Induction of Somatic Embryogenesis from Cotyledon of Oak
(Quercus castaneifolia)

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Abstract
Experiments were conducted on four species of Quercus (Q. brantii, Q. infectoria, Q. castanifolia and Q. libani). Cotyledon culture using MS medium containing BAP and IAA was applied for callus production. In the medium with BAP (1.0 mg/lit) and IAA (2.0 mg/lit) all species produced callus, however, in this medium callus was produced after 8 weeks, after 12 weeks globular embryogenic nodules were observed on the surface of Q. castanifolia calli and finally developed somatic embryos and plant regeneration. Stem and leaf culture did not produce any callus. No response was revealed in explants cultured on hormone free medium.

Keywords: Somatic embryogenesis, cotyledon culture, in vitro, Quercus.

Introduction
Several species of deciduous oaks including Quercus brantii, Q. infectoria, Q. castanifolia and Q. libani are grown in Iran. Quercus species are used as a source of economical products, and forest trees (Sharp et al. 1982). In vitro procedures for the induction and/or selection of useful mutants in plants are powerful tools for supplement conventional breeding methods (Williams & Maheswaran 1986). Quercus species require long time for crossbreeding and consequently, new technology such as somatic cell fusion (Sasmoto & Hosoi 1992) and somatic embryogenesis are shortening the breeding time (Williams & Maheswaran 1986, Mark et al. 1993).

Somatic embryogenesis is a method used for plant regeneration that potentially offers an efficient system for mass production of useful plants, e.g. transgenic plants (Das et al. 1993, Vonarnold et al. 1994). So far, somatic embryogenesis from zygotic tissue, stem, leaf segments, cell suspension and protoplasts from a few species of Quercus has been reported (Brawley et al. 1984, Gallego et al. 1997). Moreover, somatic embryos of immature embryos from several plants species, e.g. Q. acutissima (Kim et al. 1994), Q. robur (Cuenca et al. 1999), Zea mays (Armstrong & Green, 1985), and Trifolium repens (Maheswaran & Williams 1986) have been successfully obtained. However, this is the first report of somatic embryogenesis from mature cotyledons of Quercus. The objective of the present study was to evaluate the induction of somatic embryos from cotyledon explants for multiplication and possible genetic manipulation in a few Quercus species.

Materials and Methods
To evaluate the condition of embryogenesis, ten years old plants of Q. libanii, Q. infectoria, Q. castanifola and Q. brantii were chosen as source of explants. Mature seeds with a few months old were harvested during summer and sterilized with 10% NaCl solution containing 1-2 drops of Tween 80 for 20 min followed by rinsing with sterile distilled water. In the same experiments, to reduce the contamination, seeds were first sterilized with 30% H2O2 for 3 min, then with 70% Ethanol for 1 min, followed by washing in sterile water. Sterile seeds were uncoated under the air flow and zygotic embryos removed. Cotyledons were excised and cut into 8 mm segments and were transferred to Murashige and Skoog (1962) (MS) media supplemented with plant growth regulators including 1) BAP (1 mg/lit) and IAA (0.5 mg/lit), 2) BAP (1 mg/lit) and IAA (2 mg/lit), and 3) hormone free. All culture media prepared with 3% sucrose, 1% agar and pH was adjusted to 5.8. The culture media were supplemented...
with activated charcoal (0.5 mg/lit) or citric acid (20 mg/lit) for removing phenolic compounds. Explants were grown in the culture room under the 16/8 h light/dark photoperiod at approx. 25°C. Explants were transferred to the fresh medium every two weeks. In the separate experiments stem and leaf segments obtained from in vitro grown seedlings were cultured on the same media. Treatments arranged in a Randomised Block Design (RBD) with 10 replications (3-4 cotyledon segments or leaves and stems in each replication). All data were analyzed using SPSS program.

**Results**

The first step in achieving indirect somatic embryos from cotyledon explants was induction and consequently production of callus from cotyledons. In the different media supplemented with variety of plant growth regulators, one of the major problems at this stage was the releasing of browning components from explants into the culture medium. This material has been known to suppress the growth and development of the explants under in vitro culture of some plant species (Madhusudhanan & Rahiman 2000). Although, using of activated charcoal was effective in growing of the explants and reduced the concentration of the brown material but, when activated charcoal was replaced with citric acid, the responses of the explants were significantly improved and cotyledon showed better growth and development. Stem and leaf segments from germinated seedlings of four Quercus species (see section 2) failed to produce callus and somatic embryos. Callus growth was greater in medium supplemented with BAP (1 mg/lit) and IAA (0.5 mg/lit). However, cotyledons of Q. castaneifolia were the most responsive explants. Whereas, lower than 5% of explants cultured on hormone-free MS medium produced callus. About 7-8% of Q. castaneifolia cotyledons produced compact embryogenic nodules within 10-12 weeks in medium with BAP and IAA (1.0 and 2.0 mg/lit respectively) (Fig. 1). Finally, a few embryonic calli from Q. castaneifolia were developed further to plantlet (Fig. 2).

**Fig. 1- Ability of cotyledons of four Quercus species to produce callus when they were cultured in MS medium without plant growth regulators (A), 1mg/lit BAP and 0.5mg/lit IAA (B), 1mg/lit BAP and 2mg/lit IAA (C). Common letters are not significant according to Duncan test (p<0.05).**

**Discussion**

Somatic embryogenesis is one of the best known method for plant regeneration for multiplication or/and genetic manipulation in plants (Williams & Maeswaran 1986). Factors influencing somatic embryogenesis has already been reviewed for example, disruption of normal tissue interrelationship (Sharp et al. 1982), physiological state of the cells undergoing somatic embryogenesis, and ionic polarity of the cells (Williams & Maeswaran 1986).

We observed that indirect somatic embryogenesis from cotyledon culture of Quercus species followed by at least a short period of cell proliferation as callus before appearance of embryonic compact nodules. Lack of potential of callus production from stem, and leaf segments culture of four species of Quercus in this investigation indicating that leaf and stem segments failed to produce in vitro somatic embryos.

However, in contrast to our data induction of indirect somatic embryos from in vitro culture of leaf discs of red Oak (Q. rubra) has already been reported (Rancillac et al. 1996). It is indicating that somatic embryogenesis from leaf and stem in Quercus is highly depending on the genotype, combination of the medium and
environmental condition of the culture (Sharp et al. 1982).

Evaluation of different media showed that, only cotyledons of *Q. castaneifolia* were able to produce true embryonic callus on medium with BAP and IAA (0.5 and 1.0 mg/l), while cotyledon culture on medium containing 1 mg/l BAP and 2 mg/l IAA produced non-embryonic calli. The exact factor/s responsible for better response of this species is unknown but, at least adaptation of this genotype with the culture condition and expression of the gene/s responsible for somatic embryogenesis might be critical factors (Evans et al. 1978, Foulger & Jones 1986). In the present study cotyledon culture of *Q. castaneifolia* at the first step produced embryogenic nodules on the surface. Further development of embryonic granules to the globular and heart stages is showing the totipotency of this species for mass production of the somatic embryos.

The role of phytohormones in induction of somatic embryogenesis is also of interest in this context. For example, Maheswaran et al. (1986) have reported that using BAP as an only source of plant growth regulator for induction of direct somatic embryogenesis, in immature embryo culture of *Trifolium repens*. The same pattern of responses has also been reported in red clover (Gregory & Collins 1980). It appears that the type and concentration of plant growth regulators and possibly influence of external factors (Guijarro et al. 1995) may interact with factors, including ionic currents, cell activity, cell polarity and gene expression and consequently induce somatic embryos in explants (Brawley et al. 1984, Pretova et al. 1994, Gallego et al. 1997). However, factors affecting somatic embryogenesis in explants at the molecular level need to be studied in details in the future.

Fig. 2- Different stages of somatic embryo development from cotyledon culture of *Quercus castaneifolia*. A- Callus production. B- Proembryonic mass. C- Globular stage. D- Heart stage. E- Plantlet.

References
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