The effects of the physical characteristics of the culture medium on the development of red seaweeds in tissue culture

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Abstract

Explants of Gelidium versicolor, Grateloupia doryphora and Laurencia sp. were cultivated in Provasoli enriched seawater culture medium (PES) adjusted to several osmolalities (0.5, 0.7, 1.0 and 1.5 Os kg⁻¹) and solidities (agar concentration = 3, 8 and 15 g L⁻¹). Osmolality was adjusted by dilution of seawater with distilled water (50, 70 and 100% seawater) and by NaCl addition. Explants of Laurencia sp. and Grateloupia doryphora showed bud regeneration and callus formation. Explants of Gelidium versicolor only showed bud regeneration. Osmolalities of 0.5 and 1.05 Os kg⁻¹ inhibited or drastically reduced bud regeneration and callus formation. The highest callus formation and bud regeneration were observed at 0.7 to 1.0 Os kg⁻¹. An increase in the agar concentration of the culture medium was positively correlated with callus formation and negatively correlated with bud regeneration. An increase in the percentage of seawater increased the solidity of the culture medium and was positively correlated with callus formation. Glycerol was an effective carbon source for the vegetative propagation of axenic explants of Grateloupia doryphora, promoting growth and bud regeneration. An increase in glycerol concentration in the culture medium increased its osmolality, inhibiting the growth of the explants and their morphogenetic development.

Introduction

The control of cell growth and development is still a problem in seaweed cell and tissue cultures (Saga et al., 1986; Polne-Fuller & Gibor, 1987). Some authors have reported the formation of callus and callus-like structures induced by the semi-solid (agarized) state of the culture medium (Fries, 1980; Saga et al., 1982; Saga & Sakai, 1983; Polne-Fuller & Gibor, 1984; Garcia-Reina et al., 1988). The solidity of the culture medium seems to be involved in callus induction, since other gelling compounds produced the same effects as agar (Polne-Fuller & Gibor, 1987).

The role of carbohydrates in the growth and organogenesis of seaweed cell cultures remains equivocal (Lawlor et al., 1988). The addition of carbohydrates to the culture medium, without controlling osmolality, can obscure their role in seaweed growth.

The aim of the present work was to study the effects of solidity and osmolality of culture medium on callus induction and bud regeneration of several red seaweeds, as well as the effects of
glycerol addition on the osmolality of the culture medium and as a carbon source.

Materials and methods

_Gelidium versicolor_ (S. G. Gmelin) Lamouroux, _Grateloupia doryphora_ (Montagne) Howe and _Laurencia sp._, an intertidal alga similar to _Laurencia obtusa_ (Hudson) Lamouroux, were collected in Gran Canaria (Canary Islands). Voucher specimens are deposited in the herbarium of the Jardín Canario, Gran Canaria, Canary Islands, Spain (LPA), as sheets LPA 30 (_Laurencia sp._), LPA 110 (_Gelidium versicolor_) and LPA 129 (_Grateloupia doryphora_).

Two hours after collection, explants were excised from apparently epiphyte-free, highly pigmented and sterile thalli. _Laurencia sp._ explants were apical secondary branches 1 cm long; _Gelidium versicolor_ explants were cylinder-shaped fragments (0.5 cm long) excised from the middle zone of the long primary branches. _Grateloupia doryphora_ explants were disc fragments (0.3 cm diam.) excised from the middle lower thallus. Explants were cleaned following the methods described by Garcia-Reina _et al._ (1988). After cleaning, _Laurencia sp._ explants were cut again to obtain cylinder-shaped explants (0.5 cm long).

The culture medium was _PES_ (Provasoli, 1968) adjusted to the different osmalalities and solidities in Table 1. Explants (195 per treatment) were cultivated in Petri dishes (15 explants in each) with 20 mL of the culture medium. Cultures were placed in a growth chamber at 20 ± 2 °C with a day length of 18 hours and 27 μmol m⁻² s⁻¹ at the level of the Petri dishes provided by Sylvania Grolux daylight lamps.

Quantitation was made using the indices % callus (percentage of cultivated explants forming disorganized structures) and % bud (percentage of cultivated explants regenerating buds). For _Laurencia sp._, quantitation was made at 30 days of cultivation; for the other two species, at 15 days. The results were expressed as a percentage of those in the control treatment.

Axenic cultures of _Grateloupia doryphora_ were established following the methods described by Polne-Fuller _et al._ (1984). Semicircular-shaped explants (approx. 1 mm long) were excised from the axenic disc. Explants (15 semicircular explants per treatment) were cultivated in different culture media enriched with glycerol (Table 1) under the same culture conditions described before. Quantitation was made using the index 'number of buds/number of cultivated explants'. The fresh weight increase was monitored for semicircular explants cultivated in liquid _PES70_ + 0.3 M glycerol and in _PES_.

## Results

After 15 days calli (filamentous callus-like structures) and buds were regenerated by _Grateloupia doryphora_ explants. During this time, the regenerating explants of _Gelidium versicolor_ only produced buds; callus formation was not recorded even in explants cultivated for 45 days. The explants of _Laurencia sp._ developed callus (a compact cell mass in the wounded area of the explants) and buds after 30 days.

Callus and bud morphology were the same regardless of the osmalality or solidity of the culture medium.

<table>
<thead>
<tr>
<th>Culture medium notation</th>
<th>Percent seawater</th>
<th>NaCl addition mol m⁻³</th>
<th>Glycerol addition mol m⁻³</th>
<th>Osmolality Os Kg⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>PES50</td>
<td>50</td>
<td>-</td>
<td>-</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.10</td>
<td>-</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.27</td>
<td>-</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.51</td>
<td>-</td>
<td>1.5</td>
</tr>
<tr>
<td>PES70</td>
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<td>-</td>
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</tr>
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<td>-</td>
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<tr>
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<td>-</td>
<td>-</td>
<td>1.0</td>
</tr>
<tr>
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<td></td>
<td>0.25</td>
<td>-</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-</td>
<td>0.50</td>
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</table>
Effects of osmolality

The highest % callus and % bud were obtained in culture media at 0.7 to 1.0 Osm kg\(^{-1}\). Culture media at 0.5 Osm kg\(^{-1}\) or 1.5 Osm kg\(^{-1}\) inhibited or drastically reduced regenerator (Figs. 1 & 2).

Effects of solidity

The agar content of the culture media was positively correlated to callus formation (\% callus) and negatively correlated to bud regeneration (\% bud; Fig. 3).

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**Fig. 1.** Values of the index \% callus observed in explants cultivated in different osmolalities (agar concentration = 8 g L\(^{-1}\) in all media, \(n = 195\)).

**Fig. 2.** Values of the index \% bud observed in explants cultivated in different osmolalities (agar concentration = 8 g L\(^{-1}\) in all media, \(n = 195\)).
Effects of seawater

A decrease in the solidity of the culture medium was observed from PES (100% seawater) to PESS0 (50% seawater).

Callus formation in Laurencia sp. and Gratelouplia doryphora decreased from PES100 to PESS0 (Fig. 1). No direct relation between dilution of seawater and % bud formation was found, but the highest % bud formation in Gelidium versicolor and Gracidaria doryphora was observed in PES70. For Laurencia sp. (Fig. 2), the highest % bud formation was in PES100, followed by PESS0, then PES70.

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**Table 2.** Values of the index `number of buds/number of cultivated explants` (B/C) observed in explants cultivated in glycerol enriched media adjusted to different osmolalities and agar concentrations ($n = 15$).

<table>
<thead>
<tr>
<th>Osmolality (Os) kg$^{-1}$</th>
<th>Agar (g L$^{-1}$)</th>
<th>Culture medium</th>
<th>Glycerol (mol m$^{-3}$)</th>
<th>B/C index</th>
</tr>
</thead>
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<tr>
<td>0.7</td>
<td>8</td>
<td>PES70</td>
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<td>1.0</td>
<td>3</td>
<td>PES70</td>
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<td>18.25</td>
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<td>8</td>
<td>PES70</td>
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<tr>
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<td>PES70</td>
<td>0.8</td>
<td>2.60</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>PES</td>
<td>0.5</td>
<td>1.51</td>
</tr>
</tbody>
</table>
Effects of glycerol

Liquid PES70 + 0.3 M glycerol gave a greater increase in fresh weight than liquid PES alone (Fig. 4). In solid media, the best results were obtained with a PES70 + 0.3 M glycerol and 3 or 8 g L\(^{-1}\) agar; the explants became round, turned from reddish to orange, and reached a larger size than those in other culture media. Also, the highest values of the index 'number of buds/number of cultivated explants' was observed in explants cultivated in that medium (Table 2). The value of this index decreased with increasing glycerol concentration and agar content of the culture medium (Table 2).

Discussion

The best morphogenetic response of an explant (callus and bud regeneration) was obtained at the 'natural osmolality' of 1 Os kg\(^{-1}\), the osmolality of the seawater, or slightly lower, 0.7 Os kg\(^{-1}\) (Figs. 1 & 2). Variation in the osmolality of the culture medium to 0.5 and 1.5 Os kg\(^{-1}\) inhibited or drastically reduced bud regeneration and callus formation.

When the explants were cultivated in media with optimum osmolality (0.7 to 1.0 Os kg\(^{-1}\)), the solidity of the culture medium affected the development of the explant, reducing bud regeneration and increasing callus formation (Fig. 3). The solidity of the culture medium is not only involved in callus induction (Polne-Fuller & Gibor, 1987), but in fact, in switching development from organized (bud) to disorganized (callus or callus-like) structures.

Lawlor et al. (1988) reported that an increase in salinity (\% seawater) of the solid medium promotes callus formation and growth in Ecklonia. The results in Fig. 1 show the same effect of seawater on callus formation in Laurencia sp. and Grateloupia doryphora. Also, the highest \% bud formation in G. doryphora and Gelidium versicolor was observed in PES50 or PES70 medium (Fig. 2). The fact that the solidity increased with the percentage of seawater in the PES medium clearly shows that the effect of seawater on callus and bud formation is due to the alteration of the solidity.

Glycerol is an effective carbon source for the vegetative propagation of Grateloupia doryphora. Liquid PES70 culture medium enriched with 0.3 M glycerol gave an increase in fresh weight 400\% higher than PES medium alone after 45 days. This result agrees with those reviewed by Fries (1973) on the effect of glycerol as a carbon source on red seaweeds.

The inhibiting effect of high concentrations of glycerol (0.5 and 0.8 M) on morphogenesis of the explants (Table 2) is primarily due to the increase in osmolality. This result demonstrates the need to control osmolality when testing the addition of carbohydrates or other osmotically active compounds on seaweed cell and tissue cultures.

Conclusions

Osmolality and solidity are physical characteristics of the culture medium that affect the vegetative propagation of Laurencia sp., Grateloupia doryphora and Gelidium versicolor by tissue culture. The alteration of the culture medium, such as dilution of seawater or addition of glycerol modifies the response of the explants by its own effect (i.e. carbon source effect of glycerol) and also by the modification of the solidity or osmolality of the culture medium.

References