Development of an Efficient ‘One-Step Freezing’ Cryopreservation Protocol for a Georgian Provenance of Chestnut (*Castanea sativa* Mill.)

Zygotic Embryos

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Abstract

Experiments were performed to determine the influence of various dehydration and vitrification treatment times on the ‘one-step freezing’ cryopreservation of embryonic axes (EAs), composed of zygotic embryos and cotyledon residuals, from mature seeds of a Georgian provenance of chestnut (*Castanea sativa* Mill.). Dehydration was carried out in laminar flow hood from 1 to 5 h, and vitrification experiments were carried out by immersion of EAs in PVS2 vitrification solution up to 120 min, both followed by direct immersion in liquid nitrogen. Both systems resulted in inducing specimen tolerance to ultra-rapid freezing, although to a different extent. Full germination of cryo-stored EAs after 5 h of dehydration (reducing moisture content from initial 66% to 21%) has been increased from 0% to 66.7%. A pre-treatment of EAs in PVS2 vitrification solution for 30 min produced fully developed plantlets at a rate of 55.6% in post-cryopreservation. Plantlet regrowth from cryopreservation was faster in EAs that underwent the dehydration/‘one-step freezing’ procedure. All the plantlet from cryopreserved EAs could be easily acclimatized, producing healthy potted plants. Finally, the TTC test showed to be useful for a fast evaluation of specimen survival after thawing and, as a consequence, to speed up the development of optimized cryo-protocols.

Keywords: chestnut; cryopreservation; dehydration; embryonic axes; vitrification; zygotic embryos

Introduction

The forest ecosystems are of special significance for conservation of biodiversity of Georgia. The forest covers 39.9% of the territory of the country. Their greater percentage (98%) is presented by mountainous forests and they have water regulating, soil protective, climate stabilizing functions. Besides, they are an important location of many relict, endemic and threatened species of plants and animals. European chestnut or sweet chestnut (*Castanea sativa* Mill.) is the dominant of mountainous forests of Western Georgia (150-1800 m). It occupies the most percentage of areas covered with forests (approx. 75%). Dendroflora of Caucasus, issued earlier in 1961, lists 14 species within genus *Castanea*; however, only *Castanea sativa* Mill. occurs in Georgia (Gulisashvili, 1961). Chestnut forests are distributed continuously along the southern slope of the Caucasus Mountains near the Black Sea and are found in isolated populations on the north side of the Caucasus, at elevations ranging from 200 to 1400 meters (Pridnya et al., 1996). Chestnut-dominated forests comprise only a few percent of total forest cover in the Caucasus Biosphere Reserve, and usually occur in mountain valleys or coves with deep brown soil. *Castanea sativa* forests are developed in both West and East Georgia, but to the west of the country they occupy larger areas. In some localities pure stands of *C. sativa* can be found, but mainly sweet chestnut occurs as a component of oligodominant beech-sweet chestnut and hornbeam-beech-sweet chestnut forests (Gagnidze, 1975). Chestnut trees generally extend from 200 (West Georgia) up to 900–1000 meters above sea level, having the absolute upper limit at 1400 meters in sporadic locations of West and East Georgia (Nakhutsrishvili, 2013).

At present, due to low self-renewal and diseases, the big massifs of chestnut forests in Caucasus are on the verge of...
Chestnut blight was apparently introduced into the region after 1880 and continues to destroy chestnut forests today. Although chestnut seed production appears adequate, the competition with shrubs and other tree seedlings, as well as predation by herbivores and rodents, interrupts the establishment and survival of chestnut seedlings in the biosphere preserve. In addition, sweet chestnut in the Caucasus is infested by several fungal and bacterial parasites (Tavadze et al., 2012). Castanea sativa Mill. is listed in the Red List of Georgia (2006) due to the tendency to the decrease of the distribution range and habitat fragmentation. IUCN category for this taxon is evaluated as Vulnerable (VU), according to the IUCN Red List Categories and Criteria: population reduction observed, estimated, inferred, or suspected in the past where the causes of reduction may not have ceased or may not be understood or may not be reversible. According to the official IUCN list (2019) Castanea sativa has been assessed as Least Concern (Barstow and Khela, 2018).

The seeds of Castanea sativa are classified as recalcitrant (Pence, 1992; Westengen et al., 2013). In contrast to orthodox seeds, the recalcitrant seeds are shed at high water contents and are highly susceptible to desiccation and sometimes also to low temperatures (Vertucci and Farrant, 1995; Berjak and Pammenter, 1997). Such behaviour poses a major problem to seed banks for long-term conservation. Axes and cotyledons of recalcitrant seeds of Castanea sativa Mill. exhibit contrasting responses of respiration and sensitivity in relation to desiccation (Leprince et al., 1999), and their conservation at low above-freezing (0-2 °C) temperature (Girolone and Bounous, 1993) allows the maintenance of seed viability and germinability only up to 90 days (Conedera et al., 2004). This makes urgent search for alternative techniques for conservation of this vulnerable taxon.

Conservation of shoot cultures at low above-freezing temperature is one option for ex situ medium-term preservation of woody plant germplasm. As for chestnut, the technique has been investigated, allowing the conservation of micropropagated shoot cultures for a limited period of time at 4 °C (Lambardi et al., 2001), and at 8 °C (Capuana and Di Lonardo, 2013). Cryopreservation or freeze-preservation at the ultra-low temperature (-196 °C) of liquid nitrogen (LN) is the only technique currently available to ensure the safe and cost-efficient long-term conservation of germplasm of both seed and vegetatively-propagated species (Panis and Lambardi, 2006; Benelli et al., 2013). It is of particular practical interest for the long-term conservation of species characterized by sub-orthodox and non-orthodox seeds (Pritchard and Nadarajan, 2008; Walters et al., 2013). At the temperature of LN, biochemical and most physical processes are completely arrested. In this context, cryopreservation should be regarded as a sound complementary approach to the traditional seed- and in-field bank strategy of germplasm preservation, providing a real guarantee against accidental loss of plant genetic resources. As for chestnut, few reports are available, regarding the conservation in LN of shoot tips (San-José et al., 2005; Vidal et al., 2005; Vidal et al. 2010), zygotic embryo axes (Corredoira et al., 2004; San-José et al., 2005), and somatic embryos (San-José et al., 2005; Vidal et al., 2010). However, such reports always refer to seeds collected from local open-pollinated chestnut trees, and the wide applicability of selected protocols to germplasm from different and distant (in terms of geographical areas and climatic characteristics) provenances has never tested.

In order to develop an efficient strategy for the long-term conservation of threatened chestnut Georgian provenances, the present study developed a step-wise ‘one-step freezing’ protocol for the cryopreservation of excised embryonic axes (EAs) from the Caucasus region, comparing two different cryo-techniques, i.e., specimen dehydration and PVS2-vitrification, both followed by the direct immersion of EAs in LN (‘one-step freezing’). Moreover, the application of a test of seed viability after their recovery from LN has been applied for the first time in chestnut, with the aim to speed up the optimization of an efficient and repetitive cryo-procedure.

Materials and Methods

Biological material

Castanea sativa hedgehogs were collected in October-November 2018 from Tikul region in western Georgia, Imereti, at an altitude of 600-800 meters above sea level. After extraction from the hedgehogs, mature fruits were stored at 4 °C for a maximum of 45 days, until use in the cryopreservation trials.

Dehydration/’one step freezing’

For cryopreservation experiments, the fruit were washed in household detergent (2%) and rinsed three times under tap water. Then the pericarp, seed coat and lateral part of endosperm were removed. The remaining seeds were surface sterilized by successive immersion in 70% (v/v) ethanol with few drops of Tween 20 for 2 min, followed by decontamination with 10% solution of sodium hypochlorite for 20 min. After being rinsed in sterile distilled water tree times, EAs, composed by the zygotic embryos along with cotyledonal knots and 2-3 mm-long cotyledon residuals, were dissected from seeds (Fig. 5A).

To determine the moisture content (MC) of EAs, the fresh weight of single EAs, with 5 replicates for each desiccation time, was measured. After final period of desiccation time, embryos were dried at 85 °C for 24 h and re-weighted. MC was calculated as a percentage of embryo fresh weight.

As for cryopreservation trials, for each of 5 desiccation times (0, 1, 2, 3, 4, and 5 h), 50 EAs were desiccated in open Petri dishes under the air flux of a laminar flow hood. Thirty specimen were then used for determination of MC (10 EAs), for the tetrazolium chloride (TTC) test (see below; 10 EAs), and for germinability determination (10 EAs) after dehydration but without immersion in LN (LN-). Twenty EAs (10 for viability, and 10 for germinability determination) were placed in 2-ml cryovials (10 EAs in each) and immersed in LN for 24 h (LN+). For thawing, the cryovials were rapidly immersed in water bath at 40 °C for 2 min. As for EA germinability (for its determination, see below), LN+ and LN- samples were cultured in WPM (Lloyd and McCown, 1981) with 0.4 μM 6-benzylaminopurine (BAP) and 0.3% activated charcoal...
(Sigma-Aldrich), inside test tubes (20 mm × 150 mm). Sucrose was supplied at 30 g l⁻¹. The media were then solidified with 6 g l⁻¹ agar and adjusted to pH 5.7 before autoclaving. EAs were maintained in growth chamber at 24 ± 0.5 °C under 16/8 h photoperiod with an irradiance of 40 μmol m⁻² s⁻¹ from cool-white fluorescent tubes.

**PVS2-vitrification/one-step freezing**

EAs were transferred to loading solution (LS) containing 2 M glycerol and 0.4 M sucrose for 20 min at 24 °C. Then, the EAs were placed in 2-mi cryovials and treated with PVS2 vitrification solution (30% w/v glycerol, 15%, w/v DMSO, 15% w/v ethylene glycol in WPM containing 0.4 M sucrose; Sakai et al., 1990), for different times (0, 30, 60, 90, 120 min) at 0 °C, after they were plugged in LN for 24 h. For each treatment time, 40 embryos were used. A control without any treatment and LN- (10 EAs), was also included. For 0-min treatment time, EAs were only loaded in LS solution and then cultured in test tubes for germination. After immersion in LN the cryovials were thawed in a water bath at 40 °C for 2 min, and the EAs were rinsed in washing solution containing WPM liquid medium and 1.2 M sucrose (two times for 10 min each, at 25 °C), and then cultured in test tubes, as reported above.

**Determination of EAs viability after cryopreservation by the TTC test**

The viability of EAs was evaluated using 1% TTC solution. Ten EAs for each dehydration time followed by cryopreservation, and 10 dehydrated but not cryopreserved (LN-), were transferred on filter paper in 90 mm Petri dishes. The EAs were covered with the TTC solution and placed overnight in the dark at 20 °C. After staining, the solution was drained off and EAs were rinsed under tap water. Viability of each EA was interpreted according to the topographical staining pattern of the embryo and the cotyledon residuals were fully stained. Completely unstained EAs or stained in small patches were evaluated as 'non-viable'.

**Data collection and statistical analysis**

The total number of EAs used was 300 for dehydration/one-step freezing, and 240 for PVS2-vitrification/one-step freezing. Each treatment consisted in 5 replicates and all experiments were repeated 3 times. EAs germinability was assessed first time after 7 days of culture, reported as 'survival' that indicated the percentage of EAs germinability was assessed first time after 7 days of culture, 3 replicates and all experiments were repeated 3 times. After each time of dehydration under the sterile air of laminar hood, significative starting from 2 h of dehydration and reaching a final of 21.4% of MC after 5 hours (Fig. 1A and B). This dehydration trend of EAs that were not cryopreserved (LN-) didn’t affect the survival of EAs which remained 100% even at the lowest MC (Fig. 1A); however, not all these EAs completed the germination process, as plantlet regrowth from EAs dehydrated for 5 hours was reduced to 84%. EAs which were directly immersed in LN, without any previous dehydration (time 0), never survived to cryopreservation. The dehydration procedure, on the other hand, induced a clear effect of tolerance of the EAs to ultra-rapid freezing in LN. Indeed, even after only 1 h of dehydration, EAs with a MC reduced to 62.8% showed a 21.4% survival to LN. Increasing the dehydration time produced a parallel increase in the survival of EAs which was significative starting from 3-h dehydration and that reached a maximum of 80% after 5 h.

The survival of EAs refers to the first clear signs of germination (see ‘Material and Methods’), followed by shoot elongation (Fig. 5B). However, part of the apparently survived EAs did not evolve in plantlets (Fig. 1B), showing only root elongation (Fig. 5C). First evidences of plantlet regrowth (21.8%) was achieved in EAs that were dehydrated for at least 2 h, producing a reduction of their MC to 41.2%. Increasing the time of dehydration up to 5 h induced a constant increase of plantlet regrowth, up to a maximum of 66.7%, a value significantly higher than the ones achieved with lower dehydration times (Fig. 1A). Afterwards, plantlets were easily transferred in pots and acclimatized (Fig. 5E).

The TTC test showed to be an effective test for a fast evaluation of EAs viability following cryopreservation. Comparing the two lines reported in Fig. 2, the ‘viability’ line (expressing the percentages of EAs showing full red

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**Table 1.** Plantlet regrowth of *Castanea sativa* EAs subjected to various dehydration times with (LN+) or without (LN-) immersion in LN, evaluated 4 weeks after cryostorage, thawing and plating (mean values ± SE).

<table>
<thead>
<tr>
<th>Dehydration time (h)</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
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<tbody>
<tr>
<td><strong>Root length (mm)</strong></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>LN-</td>
<td>54.6±2.0</td>
<td>48.8±0.6</td>
<td>40.1±0.8</td>
<td>33.8±1.2</td>
<td>52.2±2.1</td>
<td>49.7±1.7</td>
</tr>
<tr>
<td>LN+</td>
<td>-</td>
<td>5.7±0.3</td>
<td>5.0±0.4</td>
<td>6.0±0.3</td>
<td>8.0±0.2</td>
<td>40.1±0.2</td>
</tr>
<tr>
<td><strong>Shoot length (mm)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN-</td>
<td>33.0±1.4</td>
<td>31.8±1.2</td>
<td>28.4±1.1</td>
<td>23.2±1.3</td>
<td>29.8±1.9</td>
<td>23.0±1.4</td>
</tr>
<tr>
<td>LN+</td>
<td>-</td>
<td>-</td>
<td>5.0±0.4</td>
<td>5.8±0.2</td>
<td>7.1±0.7</td>
<td>22.2±0.4</td>
</tr>
</tbody>
</table>
colour after recovery from LN and thawing) has a similar pattern to the ‘survival’ line, with a gap between the two lines of about 40% up to 4 h dehydration. Following a 5 h dehydration of EAs, the gap between the two lines was 20%.

It is interesting to note that the further development and plantlet regrowth in post-cryopreservation (LN+) of EAs dehydrated 5 h was much faster than EAs subjected to lower dehydration times, showing after 4 weeks roots longer than 4 cm and shoots longer than 2 cm (Table 1).

Fig. 1. Percentages of moisture content, survival (A) and plantlet regrowth (B) of Castanea sativa EAs from Tkibuli region, following increasing dehydration times with (LN+) or without (LN-) cryopreservation. Within each line, percentage values followed by the same letter are not significantly different at the P≤0.05 by ANOVA and LSD test (bars, SE of means)

Fig. 2. Comparison between the viability, as resulted by the TTC test, and actual survival of dehydrated and cryopreserved EAs. Within each line, percentage values followed by the same letter are not significantly different at the P≤0.05 by ANOVA and LSD test.

Cryopreservation by PVS2-vitrification of EAs

The LS, containing 2 M glycerol and 0.4 M sucrose, showed to have a limited toxic effect on the EAs (Fig 3A). Indeed, the treatment with only LS for 20 min at 0 °C, induced a small reduction of survival from 100% (control, non-treated and non-cryopreserved) to 83.4% (time 0), although significant with the untreated/LN- control. However, it induced also a tolerance to ultra-rapid freezing in LN, as EAs survival and plantlet regrowth passed from nil to almost 20%. When the EAs, following the LS treatment, were then treated with the PVS2 vitrification solution for increasing period of time and non-cryostored (LN-), EAs survival remained in the range of 75% (30 and 60 min of treatment) and 83.3% (90 and 120 min), which never showed significant differences of percentage values. Referring to EAs that were afterwards cryopreserved (LN+), a 30-min treatment with PVS2 showed to produce best result in terms of EAs survival (55.6%), significantly different from percentages obtained with both lower and higher PVS2 treatments. Indeed, when the exposure to PVS2 was increased to 60 min, a drastic decline of survival after immersion in LN and thawing was observed (25%) and reached a minimum of 20% following a PVS2 treatment of 120 min.

The treatment with PVS2 showed to negatively affect the further EAs development and regrowth in plantlets, as non-cryostored EAs (LN-) developed in plantlets from a maximum of 66.7% (30 min of PVS2 treatment) to a minimum of 40.2% (120 min of PVS2 treatment) (Fig 3B). As for cryopreserved EAs (LN+), differently from the dehydration procedure, all the EAs evolved in plantlets, confirming that the 30-min PVS2 treatment was the most effective in inducing partial tolerance to ultra-rapid freezing in LN.

The TTC test had a partially different trend from the application to the dehydration/one-step freezing procedure (Fig. 4). Indeed, starting from a PVS2-treatment time of 30 min, the overestimation of the TTC test was always in the range of 40-60%, being 40% (similarly to the value observed following the dehydration procedure) at the best treatment time of 30 min.

Full germination of EAs (expressed as plantlet regrowth) required a double time (8 weeks), in comparison to the dehydration/one-step freezing procedure. Similarly to what observed with the latter procedure, plantlet regrowth was faster in EAs that were PVS2-treated with the time achieving best result in terms of sample survival. Indeed, EAs, PVS2-treated for 30 min and cryopreserved, showed in post-cryopreservation root and shoot regrowth over 2 cm after 8 weeks from thawing and plating (Table 2).

Discussion

In the past 30 years, various explants (tissue and organs) have been successfully used in plant cryopreservation (Panis and Lambardi, 2006; Reed, 2008). Among them, sound cryopreservation procedures have been developed for orthodox (Pritchard and Nadarajan, 2008) and non-orthodox (Walters et al., 2013) by means of the storage in LN both entire seeds and excised embryonic axes (Normah and Makeen, 2008). Cryopreservation of entire seeds sometimes does not require any pre-treatment to induce
Table 2. Plantlet regrowth of *Castanea sativa* EAs subjected to various PVS2-treatment times with (LN+) or without (LN-) immersion in LN, evaluated 8 weeks after cryostorage, thawing and plating (mean values ± SE)

<table>
<thead>
<tr>
<th>PVS2-vitrification time (min)</th>
<th>Control</th>
<th>0</th>
<th>30</th>
<th>60</th>
<th>90</th>
<th>120</th>
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<tbody>
<tr>
<td>Root length (mm)</td>
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<tr>
<td>LN-</td>
<td>66.5±0.7</td>
<td>20.0±0.8</td>
<td>36.3±1.2</td>
<td>10.7±0.3</td>
<td>26.3±1.8</td>
<td>24.7±0.5</td>
</tr>
<tr>
<td>LN+</td>
<td>-</td>
<td>5.0±0.2</td>
<td>27.4±0.2</td>
<td>8.3±0.3</td>
<td>7.9±0.4</td>
<td>9.9±0.4</td>
</tr>
<tr>
<td>Shoot length (mm)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN-</td>
<td>42.0±1.4</td>
<td>30.0±0.6</td>
<td>30.0±0.6</td>
<td>21.7±1.1</td>
<td>26.0±0.4</td>
<td>24.4±1.3</td>
</tr>
<tr>
<td>LN+</td>
<td>-</td>
<td>4.0±0.1</td>
<td>23.4±0.3</td>
<td>11.2±0.2</td>
<td>9.7±0.2</td>
<td>6.7±0.2</td>
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</tbody>
</table>

Fig. 3. Percentages of moisture content, survival (A) and plantlet regrowth (B) of *Castanea sativa* EAs from Tkibuli region, following increasing PVS2 treatment times with (LN+) or without (LN-) cryopreservation. Within each line, percentage values followed by the same letter are not significantly different at the P≤0.05 by ANOVA and LSD test (bars, SE of means)

Fig. 4. Comparison between the viability, as resulted by the TTC test, and actual survival of PVS2-treated and cryopreserved EAs. Within each line, percentage values followed by the same letter are not significantly different at the P≤0.05 by ANOVA and LSD test
tolerance to ultra-rapid freezing in LN, such as in various conifers, while in many others a pre-dehydration to a MC around 20%, followed by slow cooling or by direct immersion in LN, is strictly necessary (Panis and Lambardi, 2006). However, when embryos are composed by large cotyledons, as in chestnut, some insurmountable problems (i.e., the impossibility to insert them in cryovials, the large volume occupied in the dewar when using large boxes, the difficulties in inducing homogeneous dehydration or vitrification) induce to the use of excised EAs for cryopreservation. EAs, composed of the zygotic embryo and 2-3 mm of residual cotyledons, have been used for cryopreservation in this study. The presence of a small portion of cotyledons showed to be necessary to allow a germination, differently to what observed in others species, such as peanut (Ozudogru et al., 2009) where the EA totally prived of cotyledons does not evidence any negative effect during germination. On the opposite, every reduction of cotyledons compromises Pistacia vera seed germinability in post-cryopreservation (Ozden-Tokatly et al., 2007).

Physical dehydration and vitrification cryo-procedures, both allowing the direct immersion of specimen in LN ('one-step freezing'), have been applied to the cryopreservation of embryonic axes from temperate (Pence, 1990; Ozden-Tokatly et al., 2007; Ozudogru et al., 2009) and tropical plant species (Berjak and Dumet, 1996; Berjak et al., 1999). Physical dehydration is obtained, generally, exposing the EAs to the sterile air of a laminar flow hood for an appropriate time, adequate to minimize ice-crystal formation but, at the same time, high enough to avoid reaching a minimum level, below which damages for cell plasmolysis can be produced. This minimum level should be previously determined, and it can reach very low values. In Phaseolus vulgaris and Medicago sativa, for instance, seeds can be dehydrated to 7% MC, maintaining over 90% germinibility after cryopreservation (Stanwood and Bass, 1981). Similarly, with peanut EAs, maximum germinability (100%) was still obtained following the reduction of the initial MC from 25% to 8.5% (Ozudogru et al., 2009).

The natural MC of EAs of chestnut was 66.2%, a value similar to what reported by other Authors (Pence, 1992; Correidora et al., 2004), and germinability was not affected up to 4 h dehydration (MC, 22.7%), while showed a reduction to 84% of full germination (plantlet regrowth) after 5-h dehydration (MC, 21.4%). Zygotic embryos of Correidora et al. (2004) had an initial survival of 83%, and had a beneficial effect from the dehydration treatment (2-6 h) that induced a 100% survival. In this study, survival of EAs was always 100% and was not affected by the increasing time of dehydration treatment. In accordance with what observed by Correidora et al. (2004), also in the Georgian provenance of Castanea sativa the dehydration/‘one-step freezing’ procedure showed to be very effective in inducing a clear tolerance of EAs to ultra-rapid freezing in LN. Indeed,
after 5 h dehydration, EAs achieved maximum survival in post-cryopreservation of 80%, followed by a plantlet regrowth of 66.7%. Besides the high germinability obtained with the dehydration/cryopreservation procedure, the germinated EAs could be easily acclimatized and produced healthy seedlings, without any sign of morphological alteration or callus proliferation. It is interesting to note that, in order to evaluate the efficacy of the cryo-procedure, it is not sufficient to evaluate the early germination (survival), but to determine the full germination. In Georgian chestnut plantlet regrowth (full germination) should be evaluated 4 weeks after thawing and culturing EAs in test tubes.

Also the PVS2-vitrification protocol developed in the present study showed a good applicability to the Georgian chestnut provenance, although to a lesser extent than the dehydration/one-step freezing technique. It is worthy of note that the incubation of specimens with a LS solution, prior to exposure to PVS2, was beneficial, as showed also from the partial survival and plantlet regrowth from EAs that had been only pre-treated in LS prior to cryopreservation. This is consistent with the general idea that a pre-loading of explants with cryoprotectants (at lower concentrations than they have in the vitrification solutions) is useful to minimize the possible adverse effects caused by the PVS2-vitrification solutions (Sakai, 2000). Indeed, the components of the cryoprotectant solution (glycerol, sucrose) are not permeable in the short period used for loading. However, such application causes cells to plasmolysis as result of considerable dehydration, and the presence of LS solution in the periprotoplasmic space of the plasmolyzed cells probably mitigates mechanical stress caused by severe dehydration due to exposure to PVS2 (Matsumoto et al., 1998).

The risk of toxicity of the vitrification solution is well known (Lambardi et al., 2000; Kim et al., 2011), and it is probably caused by the denaturation of proteins. The present study demonstrated that chestnut EAs tolerated well the PVS2 treatments in terms of early germinability (survival), but less in terms of further germination (plantlet regrowth), particularly starting from 60-min treatment. As a consequence, the apparently low germinability obtained with the application of the PVS2 solution to EAs prior direct immersion in LN (with a maximum of plantlet regrowth of 55.6% following a 30-min treatment) should be interpreted as the best that could be achieved, considering the almost equivalent result achieved from EAs that had been equally treated but not frozen in LN.

It has been shown in this study that only a portion of the EAs graded ‘survived’ 7 days after dehydration/PVS2-vitrification and immersion in LN were able to germinate afterwards, and 4-8 weeks had to be waited for a final evaluation of the tested cryo-procedures. Hence, in order to find a marker which could provide an early and reliable evaluation of seed germinability, the TTC test was used. The TTC test is a biochemical method, originally developed by Lakon (1949), in which viability is determined by the red colour appearing after soaking the seeds in a TTC solution for at least overnight, depending on the species. Living tissue changes the TTC to an insoluble red compound (formazan); in non-living tissue the TTC remains uncoloured. The grading of a seed into “viable” or “non-viable” depends upon the amount and location of white areas. As a consequence, correct interpretation depends upon standards worked out for specific seeds (MacKay, 1972). When the EAs of the Georgian provenance of Castanea sativa were tested just after their recovery from LN, a good correlation with their germinability was found for seeds specimen, showing completely red embryos and residual cotyledons. The general overestimation which was observed for both dehydrated/LN+ and vitrified/LN+ (starting from 30-min exposure to PVS2) EAs is in accordance with what reported, e.g., in Pistacia spp. seeds (Ozudogru et al., 2009).

Conclusions

In conclusion, the present study has clearly demonstrated that the application of both dehydration and vitrification systems for cryopreservation of chestnut zygotic embryos resulted in viable embryos capable to develop fully developed plantlets. Dehydration gave better EAs survival and plantlet regrowth, as well as a more rapid organized plant development after cryopreservation, thus proving to be more effective practice in compare to the PVS2-vitrification/one-step freezing technique. However, this lower performance of the latter procedure has been shown to be due to an intrinsic toxic effect of the PVS2 solution on the treated chestnut zygotic embryos. Research in this field should move in the direction to test for chestnut zygotic embryos other less toxic vitrification solutions (Kim et al., 2011).

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Conflict of Interest

The authors declare that there are no conflicts of interest related to this article.

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