MANUAL FOR INTENSIVE HATCHERY PRODUCTION OF ABALONE

Theory and practice for year-round, high density seed production of blacklip abalone (*Haliotis rubra*)

*Mike Heasman and Nick Savva*

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Introduction

This manual was commissioned by the Indigenous Fisheries Initiative of the NSW Government as a practical guide to commercial scale seed production of blacklip abalone (Haliotis rubra) in NSW. Methods and equipment are based on those described by Hone et al. (1997) with more recent refinements developed by Dr. Arthur Ritar and Mark Grubert in Tasmania during the course of the Fisheries Research and Development Corporation (FRDC) Project 2000/204 and in NSW during the course of additional projects supported by the FRDC, namely projects 1998/219 and 2001/033.

The manual provides specialized instruction on:
- how to collect and reproductively condition blacklip abalone broodstock
- how to induce them to spawn
- how to hatchery-rear their young through the larval and early juvenile stages to an age and size suitable for on-farming or for seeding depleted fisheries.

With minor adjustments to methods and equipment, the manual should also serve as a useful production guide for most temperate species of abalone. There is an emphasis on year-round production of spat as opposed to the seasonal production practised to date by commercial abalone farms in Australia. In view of the scarcity and high cost of coastal sites with access to marine waters in NSW, the techniques described are geared to efficient and intensive production of abalone spat using small areas of land. These techniques in some respects are quite different to those generally practised to date by commercial abalone farms in Australia.

This manual is not intended to cover all aspects of blacklip abalone biology, nor hatchery production technology. There is a large amount of such information published both in Australia and overseas in relation to blacklip and other abalone species. The manual can be used by people with a range of backgrounds. It is primarily intended for abalone farmers and technicians with a knowledge of aquaculture fundamentals; however, the information is also expected to be useful to scientists and researchers.
How to use this Manual

This manual consists of a logical sequence of chapters that provide background theory as well as practical guides of how to:

- collect wild adult blacklip abalone, domesticate them and bring them into spawning condition
- induce them to spawn
- fertilise and incubate their eggs
- produce larvae, post-larvae and juveniles of a size suitable for seeding depleted fisheries or for on-farming, in accordance with the systematic scheduling of operations illustrated in Appendix 1.

To help you apply these procedures, we have also provided step-by-step summaries that itemize key actions and essential equipment and materials. Supplementary information on the design and operation of facilities and equipment and on sub-procedures is provided as a series of appendixes. Specialist technical terms are presented in bold font and are defined the first time they appear in the text. A reminder of the meaning of these specialized words can be readily accessed in the glossary at the back of this manual.

We intend this manual not as a stand-alone text but as a companion to other publications, especially the excellent array of manuals and reports listed in ‘References and companion reading’. We strongly encourage readers to acquire and consult these and additional references as recommended from time to time through the text.
References and companion reading


Chapter 1  Abalone Biology
Chapter 1 Abalone Biology

A balone are marine snails that belong to a group of invertebrates (animals lacking a backbone or other forms of internal skeleton) called molluscs. Molluscs also include common bivalves such as scallops, oysters, mussels, pippies and cockles as well as octopus, squid and cuttlefish. Most species of abalone have a moderate to heavily calcified snail-like shell that is flattened (Figure 1). This flat body shape reduces drag forces generated by powerful waves and currents typical of their shallow (1 to 10 m deep) exposed coastal reef habitat. The shell protects the soft body tissues including a very large and undulating muscular foot that allows the abalone to remain firmly attached to rocky substrates even while moving and feeding.

Figure 1 Abalone: a marine snail distinguished by a flattened shell with a set of respiratory pores.

A series of holes (Figures 1 and 2) along the left upper margin of the flattened shell defines abalone as a particular group (genus) of snails, called Haliotis. These holes (or respiratory pores) are outlets through which seawater, partially stripped of oxygen by the gills, is exhaled together with urine, faeces, and either sperm or eggs. The head and mouth of abalone are flanked by two pairs of sensory tentacles (Figure 2); the shorter outer pair are stalked eyes. The mouth lacks teeth but instead is equipped with a long rasping tongue called a radula that is used to tear, dislodge and ingest food, which comprises mainly plant materials.

Food includes seaweed that is either still attached to rock surfaces or has already been detached by the actions of waves and currents. Detached seaweed—the
principal food source of larger juvenile and adult abalone—is commonly referred to as ‘drift’.

As already implied, abalone have separate sexes. The gonads (egg or sperm-producing organs) lie adjacent to the margin of the shell at the rear of the body on the side closest to the spire of the shell (Figure 2). The gonad can be clearly seen only by deflecting the foot out of the way (Figure 3). Male abalone can be identified by their white to cream gonads (testes) (Figure 4). Female abalone (Figure 4) have gonads (ovaries) that reflect the colour of their eggs, which varies widely from green to olive and from maroon to brown. In practice, determining the sex of blacklip abalone can be difficult because the overlying skin on the outside of the foot can be heavily pigmented or banded, ranging from pale grey to black, with the latter totally obscuring the gonad colour.

In NSW, wild blacklip abalone become sexually mature by the time they reach a shell length of about 90 mm at 3 to 5 years of age. However, their farmed counterparts become sexually mature 1 to 2 years earlier at a smaller size of 50 to 70 mm. Abalone are broadcast spawners that release sperm and eggs (Figure 5) into the surrounding seawater, where fertilisation takes place. Some populations of wild abalone form aggregations and spawn in unison to increase the chance of successful fertilisation.

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**Figure 2** Drawing of dorsal view of an abalone with shell removed to expose soft tissues (adapted from Shepherd, 1975).
**Figure 3** Simple method of deflecting the foot muscle of an abalone to determine whether it is sexually mature and, if so, its gender and breeding stage.

**Figure 4** View of soft tissues of blacklip abalone from above after removal of the shell. Note the ripe gonads of these adult abalone, characterized in the case of males (left specimen) by a large cream-coloured testis and in females (specimen on the right), by a maroon-coloured ovary. Reproduced from Liu (2005).
Figure 5  Spawning female abalone. Eggs are demersal (heavier than water) and rapidly settle on the floor of spawning containers.

In established well-run hatcheries, 60% to 80% of female abalone and 80% to 90% of males held at temperatures of 16 to 18°C— conducive to ripening of the gonads—can be expected to spawn in response to the induction protocols discussed in Chapter 4. However lower rates of success are to be expected during the first year of operation of new hatcheries because of incomplete domestication of recently captured stock; this problem is commonly compounded by staff inexperience and teething problems with the newly commissioned equipment.
Figure 6 Schematic life cycle of abalone (source Hone et al., 1997).
**Figure 7** Fully shelled second stage veliger larvae

**Figure 8** Photomicrograph of recently settled and metamorphosed post-larvae grazing on surface of crustose coralline algae coated rock (courtesy of Sabine Daume) Note: Smooth larval shell (red arrow) and sculptured post larval shell growth (yellow arrow) Black bar = 500 micron
Fertilised eggs hatch into first-stage swimming larvae called **trochophores (Figure 6)** after 18 to 24 h at an optimum temperature of about 18˚C. The time spent as **planktonic** larvae (living in the water column) differs among and within species and depends on temperature. Trochophores develop further into shelled **veliger** larvae (Figures 6 and 7) after 20 h at 18˚C. The larvae of abalone do not feed, but live on the egg yolk and by directly absorbing soluble nutrients in seawater through the skin. Blacklip abalone larvae first become competent to settle and **metamorphose** (transform) into reef-dwelling **post-larvae**. Post-larvae (Figure 8) are simply early-stage juveniles that range from 0.3 to 3 mm in shell length.

Metamorphosis of larvae into post-larvae occurs after a minimum of 6 days at an optimum temperature of 18˚C. Larvae can however, delay settlement and metamorphosis until they find a suitable habitat (Figure 9), which consists of rocky surfaces coated by a thin skin of tough, limestone-depositing **crustose coralline algae**. This is commonly known as ‘pink rock’ (Figure 10). These predominately shallow (1 to 6 m) rock habitats include wave-rubled ‘boulder fields’ and other deeper bare rock areas called **barrens**. In NSW, barrens are commonly the product of grazing by dense aggregations of the black (or purple) sea urchin (*Centrostephanus rodgersii*) (Figure 11). Urchin barrens are in fact the most common type of reef habitat in New South Wales, accounting for almost half of the 5000 ha of reef located from Port Stephens (120 km north of Sydney) south to the Victorian border.

Ready-to-settle blacklip abalone larvae ‘home in’ on these rocky habitats, being attracted by chemical cues released by the surface layer of crustose coralline algae and by recent grazing trails of juveniles and adults of their species. During and immediately following settlement, larvae and early post-larvae suffer heavy mortalities that exceed 95% during the first week alone. The principal causes are predation or accidental ingestion by a diverse array of other common and prolific reef surface grazers, especially urchins, turban shells and other marine snails, in combination with predatory free-ranging sea worms.

With progressive growth from 2 mm to about 10 mm, and further development of the rasping tongue, the main source of food for juvenile blacklip abalone progressively shifts from microscopic algae (Figure 12) towards the very abundant but tough cells of the encrusting coralline algae. Surface grazing on crustose coralline algae tissue remains an important way of obtaining food for juvenile blacklip abalone up to a shell length of about 40 mm and 18 months of age. Thereafter it is progressively replaced by a diet of seaweed that consists mainly of broken-off pieces (drift), rather than whole plants, which need to remain firmly attached to rock substrates via holdfasts in order to survive.

Post-larvae and other small juvenile blacklip abalone live on the tops and sides of submerged rocks. As they grow beyond about 5 mm, they start seeking shelter under the rocks by day, emerging only at night to feed in the absence of their most important predators. These predators include carnivorous fish and especially species such as wrasse, morwong (Figure 13) and wirra cod. Coincident with the shift in diet to seaweed, the distribution of juvenile blacklip abalone becomes more ‘patchy’. This patchiness reflects the uneven distribution
of drift, including common red seaweeds such as Gracilaria and green seaweeds such as sea lettuce (Ulva spp.). Drift is generally more abundant and accessible on the outer margins of reefs directly exposed to prevailing currents.

Red seaweeds are generally more palatable to blacklip abalone than the common brown seaweeds such as giant kelp (Macrocystis spp.), and this probably also accounts for the large variation in the carrying capacities of individual reefs for abalone—even of those within a few hundred metres of one another. Highly productive reefs in NSW are commonly characterised by fast-growing blacklip abalone with thin, oval shells. By contrast, low-productivity reefs commonly carry slow-growing abalone with thicker, heavier shells that are more circular. Many abalone within these low-productivity reefs fail to reach the minimum legal shell length, which in NSW is 115 mm. Such populations are commonly referred to as ‘stunted’.

**Figure 9** Typical shallow exposed rocky coastal habitat of blacklip abalone
Figure 10  Typical ‘pink rock’ shallow crustose coralline algae coated boulder habitat of settlement and early juvenile stages of blacklip abalone. In the centre is an early prototype mass seed-deployment device that disperses button-sized juvenile abalone. Photo taken 24 h after release. From: Heasman et al., 2004.

Figure 11  Urchin barrens - the most common type of reef habitat in NSW, accounting for about half of the State’s 5000 ha of commercially fished abalone-bearing reef
Figure 12  Examples of benthic microalgae that comprise the principal diet for early post-larvae and to juvenile abalone. These include algae eaten by abalone, especially diatoms such as species of Navicula (a & b), Nitzchia (c), Cylindrotheca (d), Ulvella (e) & crustose coralline algae (f). Reproduced from Daume, 2004.
A-E scale bar indicates 10 micron; F scale bar indicates 1 millimetre.
Figure 13 Above: Holding tank with red & banded morwong. Morwong together with common species of wrasse and other reef and demersal fish like wirra cod, are major predators of juvenile abalone 5 to 50 mm long.

Right: A collection of shells of juvenile abalone regurgitated by the morwong.
Chapter 2  Broodstock Collection and Husbandry

To run a hatchery cost-effectively it is a great advantage to be able to produce multiple batches of larvae and juveniles year-round. Unfortunately, in NSW wild blacklip abalone are not reliable sources of ripe, ready-to-spawn broodstock, even during the natural late spring/early summer breeding season. Wild adult blacklip abalone collected in apparently ripe condition (i.e. with large, swollen gonads) generally prove over-ripe or are inhibited from spawning by the stress of capture and confinement. As a consequence, abalone hatcheries located in NSW must include provision for long-term holding and year-round reproductive conditioning and spawning of captive broodstock. Fortunately, reproductive conditioning of blacklip abalone is a simple matter of feeding them a good quality manufactured diet while maintaining them in excellent (near-oceanic) quality seawater and at favourable temperatures within the range 16 to 18°C. These temperatures mimic those of sea temperatures in coastal NSW in winter, a time when blacklip abalone develop gonads at the expense of growth.

Before broodstock may be collected from the wild, a compulsory permit must be acquired from the relevant authority. In NSW this is the Fisheries Division of the Department of Primary Industries. Permits restrict size and number and the methods of collection and areas where abalone can be legally collected. All such restrictions must be strictly observed. When collecting broodstock, choose young adults in the range 90 to 120 mm that have clean, undamaged shells free of heavy biofouling and obvious infestation by shell damaging mud-worms and boring sponges. The most vigorous and healthy individuals are those with thinner and elongated shells that can rapidly right themselves when upturned and rapidly seek shade and shelter when exposed to bright light. Be aware that numbers of males and females in natural populations might not be equal and that the sex of selected stock is best checked (see methods of sexing below) as soon as possible after collection. This will ensure that sufficient numbers and the best examples of each sex are retained and that the remainder can be returned to the wild unharmed.

Special care must be taken not to injure stock when removing them off rocky surfaces. Do not use abalone knives or other sharp implements that may cut the foot of the abalone, resulting in major loss of blood, infections and eventual death. Instead, use a metal paint scraper with its corners ground off (Figure 14) or a plasterer’s plastic spatula and place the blade under the back of the foot in a rapid but smooth sliding (not levering) action to break the seal between the foot and rock surface.

During collection, broodstock can be stockpiled either in live-wells of boats or in large tubs continuously flushed with new seawater. Alternatively, broodstock can be held in mesh cages (Figure 15) or net bags hung from the stern or sides of the dive vessel. Once collected, and during transportation to the hatchery, abalone must be either kept undisturbed in flowing seawater, or stored in damp, cool, dark conditions. For both methods the abalone are encouraged to attach to smooth inert surfaces, such as sheets of plastic (Figure 15) or to each other in clusters.
If transportation time to the hatchery is likely to be more than an hour or two, the abalone should be held in seawater at ambient temperature. If they are to be temporarily held in shore-based, flow-through or chilled recirculation seawater systems, such as those used by the live abalone export trade, then water quality must be maintained at near-oceanic quality.

**Figure 14** A metal paint scraper with its corners and sharp edges ground off to minimise injury to broodstock being collected from the wild

In the case of flow-through systems, the seawater flow rate per minute should equal or exceed the total abalone biomass. Stress experienced by recently collected wild abalone is a product of both time and intensity, periods of storage and transportation should be kept to a minimum. If holding and transportation periods are likely to exceed a few hours but no more than 24 h, damp cool storage at 12 to 16ºC is recommended. Preferably this should be within sealed plastic bags in an atmosphere of pure oxygen.

On arrival at the hatchery, newly collected broodstock should be immediately inspected and any stock with injuries or other serious afflictions culled. Healthy individuals should be placed in a quarantine tank supplied with ambient, 10 µm filtered seawater. The tank should be covered with 90% shade-cloth and the abalone provided with daytime shelters. Newly acquired broodstock should be retained in quarantine for at least 2 weeks to allow them to acclimatize to captivity and for the full effects of injury and stress to become evident.
Figure 15  Top: Plastic chicken crate for stock-piling and safe transportation (following collection from the wild) Bottom: Smooth plastic sheets that safeguard attached broodstock during transportation back to the hatchery.
The shells of broodstock retained on the basis of good appearance, health and vigour are cleaned under running seawater with a wire brush or an abalone knife to remove biofouling. Utmost care is needed not to damage soft tissues or the respiratory pores of the shell. In healthy blacklip abalone, cleaning will expose the prominent red/maroon layer of the outer shell surface (see Figure 1). Once cleaned, the shell may be treated with crushed sea salt for several hours to desiccate any undetected boring mud-worms that otherwise will multiply and spread to other broodstock within the conditioning units. An option to the use of salt is to suffocate the mud-worms by coating the shell with surfboard wax. Retain the abalone in quarantine for a further 1 or 2 days before peeling off the wax and rescrubbing the shell. (See Appendix 4 for further information).

Fully conditioned (ripe) blacklip abalone suitable for the induction of spawning are those that have been held at 16ºC for 150 to 180 days (see Chapter 4 for details), either from the date of a previous spawning or from an initial state of little or no gonad development. Efficient reproductive conditioning cannot therefore be achieved without accurate and unambiguous records for individual broodstock. To do this, each broodstock must be individually marked before being measured, weighed, sexed and assessed for general health and breeding condition. Marking is done by threading commercially available tags (individually numbered and colour-coded cable ties) through the two respiratory pores closest to the front of the abalone (Figure 16). Care is needed to avoid damage to soft tissues.

Figure 16 Individually numbered or colour-coded cable-tie tags. These are inserted through the two respiratory pores closest to the front (head) of the abalone.

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**STEP BY STEP SUMMARY AND CHECK-LIST FOR BROODSTOCK COLLECTIONS**

☑ 1. Acquire all permits for collecting and holding broodstock while building and commissioning the hatchery (submit all applications at least 18 months ahead of intended start of operations).

☑ 2. Avoid injury of abalone by removing them from rocky surfaces with a rounded paint scraper or plasterer’s spatula, and transport them back to the hatchery at the prevailing sea temperature as soon as possible.

☑ 3. Provide smooth surfaces such as sheets of plastic for the abalone to adhere to during transportation to the hatchery.

☑ 4. Transport the abalone undisturbed in shaded or dark conditions in good quality seawater, or under damp, cool (less than 22 °C) conditions in air if the transportation period is less than an hour.

☑ 5. On arrival at the hatchery, allow the abalone to recover in clean, flowing seawater.

☑ 6. Remove biofouling from the shell with a wire brush or diver’s knife, taking great care not to damage the soft tissues.

☑ 7. Quarantine the newly collected abalone from existing stock for at least 2 weeks until they commence feeding (an indication of good health and habituation to captivity).

☑ 8. Double-tag, measure, and record the details of each individual on a database before introducing them into the broodstock conditioning system.
Chapter 3  Planning and Implementing Breeding Programs
Chapter 3 Planning and Implementing Breeding Programs

Important things to know

A balone broodstock-conditioning facilities are expensive to build, maintain and run. When designing a hatchery and considering issues such as the scale and timing of hatchery production, the amount of capital investment required and ongoing operational costs, it is important to know how many ripe, ready-to-spawn broodstock will be needed and when. If seed (small juveniles) abalone are to be produced for aquaculture, specific breeding lines selected for faster growth may be required. On the other hand, if seed are to be produced for restocking programs, the identity and distribution of genetically distinct populations in the depleted areas must be determined and protected.

To safeguard the genetic diversity of wild populations of abalone, a minimum number of parents (at least 30 to 40 of each sex) need to be used in the production of seed abalone for enhancing depleted wild fisheries. If selective breeding is to be practised over a number of successive generations, each generation must be derived from a large number of mating pairs to avoid inbreeding problems. Such breeding programs impose considerable additional stock management and record-keeping loads and responsibilities. They are also complex and costly. It is therefore strongly recommended that specialist advice be sought when planning and costing selective breeding programs. Such advice can be obtained from the National Abalone Selective Breeding Program, coordinated through the Abalone Aquaculture Subprogram of the Fisheries Research and Development Corporation (FRDC).

Reproductive conditioning of broodstock

In the more southern waters of Tasmania and Victoria, relatively predictable seasonal temperatures generally remain below the upper cut-off of about 19ºC for breeding in blacklip abalone. As a result, these southern stocks undergo more synchronous and predictable annual breeding patterns in which the gonads start from an initial resting stage of little or no development in autumn and end with full ripeness and spawning during the period of rapidly rising sea temperatures in late spring and summer.

In NSW, by contrast, breeding is subject to less favourable and irregular seasonal temperature regimes. As coastal sea temperatures in central NSW generally rise above the 19ºC breeding threshold by November and remain so until as late as July (Figure 17), the window for reproductive conditioning can be as short as 4 months (120 days). This is less than the minimum threshold of about 150 days at sub 19ºC temperatures required by blacklip abalone to attain ripe, ready-to-spawn condition. Regular and predictable spawning is further disrupted by large ocean eddies. These break away from the East Australian Current around Fraser Island in southern Queensland and then course slowly southward down the NSW coast. Between three and seven such eddies occur each year. If close to shore, the eddies bathe the coast in warm (up to 20ºC in winter and 25ºC in summer).
subtropical seawater, but if offshore they can bathe the coast with deeper up-welled seawater as cool as 14 to 18ºC at any time, including in summer.

![Graph showing coastal surface sea temperatures at Port Stephens from April 2002 to December 2004. Bars are standard deviations.](image)

**Figure 17** Coastal surface sea temperatures at Port Stephens in central NSW from April 2002 to December 2004. Bars are standard deviations.

An important practical implication for abalone hatcheries in NSW is that for much of the year wild blacklip abalone are asynchronous in their reproductive development. At Port Stephens, near the northern limit of the commercial abalone fishery, wild stock commonly include many females with large gonads swollen with over-ripened eggs. Even after many months of conditioning it is not possible to confidently assess whether such abalone are ripe and ready to spawn or not. In most cases they have to be subjected to repeated spawning attempts at 1 to 2 month intervals before responding. However once they have spawned they can usually be expected to spawn good-quality eggs every 150 to 200 days thereafter when held at optimal temperatures of 16 to 18ºC.

It is therefore essential that the conditioning history and spawning activity of individual broodstock be recorded and tracked. Suggested information to be recorded is provided in **Appendices 3a and 3b**. A simple diary record of the type illustrated in **Appendix 3c** is also recommended. Such records allow rapid easy tracking and collation of the information needed to maximise the spawning success in terms of yields of viable eggs and sperm. The first record-keeping operation is to enter the tag number(s) of each animal onto the spreadsheet, together with where and when it was collected and its sex. The date of each attempt to induce spawning must be linked to this record, together with the outcomes, successful or not. If spawning is successful, additional data on the quantity and quality of eggs or sperm produced and subsequent yields of fertilised eggs and of larvae put to set must also be recorded. This is particularly useful when establishing a new hatchery and for tracking down the causes of spawning failures and other problems contributing to poor or variable yields of eggs, larvae and juveniles.
These issues highlight the importance of tagging and record-keeping practices for determining which broodstock are most likely to spawn at any given time and for choosing the most opportune dates to induce spawning.

To ensure fast, efficient and reliable year-round reproductive conditioning and spawning, broodstock must be maintained with minimal stress and within a relatively narrow temperature range of 16 ± 2ºC. This requires controlled temperature-conditioning facilities and an uninterrupted supply of near-oceanic-quality seawater. Open flow-through systems consume large amounts of seawater that equate to at least the biomass of stock every minute. This is necessary to maintain broodstock within optimum ranges of water quality (including those specified in Table 1). In NSW, considerable energy may be required to chill incoming ambient seawater to the prescribed temperature of 16 ± 2ºC. Indeed, the temperature of incoming ambient seawater may need to be lowered by as much as 9ºC in summer. Incorporation of seawater recirculation in reproductive conditioning systems of the type illustrated in Figure 18 and Appendix 2 therefore offer considerable cost and energy savings. Water recirculation systems also give the broodstock considerable protection from sudden adverse changes in the quality of the source seawater, particularly at sites susceptible to the effects of storms, floods and associated pollutants and to toxic algal blooms.

### Table 1 Lethal tolerable and optimal rearing conditions

<table>
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<th>Optimal / recommended</th>
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<td>18</td>
</tr>
<tr>
<td>Larval rearing</td>
<td>?</td>
<td>?</td>
<td>18</td>
</tr>
<tr>
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<td>12-25</td>
<td>18-24</td>
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<td>Early juvenile</td>
<td>10-and 27</td>
<td>12-24</td>
<td>15-21</td>
</tr>
<tr>
<td>Late juvenile and adult</td>
<td>10 and 25</td>
<td>12-23</td>
<td>14-18</td>
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<td>32-36</td>
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<td><strong>Dissolved oxygen</strong></td>
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Some useful design and operating protocols for reproductive conditioning facilities are as follows:

- Use two or more independently operated temperature-controlled sub-units to insure against system failure and resultant gaps in year-round hatchery production.

- Operate within stocking densities and seawater exchange rates that promote high reproductive performance and general good health. In the case of the conditioning system described in Appendix 2, stocking rates of ≤ 1 kg broodstock biomass per 50 L of standing seawater volume, coupled with a net new seawater exchange rate of 3 to 5 volumes per day, are required, as are recirculation rates that equate to the entire volume of the system every 30 to 60 min. Thus, for every 1000 L of seawater in a system of the type illustrated in Figure 18 and specified in Appendix 2, seawater needs to be continuously recirculated sequentially through a solids-removal filter, a dissolved-nutrients filter (foam stripper) and a biological filter to convert ammonium to nitrite/nitrate at a minimum flow rate of 1000 L/h.

- Provide a dual-cycle air conditioner that can maintain recirculating seawater at 16 ± 2°C as a back-up in the event of seawater supply or chiller failure.

- Provide a stand-by generator capable of powering the main seawater supply pumps, air compressors, air-conditioning systems and (preferably) seawater chillers.

- Accommodate broodstock in communal tanks that provide ease of observation, access and cleaning.

- Clean and purge conditioning units of uneaten food and faeces, assess the health of abalone, and remove dead or moribund (dying, through sickness or injury) individuals at least twice weekly.

- Conduct regular (at least twice yearly) inspections of stock to detect, treat or replace those that are compromised by heavy infections of boring mud-worm and sponges.

- Apply a conditioning temperature of 16 ± 2°C that promotes rapid conditioning while also providing a reasonable buffer of time for detecting and responding to equipment and power failures. This is particularly important during the warmer half of the year, when power failures can expose broodstock to temperatures of ≥ 20°C that trigger epidemic spawning and/or resorption of developing eggs.
Figure 18  Controlled temperature reproductive conditioning system that employs recirculation and filtration of seawater to greatly reduce energy costs of heating or cooling (detailed specifications provided in Appendices 2 & 3).
**STEP BY STEP SUMMARY AND CHECK-LIST FOR CONDITIONING BLACKLIP ABALONE**

☑ 1. Handle stock as gently and as quickly as possible during all phases of collection and transportation back to the hatchery.

☑ 2. As soon after collection as practicable, measure and double-tag broodstock selected for size, health and gender.

☑ 3. Establish a data file for each batch of abalone collected; record data of the type presented in Appendix 3.

☑ 4. Place animals into a temperature-controlled conditioning facility operated at 16 ± 2°C.

☑ 5. Maintain the stock under conditions that minimise all forms of stress. This includes all forms of physical disturbance, including handling, background noise, vibration and high or variable light intensity. It also includes maintenance of high water quality (especially pH, ammonium nitrogen and dissolved oxygen), within the optimum ranges specified in Table 1.

☑ 6. Feed broodstock a conditioning diet to satiation but not to excess.

☑ 7. Siphon uneaten food and faeces out of the broodstock-holding tanks and replace with isothermal seawater at least twice weekly, but change the water filter elements and clean out the foam strippers daily.

☑ 8. After each attempted induction of spawning (see Chapter 4), record the date and the spawning success or failure of individual stock. Confirm their gender, vigour, and health, and especially rates of infestation of the shell by boring sponges and mud-worm.
Chapter 4 Induction of Spawning and Fertilisation of Eggs
Chapter 4  Induction of Spawning and Fertilisation of Eggs

Introduction

The cheapest, yet most reliable, method of inducing ripe abalone to spawn is to bathe them in seawater treated with the oxidant hydrogen peroxide (H₂O₂). An alternative method widely used by commercial abalone farms in Australia is to expose the incoming seawater to a specialised wavelength of UV (ultraviolet) light that converts oxygen dissolved in the seawater to another strong oxidant, called ozone. Both ozone-generating UV lamps and hydrogen peroxide alter the chemistry of seawater by producing free radicals that can stimulate the abalone to release a sex hormone called prostaglandin. This, in turn, triggers a series of chemical pathways that induce final ripening and subsequent spawning of eggs and sperm. We recommend hydrogen peroxide because it acts faster and has a comparable success rate to that of ozone, does not require expensive ($4,000 to $10,000) specialised UV equipment, and avoids potentially serious operator safety issues associated with UV light irradiation.

Selecting broodstock for spawning induction

The number and ratio of male to female broodstock needed for each spawning operation depends on the intended use of the juveniles produced. If they are destined for on-farming, a ratio of one male to every two females is adequate. If selective breeding is involved, or if juveniles are to be used as seed to enhance depleted fisheries, then large and equal numbers of male and female broodstock should be used to help ensure the greatest degree of genetic diversity of their offspring.

As previously discussed, sexing blacklip abalone can be difficult, especially in individuals with deeply pigmented skin that obscures the colour of the underlying gonad. Often the only reliable way to sex darkly pigmented individuals is to record their sex when they first spawn.

Spawning induction

A separate hatchery room with controlled temperature and light/dark cycles is required for spawning induction, fertilisation and incubation of eggs and for subsequent rearing of larvae. The room should have a continuous supply of freshly pumped, temperature-controlled (16 ± 2°C) seawater that is filtered to 1 μm (nominal) and UV-disinfected. Air temperature should also be maintained at 16 ± 2°C using a conventional dual-cycle air-conditioner.

It is essential to keep hatchery rooms and equipment clean and sterile through chemical disinfection, as described in Appendix 6. Two 50 to 100 L squat open-top white plastic bins, one for disinfecting, the second for rinsing off the bulk of residual disinfectant, are required. Fill the first (disinfection) bin with fresh water and make up a 100 mg/L chlorine disinfection solution by adding 1 mL of pool chlorine (10% to 13% w/w sodium hypochlorite solution) per litre. Just before
use it is good practice to hose off all equipment (spawning containers, buckets, beakers and all other containers and utensils) with 1 µm-filtered seawater immediately before dipping and draining through the disinfection and rinsing tubs. The latter will ensure the removal of all vestiges of chlorine and freshwater. These procedures also apply to all air and water supply lines, which can be disinfected by being dismantled and left to soak in the disinfection tub between successive spawning operations or by purging with 100 mg/L chlorine solution and then being allowed to drain and air dry. Always drain the rinsing bin at the end of a day’s use and leave to drain and dry before the next use.

During spawning induction it is essential that handling and other forms of disturbance to abalone are kept to a minimum. Only broodstock that meet the following criteria should be selected for spawning induction:

- have been grouped according to gender
- have undergone 150 to 180 days of conditioning at 16 ± 2°C since their previous spawning
- are in good general health.

Place from one to a maximum of three abalone in small (5 to 10 L) shallow plastic tubs of the type illustrated in Figure 19. A single broodstock per tub is recommended, as it gives certainty as to whether or not an abalone actually spawns and, if so, when it spawned and the quantity of eggs or sperm released. A further advantage of individual accommodation is that it avoids the possibility of accidental fertilisation of eggs, as may occur if an abalone is incorrectly sexed.

The spawning tubs should be mounted on shelves 0.3 to 0.6 m above floor level for ease of access, ease of handling and monitoring of broodstock, and ease of siphoning of eggs and sperm (Figure 19). The tubs should be supplied with seawater at a flow rate of at least 1 L per abalone per minute. The seawater must be freshly filtered to 1 µm, UV-disinfected, and pre-chilled or heated to the prescribed temperature of 16°C. White or clear plastic tubs are preferable for females, because the dark-coloured eggs can be seen more easily. Conversely, black tubs should be used for males, as the sperm turns the water milky even at low concentrations.

To minimise handling stress on the day of spawning, broodstock should be moved into the spawning-induction tubs on the preceding afternoon and maintained in flowing seawater at 16°C. First thing the following morning, the seawater temperature is raised by 2°C to 18°C and the broodstock left undisturbed for 2 h of ‘priming’ at the elevated temperature.

The chemical TRIS (MW = 121.14 g), which is used as a pH buffer, is added to the spawning tubs at the rate of 6.6 mL of a 2-molar (M) solution per litre of seawater to adjust the pH to 9.1. After a further 5 to 15 minutes, add hydrogen peroxide (H₂O₂) solution at the rate of 3 mL of a 6% solution per litre of seawater. The abalone are then left undisturbed in the dark for a further 3 h before the tubs are emptied and thoroughly flushed with filtered, UV-disinfected, 18°C seawater to remove residual hydrogen peroxide and TRIS buffer. The tubs
are then refilled with filtered, UV-disinfected seawater at 18°C and aerated to maintain a high level of dissolved oxygen in the absence of water exchange.

**Figure 19** Left: Stacked shelf tray system used to accommodate abalone for induction of spawning. Right: Siphoning eggs into fertilisation container.

The broodstock are again left undisturbed in the dark for a further 1 h before inspections are started at 30 minute intervals for the next 2 h, then every 15 min until spawning commences. Males usually spawn sooner than females (generally within 1 to 3 h) and continue to spawn for a longer time. A compensating delay in the induction of males can be applied to help synchronise the spawning of the two sexes. Sperm is released via the respiratory pores like plumes of white smoke that quickly disperse, turning the seawater milky. The sperm can remain viable for a few hours at a room temperature of 16°C, and for longer if they are stored in a domestic refrigerator at about 4°C. Males often produce copious amounts of sperm that are usually well in excess of that needed to fertilise all the eggs produced.

As the viability of sperm can be substantially reduced if the sperm are stored at high concentration, it is advisable to dilute dense suspensions of sperm. Copiously spawning males should be progressively moved to new tubs of seawater (three tubs may be necessary). Tubs of sperm suspension should be discarded if the time from spawning exceeds an hour. Before fertilising the eggs, estimate the density of the fresh sperm suspensions in the tubs. This can be done with a **haemocytometer**, a simple cell counter consisting of a thick glass slide with a very fine set of gridlines etched on the surface, over which a special heavy-duty glass cover slip is laid. To count the sperm, take a small (about 1 to 10 mL) sample of sperm suspension and kill the sperm by mixing in one drop of 10% solution of formaldehyde per mL of the sperm suspension. Alternatively, you can use one drop of an iodine disinfection solution per mL of sperm suspension, which both kills and stains the sperm.
Place a few drops of killed sperm suspension on the grid of the haemocytometer immediately adjacent to the cover slip and allow it to be drawn under the cover slip by capillary action. Observe the sperm at a magnification of 400x using a compound microscope fitted with a 10x eyepiece and 40x objective lens. A complete step by step account of the sperm counting procedure is provided in Appendix 5. If sperm suspensions are too dense to be easily counted, dilute the suspension with fresh filtered, UV-disinfected seawater until the appearance of the suspension is reduced to a slight milkiness.

Female blacklip abalone will start to release eggs from 3 to 5 h after the water change. Like sperm, eggs are also released via the respiratory pores. The colour of blacklip abalone eggs ranges from olive green to maroon but tends toward the latter when the broodstock are fed artificial diets. The released eggs are quite dense and rapidly sink to the floor of the spawning tubs. In preparation for fertilisation, the eggs are siphoned into 20 L white plastic buckets via coarse (1 mm) mesh screens to remove faeces and other foreign matter shed by the broodstock before spawning. Eggs should be siphoned by using a 5 to 10 mm diameter clear food-grade plastic hose.

Check the quality of the eggs by examining a sample of one to several hundred, either on a cavity slide or in a ‘weighing boat’ under a stereo or compound microscope at 50× to 100×. The eggs should be round and surrounded by a thin clear layer called the vitelline layer (Figure 20), and the yolk should be uniformly dark and dense. If the eggs are uneven in size or misshapen, or the yolks are blotchy, they are unlikely to show high levels of fertilisation or yield good quality larvae. Such eggs should therefore be discarded. As eggs do not remain viable for as long as sperm, and their viability steadily decreases over time, they should be fertilised within 1 h of spawning.

Fertilisation of eggs

Before fertilising the eggs, estimate how many good quality eggs have been collected. Next, dilute the eggs in a graduated beaker or bucket with filtered, UV-disinfected, 18°C seawater. Next, disperse the eggs into an even suspension by using a perforated-disc homogeniser, as illustrated in Figure 21.
Figure 21 Drawing: Dispersion of eggs into even suspension using a perforated disc plunging homogeniser

Figure 22 Sampling of eggs using an automatic pipette (above) and placing them into small weighing boats (right) for counting by eye or with the aide of stereo microscope at low magnification
The homogeniser must be plunged gently up and down while avoiding contact with the walls or floor of the beaker. While doing so, take from three to five 1.0-mL samples of the egg suspension with an automatic pipette (Figure 22). Dispense each sample into a small container such as a weighing boat (Figure 22). The container should be white, making the dark eggs easy to see and count. Eggs can be counted with the naked eye or with the aid of a stereomicroscope at 5× or 10×.

If there are too many eggs in the 1.0 mL samples to be easily counted, dilute the egg suspension and repeat the above sampling and counting procedures. Alternatively, dispense 1.0 mL samples onto a Sedgewick rafter slide (a specialist microscope slide with a shallow 1 mL reservoir, the floor of which is divided into 1 × 1 mm grids), where they can be counted grid by grid at 5× or 10× with the aid of a stereomicroscope fitted with a mechanical stage. After counting a minimum of three samples, calculate the average, then multiply this value by the total volume (in millilitres) of egg suspension in the bucket to estimate the total number of eggs.

Once the densities of sperm and eggs have been determined, it is time to fertilise the eggs. Before adding the sperm suspension, quickly check that the sperm are still highly motile by examining them under a microscope at 100× to 400×. If they are not, use more recently shed sperm after again determining the density as described above. For best fertilisation results, the final concentration of sperm should be 10^4 to 10^5 sperm/mL and the density of eggs around 100/mL (Figure 23). These egg and sperm densities should yield an optimum ratio of 100 to 1000 sperm for each egg.

![Figure 23](https://example.com/figure23.png)  
*Figure 23* Effect of sperm concentration on fertilisation success of eggs in suspension at 100/ ml (as indicated by yield of trochophore larvae).

A simple way to calculate the amount of sperm suspension needed to optimise fertilisation is to multiply the number of eggs to be fertilised by 1000 to give the required number of sperm and then calculate the volume of sperm suspension (of
known concentration) that will contain this many sperm. Pipette this volume of sperm suspension into the bucket containing the eggs while at the same time keeping the eggs in continuous, even suspension by using a plunging homogeniser, as previously described for egg counting.

After 2 to 3 min, resuspend the eggs uniformly, take a 1.0 mL sample, and examine it under the compound microscope at 100×. Optimal fertilisation of eggs is indicated when, by focusing on the periphery of individual eggs, three to 10 attached sperm can be seen (Figure 24). If there are too few sperm, some of the eggs will not be fertilised. However, too many sperm will dissolve the outer protective layer of the eggs or will cause polyspermy (i.e. more than one sperm fertilises the egg). Both problems will lead to lower hatch rates.

**Figure 24** Optimum sperm to egg ratio (3-10 sperm per egg).
If there are fewer than three sperm per egg on average, add a further 25% to 50% volume of the sperm suspension, wait a few minutes, and again estimate the average number of sperm attached to individual eggs. If, however, there are more than 10 sperm per egg, rinse the eggs immediately with 1μm filtered and UV-disinfected seawater. Once the optimum number of sperm per egg is observed, leave the eggs and sperm for a further 5 min. Next remove the eggs from further contact with the sperm by rinsing them with filtered, UV-disinfected seawater. To do this, gently pour or siphon the eggs onto an 80 to 100 μm mesh screen mounted in a circular shallow plastic tub pre-filled with 1μm-filtered and UV-disinfected seawater at 18ºC. The screen should have a wall height substantially greater than the depth of the weir of the tub (Figure 25) to prevent loss of eggs through spillage or overflow. The mesh floor of the screen should always remain submerged to prevent the delicate eggs from being damaged. Maintain a constant moderate flow of freshly filtered, UV-disinfected 18ºC seawater through the screen while gently raising, lowering and swirling the screen for 15 to 30 seconds to ensure the eggs are thoroughly cleansed of residual sperm.

Figure 25 Screen and tub for wet harvesting and rinsing of eggs and larvae. N.B. Wall height of screen should be substantially greater than the depth of the tub to prevent loss of eggs or larvae through spillage or accidental overflow.
**STEP BY STEP SUMMARY AND CHECK-LIST FOR SPAWNING INDUCTION**

*Preparations the day before spawning:*

☑ 1. Rinse and disinfect floors and walls of the spawning facility, spawning tanks, airlines and seawater lines with a dilute chlorine solution (see Appendix 6) and set up spawning induction tubs and air lines ready to receive broodstock.

☑ 2. Activate reverse-cycle air conditioner set at 16 °C.

☑ 3. Prepare stock solutions of the chemicals required, namely:
   - 2M TRIS solution (142 g of TRIS per litre of distilled water)
   - 6% hydrogen peroxide (H₂O₂) solution: dilute 1 volume of off-the-shelf (30% w/w) hydrogen peroxide solution with 4 volumes of distilled water. (Note: This solution should always remain stored in a sealed dark glass bottle to conserve its potency).

☑ 4. Fill spawning tubs with 16 °C, 1-µm-filtered and UV-disinfected seawater and continue to supply this seawater at a continuous flow through rate of 1 L/min.

☑ 5. Check spawning history records of stock to identify male and female abalone most likely to be ripe and ready to spawn (i.e. those that previously spawned 150 to 180 days earlier) and transfer them into individual aerated spawning tubs to acclimatise overnight.
**STEP BY STEP SUMMARY AND CHECK-LIST FOR SPAWNING INDUCTION [CONT’D]**

*On the of morning spawning induction:*

☑ 1. Drain and rinse tubs clean of faeces, then resume flow of 1-µm-filtered and UV-disinfected seawater at the rate of 1 L/min.

☑ 2. Raise water and room air temperature by 2 ºC to 18 ºC (but no higher) and leave abalone undisturbed for 2 h.

☑ 3. Stop seawater flow but provide aeration to the tubs.

☑ 4. Adjust the pH of the seawater in the tubs to 9.1 by the addition of TRIS stock solution at the rate of 6.6 mL of 2 M TRIS solution/L of seawater, then wait for 5–15 min before adding hydrogen peroxide stock solution at the rate of 3 mL of 6% peroxide (H₂O₂) solution/L of seawater.

☑ 5. Leave the abalone in darkness and do not disturb them.

☑ 6. After 3 h, empty and wash out each tub, then re-fill with freshly pumped 1-µm-filtered and UV-disinfected seawater at 18 ºC.

☑ 7. Leave the abalone in the dark, undisturbed for another hour.

☑ 8. Start checking the abalone every 30 min over the next 2 h and then, if they have not started to spawn, check every 15 min until they do. (Pre-spawning abalone commonly crawl upwards close to, or breaking, the surface, then exhibit a characteristic rearing and waving posture just before and during spawning.)

☑ 9. Collect, rinse and fertilise eggs as described in the sub-routine below.

☑ 10. At the end of spawning, record for each broodstock whether they spawned or not, and, if they did, a) the start and finish times of spawning and b) the quality (degree of clumping, and uniformity of shape and yolk density) and quantity of eggs and of sperm (% motility and amounts). Note the tag numbers of abalone that spawned, confirm the gender, and (in the case of females) record the quantity and quality of eggs.
STEP BY STEP SUMMARY AND CHECK-LIST FOR SPAWNING INDUCTION [CONT’D]

Fertilisation sub-routine:

1. After the abalone stop spawning (or sooner if you already have the required number of eggs and sperm), check the quality of each batch of eggs under a microscope at 40x. Good quality eggs suitable for fertilising are round, not irregular. They have a clear outer vitelline layer that is distinctly separated from the margin of the yolk, which should be uniformly dark (high density) and not mottled.

2. Inspect sperm under the microscope at 100x to 400x. Sperm suitable for fertilising the eggs must be highly motile (> 90% wriggling vigorously). If not, use fresher sperm that are.

3. Collect and mix approximately equal contributions of fresh sperm suspension from as many males as available in a clean beaker. Estimate the sperm density with a haemocytometer (cell counter) as described in Appendix 5.

4. Estimate the number of eggs collected (commonly 0.5 to 3 million per spawner), as follows. Harvest all good quality eggs into a single 200-L bucket and make up to a known volume with 1-µm-filtered UV-irradiated seawater.

5. Gently agitate the eggs into an even suspension with a perforated plunger/homogeniser and take three to five 1.0-mL samples.

6. Dispense each 1.0-mL sample onto a weighing boat and count by eye, or if need be, onto a Sedgewick rafter slide, and count the eggs at low magnification.

7. Multiply the average count by the volume (in millilitres) of the egg suspension to estimate the number of eggs in the bucket.

8. Calculate how much of the mixed sperm suspension is needed to provide a sperm to egg ratio of 1000:1.

9. Add the sperm suspension while continually mixing with the homogeniser (perforated plunger) to maintain an even dispersion of eggs and sperm.

10. After a few minutes examine a small sample of eggs at 40x or 100x. If there are too few (average less than three) sperm attached to the periphery of eggs, add 25% to 50% more sperm suspension, but if there are too many wash the eggs immediately in clean 1-µm-filtered and UV-disinfected seawater.

11. About 5 min after adding the sperm, transfer the fertilised eggs to a flooded 80- to 100-µm screen and rinse them for about 1 min with 1-µm-filtered, UV-disinfected seawater that has been temperature matched (18 ºC).

12. Transfer the fertilised eggs from the screen back into a clean bucket before gently pouring or ladling them into a cylindro-conical larval rearing vessel pre-filled with 1-µm-filtered, UV-disinfected and temperature-matched (18 ºC) seawater.
Chapter 5 Incubation of Eggs and Larval Rearing
Chapter 5 Incubation of Eggs and Larval Rearing

On average, about 40% of fertilised eggs spawned by well-conditioned blacklip abalone broodstock can be expected to yield 6- to 8-day-old competent (ready and able to settle) larvae. In turn, 10% to 30% of competent larvae put to set can be expected to reach a minimum weaning size of 1 mm if the hatchery and plate nursery techniques recommended below are used.

Abalone larvae are easy to rear because their planktonic larval stage is brief and does not require feeding. The most important factor for successful rearing of the larvae is strict hygiene. As described for spawning and fertilisation, all equipment used for incubating embryos (fertilised eggs undergoing successive cell divisions and cell differentiation) and rearing larvae should be washed, disinfected and thoroughly rinsed before and after use. Good survival and significant growth of post-larvae are confined to temperatures ranging from 10 to 26ºC. Maximum growth rate occurs at about 22ºC (Figure 26), while highest survival of post-larvae occurs at a lower temperature of about 19ºC (Figure 27). At a comprise temperature of 18ºC, recommended for fertilisation and incubation as well as for larval rearing, larvae are best put to set after 6 to 8 days of rearing. The final decision is based on their ‘stickiness’ that is the tendency to cling to surfaces and to each other.

Although abalone embryos and larvae can be successfully reared by using a range of alternative batch and flow-through systems described in an earlier companion manual (Hone et al. 1997), we describe only a flow-through system that has proved reliable under the stewardship of a number of hatchery managers over a total period of 7 years. Regardless of design, rearing tanks and associated equipment such as banjo screens should be thoroughly washed and disinfected with dilute (100 mg/L) chlorine solution and then thoroughly rinsed with fresh water, drained and air-dried the day before induction of spawning.

Using the flow through hatchery equipment illustrated in Figures 28, 29 and 30, initial incubation and hatching can be conducted under static or low rates of water exchange. In the latter case, fresh 1 µm filtered and UV-disinfected seawater at 18ºC is continuously supplied at a rate of about one vessel volume exchange every 4 to 6 h. A banjo filter screen fitted with fine (60 µm) polyester mesh is connected to an overflow mounted through the upper wall of the vessel to prevent loss of developing embryos. The appropriate diameter of banjo screen is 200 mm for 150 to 200 L rearing vessels and 300 mm for larger vessels up to 500 L. Overflow/wastewater discharge outlets should have an internal diameter of 20 or 25 mm.

Regardless of whether incubation and hatching are conducted under static or low flow-through conditions, aeration must be provided from the base cone of the culture vessel as moderately sized (5 to 20 mm diameter) ‘slug flow’ bubbles. This is to prevent any dead spots from developing and to ensure that the embryos and larvae are continuously and evenly dispersed through the full depth of the water column. Slug flow aeration also helps to prevent the fine-mesh banjo screens from blocking in the case of flow-through exchange and also eliminates
fine air-bubble entrapment of embryos or larvae. Slug air-flow is regulated by using a simple thumb-screw clamp on the clear silicon-rubber air supply hose at the base of the tank; this hose doubles as a drain (Figure 30).

A sample of embryos should be taken and checked at 40× to 100× under a compound microscope 18 to 20 h after fertilisation. By this time they should have completed incubation, hatched and reached the first trochophore larval stage (see Figure 6, Chapter 1). At this time the air and seawater supply should be temporarily stopped for 10 to 15 min. This is to aid the separation of unhatched embryos, empty egg cases, non-viable larvae and other organic debris from active healthy trochophore larvae. In still water, trochophores are photopositive and will swarm to the surface in pursuit of the brightest available light.

At this point the surface-swarming trochophore can either be directly ladled off and transferred into a new, clean larvae-rearing tank pre-filled with temperature-equilibrated seawater, or left for a further 24 h in the same tank operated as before, except that the original banjo screen is exchanged with a clean screen and seawater flow-through at about one exchange every 2 to 3 h is imposed.

Drain harvesting (see below) should not be practised until the larvae have reached the fully shelled veliger stage (36 to 48 h after fertilisation at 18°C; see Figure 6, Chapter 1). To harvest the larvae, a polyester 60-μm mesh flooded harvesting screen (Figure 31) is first set up as previously described for washing fertilised eggs. Immediately before harvesting, aeration is shut off by tightening the bottom drain/aeration-line clamp and allowing dead larvae and other debris to settle out for a few minutes. The clamp on the drain/aeration-line is then loosened and about 20% of the seawater and associated debris within the conical base of the rearing vessel is drained to waste. Next the larvae are gently drain-harvested onto the flooded screen.

Once all the larvae have been collected on the screen they are gently rinsed into a 20-L bucket. The bucket is then topped up with 1 μm filtered, UV-disinfected 18°C seawater to one of several reference volume points marks (5, 10, 15 or 20 L) clearly marked on the inside. The larvae are next gently agitated into suspension by using the plunging homogeniser (as described above for egg counting) until they are evenly dispersed. While they are evenly dispersed, three to five 1.0 mL samples of larval suspension are randomly collected with an automatic pipette. Larvae within these samples are counted by following the same detailed procedure, involving the use of a Sedgewick rafter slide and microscope, already described in the preceding chapter for estimating numbers of eggs.
**Figure 26** Effect of temperature on the growth (shell length) of 6-day post-settlement blacklip abalone post-larvae. From Heasman et al., 2006.

**Figure 27** Effect of temperature on survival (% yield) of 6-day post-settlement blacklip abalone post-larvae. From Heasman et al., 2006.
**Figure 28** Sketch of egg incubation and larval rearing vessels comprising 150 to 500 L cylindrical fibreglass or moulded plastic vessels with steeply sloped (45° to 60°) conical bottoms.

**Figure 29** Overhead view of incubation/larval rearing vessel showing the internal surfaces that must be smooth and scratch-resistant, and the base, that should be conical and white to aid hygiene and visual assessment of the condition of developing larvae.

**Figure 30** A thumb-screw clamp attached to a clear silicon-rubber air input and drainage hose is the simplest way to provide regulated slug air-flow within larval rearing vessels.
Finally, the larvae are restocked at up to 20 larvae/mL into a new sterile rearing vessel pre-filled with 1 μm filtered, UV-disinfected, 18 °C seawater. Continuous seawater exchange of about one vessel volume exchange every 2 to 3 h and gentle slug aeration are then resumed. These harvesting, counting and re-stocking procedures are repeated on alternate days until the larvae are ready (competent) to settle and metamorphose.

The larvae are competent to settle after 6 to 8 days of rearing, over which time they become progressively more ‘sticky’, adhering to objects they encounter, including each other, to form chains and rafts, particularly at or near the surface of the rearing vessel. However, some may begin to crawl on the base and sides of the vessel before this. Larvae should be routinely inspected in the rearing vessel and under the microscope at least once a day to assess developmental stage, level of activity, the extent of malformations and any obvious dropout of larvae from the water column or unusual aggregations. The most important malformation to look for is a breakdown of attachment of the twin retractor muscles to the inside (integument) of the larval shell (see Figures 6 and 7 Chapter 1).

Loss of one or both attachments is commonly referred to as IDS (integument/muscle detachment syndrome). This is thought most likely to be caused by a bacterial infection arising from either poor seawater quality or inadequate hygiene. The procedures recommended here place total reliance on high flushing rate and good hygiene. In particular, daily changes of banjo screens, changes of rearing vessels on alternate days, and physical filtration and UV disinfection of seawater will keep potentially harmful bacteria (especially those belonging to a group called Vibrio) at bay and will prevent IDS. In our experience the most common primary cause of IDS is malfunction or inadequate maintenance of the UV disinfection units. **Antibiotics should never be used, nor should they ever need to be.**
STEP BY STEP SUMMARY AND CHECK-LIST FOR INCUBATING EMBRYOS AND LARVAL REARING OPERATIONS

1. Wash and disinfect all equipment with dilute chlorine solution (see Appendix 6) and rinse with 1-µm-filtered, UV-disinfected seawater.

2. Stock fertilised eggs into the larvae-rearing container filled with 1-µm-filtered and UV-disinfected seawater at 18 °C.

3. Apply gentle slug aeration and either operate as a static system or apply a low seawater flow of one exchange every 4 to 6 h (about 0.5 to 1 L/min) of 1-µm-filtered, UV-disinfected 18 °C seawater.

4. After 18 to 24 h, check that trochophore larvae have hatched, stop aeration and allow 10 to 15 min to allow non-viable eggs and larvae, egg capsules and other organic debris to sink and separate from photopositive trochophores at the tank surface.

5. Either:
   - drain off the bottom 20% of seawater to remove settled debris, refill the tank and resume aeration and apply/re-apply low water exchange for a further 12 to 24 h to allow development to the fully shelled veliger larval stage
   - ladle-off surface swimming trochophores and transfer them to a new, clean rearing vessel pre-filled with 1-µm-filtered, UV-disinfected seawater at 18 °C.

6. After 36 to 48 h, drain harvested larvae into a flooded screen and rinse veligers, then transfer to a clean bucket and top-up to known volume reference mark.

7. Estimate the number of larvae by counting those in three to five 1.0-mL sub-samples and then multiplying the average larval count per millilitre by the volume of the bucket in millilitres.

8. Restock the larvae at a maximum density of 20/mL into a clean and sterile rearing vessel pre-filled with 1-µm-filtered, UV-disinfected seawater at 18 °C.

9. Repeat flooded screen harvests, counts and tank changes on alternate days. Observe larvae in the tank and examine a small sample microscopically, recording general appearance and behaviour, development, and health status every day.

10. At day 6 start checking the larvae under the microscope for competence.
Chapter 6  Nursery Production

Overview of settlement of larvae on diatom plates

There is much published information on how to settle and grow abalone post-larvae. In practice, successful settlement of larvae and rearing of post-larvae are relatively easy. As discussed in Chapter 1, competent abalone larvae settle in response to chemical cues given off by substrates that constitute their preferred habitats that can support early juvenile survival and growth. It is believed that larvae first detect the chemicals in the water column and then inspect the bottom substrate, possibly by tasting it. If the substrate has the right properties the larvae will stay attached and metamorphose. Otherwise they resume swimming and continue their search. In the case of blacklip abalone, wild competent larvae settle on rocky surfaces overgrown by crustose coralline algae, especially in shallow (1 to 5 m deep) exposed boulder areas subject to high wave energy and disturbance.

To successfully settle larvae and on-rear them as post-larvae in the nursery, settlement substrates must stimulate initial settlement and metamorphosis and also provide adequate food for early growth. Diatom-dominated biofilms are excellent for this. There are four key procedures that will ensure reliable and efficient settlement and metamorphosis of larvae and subsequent rapid growth and high survival of blacklip abalone post-larvae to a stage at which they can be reliably weaned onto formulated diets.

These procedures are:

1. Initial inoculation and colonisation of plastic nursery plates with the alga *Ulvella* *lens*. This green alga can be easily induced to develop and release motile spores that immediately attach to the surfaces of the plates and rapidly form thin plaque-like colonies of cells (Figure 12e, Chapter 1).

2. In turn, these colonies, like those of the much slower-growing crustose coralline algae, attract and promote rapid and consistently high rates (40% to 80%) of settlement and metamorphosis of post-larvae.

3. Follow-up preparation and management of plastic nursery plates to ensure that post-larvae can feed to satiation at all times through to a minimum average weaning size of about 1.5 mm on nutritious diatom-dominated biofilms.

4. Larvae are seeded onto the plates within the prescribed range of density and at favourable temperatures that help ensure consistently high yields of weanable post-larvae.

5. Post-larvae are carefully harvested off the plates and weaned onto formulated diets while they are still in peak growth and health and before they have fully exhausted the diatom-dominated biofilm on the plates.
Nursery facilities equipment and operation

In most abalone hatcheries in Australia, larval settlement plates are 600 x 300 mm and constructed from clear 1 to 1.5 mm thick PVC. These plates are mounted vertically on their long side in plastic coated or stainless steel wire baskets that hold 14 to 18 plates (Figure 32). The baskets are fully submerged in partially shaded shallow tanks commonly located under enclosed or semi-enclosed green-house like canopies (Figure 33).

Figure 32 A small-scale abalone nursery comprising baskets of plastic plates submerged in partially shaded tanks located within enclosed or semi-enclosed shade houses.

Figure 33 Standard 600 x 300 x 1.5 mm clear PVC nursery settlement plates. These plates are mounted on their long edge in plastic coated or stainless steel wire baskets that hold 14 to 18 plates.

Figure 34 Cutaway drawing of nursery tank

Figure 35 Nursery tank filter element being changed.
The tanks are fitted with standpipes to maintain a water depth of about 0.4 m. Seawater sprinkler bars are mounted centrally over each row of baskets. A high level of aeration is maintained across the surfaces of all plates continuously via floor-mounted 16 mm air diffuser hoses. Two rows of diffuser hose are anchored to the floor of the tanks beneath each row of baskets (Figure 34). Seawater entering the nursery tank is pre-filtered to 10 µm (Figure 35) to prevent the entry of zooplankton, especially tiny crustaceans called copepods that rapidly multiply and directly compete with the abalone post-larvae for food and space on the plates. Unrestricted entry of plankton will also lead to the settlement and proliferation of an undesirable epi-biota (mixed layer of plant and animal life) on the surfaces of plates. Such undesirable epi-biota commonly include filamentous and thin sheet (thallus) forms of colonial algae and an array of other invertebrates, including oysters, mussels, limpets, tube-worms, barnacles, sponges and hydroids, as well as the above-mentioned copepods.

**Preparation of diatom plates**

(For a comprehensive description of these procedures see Table 2 and the companion manual by Daume 2004).

Between 3 weeks (in summer) and 6 weeks (in winter) before starting a new hatchery/nursery production cycle, nursery plates already in use that have developed large mature patches of Ulvella lens are selected. Juvenile abalone, if present, are removed by using the standard anaesthesia techniques described below. One mature Ulvella plate is required for every basket of new plates to be seeded with abalone larvae.

The mature Ulvella plates are wiped clean with a sponge to remove diatom films and other unwanted biota before being stored for 2 weeks (summer) to 4 weeks (winter) in a separate, heavily shaded tank, where they are supplied with 1 µm filtered flow-through seawater and light aeration. This conditioning period encourages spore formation within the mature Ulvella colonies.

One week (summer) to 4 weeks (winter) before the proposed start of a hatchery/nursery production cycle, all nursery tanks and plates to be used are scrubbed clean. The tanks are filled with fresh 1-µm-filtered seawater and then chlorinated at 10 mg/L (10 ppm) active chlorine (see Appendix 6) overnight. The next morning, sodium thiosulfite solution is added and vigorous aeration applied to remove any residual active chlorine (also see Appendix 6). Next, the mature Ulvella plates are removed from the shaded conditioning tank and placed singly between each basket of new clean plates. The new tanks are then left static for 4 days without shading to enable solar heating sufficient to raise the temperature up to a range of 20 to 25°C. The resultant sudden increase in light intensity and temperature triggers spore release by mature Ulvella colonies. Spore release generally peaks after about 4 days (summer) to 8 days (winter) and starts daily 1 to 2 h after sunrise. Normal flow-through of 1 µm filtered seawater can then be resumed, the Ulvella spores having settled and germinated on the new plates.
The next critical step is to promote the establishment and growth of nutritious types of adventitious (naturally occurring) benthic diatoms (see Figures 12a, b and c Chapter 1) on the new plates. Diatom productivity on settlement plates determines their ‘carrying capacity’ in relation to abalone post-larvae. This carrying capacity in turn is determined by temperature, light and availability of nutrients. Varying grades of shade-cloth, providing from 90% down to 50% shade, can be used to adjust light intensity and hence the growth of diatoms to help match the grazing rates of the abalone post-larvae. The ideal diatom biofilm looks like a thin scum covering the surfaces of the plates (Figure 36). An accumulation of brown scum will collect on a fingertip when dragged across such plates. If the diatom film is too thin initially and the post-larvae are at particularly high densities, they can prematurely graze out the diatoms and starve before reaching minimum weaning size. Conversely, if the biofilms are too mature they will form a thick, three-dimensional layer containing unwanted and nutritionally poorer chain forming microalgae and/or seaweeds (macroalgae) (Figure 36)—especially opportunistic species such as sea lettuce (Ulva) or its filamentous counterpart, mermaids hair (Enteromorpha).

Figure 36 Left: Inspection of plates being prepared for seeding. Top Right: Photomicrograph of benthic diatoms on well-prepared nursery plate at 500x. Below Right: Close-up view of diatom plate over grown by invasive filamentous algae.
Table 2 Summarised Schedule of activities for the preparation and operation of nursery diatom plates, integrating the use of *Ulvella* to enhance rate and degree of successful settlement and metamorphosis of blacklip abalone larvae seeded at high densities (4,000-6,000 larvae per standard 300 x 600 mm plate)

<table>
<thead>
<tr>
<th>Initiation time and duration of activity conducted at warmer seasonal temperatures of 19 to 23 ºC</th>
<th>Activity</th>
<th>Initiation time and durations of activity conducted at cooler seasonal temperatures of 14 to 18 ºC</th>
</tr>
</thead>
</table>
| **Day -21 to Day - 8**  
(3 weeks before seeding of plates with competent larvae) | Select from among nursery plates in current use those extensively covered with mature colonies of *Ulvella*. One such *Ulvella* inoculation plate is required for each rack of new plates to be prepared for seeding with abalone larvae.  
- Remove residual juvenile abalone and wipe off all other extraneous biota from *Ulvella* inoculation plates before placing in a deeply shaded nursery tank with low aeration and low flow input of 10-µm-filtered seawater. Maintain this status for 4 days (warm period) to 28 days (cold period) to promote development of spores. | **Day - 42 to Day - 15**  
(6 weeks before seeding of plates with competent larvae) |
| **Day - 7 to Day - 2**  
(1 week before seeding of plates with competent larvae, and coincident with initiation of spawning induction and hatchery production of abalone competent to set larvae) | Cease seawater exchange and remove shading to promote solar heating of seawater up to the range 20 to 25 ºC. Stock *Ulvella* seeding plates at the rate of one per basket among new, clean disinfected plates in nursery tanks.  
- Spore release by *Ulvella* is promoted by temperature rises, with peak releases occurring after about 4 to 7 days, depending on temperature. New disinfected plates with settled and germinated spores soon begin to proliferate across the plates. Addition of fertilisers (Guillard’s F/2 or Aquasol) will enhance these developments. | **Day - 14 to Day - 6**  
(2 weeks before seeding of plates with competent larvae) |
| **Day - 1**  
(1 day before seeding of plates with abalone larvae) | Resume low exchange rate of 1-µm-filtered ambient seawater and maintain low aeration. Add plant fertiliser (Guillard’s F/2 media or Aquasol) to promote further proliferation of *Ulvella* colonies and adventitious benthic diatoms.  
- Inoculate with pure strain cultures of high performance diatoms (if available). | **Day - 5**  
(5 days before seeding of plates with competent larvae) |
| **Day 0**  
(induction of spawning of abalone broodstock in ripe condition and initiation of hatchery rearing cycle) | Continue as above to encourage proliferation of diatoms, adding shading once a complete film of low-profile benthic diatoms (as indicated by finger-swipe test and/or microscopic inspection of plates) is established. | **Day 0**  
(induction of spawning of abalone broodstock in ripe condition and initiation of hatchery rearing cycle) |
| **Day +1 to Day +6**  
(hatchery production cycle) | Continue as above. | **Day +1 to Day +6**  
(hatchery production cycle) |
| **Day +7**  
(seeding of plates with competent larvae) | Maintain low aeration in nursery tanks and cease seawater exchange before adding competent larvae at the rate of 4000 to 6000/plate  
- Allow larvae 12 to 24 h to achieve full settlement and metamorphosis of larvae to post-larvae. | **Day +7**  
(seeding of plates with competent larvae) |
| **Day +8 onwards**  
(on-growing of post-larvae on plates) | Resume low exchange rate of 1-µm-filtered ambient seawater and apply moderate aeration. | **Day +8 onwards**  
(on-growing of post-larvae on plates) |
The speed of development and productivity of biofilms vary with the season, being greatest in the warmer and higher light-intensity months from late spring to early autumn (November to March) and lowest in winter and early spring (July to August). Biofilm productivity may also vary among different settlement tanks according to exposure to sunlight and solar heating. The most common and nutritious naturally occurring (adventitious) diatoms are single-celled species belonging to the groups *Navicula*, and *Nitzchia*. Each type of diatom is easily recognised by its distinctive shape as illustrated in Figure 12, Chapter 1.

As already stated, installation and conditioning of settlement plates pre-seeded with *Ulvella* spores should precede spawning induction by about a week. If spawning is delayed, the nursery plates can be shaded to prevent the diatom films becoming too dense and/or rangy and overgrowing the young colonies of *Ulvella*. The net effect of the latter is reduced ability of the biofilms to attract initial settlement and metamorphosis of the abalone larvae and to support high subsequent growth and survival of the post-larvae.

As growth of diatoms is much higher on the upper surfaces of the plates (especially the sides with highest exposure to sunlight), the plates should be alternately rotated in both the horizontal and the vertical planes once a week to ensure the best and most even productivity. If the growth of diatom biofilms surges ahead of consumption by post-larvae, additional shading of the plates may be required to prevent them from becoming overgrown by filamentous algae, which may also restrict the movement of the post-larvae.

Each abalone farm site and nursery configuration has its own particular adventitious diatom production characteristics that vary with the season and weather. Optimised management of nurseries leading to consistently high and reliable yields of weaning-size juveniles within these farms is therefore likely to entail several years of keen observation and fine-tuning by technical staff.

As indicated above, a much more comprehensive description of how to manage and optimise diatom productivity on nursery plates, including the use of specially selected highly productive strains of diatoms with or without the addition of one or a combination of supplementary plant nutrients, heating and illumination of tanks, is provided in a companion manual (Daume 2004).

**Seeding of larvae onto plates at optimum densities and favourable temperatures**

As discussed above, competent blacklip abalone larvae are able to successfully settle and metamorphose over a relatively broad range of temperature (see Figure 27). At temperatures of 12 to 24°C, which span the annual range of sea temperature on the central and southern coasts of NSW, survival of blacklip abalone larvae through settlement and metamorphosis remains at or above 75% of the peak rate at 18°C. However, subsequent growth rates of early-stage post-larvae (see Figure 26) are much more affected by temperature (especially low temperature), rising steadily from zero at 12°C to a peak at 23°C, then rapidly back to zero with a further small temperature rise to 27°C. Such temperature effects render the NSW Central Coast (Sydney to Port Stephens) the most...
suitable region for year-round nursery production of blacklip abalone, with little or no need to heat or cool ambient seawater at any time. However, even within this region, the periods required by post-larvae to reach a minimum average weaning size of about 1.5 mm can vary markedly from about 30 days in the warmest summer and early autumn months up to 70 days or more during winter and early spring (Figure 37).

**Seeding density**

Blacklip abalone post-larvae consistently show exponential growth (Figure 37), even when seeded on diatom plates over a wide range of densities throughout the year. Exponential growth implies that the time taken to double in size from any starting point remains constant. Depending mainly on seasonal temperature, blacklip abalone post-larvae reared on the Central Coast of NSW double their shell length every 12 to 36 days and their weight every 6 to 12 days. An important implication of exponential growth for managing diatom plate nurseries is that daily food consumption of post-larvae increases at the same proportional rate as their weight, which equates to a massive factor of 64 times between the time of first feeding (from the first week after settlement) to the time they reach an average minimum weaning size of about 1.5 mm, 30 to 70 days later.

An important consequence of exponential growth is that if the initial larval seeding density is too high then the resultant post-larvae may run out of food and stop growing before they reach the minimum weaning size. Conversely, if post-larvae that have reached weaning size are inadvertently left and allowed to graze out the diatoms, they also will stop growing and, if left for extended periods, will begin to suffer serious consequences of starvation. As illustrated in Figure 38, stalled growth and substantial (sometimes catastrophic) losses of post-larvae can result if starvation is allowed to continue for several weeks. Such losses may be a direct consequence of starvation. However, they may also reflect an inability of nutritionally compromised post-larvae to endure the combined stresses of anaesthesia, handling, and exposure to the air and sun during transfer from nursery plates to first stage raceways for weaning onto formulated diets. Generally speaking, the higher the seeding density of larvae on diatom plates, the higher the risk of stalled growth and starvation before the larvae can attain minimum weaning size.

In commercial hatcheries, the key to achieving consistently high yields of post-larvae from diatom plates is to seed them as larvae at densities that will allow the greatest number to maintain normal exponential growth through to the minimum average weaning size of 1.5 mm but without posing undue risk of them grazing out the diatoms and starving. This task is not difficult to achieve, providing that the following conditions are met:

- The larvae are healthy and competent to settle.
- The larvae are seeded onto diatom plates at densities of 1 to 1.5/cm² (= 4000 to 6000 per standard 300 × 600 mm nursery plate).
- Temperature remains within the range 15 to 23 ºC.
Our experience is that, regardless of the initial larval seeding density, after about a month the survival of post-larvae usually stabilises at 10% to 30% of the larvae originally seeded. Provided the post-larvae on diatom plates are in good health and growing normally (exponentially), they can be successfully harvested off the plates, restocked into shallow raceways and immediately weaned onto formulated diets with relatively few losses. This can be done at an average shell length as small as 1.0 mm, but, as already stated, it is best applied at an average shell length of 1.5 mm. As illustrated in Figure 39, well prepared and managed diatom nursery plates will consistently support normal exponential growth of up to 1000 post-larvae per plate (0.3/cm²) up to the recommended average weaning size of 1.5 mm.

Figure 37 Usual exponential growth pattern of post-larvae reared diatom plates on the central coast of NSW (100 microns = 1 mm). Note: Periods required by post-larvae to reach a minimum average weaning size of 1.5 mm can vary markedly from 30 days in summer and early autumn to more than 70 days in winter and early spring (Heasman et al., 2004).
Figure 38  Effect of over-grazing of diatom plates and subsequent protracted starvation on growth of blacklip abalone post-larvae. Bars indicate standard error of means. (Heasman et al., 2004)

Figure 39  Effect of density on growth of weaning size. Bars indicate common size range (95% confidence limits). (Heasman et al., 2004)
Figure 40 Above: section of heavily grazed nursery plate. Note: Blacklip abalone post-larvae exhibiting “grazing fronts” in relation to remnant patches of benthic diatoms. Inset: Recommended method of inspecting nursery plates.
Another factor critical to achieving consistently high nursery plate yields of post-larvae is to avoid exposing them to the air while assessing their size, number and health and/or the status of diatom films. The reason is that the upper and lower short-term lethal thermal tolerance temperatures of about 30 and 5°C, respectively, of tiny (0.3 to 2 mm) post-larvae can be attained in a matter of seconds if they are exposed to air temperatures that in southern Australia can reach 35 to 45°C during summer and 0 to 5°C during winter. Plates should therefore be visually inspected in accordance with the recommendations in Table 3.

Table 3  Recommended out-of-water inspection times for nursery plates

<table>
<thead>
<tr>
<th>Air temperature (°C)</th>
<th>Max recommended air exposure times for post larvae on nursery plates</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;10</td>
<td>Nil</td>
</tr>
<tr>
<td>10-25</td>
<td>1 minute</td>
</tr>
<tr>
<td>20-25</td>
<td>30 seconds</td>
</tr>
<tr>
<td>25-30</td>
<td>5 seconds</td>
</tr>
<tr>
<td>&gt;30</td>
<td>nil</td>
</tr>
</tbody>
</table>

Even then, such brief visual inspections should be confined to the early morning or night-time. This is because the post-larvae will sustain even more extreme temperature and/or associated osmotic shock if additionally exposed to high velocity and/or low-humidity wind and/or direct sunlight.

A much more preferable practice for the safe inspection of plates is to keep them submerged in shallow (75 to 100 mm) trays filled with ambient seawater at a minimum depth of 50 mm. Such trays should be equipped with a thermometer to ensure that the temperature is always maintained within the safe limits of 10 to 25°C for blacklip abalone post-larvae.

These simple precautions will enable protracted inspection, non-invasive counting, and even measurement of post-larvae in situ at any time of the day and under most weather conditions. Fast and accurate counting and measurement of post-larvae can be made with the aid of a stereomicroscope (Figure 40) mounted on an adjustable boom and fitted with calibrated eyepiece graticules.

If the above guidelines are adhered to, then standard 300 × 600 mm nursery plates will consistently yield an average of 200 to 1000 weaning-size post-larvae (Figure 39). As previously stated, for a hatchery located on the Central Coast of NSW, the minimum average weaning size of 1.5 mm will be attained in as little as 30 days in summer at temperatures of 20 to 24°C, but in up to 70 days in winter at temperatures of 14 to 16°C. For hatcheries located farther south in NSW and elsewhere, production of weanable post-larvae during the coldest months will require heating of incoming seawater when the temperature falls below about 14°C.
STEP BY STEP SUMMARY AND CHECK-LIST FOR PREPARING AND SEEDING SETTLING PLATES
(see also Table 2)

☑ 1. Three to 8 weeks before a planned spawning and hatchery cycle, depending on the season, select sufficient nursery plates bearing large areas of mature Ulvella colonies from among an ongoing nursery operation to provide one such plate per basket of 15 to 18 plates to be seeded.

☑ 2. Remove any abalone juveniles in accordance with the procedures described in Chapter 7. Gently rub and rinse off diatom films and other extraneous biota while preserving as much of the generally tougher and strongly adhering mature Ulvella colonies as practicable.

☑ 3. Collect the plates into a separate, fully shaded tank supplied with 10-µm-filtered seawater and light aeration for 2 to 4 weeks to encourage the development of Ulvella spores.

☑ 4. Five days before attempting to induce the Ulvella colonies to release spores, set up new larval settlement tanks by placing air diffuser tubes on the bottom of the tank and starting flow of seawater filtered to 10 µm.

☑ 5. The following morning, neutralise any residual active chlorine by first adding sodium thiosulfate solution (see Appendix 6) and then apply vigorous aeration to mix and distribute the thiosulfate before using a simple test kit to verify the complete neutralisation of chlorine (also see Appendix 6).

☑ 6. Insert one mature Ulvella bearing nursery plate for every basket of new non-seeded plates, and operate the tank unshaded and static with gentle aeration for 4 days (up to and including the day of induced spore release) to facilitate solar heating and high light-intensity-mediated induction of Ulvella spore release.

☑ 7. Restore flow-through of 10-µm-filtered ambient seawater and apply moderate aeration for a further 7 to 21 days to allow inoculation and preliminary proliferation of adventitious diatoms.

☑ 8. Regularly inspect the diatom growth on the plates. A good diatom biofilm appears as a thin brown scum on the plate. The quality of the diatom film in terms of dominant types of single-cell benthic diatoms (common species of Navicula and Nitzchia as illustrated in Figure 12) can be confirmed by examining the surface of a small but representative sample of plates under a dissecting microscope. Development of a good covering of benthic diatoms generally takes from 2 to 3 weeks in summer at 20 to 24 ºC but as long as 4 to 6 weeks in winter at 12 to 16 ºC. These times can also be shortened by one or a combination of the following procedures:

- run the tanks statically (temporarily stop seawater flow) with low to moderate aeration
- heat the tanks by using clear covers, such as common bubble-film packing, that allow solar heating by day but act as heat-retaining blankets at night (capable of raising ambient temperature by 2 to 4 ºC), and/or use submersible heaters. Well-insulated tanks will require only about 100 to 200 W of heating/ m² of tank volume.
- add plant nutrients to promote propagation of diatom films
- inoculate the plates with specially selected pure cultures of highly productive benthic diatoms, available from the CSIRO (see Daume et al. 2004).
STEP BY STEP SUMMARY AND CHECK-LIST FOR PREPARING AND SETTLING LARVAE

☑ 1. Once the larvae are confirmed as being competent to settle and metamorphose into post-larvae, wet-screen harvest them (see Figure 31).

☑ 2. Gently rinse the larvae from the wet screen into a bucket and top up the bucket to a known volume with fresh 10-µm-filtered, UV-disinfected 18°C seawater.

☑ 3. Mix the larvae well in the bucket with a hand-operated homogeniser (as for counting eggs; see Figure 21) and while doing this collect three to five 1.0-mL samples of the larval suspension with an automatic pipette (see Figure 22) and dispense onto a rafter counting slide.

☑ 4. Calculate the average number of larvae per millilitre and multiply this number by the total volume of the larval suspension in millilitres to determine the total number of larvae.

☑ 5. Turn off the water flow to the settlement tank and decrease the airflow to a trickle.

☑ 6. Add between 4000 and 6000 larvae for each settlement plate (1 to 1.5/cm²) in the nursery tanks, making sure the larvae are evenly distributed among the plates.

☑ 7. After 24 h, resume water flow and full aeration.
Chapter 7  Management of Nursery Raceways and Weaning

General management procedures

For the first few weeks after they have settled, abalone post-larvae do not have a fully formed gut. During the first week they gain nutrients from the remainder of their yolk stores and by ingesting bacteria mucus and other exudates (mainly sugars) released by crustose coralline algae and diatoms. They are also able to directly absorb soluble organic compounds from seawater through the gills and skin. Beyond the first week, with progressive development of the rasping tongue and the gut, they are able to consume, digest and absorb whole benthic diatoms.

During the nursery phase, the plates have to be regularly (every 1 to 2 weeks) inspected and rotated in the vertical and horizontal planes to ensure even growth of diatoms. This method therefore requires high inputs of labour, equipment and space and consequently has high operating costs. Indeed, to meet such requirements, commercial farms commonly use 10 000 to 25 000 standard plates and nursery site areas of about 1000 m². Rearing post-larvae for long periods on diatom plates also brings uncertainty, because the growth of diatoms varies greatly with the temperature, light intensity and levels of nutrients. Most, or all, of these parameters are under only limited control by hatchery staff. Light intensity can be raised to promote greater diatom growth, as in the case of excessive grazing pressure, or lowered to inhibit excessive growth of diatoms and filamentous algae. Such adjustments are made by applying a range of alternative grades (30% to 95%) of shade-cloth that may take the form of covers or canopies.

As discussed above, blacklip abalone post-larvae can be safely harvested from the plates by the time they reach an average shell length of 1.5 mm, with most above 1 mm. On the central coast of NSW this can occur in as little as 30 days through summer but up to 70 days through winter. High-density rearing of post-larvae on the plates makes it essential that their progress is constantly monitored. As also discussed previously, there is a real risk of the post-larvae overgrazing the diatoms, leading to starvation and subsequently to losses or reduced ability to survive handling and weaning.

To avoid these hazards, a small but representative sample of plates should be inspected every few days, beginning a month after seeding (see Chapter 6 and Figure 40). If the film of diatoms covering the plates is grazed below about 10% of the plate surface, the post-larvae should be immediately harvested and transferred to raceways. This should be done even if they have not reached the desired average size of 1.5 mm, because even those with a mean size as small as 1 mm can be successfully weaned provided that they are in good general health. An option for dealing with overgrazing of individual plates before harvesting is to use back-up sets of conditioned diatom plates, batches of which should be initiated at fortnightly intervals as a supplementary food source. These can be clamped to overgrazed plates by using plastic clothes pegs.

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Pre-harvest count

From 1 to 3 days before transfer of the post-larvae to raceways, the total number of post-larvae can be estimated by counting those on at least one plate selected at random from each basket of plates. This can be done by two alternative methods. The first is direct counting of undisturbed post-larvae in situ (i.e. still attached to the plates). As already discussed, direct counts should always be conducted on plates fully submerged in seawater in shallow (75 to 100 mm) clear plastic trays (see Figure 40) maintained at or below 23°C. Counting can be facilitated by incorporating a prominent 2 x 2 cm black-line grid on the floor of the trays. Additionally, transparent or translucent plastic trays can be lit from below to silhouette both the counting grids and the post-larvae.

The second option is to pre-harvest and estimate the average numbers of post-larvae per plate (and hence the total numbers of post-larvae available for harvesting) from the representative sub-sample of plates. These sub-sampled plates should be pooled into a single basket before being subjected to the same step-by-step procedures described below for the main harvest. The combined settled volume of these sub-sampled post-larvae can then be determined by transferring them into a seawater-filled measuring cylinder. This volume is finally multiplied by the average count of post-larvae in a minimum of three 1.0-mL settled sub-samples and the result divided by the number of sub-sampled plates to estimate the average count per plate and hence the total number of post-larvae available for harvest. To avoid double-handling, the sub-sampled post-larvae should be immediately transferred to a raceway ahead of the main harvest.

Harvesting and transfer of post-larvae to raceways

Individual baskets of plates are placed in a dedicated tub where the post-larvae are anaesthetised. The tub should be large enough to easily accommodate, submerge and manipulate a full basket of plates in a vigorously aerated anaesthetising solution of benzocaine at 50 mg/L seawater (see step-by-step summary for preparation details). The post-larvae are detached from the plates after 5 to 10 min of anaesthesia by using a soft-bristle paintbrush aided by continuous flushing with seawater. Post-larvae can be detached from the plates into the tub and then drain-harvested onto a submersed screen for transfer to first stage raceways. Alternatively, the post-larvae can be detached plate by plate directly into recipient raceways. The latter is preferable in the sense that double-handling is avoided and the duration of the anaesthesia minimized.

To allow good quality films of diatoms to develop, seawater flow through the nursery raceways should be started about 2 weeks before stocking with post-larvae. To expedite biofilm development, raceways can be inoculated with diatoms scraped from conditioned nursery plates. Daytime hides such as concrete pavers should also be installed in the raceways 1 to 2 weeks before stocking with post-larvae. Although the hides are not used as shelters by the post-larvae for the first few weeks, their initial presence avoids the risk of post-larvae being crushed during their installation. Healthy post-larvae are able to recommence feeding on diatoms the first night after transfer, and the presence of diatom films reduces or
eliminates growth checks otherwise commonly associated with transfer to raceways and weaning onto fine-grained (powdered) formulated diets.

The amount of supplementary formulated food provided over the first week is a matter of trial and error. If there is a lot of uneaten food left on the first morning after transfer, reduce the ration by about 25%, then repeat the exercise until food supply and consumption are matched. If, however, there is no food left over, increase the ration by 25% each night until some residual food is apparent. Avoid over-feeding, which results in both wastage and reduced water quality and hence impaired growth performance and health.
**STEP BY STEP SUMMARY AND CHECK-LIST FOR DIATOM PLATE NURSERY REARING AND EARLY WEANING**

1. Once plates have been seeded with post-larvae, inspect regularly to monitor the percentage cover of the diatom bio-film and the growth of the post-larvae. Inspections should be made fortnightly for the first 4 weeks, then on alternate days or daily once plates with a diatom bio-film coverage below about 10% are first detected and/or as post-larvae approach the minimum average weaning size of 1.5 mm.

2. If the biofilm on a majority of plates drops below 10% coverage, wean the post-larvae immediately, whether or not an average size of 1.5 mm has been reached.

3. When the post-larvae reach an average shell-length of 1.5 mm (a process that will take from as little as 30 days in summer but as long as 70 days in winter), wean them regardless of the remaining biofilm coverage.

4. Set up the nursery raceways with flow-through seawater up 2 weeks before the anticipated first date of harvest and weaning to promote development of diatom films on the floor and walls of the raceways.

5. Estimate the total numbers of post-larvae 1 to 3 days before transferring them to raceways, by counting those on a plate randomly selected from each basket to be harvested. There are two alternative ways of doing this:
   - Count the numbers of post-larvae on the plates in situ. This should always be done on plates fully submerged in seawater in shallow (75 to 100 mm) clear or translucent plastic trays maintained at or below 23 ºC. This method can be facilitated by incorporating a prominent 2 x 2-cm black-line grid on the floor of the trays and/or by under-lighting them to silhouette both the grids and the post-larvae; or
   - Harvest and count post-larvae from a representative sub-sample of plates by using the techniques described below.

6. On the day of harvesting and transfer of post-larvae to the raceways, fill a tub large enough to accommodate a full basket of plates with 10-µm-filtered seawater and add anaesthetic stock solution as follows:
   - Make up a stock solution of benzocaine by adding 50 g of ethyl-p-amino-benzoate in 500 mL of 95% ethyl alcohol (0.4 M solution).
   - Add 0.5 mL of this solution per litre of seawater to the tub1.

7. Leave the plates in this solution for about 5 min, after which time you will able to easily dislodge the post-larvae with seawater by rinsing aided by gentle use of a soft-bristle brush. Post-larvae from plates can either be:
   - dislodged into the tub for collective harvesting into a flooded 10-µm-mesh screen and then transferred into a nursery raceway; or preferably
   - dislodged as described, but directly off the plates into prepared raceways.

8. In either case, raceways must be supplied with flowing seawater and vigorous aeration to speed the recovery of the post-larvae from anaesthesia2.

---

1 Do not attempt to handle or to anaesthetize post-larvae or small juveniles when heat-stressed at temperatures above 23°C.

2 Total time of exposure of post-larvae to anaesthesia should be kept as brief as possible and less than 15 min.
Chapter 8  Intensive on Rearing in Shallow Raceways
Chapter 8 Intensive On-Rearing in Shallow Raceways

General

Shallow raceway nursery rearing of post-larvae from 1 to 2 mm to juveniles in the range 7 to 15 mm and suitable for on-farming or fisheries reseeding operations is best completed in two stages. First-stage raceways of the type illustrated in Figure 41 may be as small as 1 m² (2 × 0.5 m) and no more than 15 cm deep.

Cascade systems can be used to improve floor space efficiency, but the minimum seawater exchange needed ensure maintenance of optimum seawater quality (≥ 1 L seawater/kg biomass/min) must be adjusted accordingly. Likewise, care needs to be taken to ensure that the longer seawater flow-paths do not result in the abalone being exposed to temperatures outside their tolerable range of 10 to 23°C.

Stocking rates

First-stage raceways can be stocked with 1 to 2 mm ex-plate post-larvae at initial densities of up to 50 000/m². However, even with good husbandry, numbers of juveniles are likely to progressively fall by an average of 30% to 50% over the subsequent 2 to 3 months. By this time they should have grown to an average size of 4 to 5 mm, ready to shift to second-stage raceways. A second important criterion for adjusting stocking density is that the collective shell area of stock should not occupy more than about 60% of the floor area of the raceways. The exponential shell growth of post-larvae reared on diatom plates immediately shifts to a constant (linear) rate (Figure 42) following transfer of the post-larvae to shallow raceways and weaning onto formulated diets.

At 4 to 5 mm, juveniles are stocked at initial densities up to 20 000/m² into the larger second-stage raceways of the type illustrated (Figure 43). Here they soon adopt a strictly nocturnal pattern of activity (Figure 44), remaining cryptic and inactive by day. As in first stage raceways, densities commonly fall by 30% to 50% as the juveniles grow through to a ‘button’ size range of 7 to 15 mm over the ensuing 2 to 3 months. At this size they are suitable either for fisheries seeding operations or for on-farming to a final market size, most commonly in the range 80 to 90 mm.

In raceways the average monthly shell growth of seed from the time of weaning to a final mean size of 7 to 15 mm can vary from 45 to 80 µm/day, depending mainly on the seasonal temperature. Farming methods include offshore cages located either on the seabed or suspended from mid-water long-lines, or (more commonly) land-based raceway tank systems. These shallow raceways or ponds are provided with either hides or riffle ridges incorporated into the floor. Various types of newer-generation grow-out tanks commonly used throughout southern Australia are illustrated (Figure 45).
Stocking rates up to harvest size (80 to 90 mm and 80 to 100 g) continue to be limited to 60% of the available tank floor area, which amounts to 200 abalone/m² or a biomass of about 12 kg/m².

**Figure 41** Top left: First stage nursery raceway under aeration

Below central: Same raceway with aeration turned off to reveal air diffuser tube and day time hides.

Bottom right: Close-up of 1 to 3 mm recently weaned post-larvae and 20 mm aeration diffuser tube
**Figure 42** Constant (linear) shell growth of weaned raceway reared juveniles. (From Heasman et al., 2004) Note the contrast with earlier exponential growth pattern of diatom plate reared post-larvae (see Figure 37).
Figure 43 Larger (3 to 4 m²) second-stage raceways operated at a minimum recommended depth of 5 cm. Note: These are used to on-rear the juveniles from 4 to 5 mm up to 7 to 15 mm at densities of 20 000/m² (initial) to 10 000/m² (final).
**Figure 44** Top: Daytime photograph of secondary nursery system with aeration off to show juveniles inactive and sheltering under artificial hides

Bottom: Same system photographed 1 h after nightfall, revealing active nocturnal foraging
**Figure 45** Various types of land based abalone grow-out farm systems used in Australia

Left and above: Greenlip abalone in "Slab" tanks

Middle left: Large outdoor circular tanks

Bottom left: Maze tanks

Below & bottom right: Intermediate pipe tanks being used to rear greenlip abalone
Water quality and temperature

Juvenile blacklip abalone, especially in the larger 1+ to 3+ year old age-classes, should always be maintained at temperatures below 24 °C and in continuously flowing, oxygen-saturated seawater. As already stated, a useful guide to the minimum seawater exchange rates required to maintain optimum growth and health is that they should match or exceed the combined body weight of abalone every minute (1 L of seawater for every kg of abalone). Assessment of density for blacklip abalone should be made during the period of peak activity 1 to 4 h after nightfall. In this regard it is important to know that the older and larger the juveniles, the less tolerant they are to common forms of stress, including reduced seawater quality parameters such as pH, salinity, dissolved oxygen and temperature extremes (especially high temperatures). It is also important to appreciate that any form of stress will lower resistance to other stress factors, including infectious diseases, starvation and nutritional deficiencies.

Feeding

Dry formulated feeds should be ordered in regularly (monthly) to avoid extended storage times and associated deterioration. Even then, they are best stored deep-frozen (-18°C) to preserve their nutritional integrity—especially that of the volatile/labile components, including some essential vitamins and fatty acids. Frozen storage also minimises microbial spoilage, including the production of fungal-generated toxins (aflatoxins). Food consumption and feed application rates should be monitored and matched on a routine basis to achieve satiation feeding. Satiation feeding maximises growth and water quality (and hence health) while minimising the cost of wasted food. This is a simple process of making small (5%) upward or downward adjustments in the weight of formulated food rations fed to each raceway. Thus at the beginning of each week the ration fed to particular raceways will be raised if very little or no residual (uneaten) food has been logged over the previous seven mornings. Likewise, raceway rations should be left unchanged if near-satiation feeding has been achieved, or reduced by 5% if low to moderate levels of uneaten food have been consistently observed over the previous week.
Appendices

Appendix 1 Schedule of hatchery operations

**Broodstock**

- Days -10 to 14: Induce spawning
- Day 0: Select and precondition at ~18°C

**Larvae**

- Day 1: Setup incubation larval rearing tanks
- Day 1: Fertilisation and incubation
- Day 6 8: Transfer larvae to clean isolation tank
- Day 6 8: Settlement of larvae on plates
- Day 20 (mid summer): Harvest post-larvae at 1-2 mm transfer to raceway
- Day 30 (mid summer): Attainment of button size (10 mm) for seeding or on-growing in commercial raceways

**Post-larvae on plates**

- Day 0: Setup diatom plates in settlement tanks
- Day 1: Culture of benthic diatoms on plates

**Juveniles**

- Day 6 8: Larvae rearing
- Day 6 8: Set-up diatom plates in settlement tanks
- Day 20 (mid summer): Setup 1st stage raceway and hides
- Day 30 (mid summer): Harvest post-larvae at 1-2 mm transfer to raceway
- 5 months: Stock into raceway for weaning
- 5 months: Transfer into stage 2 raceways at 5 mm
- 6 8 months: Attainment of button size (10 mm) for seeding or on-growing in commercial raceways
- 6 8 months: Market size 80 mm 25 mm/year
- 3 - 5 years: Market size 80 mm 25 mm/year
Appendix 2 Design and operating specifications of a relatively simple and inexpensive abalone broodstock conditioning system

As insurance against system failure at least two independent units of the type illustrated in this Appendix should be used for year-round temperature-controlled reproductive conditioning of abalone. Each unit needs to be separately housed within an insulated room. The units illustrated in Figure A1 are hybrid systems incorporating both continuous low-level exchange (1 to 3 total system volumes/day) with a much higher rate of seawater recirculation (1 to 1.5 total system volumes/h). Recirculation needs to incorporate continuous or semi-continuous removal of solids by physical filtration, removal of dissolved organics—especially proteinaceous and oily (lipid) materials—by using foam (fractionation), and removal/detoxification of excreted ammonia by using biological and/or denitrification filtration.

**Holding tank:**
FRP (fibreglass)-constructed rectangular tank, 1500 L nominal capacity.
Dimensions: 2700 × 1000 base × 500 mm tapered wall.
BU1: Open tank with hides of fibre cement sheets.
BU2: Buckets. Abalone held in fifteen to eighteen 30 L drums with five to eight abalone per drum. (The drum system follows that used by Moss [pers. comm.], Mahunga Bay, New Zealand for *H. australis*.)

**Header tank:** 1000-L nominal capacity.

**Physical filter:** Dacron (Aquahort) or similar.

**Biofilter:** Layer of Dacron over 10 L of coral debris over 100 L of bioballs.

**Pumps:** Davey Models XF171 and XF221.

**Foam fractionator (Figure A2):**
Diameter: Downtube 150 mm PVC class-18 pressure pipe.
Height: 1890 mm.
Foam effluent: Transparent cone with 19 mm poly waste line.
Venturi: Mazzel injector model 484 (20 mm).
Flow-through fractionator, 1 L/s.

**Temperature control:**
Air-conditioner, Email GM 155H dual cycle, 1.5 kW or similar
Chillers. Carrier 30ZQ024 on header tank. Ozsea 2 hp on recirculation.
Temperature maintained at 15.0 or 16.0°C ± 1.0°C, monitored by temperature dataloggers (Hastings).
The header tank also supplies temperature-controlled water to the spawning/larvae-rearing unit.

**Aeration:**
Gentle aeration through 3-mm CVT (clear vinyl tube) to each bucket.
Flow rates:
New water throughput, 2 to 4 L/min. (2 to 4 total system volume exchanges/day.)
Recirculation through chiller, foam fractionator and biofilter is 0.5 to 1 L/s.
(using Davey XF 171 pump), representing 1 to 2 total volume turnovers/h.

Stocking rate:
Each system is designed to accommodate a total of about 100 adult abalone (total
biomass 30 kg)

Feed rates:
0.75% to 1% of body weight/day of formulated feed (Adam and Amos).

Ammonia levels:
Routinely less than 0.2 mg/L total ammonia nitrogen, pH range 7.9 to 8.2

Light levels:
Dim light. Constant photoperiod of 10 h light, 14 h dark.

Maintenance:
Daily. Feeding, manual temperature check and visual check of system and
abalone.
Each 2 to 3 days. Full drain-down, water change and cleaning of holding tank to
remove detritus and surface biofilms. Cartridge filter cleaned.
Foam fractionator and Dacron solids-removal filter cleaned.
Each month. Wash half of biofilter. Test for ammonium (Merck, Aquamerck
ammonium test) and pH.

Alarm system:
Notification of system failures via phone link (Telstra SMS).
Temperature alarm if outside ± 2.0°C.
Header tank low-water level alarm.


A abalone broodstock conditioning unit
NSW Fisheries / FRDC, Abalone enhancement project.

Nick Savva, June 2000.

**Figure A1 Example of the broodstock conditioning unit - schematic design**
Not to scale. All dimensions in mm. Motive flow, Davey XF 171. Flow rate, 1

Figure A2 Detailed design of foam fractionator
Appendix 3(a) Data to be recorded for broodstock, their collection and conditioning history and spawning performance

Collection:

- Where collected
- Depth
- Time/date/ambient sea temperature
- Allocated Tag and Code numbers
  
  e.g. Tag number a-012 = Code number BH - 230203-(2F) where: BH is letter code of collection site (Boat Harbour); 230203 is date (ddmmyy) of collection and 2F is designated number and sex of individual
- Conditioning tank number allocated to after quarantine

Conditioning protocol (per tank):

- Temperature (maximum/minimum monitored daily)
- O₂ (monitored weekly)
- N as ammonia and nitrite (monitored weekly)
- Net exchange and recirculation of flow rates (monitored weekly)
- Date of cleaning
- Mortality (monitored daily)
- Feeding rate recorded daily (normally 1% body weight per day)

Animal condition and performance:

- Weight and length when initially transferred to conditioning unit and on all subsequent spawning induction occasions
- Gonad index when initially transferred to conditioning unit and on all subsequent attempted spawning induction occasions
- Spawning response including egg numbers and quality in the case of females
Appendix 3(b) Data to be recorded for all attempted inductions of spawning

Batch code/identification:
- date
- person responsible for spawning

Spawning:
- tank setup - including how many abalone per tank
- time UV was turned on
- time males spawned
- time females spawned
- temperature and any alterations to temperature
- comments on egg quality (colour, form/shape, clumped/uniform/layer)

Fertilisation:
- number of females and males spawned fully
- approximate number of eggs per female
- sperm concentration and motility
- time fertilised
- time washed

Hatchout phase:
- system setup
- temperature of water
- time to hatchout
- percent hatched

Larval phase:
- system setup
- water temperature
- water flow/exchange rate
- larval density each day
- cleaning routine
- time to shell development
- time to settlement

Juvenile/adult phase:
- growth rate
- survival rate
## Appendix 3(c) Example of data collection for spawning operations

### Spawning Attempt 5/4/01

Broodstock collected from Broughton Is. 15/4/99

<table>
<thead>
<tr>
<th>Rivet Tag</th>
<th>Glued Tag A</th>
<th>Visual gonad index</th>
<th>Tub</th>
<th>Eggs?</th>
<th>Comments</th>
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<tr>
<td>22 FEMALES</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>1001</td>
<td>502</td>
<td>3</td>
<td>7</td>
<td>0.91</td>
<td></td>
</tr>
<tr>
<td>1002</td>
<td>504</td>
<td>2</td>
<td>9</td>
<td>1.0</td>
<td>NB. Except for 1001, nos. of eggs are averaged across multiple spawners.</td>
</tr>
<tr>
<td>1004</td>
<td>506</td>
<td>1.5</td>
<td>5</td>
<td></td>
<td></td>
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<td>507</td>
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<td>8</td>
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<tr>
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<td>512</td>
<td>1</td>
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<td>1.0</td>
<td></td>
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<tr>
<td>1010</td>
<td>513</td>
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<td>6</td>
<td>1.0?</td>
<td>Unsure which or both in tub 6 spawned.</td>
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<td>2</td>
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<tr>
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<tr>
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<td>521</td>
<td>1</td>
<td>6</td>
<td>1.0?</td>
<td>Unsure which or both in tub 6 spawned.</td>
</tr>
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</tr>
<tr>
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<td>522</td>
<td>1</td>
<td>11</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Appendix 3(c) cont’d. Example of Companion Diary Notes of Spawning Operation

0850 Abs transferred from BU 2 to spawning containers in hatchery.

0905 Tris added at 6.6ml/L

0915 Peroxide added at 3ml/L.

1015 +1hr. All quiet

1115 +2hrs. All quiet, plenty of bubbles.

1215 +3hrs. FWC. Flow on. Water temp.

1310 Flow off. Air on. 17.0°C.

1320 Sperm released in tubs 11 and 12 (only light so far.) very small # of eggs in Tubs 7 and 2. Eggs from tub 7 are of poor quality and are possibly atretic. Eggs from tub 2 appear fine for fertilisation. Will wait for hopefully more eggs before acting.

1345 Rapid succession of spawning females over past 35 mins
Coll’n 1. Tub 1, #1014 and 1012. 1.05 million.
Coll’n 2. Tub 2, #1015 (lost tag). 1.4 million.
Coll’n 3. Pooled tubs. 1,2,6,8. 1.38 million.
Coll’n 4. Tub, 7. #1001. 0.91 million, questionable quality.
Coll’n 5. Tubs, 1,2,6,8,9. 2.64 million.
Coll’n 6 Tubs, 1,2,6,8,9. 2.84 million.
All eggs except Tub 7, to LT2. 9.3 million.
Eggs from tub 7, #1001 to LT100:2. 0.91 million.

TOTAL EGGS SPAWNED 10.2 million.

Abs returned to BU 1.

Batch designated as 02-0401.
Appendix 4 Prevention and treatment of mud-worm infestation (adapted from M. Lleonart, 2001).

Mud-worm infestation of the shells of conditioning broodstock can and does seriously reduce productive performance of and indeed can cause the death of these valuable stock if left untreated. The threat to broodstock abalone is particularly high because of the ability of mud-worms to breed asexually, and hence to rapidly and heavily re-infect both their hosts and other previously un-infested broodstock in close proximity within controlled temperature conditioning systems. The first and most important way of preventing and controlling mud-worm infestation is therefore to cull heavily infested stock at the time of their collection from the wild. Effective treatment of new lightly infected stock can be accomplished by use of either shell drying shell salting or shell waxing. For shell drying, the abalone need to be removed from water and their shells allowed to dry out in accordance with the following conditions.

- 2-4 hours out of water
- Minimum temperature of 15°C & recommended maximum of 21-22 °C
- Air humidity of less than 64%

If shells of stock are not apparently drying within ~30 minutes of air exposure the conditions are probably unsuitable. Merely removing abalone from water, even for longer periods of time and/or higher temperatures, will not kill mudworms if shell surfaces do not dry properly. Air humidity can be measured simply and cheaply with a wet & dry bulb hygrometer and accompanying conversion tables (scientific equipment supply companies ~$30). Typically, days with minimal cloud cover will be sufficiently dry for treatment in southern Australia. Abalone should be dried in shade, not direct sunlight. Depending on-rearing vessel design and degree of fouling of these, it may be possible to dry stock within rearing vessels. If not, abalone will have to be removed and placed on a suitable substrate such as sheet plastic. It may be appropriate to combine treatment activities with grading, thinning or other activities requiring handling.

Mud-worm kill rates of up to 90% may be achieved by shell drying provided the worms treated are small (less than 6 months old). Reduction in worm numbers may be possible after this time but the result will vary with period of infestation, extent and depth of blisters, size of abalone and degree of shell fouling. The treatment drying conditions referred to are generally safe for blacklip abalone. However drying can significantly suppress growth and probably gonad development as well. Moreover as exposure of conditioning broodstock to temperatures of 19°C or higher will either promote unplanned spawning of ripe eggs or atresia (regression) of developing eggs, drying should not be applied to conditioning broodstock except for individual stock in the period immediately following successful induction of spawning.

The alternative treatment of shell waxing involves the manual application of a thick (2 to 5mm) layer of pliable non-toxic surfboard wax. Shell waxing, which is best preceded by blot drying with paper toweling and swabbing with 90% ethanol, is far more labour intensive and therefore expensive than shell drying. However it has a number of major advantages over shell drying namely:
- it is capable of suffocating all mud-worms and their young if applied diligently and left for a minimum of 2 days
- broodstock suffer much lower stress in the absence of desiccation and much shorter periods of air exposure (less than 30 minutes)
- it can also be safely applied to conditioning stock at air temperatures of 10-15°C and the broodstock immediately returned to the conditioning system thereby eliminating risk inadvertent induction of spawning of ripe eggs or regression of developing eggs

Shell salting, the simplest and cheapest of the three alternative treatments. It is also the quickest entailing a one hour application of copious amounts of damp sticky rock salt to the drained damp shell of broodstock while strongly attached to a smooth flat horizontal surfaces such as those provided by clean diatom nursery plates. Salt coating has the same advantages of waxing in application to conditioning broodstock in being able to be applied within the safe temperature range of 10-15°C and without the risk of undue physiological stress associated with shell drying. However the killing efficiency of salting is probably lower than both drying or waxing as mudworm occurring on the curved peripheral margin of the shell and adjacent to the respiratory pores may escape exposure to lethal levels of osmotic stress. This is because of the poorly adherence characteristics of the salt and the need to prevent salt from coming in direct contact with soft delicate tissues of the foot and mantle on the underside of the abalone and with the gills and neighboring tissues in the mantle cavity via the respiratory pores.
Appendix 5 Determination of sperm concentrations using haematocytometer

Haemocytometer slides have 2 rafters allowing for 2 sub-samples to be examined. The sampling and counting procedures are as follows:

1. dilute sperm sample if needed (use formalin 4% to fixate moving sperm) clean slide and cover-glass with Kleenex-paper
2. press cover glass onto the slide until the Newton diffraction rings appear
3. fill both slides of the counting chamber under the cover-glass with a single smooth flow of suspension using a Pasteur pipette (avoid air bubbles)
4. count sperm in, 20 small squares under a microscope (objective 40 x). Count cells which touch the upper and left border but not those which touch the lower and right borders (see schematic diagram)
5. the sub-sample on the other side of the chamber is counted in the same way

- calculation of density per ml.

numbers of cells per ml = \( \frac{n_1 + n_2}{2 \times 20} \times 250 \times 10^3 \times d = \frac{(n_1 + n_2)}{160} \times 10^6 \times d \)

where:

- \( n_1 \) = number of sperm counted in upper rafter
- \( n_2 \) = number of cells sperm counted in lower rafter
- \( d \) = dilution factor

NB For greater accuracy make 3 duplicate counts (3 separate dilutions each counted in two rafters).

Figure A3 Counting directions (follow arrow), count the sperm cells in the square and those which touch the top and left borders (•). Do not count the ones touching the right and bottom borders (○).
### Appendix 6 Chlorine disinfection and neutralization protocols

<table>
<thead>
<tr>
<th>Disinfection Task</th>
<th>Procedures</th>
<th>Amount of stock chlorine to use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nets/buckets/boots:</td>
<td>0.25ml stock chlorine solution(^1) per litre of freshwater for ≥ 24hours</td>
<td>25ml/1000L</td>
</tr>
<tr>
<td>Transport containers and</td>
<td>0.8ml stock chlorine solution per litre of freshwater for ≥ 24hours</td>
<td>80ml/1000L</td>
</tr>
<tr>
<td>Rearing vessels</td>
<td>1.25ml stock chlorine solution per litre freshwater for ≥ 24hours</td>
<td>125ml/1000L</td>
</tr>
<tr>
<td>Nursery plates and seawater prior inoculation with Ulvella spores</td>
<td>0.8ml stock chlorine solution per litre of 1(^{\circ})m filtered seawater for ≥ 24hours followed by the addition of 0.02 ml/L of stock thio(^2) solution and heavy aeration (air stripping) and finally test for traces of residual chlorine(^3)</td>
<td>80ml/1000L</td>
</tr>
</tbody>
</table>

---

1. **Stock hypochlorite solution** contains 10-12% (100 to 120 g/L) active chlorine

2. **Stock thio** (chlorine neutralising) solution 2 molar (316g/L) solution of sodium thiosulphite in freshwater

3. **Residual chlorine test** - add 1 DPD No 4 chlorine test tablet per 10-20ml sub-sample of thio neutralised and air stripped seawater
Glossary

adventitious, naturally occurring; used here to describe types of wild microalgae and macroalgae that opportunistically colonise bare substrates

banjo filter, a filter used in flow-through larval culture to stop loss of larvae via the outflow

barrens, bare (or, more commonly, crustose coralline algae-coated) rock areas commonly created by dense aggregations of grazing animals, especially sea urchins and (in NSW) the black (or purple) urchin (Centrostephanus rodgersii)

benthic, bottom-living

benzocaine, a chemical (ethyl-p-amino-benzoate) used to anaesthetise abalone before handling or examination for health and breeding status

biofilms, thin films of micro-organisms (initially bacteria and fungi) that colonise bare substrates

competence, the ability and urge of larvae to actively seek out and settle on suitable habitats and to be able to metamorphose (transform) into reef-dwelling post-larvae

copepods, very small planktonic and surface-living crustaceans

crustose coralline algae, colonial algae that form a tough, limestone depositing skin on rocky surfaces that constitute the principal settlement substrate and early juvenile habitat of blacklip abalone; commonly known as ‘pink rock’

diatoms, a special group of microalgae with an outer cell-coat of silica

demersal, found near or on the sea bed

drift, broken-off and drifting pieces of seaweed, including common red seaweeds such as nori (Porphyra spp.) and Gracilaria and green seaweeds such as sea lettuce (Ulva spp.)

embryos, fertilised eggs undergoing successive cell divisions and cell differentiation

epi-biota, mixture of plants and animals attached to surfaces

exponential growth, implies that the time it takes to double in size from any starting point remains constant

gonads, the reproductive organs of abalone and other animals, being the egg-producing ovaries in females and sperm-producing testes in males

haemocytometer, a simple cell counter consisting of a thick glass slide with a very fine set of gridlines etched on the surface, over which a special heavy-duty glass cover-slip is laid

IDS (integument/muscle detachment syndrome), the most important affliction of abalone veliger larvae, whereby there is a breakdown of attachment of the twin retractor muscles to the inside (integument) of the shell. This loss of one or both attachments is commonly considered a symptom of bacterial infection (usually by Vibrio) arising from either poor quality of seawater or inadequate hygiene.

macroalgae, large filamentous or thallus-forming seaweeds

metamorphose /metamorphosis, transformation of planktonic veliger larvae into reef-dwelling benthic post-larvae

microalgae, microscopic single or chain-forming algal cells

moribund, sick, weakened and dying as consequence of injury, disease, malnutrition or age

photopositive, attracted to light
**plankton(ic)**, small aquatic animals (zooplankton) and plants (phytoplankton) that live permanently in the water column

**polyspermy**, deleterious effects of very high sperm to egg ratio during fertilisation, when more than one sperm fertilises an egg

**post-larvae**, name given to the juvenile form of animal that in abalone spans the period after metamorphosis up to 3 mm shell length

**radula**, rasping tongue of the abalone

**Sedgewick rafter slide**, a special type of microscope slide with a shallow 1-mL reservoir, the floor of which is divided into 1 × 1-mm grids where numbers of small particles such as larvae or microalgae sampled from a dispersed suspension can be systematically counted and measured grid by grid at 5× or 10× with the aid of a stereomicroscope fitted with a mechanical stage.

**trochophore**, first-stage swimming but non-feeding planktonic larval stage of abalone that develops about 16 h after fertilisation and lacks a shell

**veliger**, second and final fully-shelled larval stage of abalone and many other types of mollusc