Genetic stability of wild pear (*Pyrus pyraster*, Burgsd) after cryopreservation by encapsulation dehydration

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Shoot tips of *Pyrus pyraster* were successfully cryopreserved by encapsulation-dehydration. Na--alginate beads each containing one shoot tip, dehydrated for 2 days in 0.75M sucrose and desiccated to 20% moisture content (fresh weight basis), gave 60% recovery after exposure to liquid nitrogen. Regenerated shoots showed no differences in length and leaf shape compared to the mother plant. Multiplication rate and rooting ability of cryopreserved shoots were lower than those of untreated controls after one subculture, but were completely restored following the third subculture. Fifteen cryopreserved lines derived from single buds were used for genetic analyses by RAPDs and SSRs, in comparison with the mother plant. In RAPD analysis, of a total of 24 primers, only 15 showed reproducible and well resolved bands and were further used. These primers produced a total of 66 fragments ranging from about 500 to 2500 base pair size. SSR (microsatellite) marker amplification was performed using 19 primers which produced 57 reproducible fragments. Microsatellites fragments ranged from 60 to 600 base pairs. Both RAPDs and SSRs did not reveal any polymorphism between cryopreserved lines and the original genotype, suggesting that cryopreservation, using encapsulation-dehydration, does not affect genetic stability of wild pear.

Key-words: germplasm preservation, multiplication ability, RAPDs, rooting, somaclonal variation, SSRs, tissue culture.
Introduction

In situ plant germplasm preservation plays an important role in the maintenance of biodiversity and the avoidance of genetic erosion, but the preservation of woody species in field gene banks requires huge land areas and it is expensive (Panis and Lambardi 2005). Cryopreservation is an alternative choice for the long-term conservation of germplasm including woody fruit species and their wild relatives (Engelmann 2004). In recent years, several new techniques have been developed for the cryopreservation of shoot tips of tropical and temperate plant species. The encapsulation-dehydration method, originally described for cryopreservation of *Solanum* shoot tips (Fabre and Dereuddre 1990), has been successfully applied to cryopreserve shoot tips of several fruit tree species (Gonzalez-Arnao and Engelmann 2006), including *Pyrus* (Dereuddre et al. 1990, Niino and Sakai 1992), *Prunus* (Shatnawi et al. 1999), *Malus* (Niino and Sakai 1992, Paul et al. 2000, Wu et al. 2001b), *Vitis* and *Actinidia* (Plessis et al. 1993, Wu et al. 2001a).

*Pyrus pyraster* is considered an important wild relative of cultivated pear (*Pyrus communis* L.). The tree is considerable in size and diameter and its high quality wood makes this species interesting for reforestation of marginal farmland and for the production of highly valued timber (Kleinschmitt et al. 1998). In cultivated regions with calcareous soils, where Fe-chlorosis is a serious problem, wild pear can be also preferred as rootstock for pear cultivars. The species is indigenous in nearly all Europe, except in the northern countries, but it is now seriously endangered (Kleinschmitt et al. 1998). In in vitro propagated cultures, obtained from a single axillary bud of a wild pear (*Pyrus pyraster*, Burgsd) genotype, were established according to Caboni et al. (1999).

Shoots in the proliferation phase were subcultured every 21 days on a medium (“LPmod” medium) consisting of LP (Quoirin and Lepoirre 1977) salts and the following organics: 0.5 mg L\(^{-1}\) nicotinic acid, 0.5 mg L\(^{-1}\) pyridoxine, 2.0 mg L\(^{-1}\) glycine, 0.5 mg L\(^{-1}\) thiamine-HCl, 150 mg L\(^{-1}\) myo-inositol, 1.0 mg L\(^{-1}\) Ca-pantothenate, 0.1 mg L\(^{-1}\) biotin and 0.5 mg L\(^{-1}\) riboflavin, according to Caboni et al. (1999). The medium was also supplemented with 1.78 µM benzyladenine (BA), 0.25 µM indole-3-butyric acid (IBA), 20 g L\(^{-1}\) sucrose and 6 g L\(^{-1}\) agar (B & V - Italy). The pH was adjusted to 5.7 before sterilization and cultures were
maintained at 25°C under a 16 h photoperiod and a light intensity of 40 µmol m⁻² s⁻¹ provided by cool white fluorescent tubes (Philips TLD - France).

Cryopreservation protocol

Mother plants were cold acclimatized for 2 weeks in darkness at 5 °C. Subsequently, shoot tips (2–4 mm in length) were excised and sub-cultured for 24 h on the LPmod medium containing 0.3M sucrose. Apices were then transferred to MS (Murashige and Skoog 1962) calcium free liquid medium, supplemented with 3% alginate (Sigma). Beads were prepared by dispensing drops of alginate medium, each one containing one shoot tip, in a 100 mM CaCl₂ MS liquid medium. The beads formed were cultured in liquid LPmod containing 0.75M sucrose for 2 days, desiccated in vessel containing silica gel (5 beads in 18g) to a bead moisture content of 20% fresh weight, placed in 1ml cryo-vials (Nalgene, 10 beads in each cryovial) and immersed in liquid nitrogen where they were kept for 1 week. Beads were then directly transferred to Petri dishes containing 0.3M sucrose enriched LPmod medium in darkness at room temperature and finally transferred to the standard multiplication medium and sub-cultured regularly, as reported above.

Morphological observations, multiplication and rooting performance evaluation

Thirty days after recovery, necessary to overcome an initial lag phase in growth after storage in liquid nitrogen, 30 single shoots were transferred to standard proliferation conditions reported above for morphological and multiplication ability evaluation, in comparison with 30 shoots obtained from shoot tips of the mother plant. Multiplication rates (final number of shoots – initial number of shoots divided by initial number of shoots) were calculated and shoot length and leaf shape were evaluated in newly formed shoots. Data were collected at the end of the 1st and 3rd subculture.

For rooting experiments, 30 microcuttings were immersed for 5 days in a 2 mg L⁻¹ IBA solution plus 20 g L⁻¹ sucrose in the darkness and then transferred to a hormone free LPmod medium and to the light. Data were collected 30 days after the beginning of the root induction treatment. The experiment was repeated after three subcultures. Data (percentages were transformed to arc-sin root before analysis) were subjected to analysis of variance and differences among means were compared by Fisher’s test.

RAPD and SSR analysis

Fifteen of the recovered shoots were cultured separately and these lines were used for molecular analyses. Genetic stability of the 15 lines was tested, in comparison with the mother plant, using 24 RAPD primers, designed according to Williams et al. (1990) and 19 SSR primer pairs, previously selected in Yamamoto et al. (2002) for giving unambiguous and reproducible fragment patterns in pear. Total DNA was extracted from 100 mg of plant tissue with DNeasy Plant Mini Kit (Quiagen). Two independent extractions were performed for each line and for the control (mother plant). RAPD reactions were carried out in a volume of 30 µl containing 25 ng total DNA, 1X PCR buffer (Qiagen), 1.5 mM MgCl₂, 200 µM dNTPs, 0.4 µM 10-mer oligonucleotide primer (Invitrogen) and 1U Taq polymerase (Qiagen). SSR amplifications were performed in a volume of 30 µl containing 25 ng total DNA, 1X PCR buffer (Qiagen), 2.3 mM MgCl₂, 200 µM dNTPs, 0.3 µM oligodeoxynucleotide primers (Invitrogen) and 1U Taq polymerase (Qiagen). DNA amplifications were performed in a Biometra T thermal cycler with a preliminary step of 5 min at 94 °C, 45 cycles of 60 s at 94 °C, 60 s at 36 °C and 2 min at 72 °C and a final step of 5 min at 72 °C for RAPDs. For SSRs, an initial step of 5 min at 94 °C, followed by 35 cycles of 1 min at 94 °C, 1 min at 55–58 °C and 2 min at 72 °C and a final 5 min extension at 72 °C were performed. In order to obtain reproducible and clear DNA fragment patterns, each amplification was repeated twice. RAPD amplification products were separated in a 1.5% agarose (Duchefa – NL) gel using
SSR fragments were analysed on 3.5% MetaPhor agarose (Cambrex Bio Science - USA) gel in 1X TBE buffer and stained with ethidium bromide.

**Results**

After cryopreservation the survival rate of the explants was 60% (Fig. 1), they grew well, developed normally and, compared with the control, no morphological differences were observed (Fig. 2; Table 1). Multiplication rate and rooting ability of the cryopreserved shoots were lower than those of the control after one subculture, but were completely restored after three subcultures (Table 1).

In order to evaluate if the encapsulation dehydration method preserves genetic integrity in wild pear, we used RAPD and SSR markers for the molecular analysis. To increase the confidence of the analysis, we selected only those primers which gave very reproducible bands. In RAPD analysis, a total of 24 primers were firstly used to amplify DNA of all genotypes and with 15 of them we obtained reproducible and well resolved bands and they were selected for further use (Table 2). These primers produced a total of 66 fragments ranging from about 500 to 2500 base pairs in size. The highest number of analysable bands was obtained with the primers 70.13 and 70.20 (six fragments each), the lowest with the primer 70.15 (one fragment).

SSR marker amplification was performed using 19 primers that produced 57 reproducible fragments (Table 3). Fragments ranged from 60 to 600 base pairs; the highest number of analyzable bands was obtained with the primer pair NH020 (six fragments), the minimum with primer pair NB103 (one fragment). A summary of the results of RAPD and SSR marker analysis is given in Tables 2 and 3. The total number of fragments scored for the whole plant material analysed was 1056 (66 fragments

Table 1. Shoot length, leaf shape, proliferation (MR, multiplication rate) and rooting (% of rooted explants) ability in cryopreserved and control (mother plant) shoots after one or three sub-cultures (sc)

| Shoot length | Leaf shape | MR | Rooting % |
|--------------|------------|----|-----------|
| 1sc | 3sc | 1sc | 3sc | 1sc | 3sc | 1sc | 3sc |
| Cryo-shoots  | 3.1a | 3.3a | stand* | stand | 2.4a | 5.6a | 24.4a | 66.6a |
| Control      | 3.4a | 3.2a | stand | stand | 5.4b | 5.6a | 62.4b | 68.7a |

*Stand, expanded leaves of standard obvoidal form. Means on the column followed by the same letters are not significantly different at p=0.05. Percentage data were transformed to arc-sin root before statistical analysis.
× 16 samples, included the control) and 912 (57 fragments × 16 samples, included the control) for RAPD and SSR markers, respectively. Identical patterns were obtained with both markers when cryopreserved plantlets were compared with the mother plant (control plant) (Fig. 3 and 4).

Table 2. RAPD primers used for DNA amplification and total fragments scored.

| RAPD primers | Sequence 5’–3’ | Total fragments scored |
|--------------|----------------|-----------------------|
| 70.2         | CACAGGC CGGA    | 4                     |
| 70.4         | CCCGCTACAC      | 5                     |
| 70.5         | CAAAGGGCGGG     | 5                     |
| 70.7         | AAGTGCAACGG     | 4                     |
| 70.9         | AACGCCC CGTC    | 4                     |
| 70.12        | GGCCTACTCG      | 3                     |
| 70.13        | GTGTGCAGAG      | 6                     |
| 70.15        | GCCCTCTTCG      | 1                     |
| 70.17        | GAGACCTCCG      | 5                     |
| 70.19        | GCTCTCACCAG     | 5                     |
| 70.20        | TGCACGGACG      | 6                     |
| 70.22        | GTGGCCTCA       | 5                     |
| 70.23        | TTGGCACCGG      | 3                     |
| 70.24        | GTGTGCCCTCCA    | 5                     |
| 70.30        | CGCGCTACGT      | 5                     |

Table 3. SSR primers, annealing temperatures used for DNA amplification and the total number of fragments scored.

| SSR primer couples | Annealing Temperature (°C) | Total of scored fragments |
|--------------------|---------------------------|--------------------------|
| NB102              | 55                        | 2                        |
| NB103              | 55                        | 1                        |
| NB105              | 55                        | 5                        |
| NB106              | 55                        | 3                        |
| NB109              | 55                        | 4                        |
| NB110              | 55                        | 2                        |
| NB111              | 55                        | 4                        |
| NB113              | 58                        | 2                        |
| NB019              | 55                        | 2                        |
| NB020              | 55                        | 6                        |
| NB021              | 55                        | 3                        |
| NB022              | 58                        | 2                        |
| NB023              | 55                        | 4                        |
| NB024              | 55                        | 3                        |
| NB025              | 58                        | 3                        |
| NB026              | 55                        | 3                        |
| NB027              | 55                        | 2                        |
| NB029              | 58                        | 3                        |
| NB030              | 58                        | 3                        |

Fig. 3. RAPD banding profiles of DNA samples from mother plant (control, C) and cryopreserved shoots (1–15) of Pyrus pyraster. Amplification products were generated by primer 70.8. M: HyperLadder II (Bioline) marker.

Fig. 4. SSR profiles of DNA samples from mother plant (control, C) and cryopreserved shoots (1–15) of Pyrus pyraster. Amplification products were generated by primer couple NH030 FW-NH030 RW. Ma: 50 bp ladder marker and Mb: 100 bp ladder (Amersham-Pharmacia) marker.
Discussion

Conservation of plant genetic resources has to rely on methods that not only allow to obtain good survival, but also to guarantee that material remains genetically stable after conservation. Some of the steps involved in cryopreservation, putatively causing stress responses, may induce genetic instability (Engelmann 2004). For this reason, evaluation of genetic variation in cryopreserved material is an essential step before the large scale use of the established storage protocols.

In this study we used morphological, physiological and molecular markers to evaluate stability of preserved material. Similarly to Liu and co-workers (2004), who evaluated cryopreserved apple, we did not observe any morphological difference between leaf shape of the mother plant and of the cryopreserved wild pear shoots and, while multiplication and rooting ability was lower after one sub-culture, they were fully restored after three subcultures. Rooted microcuttings were acclimatized and they are now under further observation in the greenhouse.

Most of the studies previously performed have shown that the cryopreservation process does not affect genetic stability of the stored lines. Nevertheless, genetic variation was recently shown to be inducible by the encapsulation-dehydration method in *Dendrathema grandiflora* (Martin and González-Benito 2005) suggesting that attention should be paid to the evaluation of the genetic stability in cryopreserved lines.

In this study, we used RAPDs and SSRs to evaluate the genetic stability of the cryopreserved lines. These markers, both visualised by PCR (Polymerase Chain Reaction) and agarose-based electrophoresis, offer the advantage of being less expensive and quicker to be performed than RFLP or AFLP (Lanham and Brennan 1999, Powell et al. 1996). RAPDs, in particular, have been widely used to evaluate genetic stability in tissue cultures (Carvalho et al. 2004, Palombi et al. 2007 and references therein) and they have been also adopted to evaluate stability of cryopreserved material in various species (Hao et al. 2002, Ryynanen and Aronen, 2005, Zhai et al. 2003, Ventkatachalam et al. 2007). SSR markers allow screening of different regions of the genome than RAPDs, including repetitive and hypervariable DNA regions and they were shown to be valuable molecular tools for determining somaclonal variation in tissue culture (Rahman and Rajora 2001) and for genetic fingerprinting of fruit tree species, pear included (Yamamoto et al. 2002). Thus, RAPDs and SSRs, showing a different polymorphism capability, can be conveniently used, in combination, to evaluate somaclonal variability induced by tissue culture (Palombi and Damiano 2002).

In our study we used 15 RAPD primers to amplify DNA of 15 cryopreserved lines and of the mother plant. These primers produced a total of 66 fragments and the total number of bands considering all the plant material analysed was 1056. No differences were found between the wild pear lines and the mother plant in the number of fragments obtained, as also found in most of the works performed on genetic stability of cryopreserved material (Dixit et al. 2003, Sales et al. 2001, Zhai et al. 2003).

SSR marker amplification was also performed using 19 primer pairs that produced 57 reproducible bands corresponding to a total number of 912 fragments in the analysed material. Also this method showed no differences between the cryopreserved lines and the mother plant.

This is, to our knowledge, the first report on the combined application of RAPDs and SSRs for evaluation of genetic stability in cryopreserved lines of fruit trees. We analysed a total of 1968 fragments (from RAPD and SSR markers) without observing any genetic variation. This number of analysed fragments, can be considered to be informative, as reported in other studies performed on genetic stability of cryopreserved materials (Dixit et al. 2003, Sales et al. 2001, Zhai et al. 2003) and also in micropropagation (Kawiak and Lojkowska 2004).

However, due to the relatively low fraction of the genome screened with the molecular markers and the moderate sized test population used, these results cannot be interpreted as a final proof that no somaclonal variation has occurred. However, they
give us a preliminary information on the genetic stability of the wild pear cryopreserved material and let us to be confident of the possibility of using routinely the encapsulation-dehydration method for long term conservation of *Pyrus pyraster*.

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