INTRODUCTION

Red (*Trifolium pratense*), alsike (*Trifolium hybridum*) clovers and alfalfa (*Medicago sativa*) are the most popular fodder legumes suitable for growing in agronomical and ecological conditions of Latvia (Jansone, 1997). Clover breeding has been performed in Latvia since the 1920s and several widely used varieties were bred. Currently, only one alfalfa variety bred in Latvia (‘Skrîveru’) is registered for cultivation in Latvia (Jansone, 2008). For breeding of new varieties it is very important to have genetic resources with the necessary genetic variability. In the former Soviet Union, plant genetic resources activities were concentrated only in the N. Vavilov All-Union Institute of Plant Industry, now the N. Vavilov All-Russian Institute of Plant Industry, St. Petersburg, Russia (VIR). In 1993, when in Latvia plant genetic resources activities were started, it was recognised that many local varieties, as well as landraces and wild or semi-wild forms, were lost. Several accessions of Latvian origin were repatriated from the VIR and other gene banks. Seeds of the VIR collections were kept at room temperature. The majority of repatriated clover accessions were produced more than 20 years ago, some alfalfa accessions even 44 years ago, and therefore the seeds did not successfully germinate in soil.

Germination of seeds in *in vitro* culture conditions has been reported for many plants species, such as citrus *Citrus sinensis* (Niedz, 2008), strawberry (Hamdouni *et al*., 2001), almond (San and Yildirim, 2009) and others. Germination of old seeds is important for seed banking and conservation, for restoring genetic resources, to allow to improve modern plant breeding and cultivation, and to provide insight on the history of plant domestication and breeding. Special attention is paid on elaboration of methods for viability restoration of aged seeds, non-germinating in soil. In this respect, germination of an 1300-year-old seed is an unique case (Sallon *et al*., 2008).

There are known numerous factors that influence seed germination rate and seedlings growth: seed age and storage conditions (Rice and Dyer, 2001; Grauda *et al*., 2006; Val-leriani and Tielbörger, 2006), age of mother plants (Raja *et al*., 2004; Espahbodi *et al*., 2007), heredity (Revilla *et al*.,...
2009), conditions of seed germination and plantlet growth (Bhattacharya and Khuspe, 2001). Several tricks can improve aged seed germination ability: seed pre-treatment by temperature, light or phytohormones (Takayanagi and Harprove, 1971; Bhattacharya and Khuspe, 2001), and adding different phytohormones to cultivation medium (Bhattacharya and Khuspe, 2001; Soylar and Khawar, 2007). For various plant species, positive influence of AgNO$_3$ on seed germination and plantlet formation in vitro has been shown (Ornicāne and Rashal, 1997); similar effects were found also for KMnO$_4$ (Grauda et al., 2006), KH$_2$PO$_4$ (Sathish et al., 2011), activated carbon (Pacek-Bieniek et al., 2010) and TiO$_2$ nanoparticles (Zheng et al., 2005). The goal of this study was to obtain mature plants from aged red and alsike clover and alfalfa seeds of accessions of Latvian origin by using in vitro methods.

MATERIAL AND METHODS

Seeds of alfalfa and clover accessions of Latvian origin that were repatriated from the N. Vavilov All-Russian Institute of Plant Industry (VIR) or were kept in the Latvian Research Institute of Agriculture (RIA) were used for the germination experiment (Table 1). The accessions had not been multiplied for a long time and all available seeds had lost germinating ability in the soil.

The in vitro germination experiment was conducted in two stages. In the first stage the best conditions of seed germination were determined for a sample of accessions; in the second stage, those were applied to the entire set of accessions.

Clover. For determination of the best concentration of KMnO$_4$, in the first stage seeds of four clover accessions (‘Vietējais agrais’, ‘Stendes vēlais 2’, ‘Priekuļu tetraploidāsis’, ‘Vietējais’) were soaked for 40 minutes both in 0.05% and 0.1% of KMnO$_4$. For the remaining accessions, in the second stage of experiment only 0.1% concentration of KMnO$_4$ was used. Seeds were sterilised in 50% solution of commercial bleach “Belizna” for 20 minutes, which had been found earlier to be optimal (Ornicāne and Rashal, 1997). The bleach solution was cooled to 10 °C before use, according to Yildiz and Er (2002).

To induce germination, 100–400 seeds from each accession were placed in Petri dishes on Murashige and Skoog (MS) basal medium (Murashige and Skoog, 1962) with 2 mg/l kinetin and 0.5 mg/l NAA, or on the MS basal medium without phytohormones (15–30 seeds in a Petri dish). Seeds were cultivated in darkness at 26 °C. Germination rate was determined as number of germinated seeds to the total number of placed seeds.

Germinated seeds were placed on the MS basal medium without phytohormones and moved into light conditions at 24 °C (16 h day). Developed clover plantlets were multiplied by cuttings of different parts (stem segments, leaf petiole segments, leaf segments and axillary buds), placed on MS basal medium without phytohormones, and cultivated in light conditions at 24 °C (16 h day). Plantlets with good roots at stage of 3–4 leaves were planted in a plastic (100 ml) pot and covered with plastic foil for better acclimatisation, and cultivated in a growing room (16 h day, 60% humidity, temperature 18–20 °C). After several weeks of cultivation in the growing room, plants were moved to a greenhouse and grown for 2–3 months. Well developed plants were transferred for field trials and seed production to the Latvian Research Institute of Agriculture (RIA).

Alfalfa. Because only 1.0–1.5 g of seeds were available for accessions from the VIR, only alfalfa seeds of accessions from the Latvian Research Institute of Agriculture were used to determine the best pre-treatment and cultivation conditions. In the first stage, seed pre-treatment with both 0.07% and 0.1% KMnO$_4$ solution were performed for 40 or 60 minutes (Ornicāne and Rashal, 1997). Then seeds were sterilised by 50% solution of commercial bleach “Belizna” for 20 minutes. After pre-treatment, seeds were placed in Petri dishes on MS basal medium, or on the MS basal medium supplemented by 10 mg/l AgNO$_3$ or on the MS basal medium supplemented by 1 g/l activated carbon. In the second stage, seeds of accessions from the VIR were germinated using pre-treatment by 0.1% KMnO$_4$ for 60 minutes and MS basal medium without supplements.

<table>
<thead>
<tr>
<th>Accession name</th>
<th>Species</th>
<th>Age of seeds</th>
<th>Seeds</th>
<th>GM rate ( \text{in vitro}^* )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local k-31068</td>
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<td>44</td>
<td>VIR</td>
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<td>VIR</td>
<td>62.0</td>
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<td>Trifolium pratense</td>
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<td>33</td>
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<td>Trifolium hybridum</td>
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<td>VIR</td>
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<td>29</td>
<td>VIR</td>
<td>14.0</td>
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<tr>
<td>Vietējais</td>
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<td>Medicago sp.</td>
<td>14</td>
<td>RIA</td>
<td>18.0</td>
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</table>

* – seeds of all accessions did not germinate in soil
All seeds were cultivated at 20–26 °C (16 h day) for 3–4 weeks. Plantlets with good developed roots and 3–4 leaves were planted in soil in plastic pots (100 ml) and cultivated in a greenhouse. For better acclimatisation, pots with plantlets were covered by plastic film. After acclimatisation, the plastic film was removed and plants were grown for 2–3 months. Good developed plants were planted in field conditions for seed production.

RESULTS

Seeds germination started after 10–14 days of cultivation in vitro and continued for the next 15 days. All viable seeds germinated within four weeks after initiation.

The seeds of different accessions showed various effects (Fig. 1) of the used KMnO4 concentrations for seeds soaking. For alsike clover, best results were for soaking in 0.1 % KMnO4 solution, but on red clover the results obtained were similar for both used concentrations. Soaking in 0.1% KMnO4 solution decreased the percentage of infected seeds, depending on accession (data not shown).

For alfalfa, MS medium supplementation with AgNO3 or activated carbon did not significantly influence germination rate (Fig. 2). In addition, many plantlets obtained on the MS media supplemented with AgNO3 were soft, without roots and/or leaves and with tendency to calli formation (Fig. 3).

The highest germination rate for alfalfa was observed after pre-treatment with 0.1% KMnO4 solution for 40 minutes (Fig. 4). Accession ‘Lucerna Nr. 2’ had the highest germination rate after pre-treatment with 0.07% KMnO4 for 60 minutes, but in this case some of the seeds were contaminated. Therefore, pre-treatment with the highest concentration of KMnO4 (0.1%) for 60 minutes and MS basal medium without supplement was chosen as the best conditions for in vitro germination of aged alfalfa seed.

The percentage of germinated seeds differed significantly depending on accessions, and was not always correlated with seeds age (Table 1). The germination rate reached 71.8% (clover) and 62% (alfalfa). Tolerance to long-term storage at room temperature varied both among and within species — red clover ‘Vietējais vēlais’ and alsike clover ‘Savvaļas’, ‘Vietējais’ (k-33239) and ‘Vietējais’ (k-33240) had the period of storage (29 years), but their germination rates differed by almost up to six times. The germination rate for both 44-year-old alfalfa seed accessions Local k-31068 and Local k-31069 was similar, but it is interesting that the germination rate for these accessions was significantly higher than for seeds with storage age 14–20 years.

Most germinated seeds formed phenotypically normal plantlets with all organs. An exception was the variety ‘Stendes agrais’: 23% plantlets did not form normal leaves, but only roots and stem. The use of phytohormones did not improve seed germination and had a negative influence on plantlets quality — after transfer into soil only 29% of plantlets started to grow, while 86% of plantlets grew after cultivation on medium without phytohormones.

After four weeks of in vitro cultivation, clover plantlets (stage of 3–4 leaves) were used for micropropagation. It was
found that best results could be achieved by micropropagation of stem segments that were approximately 2 mm in length. Alsike accession ‘Viètejās’ developed axillary buds that were excellent explants for the second multiplication. From a plantlet, 2–7 new plantlets were obtained depending of genotype. After one month of cultivation, plantlets were ready for planting in soil or for the new cycle of multiplication. Positive results for micropropagation were not obtained only for accession ‘Stendes agrais’: stem segments started to grow but formed only non-regenerable calli.

DISCUSSION

The influence of different factors on in vitro germination rate of aged seed is known. One of the most important factors is seed sterilisation; the optimal method of sterilisation can considerably increase the seed ability for plant formation (de Sousa et al., 1999; Hamdouni, 2001; Yildiz and Er, 2002). It is also important to determine the optimal method of pre-treatment (Ornicãne and Rashal, 1997; Grauda and Rashal, 2004, Niedz, 2008; Sathish et al., 2011). Our results suggested that pre-treatment with potassium permanganate was effective for both seed sterilisation and germination stimulation; influence of genotype was observed in all used species. Cultivation medium and conditions are usually mentioned as main factors for seed in vitro germination and obtaining plantlets (Bhattacharya and Khuspe, 2001; Mauromicelle and Licandro, 2002; Klcová and Gubišová, 2003; Dutra et al., 2008; Greer et al., 2009; San and Yildirim, 2009; Mahamoodzadeh et al., 2010). Different effect of addition of silver ions or activated carbon to cultivation medium on the used genotypes was observed in our experiment.

Germination was successful for all accessions, even in the case of 44 years old seeds (alfalfa). It has been known for a long time that seed germination differs between genotype (Rowley, 1956), as also observed for maize (Revilla et al, 2007; Pereira et al., 2008; San and Greer, 2009). In vitro germination and dormancy responses of Hydrangea macrophylla and Hydrangea paniculata seeds to ethyl methane sulfonate and cold treatment. HortScience, 44 (3), 764–769. 


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TAURIŽIEŽU LATVIJAS IZCELSMES ĶEŅĒTİSKO RESURSU ATJAUŅOŠANA AR AUDU KULTŪRAS PALĪDZĪBU

Audu kultūras apstākļos panākta ūdu vecu aboliņu un lucernas šeiku izdīgšana, kuriem bija zaudēta diģstspēja parastās apstākļos. Iegūti fertili augi, kuri papildina pieejamos ķeņētiskos resursus attiecīgo kultūraugu selekcijas vajadzības.