0.03            | 25.25           | 3.73  | 77.6 | 42.5 |
| 24        | 87              | 191             | 2.72            | 0.92            | 0.19            | 0.05            | 25.27           | 5.3   | 58.2 | 48.5 |
| 25        | 191             | 325             | 3.28            | 0.19            | 0.46            | 0.01            | 29.66           | 0.55  | 50.6 | 50 * |
| 26        | 246             | 377             | 2.42            | 0.33            | 0.27            | 0.02            | 16.36           | 0.02  | 50.5 | 50 * |

This result is confirmed by the dendrogram within the heatmap (Figure 7A,B) as well as the PCA biplot (Figure 7C,D), which revealed a high positive correlation between E, gs, and A with relative water content and water potential (Figure 7B). Moreover, a negative correlation was found between these parameters and ci. On most of the plants, water potential was below the detection limit (−50 MPa). The PCA analysis has also revealed a very diverse behavior of plants, regardless of their provenience either on t0 or t3 (Figure 7C,D). Thus, plants with the best and worst performance under water stress are from both proveniences. On t0, principal component 1 (PC1) contributes 67.8% to the total variance and A, gs, and E are the parameters with a higher weight on this component, whereas PC2 contributes 29.8% to the total variance, and ci is the variable that most contributes to this variance. The genotypes identified due to a better performance (3, 5, 13, 21, and 25) are grouped (Figure 7C). On t3, principal component 1 (PC1) contributes 76.5%
to the total variance. $A$, $gs$, $E$, relative water content, and water potential are the parameters with a higher weight on this component. PC2 contributes 14.6% to the total variance. As mentioned before, some of the tested plants showed a better overall performance in terms of net CO$_2$ assimilation under drought. Thus, genotypes 12, 13, 14, 15, 17, and 18 are grouped together by the influence of some gas exchange parameters ($gs$, $E$, and $A$), relative water content, and water potential. Genotype 19 is also on this cluster, as it was able to maintain relatively high values of relative water content and water content, in spite of its low performance in terms of net CO$_2$ assimilation. Plants with a worst overall performance are grouped together by the influence of $ci$ (Figure 7D).

Figure 7. Physiological parameters measured on 26 hybrid plants from populations C1 and C2 on $t_0$ and $t_3$ (3 weeks under water deficit): (a,b) heatmap with dendrogram on $t_0$ and $t_3$; (c,d) principal component analysis (PCA) on $t_0$ and $t_3$. $ci$—intercellular CO$_2$ concentration, $E$—transpiration rate, $gs$—stomatal conductance, $A$—net CO$_2$ assimilation rate, $RWC$—relative water content, $WP$—water potential.

When comparing the two groups (C1 and C2), although no statistical differences were found between groups (Figure 8A), transpiration rates greatly decreased after 3 weeks on both groups (Figure 8B), as well as stomatal conductance (Figure 8C) and net CO$_2$ assimilation rate (Figure 8D). No statistical difference was found for relative water content as well, with values of $62.4 \pm 13.0\%$ on C1 and $65.0 \pm 14.7\%$ on group C2.
Figure 8. Gas exchange-related parameters of hybrid plants from populations C1 and C2 on t0 (plants watered to full field capacity) and t3 (3 weeks under water deficit): (a) intercellular CO₂ concentration; (b) transpiration rate; (c) stomatal conductance; (d) net CO₂ assimilation rate (D). Values are means ± SDs, n = 13, different letters indicate significant differences between treatments at p ≤ 0.05.

4. Discussion

The long phenological cycle observed on the trees analyzed on this work is similar to the one that has been reported by Villa [10]. The inversion process of the anthers observed, as well as the development of the two appendages, has been described as a characteristic feature of the Ericaceae family [29]. The pollen morphology observed is similar to data reported by Villa [10], but it slightly differs from that described by Mateus [11]. According to this author, the endoapertures of strawberry tree pollen are endocolpus. In contrast, in this study, the endoapertures observed were endopores. The size of the pollen tetrads is slightly different as well: the diameter of the tetrads (D) determined on this work range from 42 to 67 µm, compared to 49–66 µm obtained by Mateus [11], while the size of single pollen grains (d) range from 22 to 36 µm compared to 33–41 µm.

The pollen germination rates obtained on the germination medium with 12% sucrose (89.40%) are similar to the ones obtained on other species, such as *Prunus domestica* L. [30] and *Pistacia* spp. L. [31], and they are much higher than the germination rates obtained on *Annona cherimola* Mill. [32] and *Olea europaea* L. [33]. Therefore, the germination medium used is adequate for strawberry trees and should be used for pollen viability tests. Although it has been reported by Cane [34] that 90% of the tetrads generated 3–4 pollen tubes on another member of the Ericaceae (*Vaccinium macrocarpum* Aiton), we found that most of the grains on the pollen tetrads were found aborted, even in the initial developmental stages. The inhibition of pollen germination caused by fructose has been reported on the literature by Okusaka and Hiratsuka [35]. According to these authors, fructose completely inhibited pear pollen germination but without pollen viability loss. Thus, fructose is not an adequate sugar for strawberry tree pollen germination, and other carbon sources should be used instead, preferably sucrose. Plant growth regulators (PGRs) can be an extremely useful tool for plant breeders, either as gametocides or on the contrary by promoting pollen germination and eventually increasing fruit seed-set. Different PGRs have been tested on diverse crop species including rye [36], barley [37], onion, tomato, eggplant, pepper, watermelon [38], and wheat [39], on most cases in concentrations similar.
to the ones tested on this study (10–500 mg L\(^{-1}\)). Our results showed an inhibitory effect of IBA and NAA on strawberry tree pollen germination in similar concentrations to those reported in the literature: the application of NAA (50 mg L\(^{-1}\)) on eggplant and IBA or NAA (10–100 mg L\(^{-1}\)) on onion proved to have an efficient gametocide effect [38]. On the other hand, the applications of at least 100 mg L\(^{-1}\) of GA\(_3\) greatly promoted pollen germination, which is an effect that has also been observed on strawberry with a similar concentration (50 mg L\(^{-1}\)) [40] and blueberry where the application of GA\(_3\) on flowers (on concentrations approximately between 30 and 500 mg L\(^{-1}\)) lead to an increased fruit set [41]. For this reason, the effect of GA\(_3\) on strawberry tree pollen germination and fruit set should be further tested due to its potential as a breeding aiding tool.

Although a previous study had suggested that pollen tubes growth speed is slower on self-pollination due to higher rates of attrition [42], this was not the case in strawberry trees, and similar pollen tube growth patterns on self- and cross-pollinations was observed. Moreover, no difference was found between the effectiveness of self- and cross-pollinations. The observed accumulation of callose on pollen tubes of a strawberry tree has also been observed in *Chaenomeles japonica* (Thunb.) Lindl. ex Spach [43]. Only a small portion of the hand-pollinated flowers on the field were able to complete the long development process and bear fruits. Due to the long development process that takes a year to be completed, strawberry tree pollinated flowers and fruits under development are subjected to a wide range of environmental conditions and interferences, and only a small portion of fruits is able to complete its development. This might help explain the low success rate of hand-pollinations along with the high manipulation required to carry out the pollination procedure. In fact, it has been reported that fruit production on *Vaccinium* spp. is lower when plants were cross- or self-pollinated by hand than when natural pollination occurs [15]. Thus, the improvement of pollinations conditions is something to be pursued in the near future in order to increase success rates of hand-pollinations. The increase of the amount of pollen placed on the stigma and/or the use of PGRs (e.g., GA\(_3\)) should be considered, as well as the implementation of open pollinated seed orchards. The size and seediness of fruits may also be affected when hand-pollination is carried out, as referred by Usui et al. (2005). In this work, the average of viable seeds obtained on hand-pollinated fruits was 50%, which is much lower than the 77% obtained on open pollinated trees. Nonetheless, due to the low amount of hybrid fruits obtained, these results are not significant, and further analyses should be carried out in the future. The high germination rates obtained are similar to other works [44–46] indicating that the germination ability of the hybrid seeds is not compromised. However, such rates were obtained after cold stratification, which is a procedure that should be followed in order to break seed dormancy. In fact, it has been reported that fruit production on *Vaccinium* spp. is lower when plants were cross- or self-pollinated by hand than when natural pollination occurs [15]. Thus, the improvement of pollinations conditions is something to be pursued in the near future in order to increase success rates of hand-pollinations. The increase of the amount of pollen placed on the stigma and/or the use of PGRs (e.g., GA\(_3\)) should be considered, as well as the implementation of open pollinated seed orchards. 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on this work, which is probably related with the period under water stress as well as the age of the plants that might have more lignified tissues and a higher resistance to cavitation and low water potential. Still, this hypothesis should be further tested and confirmed on future analysis.

Although most of the tested plants showed a poor performance under drought stress, we successfully identified two groups of plants that followed a different strategy to cope with water deficit and were able to maintain a high stomatal conductance and consequently higher net CO$_2$ assimilation rates. Genotypes 14, 15, and 18 were able to maintain their basal levels of photosynthesis, which was accomplished by maintaining stomata open, as these plants were able to maintain relatively high levels of water (relative water content and water potential). Genotypes 12, 13, and 17 were shown to have an intermediate performance under drought stress. Although these plants were able to maintain the photosynthetic mechanisms active after 3 weeks under water-deficit conditions, they were already probably close to their resistance limits, and a significant drop on net CO$_2$ assimilation rates was expected after a few more days under stress. Finally, some genotypes (e.g., 19) were able to maintain relatively high levels of water (relative water content and water potential), but they were unable to maintain satisfactory levels of net CO$_2$ assimilation, which might be due to the biochemical limitation of photosynthesis rather than stomatal constraints. From the genotypes identified on t0 that have higher net CO$_2$ assimilation rates, only genotype 13 was able to maintain a similar performance under stress. On the other hand, genotypes with lower net CO$_2$ assimilation rates on t0 (e.g., 17) were able to cope with drought stress and maintain the levels measured at t0. The genotypes identified on t0 for its high net CO$_2$ assimilation rates (3, 5, 13, 21, and 25) might have high potential, and their productivity should be evaluated. Nonetheless, on the water restriction scenario we hypothesize on this work, with the exception of genotype 13, they generally fail to cope with water stress.

In addition to revealing the importance of genotype on strawberry tree physiological performance and response to drought, these results show that strawberry tree plants have a high phenotypic plasticity and are able to adjust differential strategies to cope with stress. In order to facilitate and considerably reduce the necessary required time for selection, the identification of other adequate selection parameters should be pursued. In particular, metabolites such as phenols, proline, chlorophyll, anthocyanins, and several hormones (e.g., abscisic acid, jasmonic acid, and salicylic acid) that are known to be essential on plant response mechanisms to drought stress, might be used as markers to identify plants with a better appetite to undergo extreme drought events.

Although a great variance was observed between individuals from the same population, no differences were observed between populations on all the tested parameters. These results suggest that intra-population variation should be take into account and prioritized over inter-population on future selection endeavors, and a large number of individuals from within a population must be sampled. In contrast, results obtained by Vasques et al. [49] showed that seedlings provenience might influence the tolerance of plants under water stress, thus suggesting local adaptations of plant populations, which reinforces the importance of inter-population variance on plant behavior. Due to its implication on plant selection, this hypothesis should be further investigated. Overall, the obtained results will have important repercussions on strawberry tree phenotyping and early plant selection as well as breeding toward the obtention of drought stress resistant genotypes.

5. Conclusions

As a basis for any breeding program that includes plant hybridization, a deep knowledge of the plant reproduction system is necessary. This work provides the first insights of a strawberry tree reproduction system, which will be crucial on future breeding attempts. As a tool for plant selection, the physiological parameters used in this study proved to be adequate. However, the analysis of biochemical parameters could not only elucidate the tolerance mechanism of A. unedo but also identify key metabolites (e.g., phenols, hormones,
and pigments) that could be used as markers for early plant selection. Three genotypes (14, 15, and 18) showed a particular aptitude to cope with water stress and may be the basis for a future breeding program. However, due to the influence of genotype on plant response to water stress and the observed phenotypic plasticity, the analysis of a large number of individuals should be carried out in order to develop a long-term breeding program. The selection and breeding of strawberry tree genotypes more tolerant to drought stress is essential in order to maintain species sustainability and our promising results are a step forward in order to ameliorate strawberry tree adaptation while preserving productivity on drought prone areas.

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