Cryopreservation of Ectocarpus

PREAMBLE. The method outlined below is based on Heesch et al. (2012).

For all cryopreservation procedures it is normally optimal to employ a healthy, vigorous, relatively dense, late log-phase, or early stationary phase culture. Where practicable axenic strains, or those with low levels of bacterial contaminants should be employed and those contaminated with eukaryotes avoided.

1. One to two weeks prior to cryopreservation separate thalli into smaller pieces (approx. 1-2 mm in length). Transfer to Erlenmeyer flasks containing 100 ml of culture medium, under the same culture conditions.

   Exchange culture medium once a week prior to cryopreservation, to maximise culture quality and minimise the bacterial numbers in non-axenic strains.

2. Prepare 10-ml aliquots of cryoprotectant solution [10% (v/v) DMSO & 9% (v/v) D-sorbitol in the appropriate culture medium] and filter-sterilise into a sterile Universal tube.

3. Aseptically transfer 1-ml aliquots of sterilised cryoprotectant solution into 2-ml cryovials.

4. Aseptically transfer intact sections of *Ectocarpus* thalli to each cryovial, seal the cryovials and incubate for 15 – 30 min. at room temperature.

   At SAMS 15 vials are normally filled and processed (see below).

5. Programme the controlled cooler: Start temperature 20°C; ramp, cool at -1°C min.\(^{-1}\) to -40°C; dwell, hold at -40°C for 10 min.

   In general mechanical seeding of ice is not used for algal cryopreservation, but if it is an option in the system employed, empirical experimentation may be used to ascertain if it results in higher post-thaw viability.

6. Start the programme to purge the system with nitrogen vapour and to allow the system to stabilise at the start temperature.

7. On reaching the start temperature, (most systems have an audible alarm) transfer the cryovials to the cooling chamber of the programmable cooler and initiate the cooling ramp.

8. After the end of the programme an alarm will sound, rapidly transfer the cryovials to a small dewar containing liquid nitrogen using long forceps.
It is essential that the cryovials are not allowed to warm up prior to plunging into liquid nitrogen.

9. Samples for storage should be transferred to the cryostat (ultra-cold freezer) in the liquid nitrogen containing dewar. Transfer of cryovials to the storage system should be done rapidly using long forceps.

Storage temperature is critical and should be <-130°C. It is advisable that replicate samples are stored in at least two separate “refrigerators”. At SAMS 10 cryovials are normally stored in a working bank, 2 vials are stored in a master/back-up bank and 3 are used to check viability/efficacy of the protocol.

10. To recover cultures, vials are thawed by placing in a pre-heated water-bath (40°C) and agitated until the last ice crystal has just melted.

For most marine taxa it is important not to prolong their incubation at 40°C. Alternative, slower warming e.g. in a 25°C water-bath may be beneficial for some strains, but in general rapid warming is optimal as it avoids/minimises ice crystal regrowth.

11. On thawing rapidly transfer to a laminar flow/biological safety cabinet and wipe the outside of the vial with 70% (v/v) ethanol.

Note: there may be high levels of viable bacterial and fungal spores in liquid nitrogen that may contaminate recovered cultures.

12. Using a disposable plastic pipette the cryoprotectant solution is aseptically removed, discarded and replaced by 1-ml of fresh sterile medium. The thallus is then aseptically transferred to a labelled Petri-dish containing 10-ml of an appropriate medium. Cover in aluminium foil and re-label with strain designation and date.

It is optimal to prevent further biochemical-based injuries to incubate for a period in the dark.

13. Incubate at standard culturing temperature for the cryopreserved organism, after 24h partially remove the foil and after a further 24-96 hours remove all the foil covering.

It is important to ensure that cells are not subjected to light levels likely to induce photo-oxidative stress during the recovery phase. It is worth noting that cell numbers are generally very low and, particularly for strains where recovery levels may be low, that there is little self-shading.

14. After one week, using standard aseptic techniques, replace the medium.

Because of the lysis of cells killed by the cryopreservation procedure it is necessary to exchange culture medium after a week to reduce the bacterial numbers in non-axenic strains. For more heavily bacterised cultures this should be done more frequently to maximise the possibility of recovering a healthy culture.

15. After an appropriate period (2-8 weeks, depending of the strain) a normal culture should be obtained. This may be maintained by routine serial transfer, or employed for experimental use.