MORPHOGENIC PROCESSES IN CALLUS TISSUE CULTURES AND DE NOVO REGENERATION OF PLANTS IN ACTINIDIA CHINENSIS PLANCH.

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ABSTRACT

Our experiments have confirmed the considerable disposition of leaf explants of Actinidia chinensis Planch. for induction and intensive proliferation of callus cultures, as well as, a possibility to regulate morphogenesis in in vitro conditions. Under specific culture conditions the morphogenic potential of callus cells of Actinidia chinensis was manifested both in organogenesis and somatic embryogenesis. Organogenesis was represented by induction of adventitious buds and regeneration shoots on the modified MS culture medium (Murashige and Skoog 1962) with BAP in combination with GA3 (each 1.0 mg. 1-1). Rooting of shoots was successful on modified MS medium containing IBA (0.5-1.0 mg. 1-1).

Histological studies of callus tissues revealed their structural heterogeneity. Morphogenic processes in the callus were characterized by the appearance of meristematic zones and vascular elements. The formation of apical meristem, leaf primordia and finally shoot development proved de novo regeneration in callus culture.

The results demonstrate a possibility of plant regeneration through indirect organogenesis, which can be used for propagation of Actinidia chinensis Planch.

KEY WORDS: Actinidia chinensis, leaf explants, callus culture, morphogenic responses.

INTRODUCTION

Callus cultures are excellent objects for the study of differentiation of cells and tissues. An important aspect of callus cells and tissues is the ability to regenerate plants. Many woody species are difficult to regenerate from callus cultures and are propagated for commercial purpose by apical and axillary buds proliferation. Certain plant species are able to realize the in vitro process of dedifferentiation and redifferentiation, but this ability is often limited, especially in forest trees.

The successful callus induction and realization of morphogenic responses (organogenesis, somatic embryogenesis) in callus tissue cultures depend on the plant species and on the genotype of the donor plant and even on an explant type. Composition of the culture medium, mainly the content of growth regulators influence the mentioned processes to a considerable degree. Endogenous phytohormones present in explants play also a great role. Many authors (Opatrný 1983, Bonga, Aderkas 1992, Georgie 1993 and others) emphasized the mechanism of growth regulators effect on the regulation of the morphogenesis in tissue cultures.

Several recent studies have confirmed the morphogenic potential of callus cultures derived from leaves of trees (Jones 1993) and also presented an efficient adventitious shoots development from callus cultures (Fasolo et al. 1989, Escalettes, Desba 1993, Economou and Maloupa 1995).

This paper is intended to present the effect of growth regulators on induction of callus culture, observations of the morphogenetic responses in callus cultures derived from leaf explants of Actinidia chinensis and also the de novo in vitro plant regeneration. Histological investigations of the structural changes in callus tissue cultures and alternatives of morphogenetic response are illustrated.

MATERIAL AND METHODS

Leaf explants were separated from shoot cultures obtained from axillary buds of about 5-yr old plants (from the Botanical Garden of the Slovak Agricultural University in Nitra). Leaf segments were placed with overleaf into Petri dishes (25 ml/dish) with a medium. Basal modified medium of Murashige and Skoog (1962) (with a half concentration of macroelements) supplemented with 3% (w/v) of sucrose and 6% (w/v) of agar was used. Experimental variants the MS medium were supplemented with BAP, 2,4-D, GA3 (each 1.0
mg. 1⁻¹) as sole growth regulators or combinations of BAP with 2.4-D, BAP with GA₃ and 2.4-D with GA₃ (each 1.0 mg. 1⁻¹) were used. pH of culture media was adjusted to 5.6-5.7. Cultures were incubated at 25-28°C and a 16/8 hour photoperiod with light intensity of 60 µE.m⁻².s⁻¹. After one month, when callus was induced, the explants with callus were transferred into a 100 ml conical flasks (containing 25 ml of the medium). The cultures were further monthly subcultured on fresh media.

Shoots regenerated from callus tissue cultures were transferred on a rooting medium containing IBA (0.5-1.0 mg. 1⁻¹). The regenerated plantlets were transferred into a substrate consisting of perlite and peat (1:2) and placed in a greenhouse.

For histological studies the callus tissue samples (derived on the medium with BAP and GA₃) after one month and after three months of culture were fixed in FAA (70 w/v ethanol; glacial acetic acid; formalin – 18:1:1), dehydrated (ethanol-xylool) and embedded in paraffin wax. Sections (10-13 µm thick) were stained with Ehrlich’s hematoxylin.

RESULTS AND DISCUSSION

The micropropagation of the species Actinidia chinensis Planche was obtained in in vitro conditions by cultivation of different explants - root, zygotic embryos, stamens and first of all apical and axillary buds (Harada 1975, Wang et al. 1982, Brossard-Chiriqui, Tripathi 1984, Standardi 1983, Monette 1986, Kamenická, Rypák 1989, Cholaková 1990, Ostrovčuk 1993). The results of the above authors and our own results have confirmed a high regeneration potential and considerable flexibility of Actinidia chinensis in morphogenic responses in comparison to other woody plants.

The presence of growth regulators in the culture medium has considerably affected the morphogenic processes. The importance of growth regulators for the induction of cell morphogenic response was confirmed by the fact, that leaf segments of Actinidia chinensis cultured on a modified MS medium without growth regulators failed to induce callus and gradually necrotized. The effects of individual growth substances and their combinations (in concentration of 1 mg. 1⁻¹) on callus induction and on the character of morphogenic processes were different (Table 1).

The initiation of callus induction occurred on the overleaf segments and on their margins on the 6th-10th day. The callus overgrew the whole surface of the primary explant. Callus formation was lower on the medium containing sole growth regulator (with BAP - 30%, with GA₃ - 68%) (Table 1). The 30% frequency of initiation on BAP was the lowest one obtained in our experiments. The combination BAP and GA₃ gave a higher initiation frequency (86%). Cultures growing on the medium with combined BAP and GA₃ retained their viability for a long time what can be ascribed as a positive effect of GA₃. The medium with combined BAP and 2.4-D appeared to be the most efficient (98%) in callus induction (Fig. 1). Similarly some authors have found that callus induction depends on the presence of auxins in combination with cytokinins in culture media (Kamenická, Rypák 1989, Bonga, Aderkas 1992). Monette (1986) reported that callus proliferation in Actinidia chinensis was markedly stimulated by BAP in combination with NAA.

At the same time we observed a specific effect of individual growth substances on the callus consistence and colour (Table 1). Callus on media containing BAP, GA₃ and BAP in combination with 2.4-D was white, of compact consistence. On a medium with 2.4-D, BAP in combination with GA₃ and 2.4-D with GA₃ callus was white, later light green to green and friable consistence. On a medium with BAP and GA₃ reddish areas appeared on the callus which is supposed to be anthocyanins synthesis manifestation. We found that the callus consistence can change after changing the medium composition and longer culture. For instance the friable callus induced on the medium 2.4-D became compact after three months of culture on the medium with BAP and 2.4-D. A regular subculturing

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### Table 1. The callus initiation and variability of morphogenic response in the callus cultures of Actinidia chinensis

<table>
<thead>
<tr>
<th>Morphogenic response</th>
<th>Variants of culture media</th>
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<tr>
<td></td>
<td>Growth substances and their combinations (each 1.0 mg = mg. 1⁻¹)</td>
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<td>BAP</td>
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<td>Callogenesis</td>
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<td>The consistency of callus</td>
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<td>The colour of callus</td>
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<td>Other responses</td>
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<td>Organogenesis</td>
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<td>Somatic embryogenesis</td>
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A - number of primary explants (leaf segments)
B - primary explants with callus initiation (%)
C - MS medium without growth substances
CC - callus of compact consistency
FC - callus of friable consistency
W - callus of white colour
LG - callus of light green colour
G - callus of green colour
AN - anthocyanins synthesis
R - differentiation of roots
S - differentiation of shoots
E - embryogenic structures
Fig. 1. Induction of callus from leaf segments on modified MS medium supplemented with BAP and 2,4-D (each 1 mg. l\(^{-1}\)) after 30 days cultivation.

Fig. 2. Callus mass derived from leaf segments on modified MS medium with BAP and GA\(_3\) (each 1.0 mg. l\(^{-1}\)) after three months of culture and formation of adventitious buds in callus.

Figs 3, 4. Shoot differentiation from leaf callus and their development on MS medium containing BAP a GA\(_3\) (each 1.0 mg. l\(^{-1}\)).

Fig. 5. The rooted plantlets two months after transplantation into substrate.

Fig. 6. Direct differentiation of roots on the main vein of leaf explant on modified MS medium with BAP and 2,4-D and GA\(_3\) (each 1.0 mg. l\(^{-1}\)).

Fig. 7. Embryogenic callus derived from leaf segments on modified MS medium supplemented with BAP a 2,4-D (each 1.0 mg. l\(^{-1}\)).

Fig. 8. Further development of embryogenic callus – numerous globular somatic embryos.
Morpho-histological study of organogenic callus induced from leaf segments on modified MS medium with BAP and GA₃ (each 1.0 mg. L⁻¹) (Figs 9-17).

Fig. 9. Vacuolated parenchymatic cells of irregular shape in callus after 30 days of cultivation.
Fig. 10. Morphological variability of parenchymatic cells with conspicuous nuclei observed in callus after three months of cultivation.
Fig. 11. Accumulation of starch grains (S) in callus cells.
Figs 12-13. Formation of vascular elements (V) in callus tissue.
Figs 14-15. Indirect organogenesis is illustrated by occurrence of meristematic zones (M), formation of apical meristem (M) and of leaf primordia (L).
Fig. 16. Development of shoot of adventitious origin.
Fig. 17. Differentiation of multicellular globular structures.
(every 30 days) favourably affected the continuous growth of the callus culture in all variants of culture media.

A possibility to regulate morphogenesis in in vitro conditions has been confirmed in our experiments (Table 1). The leaf explants of the species Actinidia chinesis manifested an ability not only for initiation of the callus culture, but also for redifferentiation of induced callus. Under certain culture conditions the callus cells morphogenetic potential was manifested both towards organogenesis (Figs 1-4) and somatic embryogenesis (Figs 6-8). A favourable effect of growth regulators in their synergic activity was confirmed. In some parts of the callus on the medium BAP in combination with 2,4-D root formation occurred and also differentiation of roots have been observed on the medium with GA3 and 2,4-D, directly along the main midrib (Fig. 5). During root growth a polarity disturbance – negative geotropism was observed, which was also recorded by Kamenická and Rypák (1989) in Castanea sativa. We think that the stimulatory effect on the rhizogenesis induction can be ascribed to 2,4-D. Vieitez et al. (1978) reported a positive effect of BAP on the rhizogenesis process in Castanea sativa.

The morphogenic potential and regeneration ability of callus culture enabled adventitious buds induction and subsequent regeneration of shoots of fruit trees and other woody species (Miguel et al., 1996; Vieitez, San-José, 1996). The callus culture of Actinidia derived from leaves can also be used for propagation because it is very effective for induction of adventitious shoots. Our experiments revealed that numerous buds and adventive shoots were developed on a medium containing BAP and GA3 (Figs 3, 4). We recorded an efficient proliferation of shoots in the callus culture of Actinidia chinesis. Each callus produced 15-20 shoots. Kamenická and Rypák (1989) reached multiplication of Actinidia shoots on a medium containing BAP (1.0 mg, 1%) and GA3 (2.0 mg, 1%) through a direct organogenesis. Cholvadová (1990) points to an efficient effect of zeatin in the organogenesis process in Actinidia.

Regenerated shoots (from 2 to 2.5 cm in height) were excised and transferred on a rooting medium (MS with a half concentration of macroelements) containing IBA (0.5-1.0 mg, 1%) where the rooting was successful (to 92-98%). The obtained regenerated plants are able of adaptation and they also grow intensively in in vivo conditions (Fig. 5).

The proliferation of embryogenic callus and differentiation of embryogenic structures (Figs 7, 8) were observed on the medium with BAP in combination with 2,4-D. Embryogenic structures on the level of globular and torpedo stages were attacked by an infection after subculturing in consequence of which we did not succeed in obtaining somatic embryos. The positive BAP and 2,4-D effect on differentiation of somatic embryos has been recorded e. g. in oak (Ostrolucká and Krajmerová 1996).

The obtained results demonstrate a reproducible plant regeneration system of indirect organogenesis and the Actinidia callus culture ability of redifferentiation and de novo regeneration. This system enables a high plant production, but on the other hand does not guarantee the genetic stability of regenerants.

Histological investigations of callus tissue induced on BAP medium in combination with GA3 showed that after 30 days the callus consisted of parenchymatous cells with intercellular spaces. Parenchymatous cells were irregularly shaped, of different size and highly vacuolated (Fig. 9).

The studies on callus tissues revealed their structural heterogeneity. Different parts of the callus gave different histological patterns. Parenchymatous cells of different size, with one and sometimes even with two nuclei appeared what can indicate a different degree of their ploidy. Strongly vacuolated cells were also observed (Fig. 10).

Further observations of intensively proliferating callus (carried out three months later) on BAP medium with GA3 have confirmed the heterogenous morphogenic activity of some parts of the callus, whereas other parts remained amorphous without organogenic manifestations. In some callus cells many starch grains appeared (Fig. 11). We suppose that these starch grains are a source of reserve substances and are important in the process of organogenesis. The anatomical studies of shoot development of kiwifruit long term callus culture were carried out by Jäskil and Lux (1992). They have recorded in detail the starch synthesis in the organogenic callus.

Morphogenic processes were correlated with the changes in callus culture. The differentiation of callus cells led to the formation of numerous tracheads of different length and tracheal zones (Figs 12, 13) as well as of meristematic centres (Fig. 14). The presence of vascular elements in a non-organized parenchymatous tissue enables the transport of growth regulators and nutrients and supports local activation of cell division. Opatrný (1977) reported that a distinct differentiation of tracheads occurs during the so-called process of an active state of the tissue and the appearance of trachead elements can be the first degree of regeneration of organs in a non-organized tissue. According to the author, the active state of the tissue – the cell division is preceded by utilization of the starch grains in cells.

The formation of apical meristem, leaf primordia and multiple shoots illustrates the regeneration in callus cultures (Figs 15, 16, 17).

LITERATURE CITED


PROCESY MORFOGENETYCZNE W KULTURACH TKANKOWYCH KALUSA I REGENERACJA ROŚLIN U ACTINIDIA CHINENSIS PLANCH.

STRESZCZENIE

Stwierdzono wysoką zdolność regeneracyjną Actinidia chinensis w warunkach in vitro. W niniejszych badaniach wykazano: dużą zdolność eksplantatów z liści Actinidia chinensis do indukcji, intensywne rozmnazanie się kultur kalusa oraz możliwość regulowania morfogenety z warunków in vitro. W określonych warunkach kultury potencjał morfogenetyczny kalusa. Actinidia chinensis przejawiał się zarówno w organogenezie, jak i somatycznej embriogenezie. Organogeneza polegała na indukcji pąków przybyszowych i regeneracji pedów na zmodyfikowanej pożywce MS (Murashige i Skoog 1962) z dodatkiem BAP w kombinacji z GA3 (po 1,0 mg. l⁻¹). Ukorzenianie się pedów było możliwe na zmodyfikowanej pożywce MS zawierającej IBA (0,5-1,0 mg. l⁻¹). Badania histologiczne tkanki kalusa wykazały ich strukturalną niejednorodność. Procesy morfogenetyczne charakteryzowały się pojawianiem się stref merystematycznych i elementów naczyniowych. Tworzenie się merystemu wierzchołkowego, zawiaszków liści oraz rozwój pedów potwierdziło zachodzenie regeneracji w kulturze kalusa. Uzskształcone rezultaty prezentują wydajną system regeneracji roślin poprzez poszczególną organogenezę, który wykorzystać można do rozmnazania Actinidia chinensis.

SŁOWA KLUCZOWE: Actinidia chinensis, eksplantat z liści, kultura kalusowa, reakcje morfogenetyczne.