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Production and Bioassay of a Diffusible Factor that Induces Gametophyte-to-Sporophyte Developmental Reprogramming in the Brown Alga *Ectocarpus*

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**[Abstract]** The brown alga *Ectocarpus* has a haploid-diploid life cycle that involves alternation between two multicellular generations, the sporophyte and the gametophyte. Life cycle generation is not determined by ploidy but by a genetic system that includes two different three amino acid loop extension homeodomain transcription factors called OUROBOROS and SAMSARA. In addition, sporophytes have been shown to secrete a diffusible factor into the medium that can induce gametophyte initial cells to switch from the gametophyte to the sporophyte developmental program. The protocol presented here describes how to produce sporophyte-conditioned medium containing the diffusible sporophyte-inducing factor and how to assay for activity of the factor using a meio-spore-based bioassay. The protocol, which describes how several steps of these procedures can be optimised, will represent a useful tool for future work aimed at characterising the diffusible factor and investigating its mode of action.

**Keywords:** Brown algae, Diffusible sporophyte-inducing factor, *Ectocarpus*, Gamete, Gametophyte, Life cycle, Meio-spore, Sporophyte

**[Background]** *Ectocarpus* is a small, filamentous brown alga that has been adopted as a genetic model organism for this group of seaweeds (Peters *et al.*, 2004; Coelho *et al.*, 2012a). The brown algae exhibit a broad range of different life cycles (Cock *et al.*, 2014). *Ectocarpus* has a haploid-diploid life cycle (Figure 1) that involves alternation between two multicellular generations, the gametophyte and the sporophyte (Müller, 1967; Bothwell *et al.*, 2010). Transition from the sporophyte to the gametophyte generation involves the production of meio-spores, which are released into the surrounding environment and germinate to produce separate male and female, haploid gametophytes. To complete the life cycle, mature female and male gametophytes produce gametes, which fuse to produce a zygote, the initial cell of the diploid sporophyte generation. Life cycle generation (*i.e.*, sporophyte or gametophyte identity) is not determined by ploidy because single gametes are able to develop parthenogenetically, without fusing to produce a zygote, leading to the production of haploid partheno-sporophytes that are
phenotypically indistinguishable from diploid sporophytes derived from a zygote (Müller, 1967; Bothwell et al., 2010). The gametophyte and sporophyte generations of *Ectocarpus* are free-living, exhibit distinct morphologies and functions and show spatio-temporal independence, facilitating the study of the developmental program of each generation and their relationship with the life cycle.

**Figure 1. The *Ectocarpus* life cycle.** The diploid sporophyte produces meio-spores via a single meiotic cell division (*R! for reduction*) followed by multiple mitotic divisions in each unilocular sporangium (US). The meio-spores are released and develop as gametophytes. Gametophytes are either male or female (dioicy) and produce either male or female gametes in plurilocular gametangia (PG). Pairs of male and female gametes fuse (*F! for fusion*) to form zygotes, which develop as diploid sporophytes to complete the sexual cycle (left). Unfused gametes can undergo parthenogenesis to produce partheno-sporophytes. Partheno-sporophytes produce spores in unilocular sporangia, which develop as gametophytes to complete the parthenogenetic cycle (right).

In addition, both diploid sporophytes and partheno-sporophytes produce mito-spores in plurilocular sporangia (PS), which germinate to produce a new sporophyte generation with the same ploidy as its parent (dotted line). Treatment with sporophyte-conditioned medium (+ SCM) induces a subset of meio-spores to switch from the gametophyte to the sporophyte developmental program (grey text). Adapted from Peters et al. (2008).

Alternation of generations in *Ectocarpus* has been shown to be controlled by two homeodomain transcription factors (HD TFs) of the three amino acid loop extension (TALE) class, OUROBOROS (ORO) and SAMSARA (SAM), which are necessary for the initiation of the sporophyte program (Coelho et al., 2011; Arun et al., 2019). In addition, the sporophyte has been shown to secrete a non-cell-autonomous, diffusible factor that can cause receptive cells to switch from the gametophyte to the sporophyte developmental program (Arun et al., 2013 and 2019). This diffusible sporophyte-inducing factor is only effective on the single-cell stage of the gametophyte generation. Developing meio-spores became resistant to the factor at the same point in time as they synthesise a cell wall, about 24-48 h after release from unilocular sporangia (Arun et al., 2013). This observation suggests that the cell wall may play a
role in locking the individual into the developmental program that has been initiated. *oro* and *sam* mutants do not respond to treatment with the diffusible factor, suggesting that ORO and SAM are part of the signalling network that detects the factor.

Preparations of the diffusible sporophyte-inducing factor are produced by filtering seawater medium in which *Ectocarpus* sporophytes have been cultivated, resulting in cell-free, sporophyte-conditioned medium (SCM). Procedures to produce and assay the diffusible factor were described briefly in Arun et al. (2013) and Arun et al. (2019). Activity of the diffusible factor can be assayed either by treating freshly released meio-spores or by treating gametophyte-derived protoplasts (Arun et al., 2013). The protocol below describes treatment of meio-spores, which are easier to obtain than gametophyte-derived protoplasts. For information about the method using gametophyte-derived protoplasts, please refer to Arun et al. (2013). The meio-spore-based procedure is described in detail and some specific tips to improve both SCM production and the bioassay for the diffusible factor are provided. This detailed protocol is currently being employed in experiments aimed at biochemically characterising the diffusible factor.

**Materials and Reagents**

1. 55-mm diameter Petri dishes (Corning, catalogue number: CLS430589)
2. 150-mm diameter Petri dishes (Corning, catalogue number: CLS430597)
3. 10 L polycarbonate carboys (Nalgene, catalogue number: 2251-0020)
4. Parafilm (Sigma-Aldrich, catalogue number P7793-1EA)
5. Coverslips (Knittel, catalogue number: MS0009)
6. Large size cell strainer (for example a 12.5 cm Finlandek permanent coffee filter)
7. 40 µm nylon cell strainer (Falcon, product number: 352340)
8. 0.22 µm Millex-GP polyethersulphone membrane filter (Millipore, catalogue number: SLGP033RS)
9. Glass bottles (Fisherbrand, catalogue number: 15476113)
10. A closed (e.g., metal) box to keep Petri dishes in the dark
11. Sterile, plugged, glass Pasteur pipettes (Sigma-Aldrich, catalogue number: S6143)
12. Sterile, 10 ml plastic pipettes (Corning, catalogue number: 4488)
13. 22 x 30 mm coverslips
14. Sporophytes and gametophytes of *Ectocarpus* male strain Ec32 (reference CCAP 1310/4 in the Culture Collection of Algae and Protozoa, Oban, UK)
15. H3BO3
16. FeCl3
17. MnSO4·H2O
18. ZnSO4·7H2O
19. CoSO4·7H2O
20. EDTA
21. Vitamin B12
22. TRIS (Trisma base)
23. (NH₄)₂Fe(SO₄)₂·6H₂O
24. NaNO₃
25. C₃H₇Na₂O₆P·5H₂O
26. Provasoli solution (see Recipes)
   Solution 1
   Solution 2
   Solution 3
   Solution 4
   Solution 5
27. Provasoli-enriched natural seawater (PES) (see Recipes)

**Equipment**

1. Neubauer chamber Bürker (0.1 mm depth, 0.0025 mm²)
2. Fine dissecting forceps (Dumont, catalogue number: 0103-27-PO)
3. Thermostatically-controlled, illuminated growth cabinet (Pol-Eko-Aparatura, model: KK 240 FIT P) or growth room
4. Binocular dissecting microscope (Olympus, model: SZ61)
5. Inverted light microscope (Olympus, model: CKX41)

**Software**

1. Excel, version 15.32 (Microsoft)

**Procedure**

A. Preparation of SCM

1. Preparation of *Ectocarpus* strains
   a. Prepare sporophytes and gametophytes of *Ectocarpus* male strain Ec32.
   b. Maintain the sporophytes and gametophytes separately by growing in 10 ml of Provasoli-enriched seawater (PES, *i.e.*, natural seawater enriched with Provasoli supplement; Starr and Zeikus, 1993) in 55-mm diameter Petri dishes or in 100 ml PES in 150-mm diameter Petri dishes under standard growth conditions: 13 °C, 12 h:12 h (light:dark) and 20 μM photons/m²/s (Coelho et al., 2012b). The edge of the Petri dish is wrapped with a band of Parafilm, which allows sufficient passive aeration for algal growth.
2. Growth of partheno-sporophytes for the production of SCM
   a. Grow cultures of *Ectocarpus* strain Ec32 gametophytes in 150 mm Petri dishes (about
0.13 g per Petri dish) for 2 weeks under low light conditions (2-3 µM photons/m²/s, 12 h:12 h light: dark) in PES at 13 °C.

b. Induce the release of gametes by grouping 20 to 30 gametophytes together in a small volume of medium (to simulate low tide conditions) and incubate in the dark in a closed box for four hours at 13 °C (Figure 2).

Figure 2. Induction of gamete release. A. Gametophytes grouped into a small volume of PES. B and C. Removal of excess PES with a pipette. D. Gametophytes after removal of excess PES. E. After incubation in the dark for four hours, 300 µl of PES is added. F. Gametophytes are placed under strong light to induce gamete release.

c. Add 300 µl PES and place under strong light (25-30 µM photons/m²/s) to induce gamete release. Synchronous release of large numbers of gametes should be observed within ten minutes of illumination with strong light (Video 1).

Video 1. Release of Ectocarpus gametes from a plurilocular gametangium and swarming gametes

d. Estimate the number of gametes released using a Neubauer chamber. Mix 9 µl of the
released gametes in PES with 1 μl of 10% glutaraldehyde. After 10 min at room temperature pipette the mix onto the grid of the Neubauer chamber and cover with a coverslip. Count the number of gametes in a 0.04 mm² square on the Neubauer chamber grid, then repeat the counting at least nine times and calculate the average of these counts (= X). The number of gametes per μL is calculate by dividing the average number of gametes counted per 0.04 mm² square by the surface area (mm²) x chamber depth (mm) and then correcting for the dilution (i.e., X/(0.04 x 0.1) x 10/9).

e. Use each batch of released gametes to inoculate two 150 mm Petri dishes (about 10⁶ gametes per Petri dish) and cultivate the resulting partheno-sporophyte germlings under low light conditions for 14 days.

f. Inoculate a 10 L carboy of PES with 0.5 g of partheno-sporophyte thalli and grow for between four and eleven weeks at 13 °C under low light conditions. The carboy culture should be aerated by pumping air through a 0.22 μm filter. The entry and exit tubes for the air pass through holes in the top of the carboy and a 10 ml sterile pipette is used to deliver the air bubbles to the bottom of the carboy (Video 2).

Video 2. Culture of Ectocarpus partheno-sporophytes in a 10 L carboy

3. Collection and storage of the SCM
   a. Use a large size cell strainer to remove the bulk of the partheno-sporophyte tissue from the culture and then filter the SCM through a Falcon 40 μm cell strainer and a 0.22 μm filter to remove any remaining cells.
   b. Store the SCM in glass bottles (prepared beforehand by cleaning with 70% ethanol) in a cold room at 4 °C. The SCM can be stored for at least eight weeks at 4 °C without loss of activity.

B. Bioassay of the diffusible sporophyte-inducing factor
   1. Production of unilocular sporangia containing meio-spores
      a. The bioassay of the diffusible sporophyte-inducing factor is carried out using meio-spores
produced by fertile sporophytes. Depending on the strain of *Ectocarpus*, meio-spores can exhibit different levels of heteroblasty, i.e., spontaneous initiation of the sporophyte program instead of the gametophyte program (Müller, 1967). To assay the activity of the diffusible factor, it is important to use a strain that exhibits a low level of spontaneous heteroblasty, such as the strain Ec32.

b. Induce the release of gametes from 45 to 60 mature gametophytes by grouping the material together in a small volume of PES (to simulate low tide conditions) and incubating in the dark at 13 °C for four hours (follow the procedure described in Steps A2a-A2c above).

c. Allow the gametes to germinate parthenogenetically to produce partheno-sporophytes under standard growth conditions: light intensity of 20 µM photons/m²/s, 12 h:12 h light:dark at 13 °C in 150-mm Petri dishes (Coelho et al., 2012b).

d. Change the culture medium regularly (once every 2 weeks) until the partheno-sporophyte filaments produce upright filaments. When upright filaments are produced, remove individual partheno-sporophytes and culture separately. Note that cultures grown at high density will not produce unilocular sporangia. Plurilocular sporangia should develop after about 1 to 2 weeks and the unilocular sporangia appear about one week later.

e. Alternatively, unilocular sporangia can be produced on cultured upright filaments, for example by transferring previously dissected upright filaments with unilocular sporangia that have released meio-spores back into culture. These filaments will adhere to the bottom of a 55-mm Petri dish and produce new upright filaments in a few days under standard culture conditions. The upright filaments will produce many unilocular sporangia after about one week. Cultivation of upright filaments results in the production of fewer (or no) plurilocular sporangia than cultivation of partheno-sporophytes (reducing the risk of contaminating meio-spore preparations with mito-spores). This method of producing unilocular sporangia is more rapid than cultivation of whole partheno-sporophytes.

2. Bioassay of the diffusible factor by treatment of meio-spores

   a. Under a binocular microscope, dissect a piece of sporophyte upright filament that bears one or more unilocular sporangia using a sterile glass Pasteur pipette that has been broken to create a sharp, cutting point (Figure 3).
Figure 3. Dissection of unilocular sporangia and meio-spore release. A. *Ectocarpus* partheno-sporophyte bearing unilocular sporangia. B. Dissection of a filament fragment bearing a unilocular sporangium (arrowhead) using a sterile Pasteur pipette. C. Cut filament fragment with unilocular sporangium (arrowhead). D. Drop of PES on a coverslip in a 55-mm Petri dish humidified with drops of PES at the periphery. E. Filament fragment bearing unilocular sporangium transferred to the drop of PES. The inset shows the filament fragment in the drop of PES. F. Unilocular sporangium before meio-spore release. G. Unilocular sporangium after meio-spore release. Scale bars: 1 mm (A), 500 µm (B, C), 1 cm (D), 2.5 mm (E), 400 µm (E inset), 20 µm (F, G).

b. Wash the fragment twice in PES by transferring sequentially to two 55-mm Petri dishes, each containing 10 ml of PES (to ensure that there is no carry-over of possible contaminating mito-spores), then transfer into a drop of 300 µl of SCM (Figure 2) on a coverslip. The coverslip should be fixed to the bottom of the Petri dish by placing it on a drop of 10 µl of PES. Place three drops of PES around the edge of the Petri dish to keep the chamber moist and seal the Petri dish with parafilm. Note that the SCM can be replaced by any other seawater-based sample that you would like to test for diffusible factor activity.

c. In parallel, transfer unilocular sporangia to drops of PES, using the same set-up, as negative controls (i.e., incubation in seawater that does not contain any active factors).

d. Place the unilocular sporangia in strong light conditions (30 μM photons/m²/s) overnight to induce release of the meio-spores (100-200 meio-spores per unilocular sporangium) directly into the 300 µl drop of test or control solution. Release of the meio-spores from the unilocular sporangia should take less than 48 h. Discard any plates in which release has not occurred within this time frame.

e. After meio-spore release, remove the piece of upright filament with forceps.

f. After 72 h, gently add 10 ml of PES to the Petri dish. The meio-spore-derived germlings will
be weakly attached to the coverslip at this stage and the added medium assures that they grow under optimal conditions.

g. After an additional three or four days examine the Petri dish under an inverted microscope to score the numbers of gametophyte and sporophyte individuals (Figure 4). Sporophytes can be distinguished from gametophytes based on a symmetrical pattern of initial cell division and the presence of thick walled round cells rather than wavy rhizoid cells. Note that, if meio-spores are released within 48 h but then germinate slowly, the germination process can be accelerated by adding an additional 300 µl of the test medium to the 300 µl drop. This will allow the germlings to grow and attach to the coverslip before the plate is flooded with the 10 ml of PES.

Figure 4. Morphological differences between gametophytes (normal development, left) and sporophytes (modified development, right) derived from SCM-treated meio-spores. Meio-spores were incubated in SCM for three days and then grown in PES. A. and E. two-cell stages of the gametophyte (asymmetrical cell division) and the sporophyte (symmetrical cell division), respectively. B. and F. gametophyte and sporophyte germlings, respectively, after three days in SCM. C. and G. gametophyte and sporophyte germlings, respectively, after five days in SCM. D. and H. gametophyte and sporophyte germlings, respectively, after 14 days in SCM. Arrows indicate wavy rhizoid cells typical of the gametophyte germling, arrowheads indicate round cells typical of the sporophyte germling. Scale bars: 20 µm.
h. When working with active preparations of SCM, expect between 2% and 30% of the meio-
spores to be switched from gametophyte to sporophyte identity but note that the percentage 
of switching can be highly variable between assays. It is therefore preferable to carry out at 
least three assays for each test to obtain statistically robust estimations of diffusible factor 
activity.

Data analysis

Use the Wilcoxon rank sum test with Holm-Bonferroni \( P \)-value adjustment to determine whether test 
samples produce significantly more sporophyte individuals than the PES control treatments (Figure 
5).

![Figure 5. Example of data obtained using the diffusible factor bioassay. Percent of 
germlings that exhibited sporophyte morphology following the release of meio-spores from 
unilocular sporangia treated with either Provasoli-enriched natural seawater (PES, 69 unilocular 
sporangia, 7040 meio-spores) or sporophyte-condition medium (SCM, 103 unilocular sporangia, 
9106 meio-spores). Error bars indicate standard error of the mean, letters above bars indicate 
significant differences (\( P \)-value < 0.05).]

Notes

One difficulty with this procedure is that the percentage of meio-spores that switch from the 
gametophyte to the sporophyte developmental program can be very variable (between 2% and 30%). 
This appears to due to two factors, variation in the quantity of diffusible factor in different batches of 
SCM and variation in the capacity of meio-spores to respond to the diffusible factor. The exact cause 
of this variation has not been determined but it is important that the operator be prepared to carry 
out additional replicate tests if the effect of the SCM is weak.
Recipes

1. Provasoli solution

**Solution 1**
1.9 g H$_3$BO$_3$
0.05 g FeCl$_3$
0.273 g MnSO$_4$·H$_2$O
0.0367 g ZnSO$_4$·7H$_2$O
0.008 g CoSO$_4$·7H$_2$O
11.4 ml EDTA (0.5 M, pH 8)
ad 1,000 ml ddH$_2$O

**Solution 2**
3.35 mg Vitamin B12 (cyanocobalamine)
165 mg Thiamine hydrochloride (vitamin B1)
1.65 mg Biotin C$_{10}$H$_{16}$N$_2$O$_3$S
166.5 g TRIS (Trisma base) C$_{6}$H$_{11}$NO$_3$
ad 500 ml ddH$_2$O

**Solution 3**
1.17 g (NH$_4$)$_2$Fe(SO$_4$)$_2$·6H$_2$O
6.8 mL EDTA (0.5 M, pH 8)
ad 1,000 ml ddH$_2$O

**Solution 4**
23 g NaNO$_3$
ad 1,000 ml ddH$_2$O

**Solution 5**
3.33 g C$_3$H$_7$Na$_2$O$_6$P·5H$_2$O “glycerophosphate”
ad 1,000 ml ddH$_2$O

Prepare each stock solution separately, autoclave, and store in glass bottles at 4 °C. Use a dark bottle for Solution 2. For 1 L of Provasoli solution, add 100 ml of each of Solutions 1, 3, 4, and 5 plus 10 ml of Solution 2 to ddH$_2$O (the starting pH should be between 9.6 and 9.8). Adjust to pH 7.8 with concentrated HCl (37%) and adjust the volume to 1 L with ddH$_2$O. Aliquot into small glass bottles (20, 50, 100, or 200 ml), autoclave, and store at 4 °C.
2. Provasoli-enriched natural seawater (PES)
   If possible, collect natural seawater offshore.
   a. Prepare 1 L of natural seawater by filtering through a 5 µm mesh and autoclaving in a plastic bottle
   b. Add 20 ml of Provasoli solution
   c. Autoclaved natural seawater can be stored at 13 °C before addition of the Provasoli solution and PES can be stored at 4 °C

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Competing interests

The authors have no competing interests.

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