Plant Regeneration from Protoplasts Isolated from Callus of Taro

(Colocasia esculenta Schott)

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Summary
A system of protoplast isolation, callus formation, and plant regeneration in taro (Colocasia esculenta Schott cv. Eguimo) was established.

1. Friable calli were induced by culturing etiolated stem segments of taro on Murashige and Skoog (MS) medium supplemented with 30 g liter\(^{-1}\) sucrose, 2 mg liter\(^{-1}\) 2,4-D and 2 mg liter\(^{-1}\) 2ip. Calli were maintained by subculturing on a fresh MS medium.

2. Protoplasts were easily isolated from suspension cells derived from the friable calli. The component of the enzyme solution for the isolation consisted of 1 g liter\(^{-1}\) Pectolyase Y-23, 5 g liter\(^{-1}\) Cellulase Onozuka RS, 5 mM MES, 5 mM CaCl\(_2\)•2H\(_2\)O, and 0.5 M mannitol.

3. Isolated protoplasts were cultured in the liquid media consisting of half strength MS inorganic salts, Kao and Michayluk's (1975) organic substances, various levels of NAA and BA, 0.1 M glucose and 0.3 M mannitol. Numerous colonies were formed in the medium containing 2 mg liter\(^{-1}\) BA.

4. Shoot regeneration from protoplast-derived calli occurred on solid MS medium with 0.2 mg liter\(^{-1}\) NAA and 2 mg liter\(^{-1}\) BA. The shoots initiated roots after being transferred to a MS medium without phytohormones.

The protoplast culture system established in this study might be useful for cell fusion and electroporation of genes as an approach to breeding of taro.

Chemical names used: 2-N-Morpholino ethanesulfonic acid (MES); 1-naphthanlenacetic acid (NAA); 2,4-dichlorophenoxy acetic acid (2,4-D); N-phenylmethyl-1H-purin-6-amine (BA); N-3-methyl-2-butenyl-1H-purin-6-amine (2ip).

Introduction

Taro, Colocasia esculenta Schott, is an important vegetable crop and staple food in Asia, Africa, and in the Pacific Islands. The use of cross-breeding method has been limited in this species, because most of the cultivars are triploid and, even in diploid plants, flower initiation is very difficult. Therefore, somatic hybridization through protoplast fusion seems to be a possible means of genetic improvement. A prerequisite for a cell fusion technique is the establishment of a culture system for plant regeneration from protoplasts. However, taro is one of the most difficult crop species for protoplast culture. Oosawa and Takayanagi (1984) first reported culture of protoplasts isolated from taro leaf, but callus formation and plant regeneration from protoplasts have not been successful.

In monocotyledonous plants, calli or suspension cultured cells have been frequently used as a source of protoplasts. In taro, the growth of callus derived from shoot-tip was limited (Jackson et al., 1977; Nyman et al., 1983), whereas callus obtained from etiolated stem segments grew vigorously (Murakami et al., 1992). Therefore, we used calli obtained from stem segments. In the present paper, we describe a system of protoplast culture, including donor callus formation, protoplast isolation and culture, and plant regeneration.
Materials and Methods

Callus initiation and culture

Shoot apices of taro cv. 'Eguno' were cultured on a basal medium supplemented with 0.2 mg·liter⁻¹ NAA. Etiolated stems were obtained by culturing these aseptic plantlets on the medium for 30 ~ 60 days in the dark (Fig. 1). The basal medium consisted of inorganic salts (Murashige and Skoog (MS), 1962) plus (as mg·liter⁻¹): 100 myo-inositol, 2 glycine, 0.5 nicotinic acid, 0.5 pyridoxine·HCl, 0.1 thiamine·HCl, 30,000 sucrose, and 2,000 Gelrite. Stem segments of 5 mm were cultured on a basal medium containing 2 mg·liter⁻¹ 2,4-D plus 2 mg·liter⁻¹ 2ip at 25°C and 16 hr daylength with a light intensity of 20 μ mol·sec⁻¹·m⁻² from a fluorescent light to obtain friable calli (Murakami et al., 1992). Vigorous and friable calli were selected and subcultured after 60 days on a fresh medium for further proliferation. Callus was then subcultured monthly on a fresh medium for 8 months.

Protoplast isolation and culture

Two to three grams of calli were cultured in a 30-ml liquid medium containing 2 mg·liter⁻¹ 2,4-D and 2 mg·liter⁻¹ 2ip under dim light conditions on a rotary shaker adjusted to 100 rpm at 25°C. Suspended cell aggregates which formed after 10 ~ 15 days were filtered through a 500 μm mesh stainless sieve, and isolated on a 45 μm stainless sieve.

These cell aggregates of 45 ~ 500 μm were re-suspended in a 20-ml reaction mixture containing 1 g·liter⁻¹ Pectolyase Y-23, 5 g·liter⁻¹ Cellulase Onozuka RS, 5 mM 2-N-Morpholino ethanesulfonic acid (MES), 5 mM CaCl₂·2H₂O and 0.5 M mannitol. The suspension was shaken on a rotary shaker adjusted to 100 rpm for 3 hours at 25°C. Undigested cell clumps were removed by passing the solution through a 63 μm stainless sieve. The protoplasts were collected by centrifugation and washed three times with a solution of 5 mM MES, 5 mM CaCl₂·2H₂O and 0.5 M mannitol.

A protoplast culture medium consisting of a half strength MS inorganic salts, Kao and Michayluk's (1975) (KM) organic substances, various phytohormones, 0.3 M mannitol and 0.1 M glucose were used, because KM organic substances were found to be suitable for taro protoplast culture (Murakami and Matsubara, 1992), and shoot regeneration from callus was observed by subculturing them on MS medium supplemented with BA or with NAA and 2ip (Murakami et al., 1992). In the present study, therefore, the effects of BA and 2ip or the combination of NAA and BA at various concentrations were tested.

The two experiments were conducted in triplicate. In Experiment 1, isolated protoplasts were plated on media with BA or 2ip at 0.2, 2, 5 mg·liter⁻¹. In Experiment 2, the protoplasts were plated on media containing 0, 0.2, 1, or 5 mg·liter⁻¹ NAA plus 2 mg·liter⁻¹ BA. The media were sterilized by filtering through 0.2 μm cellulose nitrate membranes. The isolated protoplast equivalent to 2 × 10⁵ cells per ml were plated on 2 ml liquid media in 60 mm diameter plastic petri-dishes. These petri-dishes were sealed with Para-
film and kept in the dark at 25 °C for 40 days.

Two weeks after plating, 10 microscopic fields selected at random per petri dish were examined and the plating efficiency (percent of divided cells per total number of protoplasts) was recorded.

Regeneration of plant

Calli proliferated from protoplasts on the medium supplemented with 2 g·liter⁻¹ BA were transferred to a basal medium containing 2 mg·liter⁻¹ BA plus 0, 0.2 or 2 mg·liter⁻¹ NAA. Formation of green protocorm-like structures was observed 60 days later. Regenerated shoots were transferred to a basal medium without phytohormone to promote shoot growth and root formation.

Results

Protoplast isolation and the plating efficiency

The fine and friable calli which were obtained after subculturing (Fig. 2) were gradually dispersed in a liquid medium by shaking for 10–15 days. The average yield of protoplasts was 3.8 × 10⁶ per g of callus. Considerable variation was observed in the size of protoplasts (Fig. 3), but most of them contained many small granules, similar to the protoplasts of asparagus (Elmer et al., 1989) and rice (Thomson et al., 1986) derived from suspension cells.

The tests of various levels on BA and 2ip concentrations on the plating efficiency (Exp. 1) revealed that 0.2 mg·liter⁻¹ BA was optimum (Table 1), but that the subsequent colony growth on this medium was poor (data not shown). Good colony growth was obtained on the medium with 2 mg·liter⁻¹ BA. 2ip was generally less effective on plating efficiency than BA. Combining NAA at various levels with 2 mg·liter⁻¹ BA (Exp. 2) revealed that plating efficiency slightly increased by adding 0.2 mg·liter⁻¹ NAA, compared to a medium with 2 mg·liter⁻¹ BA alone (Table 2). However, increasing the levels of NAA lowered the efficiency.

Table 1. Effects of BA and 2ip on the division of taro protoplasts (Experiment 1).

<table>
<thead>
<tr>
<th>Phytohormone treatment²</th>
<th>Plating efficiency³ (%)</th>
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<tbody>
<tr>
<td>BA (mg·liter⁻¹)</td>
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<tr>
<td>0.2</td>
<td>1.6</td>
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<tr>
<td>2.0</td>
<td>1.3</td>
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<tr>
<td>5.0</td>
<td>0.9</td>
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<tr>
<td>2ip (mg·liter⁻¹)</td>
<td></td>
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<tr>
<td>0.2</td>
<td>0.5</td>
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<tr>
<td>2.0</td>
<td>0.9</td>
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<tr>
<td>5.0</td>
<td>0.8</td>
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² NAA was not added to all media.
³ Percent of dividing cells per total protoplasts after 14 days of culture.

Table 2. Effects of NAA and BA on the division of taro protoplasts (Experiment 2).

<table>
<thead>
<tr>
<th>Phytohormone treatment</th>
<th>Plating efficiency³ (%)</th>
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<tbody>
<tr>
<td>NAA (mg·liter⁻¹)</td>
<td></td>
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<tr>
<td>0</td>
<td>2.0</td>
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<tr>
<td>0.2</td>
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<tr>
<td>1.0</td>
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<tr>
<td>5.0</td>
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² See Table 1.

Fig. 2. Friable callus cultured on MS medium supplemented with 2 mg·liter⁻¹ 2,4-D and 2 mg·liter⁻¹ 2ip.

Fig. 3. Freshly isolated protoplasts.
Plating efficiency on the medium with 2 mg·liter⁻¹ BA was higher than that of the same medium in Exp. 1. This may be due to the morphological difference of donor callus between two experiments. Donor callus used for Exp. 2 was softer and more homogenous than that used for Exp. 1.

**Plant regeneration**

Small colonies were formed after 21 days of culture (Fig. 4) which grew to 1~2 mm in diameter after 40 days (Fig. 5). Green protocorm-like structures were formed after transferring the calli to solid regeneration medium. More than half of the calli on 0.2 mg·liter⁻¹ NAA plus 2 mg·liter⁻¹ BA produced protocorm-like structures (Table 3). After these protocorm-like structures were transferred to a fresh medium several times, 20% of them formed adventitious shoots. Thus, about 10% of the transferred calli developed adventitious shoots of which 40% regenerated additional multiple shoots upon subculturing (Fig. 6). Each shoots were separated from multiple shoots and transferred to the basal medium. They grew slowly after transfer, and 3 of the 20 transferred shoots (15%) formed roots (Fig. 7). It took almost 1 year to obtain plantlets since initial culture of protoplasts.

**Discussion**

The type of callus for protoplast source influences the efficiency of protoplast culture rather than the cultural conditions after the isolation (He et al., 1992; Kasem and Sagi, 1993; Morocz et al., 1990; Vasil, 1987). Callus for protoplast source is required to have the following characteristics: fast-growing, friable and regenerative. Protoplasts

<table>
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<th>Phytohormone treatment</th>
<th>Calli forming green protocorm-like structures (%)</th>
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</thead>
<tbody>
<tr>
<td>NAA</td>
<td>BA (mg·liter⁻¹)</td>
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<tr>
<td>0</td>
<td>2</td>
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<td>0.2</td>
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*Calli developed from protoplasts on the medium supplemented with 2 mg·liter⁻¹ BA were used as explants.*
isolated from fast-growing callus develop colonies at a high rate. Friable callus easily disperses in liquid medium or enzyme solution. The regenerating ability of callus is the most important character to obtain plantlets. Vasil (1987) reported that regenerative protoplasts were isolated from regenerable calli or suspension cells (Vasil, 1987). In the present study, we obtained fast-growing and friable calli which we used for isolating protoplasts. Although induction of such vigorous callus from shoot-tip has been difficult, we succeeded in its induction by using etiolated stem segments as the initial callus explants.

Our system of protoplast culture was a simple, the protoplasts being cultured in liquid medium. Nurse culture technique, conditioned medium, agarose and other gelling agents, which were developed for the efficient protoplast culture, were not used. Nevertheless, colonies were easily developed from protoplasts in this medium which suggests that the source of the protoplasts is more important than the cultural conditions.

In the present study, however, it took an extended period to regenerate plants from protoplasts and the frequency of regeneration was low. This might be due to low regenerative ability of the callus used as protoplast source. Continuous selection of faster growing and non-regenerative callus for a long period may cause a decline of regeneration ability. Such a repeated long-term subculture also may induce gene mutations. Therefore, further study is necessary to reduce the period for callus establishment and to prevent a decline of regeneration ability. Recently, formation of embryogenic callus became possible in taro (Karube et al., 1992). Use of the embryogenic callus for protoplast source may be an approach to increase the frequency of regeneration. On the other hand, there are some reports concerning the promotion of regeneration by pretreatments of callus. For example, culturing callus in hormone-free medium promoted regeneration from protoplasts in asparagus (Kunitake and Mii, 1990) and Phalenopsis (Kobayashi et al., 1993). Thus, it would be necessary to examine various callus pretreatments for promotion of regeneration also in case of taro. By improving the frequency of plant regeneration from protoplast in the future research, the protoplast culture system described here may be applicable to the breeding of taro using the techniques such as cell fusion and electroporation.

Acknowledgements

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Literature Cited


