Abalone, *Haliotis mariae* (Wood, 1828), Hatchery and Seed Production Trials in Oman

Khalfan M. Al-Rashdi*1 and Tsueno Iwao2

1Ministry of Fisheries Wealth, Aquaculture Center, P.O. Box 427, PC 100, Muscat, Sultanate of Oman
2Japan International Cooperation Agency, Fisheries Department, Japan

Introduction

The abalone *Haliotis mariae* (locally called As’sufailah) is endemic to the subtropical Arabian Sea coast of southern Oman, where it inhabits inter- and subtidal rocky substrates down to 20 m depth (Bosch and Bosch, 1982; Johnson et al., 1992). In this habitat, abalone shelter among rocks and in crevices during daytime and forage nocturnally. Larger individuals...
occur in deeper areas than smaller animals (Anon, 1984).

The life cycle of *H. mariae* is related to the southwest monsoon driven upwelling that occurs from June to August (Johnson *et al.*, 1992), which decreases seawater temperature and increases nutrient concentrations (Barratt *et al.*, 1986). During the pre-monsoon (March-May) and monsoon period, the brown and red algae on which *H. mariae* predominantly feed are scarce or absent (Barratt *et al.*, 1984; Jupp, 2002). In the post-monsoon months of September to February, abalone move into shallower waters to reproduce and feed (Al-Hafidh, 2006). Females appear to spawn once a year with a peak spawning during December and January, and a gradual decrease during February and March as algae become scarce (Al-Hafidh, 2006). Sea urchins appear to be a competitor for food (*Tripneutes gratilla*) and living space (*Diadema* spp.), and natural predators on abalone include scalloped spiny lobster (*Panulirus homarus*), seastars (*Asterias* spp.), octopus (*Octopus aegina*), cuttlefish (*Sepia* spp.) and finfishes like morays (*Gymnothorax* spp.).

A fishery for *H. mariae* is concentrated around Sadah, with catches declining towards Sharbithat in the East and Mirbat to the West (Fig. 1). The annual production of abalone from a two-month fishing season (Table 1) was estimated at 50 t in 2006, valued at more than US$ 8 million (MAF, 2007).

**Figure 1.** Map of Oman with important locations for abalone fisheries at Dhofar Region.
Economically, it has the highest yield per kg of all Omani fisheries products. Intense fishing by locals using free diving methods have led to sharp decline in catches, and abalone stock is presently regarded as overfished (Al-Hafidh, 2006).

The aquaculture of abalone on a commercial scale has developed rapidly in several countries including the USA, Mexico, South Africa, Australia, New Zealand, Japan, China, Taiwan, Ireland and Iceland (Hahn, 1989; Gordon and Cook, 2001). Specific culture techniques need to be developed or adapted for each abalone species. The geography and subtropical climate of Oman with warm clean coastal waters lends itself to aquaculture, and *H. mariae* furthermore has a unique yellow foot and excellent meat texture. Given the high value of abalone products on international markets and the decline of wild stocks, the aquaculture of *H. mariae* in Oman was investigated in a series of projects since 1994, the resources of which are available in several internal- and unpublished reports (Ogawa, 1994, 1997; Iwao, 2000; Al-Rashdi, 2001; Endo, 2005), but have not appeared in the peer-reviewed literature. This paper reviews the progress of the abalone aquaculture trials in Oman, with emphasis on the development of the culture facilities, hatchery and handling techniques, and constrains to the culture of *H. mariae*.

### Materials and Methods

#### Study site and culture system

Mirbat bay was selected as the site for the Mirbat Abalone Seed Production Station (MASPS) based on several criteria. The bay is characterized as semi-closed, opens to the Arabian Sea in an eastward direction, and is only moderately affected by wind and adverse sea conditions during the monsoon season. It has clean and unpolluted coastal waters. Several seaweed species are available in the vicinity as abalone food, and Mirbat town is close to markets and other existing infrastructure.

A Land-Based (L-B) culture system with seawater pumped and filtered on a 24 h basis from 270 m offshore at a maximum capacity of about 120 t.h\(^{-1}\) (only 20 t.h\(^{-1}\) utilized) was installed. Water was filtered to 40 µm to prevent invasion of fouling organisms.

#### Broodstock collection and conditioning

*H. mariae* broodstock of 60–120 mm shell length (SL) were collected from the wild in May (pre-monsoon) and September (post-monsoon) by SCUBA diving during the daytime at depths of 5–20 m. Care was taken not to damage the foot while dislodging them from the substratum. Animals were transported to the hatchery within 4h using a cool box containing a wet sponge to absorb excretory products and ice to reduce temperature and metabolic rates. The total SL and weight of each abalone were measured using a vernier caliper (±0.1 mm) and electronic balance (±0.1 g).

Artificial conditioning trials were carried out in 1000 l capacity rectangular fiberglass tanks over 4-month and 6-month periods, respectively for abalone collected before and after the monsoon period. Densities in tanks were limited to 25-60 individuals per tank. Inverted V-shaped Glass Reinforced Plastic (GRP) structures were placed at the bottom of the tanks as shelters. The water was filtered through three layers of sand and then recycled at an ambient temperature of 26–28°C. The water was re-circulated and re-filtered to allow microorganisms to eliminate ammonia, and aeration was provided. An artificial photoperiod (12L:12D) was maintained with a light intensity of approximately 150 lux at the bottom of the tank during the daylight hours. Broodstock were fed daily with green alga, *Ulva fasciata*, at a rate of 10% of the total body weight. A gonad maturity stage was defined by visual assessment as follows: stage (0) - no gonadal development visible; stage (1) - immature, with the level of the shell edge higher than the gonad coverage; stage (2) - mature, with the level of the shell edge equal to the level of the gonad coverage; and stage (3) - fully mature, with the shell edge lower than the gonad coverage (Fig. 2). The collected broodstock had mostly immature gonads (stage 1) at the onset of conditioning. A maturity index was calculated as the average of individual gonad maturity stages in each month.
**Spawning induction and fertilization**

Spawning was induced following the method developed by Kikuchi and Uki (1975) for *H. discus hannai* in Japan. Broodstock were first exposed to the air for an hour and then immersed in filtered irradiated sea water with ultraviolet (UV) light at slightly elevated temperature (+3°C). The UV irradiation dose level used was 2400 m.W.h.l⁻¹ at flow rates of 200 ml.min⁻¹, distributed equally to ten 20 l aquaria. The broodstock were then left undisturbed and checked for spawning every hour for the first 3 h and then for every 30 min for a further 5 h.

After spawning, eggs were siphoned out of the aquaria and filtered through a 300 µ mesh to remove unwanted particles, and the number of eggs was counted under binocular microscope by taking 5 ml sub-samples from egg water aquaria. Fertilization was carried out according to Hahn’s (1989) technique. Eggs were maintained at the bottom of 20 l aquarium in a single layer (approximately 400,000 eggs in 2 l of water) and then fertilized with a mixture of sperms (100,000 sperm per ml) collected from several spawning males. The eggs and sperms were left for 15 min for fertilization. After all the fertilized eggs settled at the bottom, the upper water layer was decanted off and the fertilized eggs aquarium was refilled with filtered irradiated sea water. This washing process was repeated 10 times at intervals of 20 min (Hahn, 1989). The fertilization rates were estimated by aliquot sampling (Ebert and Houk, 1984). Aquaria were kept in the dark in an air-conditioned room until hatching.

**Larval rearing**

The newly hatched trochophore larvae swim to the surface, leaving the egg debris and aberrant larvae at the bottom. The healthy trochophores were decanted immediately into separate sterilized aquaria to avoid bacterial contamination (Hahn, 1989). The water level in the aquaria was made up to 15 l with filtered sterilized seawater. The rearing was undertaken until the larval shell was completely formed.

**Primary film formation and larval settlement**

Settlement and metamorphosis of *H. mariae* larvae were carried out on 0.33 m × 0.33 m polycarbonate transparent flat culture plates set as artificial substrates. The plates were first prepared by exposing them to a continuous through flow of sand-filtered seawater in outdoor tanks for less than 60 days. This process covers the plate surfaces with a primary film of organic material with a complex microbial community, mainly diatoms (Hahn, 1989).

Five sets of 60 plates each were suspended vertically in two indoor rectangular fiberglass tanks (2.2 m × 1.2 m × 0.5 m) filled with filtered UV sterilized seawater, and the water level was adjusted to cover all plate surfaces. The metamorphic larvae were stocked at a density of 300 to 480 larvae per plate after which the water circulation was initiated at 300
Abalone, hatchery and seed production trials in Oman

1.0 \text{ l.h}^{-1} \text{ through a perforated pipe along the tank bottom, and the water was aerated. Changes in larval external morphology were studied by regularly cutting off small pieces of plates from 3 sets and examining the developing larvae under the microscope. After 7–15 days the settlement plates were transferred to outdoor circular tanks for further rearing. Total survival was determined after one month by counting surviving juveniles.}

Handling of juveniles and grow-out
To detach thin-shelled juveniles for transfer between tanks or other purposes (Hahn, 1989), 70% ethanol was diluted to concentrations of 1%, 2% and 4%, and the drop-off and recovery times of 10 juveniles (10–20 mm in size) were measured in each dilution. ANOVA was used to compare the drop-off and recovery times between the three treatments (Zar, 1996). Juveniles produced at MASPS were maintained for 13 months at ambient temperature to assess growth rates.

Results and Discussion
The culture of abalone is a high priority project in Oman, owing to the high value product and demand on international markets. Wild stocks are overfished and food-limited during the monsoon season, and faster growth rates than in nature may be achievable through culture (Johnson et al., 1992; Shepherd et al., 1995).

The maturity index of the broodstock collected in the pre-monsoon (May) increased between June and August, and all the individuals had mature gonads by the end of the period in December over a period of 6 months (Fig. 3). The maturity index of breeders in the post-monsoon (September) samples increased over a shorter period and reached higher value in December over 60 days (Fig. 3). After the spawning inducement in December, the samples were returned into conditioning tank and regained full maturity in February, over another 60 days (Fig. 3). The post-monsoon group was collected from the wild at the onset of the reproductive season, and was presumably physiologically ready for maturation. Regardless of

Figure 3. Changes in the maturity index of H. mariae from the pre-monsoon (May) and post-monsoon (September) samples, and conditions for six months and two months, respectively, prior to spawning.
the controlled conditioning methods applied in various abalone species, the result of shorter conditioning period achieved in our experiment (60 days) was better than that obtained with several abalone species (Table 2). Therefore, it is advantageous to collect broodstock in September, because the shorter conditioning time required will reduce costs.

Four spawning inducement trials were carried out successfully between December 1999 and April 2000 (Table 3). Spawning occurred mainly during nighttime at a temperature of 25–27°C. The pattern was similar to tropical abalone *H. asinina* in the Philippines (Capinpin and Hosoya, 1995; Fermin *et al*., 2000) and *H. varia* in India (Najmudeen and Victor, 2004). Spawning in males occurred 3 h after immersion in UV-irradiated seawater; whereas, in females it was after 5 h. Males spawned more frequently than females, which facilitated higher fertilization rates. The success rates of male and female spawning induction averaged about 63% and 11% respectively (see Table 3). However, spawning success rate of 50% or more was obtained with female *H. ruber* and *H. laevigata*, using UV irradiation and temperature shock methods (Grubert and Ritar, 2005; Daume, 2007). The low rate may have resulted from incorrect technique of UV stimulation (Hahn, 1989) or the female broodstock not being sufficiently ripe for these stimulus (Morse, 1984), thus technical modifications on female *H. mariae* spawning induction and further work on testing other spawning methods are required.

The total number of eggs spawned in the four trials ranged from 0.59–2.8 million per female with a shell length of 60–80 mm. Al-Rashdi (2001) suggested that larger female *H. mariae* (>110 mm SL) can spawn as much as 5 million eggs. This compares well with other species: 5.2 million eggs in *H. gigantea* (Yoo, 1989), 0.3 million in *H. discus hannai* (Yoo, 1989), 0.2 million in *H. diversicolor supertexta* (Chen, 1989) and 0.08 million in *H. varia* (Najmudeen and Victor, 2004). Fertilized eggs were spherical and 200 μm in diameter (Fig. 4). The fertilization rates varied between 72% and 98.6% (see Table 3).

Hatching of *H. mariae* occurred 8 h and 25 min after fertilization at 23.6°C and 6 h and 30 min at 24.7°C (Ogawa, 1997), and the external morphology of larvae

### Table 2. Conditioning period of various abalone species.

<table>
<thead>
<tr>
<th>Abalone species</th>
<th>Conditioning period (days)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Haliotis mariae</em></td>
<td>60</td>
<td>This study</td>
</tr>
<tr>
<td><em>Haliotis discus hannai</em></td>
<td>80</td>
<td>(Uki and Kikuchi, 1984)</td>
</tr>
<tr>
<td><em>Haliotis rufescens</em></td>
<td>90</td>
<td>(Ault, 1985)</td>
</tr>
<tr>
<td><em>Haliotis discus</em></td>
<td>160</td>
<td>(Kafuku and Ikenoue, 1983)</td>
</tr>
<tr>
<td><em>Haliotis laevigata</em></td>
<td>90-120</td>
<td>(Grove-Jones, 1996)</td>
</tr>
<tr>
<td><em>Haliotis roei</em></td>
<td>&gt; 180</td>
<td>(Freeman, 2001)</td>
</tr>
</tbody>
</table>

### Table 3. Data on the spawning success of *Haliotis mariae* broodstock from the various trials. The number in parenthesis indicate the number of broodstock tested for each spawning trial.

<table>
<thead>
<tr>
<th>Trial No.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date</td>
<td>13.12.99</td>
<td>16.12.99</td>
<td>23.01.00</td>
<td>01.04.00</td>
</tr>
<tr>
<td>Water temperature (°C)</td>
<td>26</td>
<td>25</td>
<td>26.5</td>
<td>27</td>
</tr>
<tr>
<td>♂ 2 (15)</td>
<td>♂ 1 (17)</td>
<td>♂ 2 (15)</td>
<td>♂ 2 (17)</td>
<td></td>
</tr>
<tr>
<td>♂ 9 (11)</td>
<td>♂ 7 (12)</td>
<td>♂ 7 (12)</td>
<td>♂ 6 (11)</td>
<td></td>
</tr>
<tr>
<td>No. of spawners</td>
<td>282</td>
<td>59</td>
<td>68.7</td>
<td>500</td>
</tr>
<tr>
<td>No. of eggs (10⁶)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fertilization rate (%)</td>
<td>72.5</td>
<td>98.6</td>
<td>88.7</td>
<td>90.4</td>
</tr>
</tbody>
</table>
over the next 30 days, from hatching to plantigrade veliger stage, is shown in Fig. 4. The laboratory observation of the *H. mariae* larval development indicated that there were no obvious differences between the external morphology of *H. mariae* larvae compared to *H. discus hannai* larvae (Seki and Kanno, 1977), except that *H. mariae* developed faster, presumably because of higher water temperatures (S.A. Shepherd, personal communication). The survival rates from trochophore to plantigrade veliger

![Figure 4](image1.png)

**Figure 4.** Embryonic development of *H. mariae* from eggs to seed production {1: fertilized egg (200µm), 2: newly hatched out larva (trochophore, 210µm), 3: pre-settlement veliger larva, 4: metamorphosed larva, 5: newly settled larvae, 6: post-settlement larva with shell formation, 7: early juvenile with first respiratory pore in the shell (2.5mm, 30 days), and 8: a produced seed in the hatchery (25mm, 120 days)}.

![Figure 5](image2.png)

**Figure 5.** The number of *H. mariae* veliger larvae and their survival rates from larval development to larval pre-settlement.
larvae obtained from the four spawning trials and introduced to the settlement plates were low and varied between 35.9% and 73.7% (Fig. 5). However, 90% has been recorded in several abalone species (Hahn, 1989). It is presumed that the mortalities are due to suboptimal water management, handling methods and temperature fluctuations (Hahn, 1989). At 25°C to 27°C, the trochophore larvae metamorphosed after 56 h and settled after 72 h from hatching (Fig. 4).

The survival of settled larvae is affected by the ingestibility and digestibility of the diatom which, in turn, depends on the species dominated in the biofilm (Roberts et al., 1999). Survival rates during the postsettlement period were generally low and variable (Searcy-Bernal et al., 1992). However, the survival of the settled larvae recorded in all our four trials ranged from 0.1–3.6% after one month (Fig. 6). These low survival rates are generally a major problem in abalone seed production (Ebert and Houk, 1984; Hahn, 1989), and were attributed to the inability to maintain the quality of diatoms (Nitzschia, Navicula, and Cocconeis) species on culture plates for abalone settlement and subsequent feeding (Mcshane, 1992; Daume et al., 2004). Growth of filamentous green algae and encrusted coralline algae on the culture plates and tank sides reduced diatom growth and quality in terms of space competition. Similar observations have been reported in tropical species such as H. varia in India (Najmudeen and Victor, 2004), perhaps as a result of high light intensity (Daume et al., 2004), which is also characteristic of Oman.

Growth of juvenile abalone was enhanced at elevated temperatures, and had reached an average length of 2.5 mm.mo⁻¹ at 26°C, in which its first respiratory pore was fully formed and visible to the naked eye (Fig. 4). Similarly, the first respiratory pore was visible after 27 days in H. varia at 27°C (Najmudeen and Victor, 2004) and 28 days in H. assinina at 27°C (Singhagraiwan and Sasaki, 1991), but after 8 weeks in H. rufescens at 15°C (Ebert and Houk, 1984) and after 43 days in H. midae (Genade et al., 1988).

Anesthetized juveniles dropped off the culture plates after an exposure time of 10–14 min to 1% ethanol (12.5 ± 1.3 min, n = 10) but this drop-off period shortened significantly to 2–4 min at a 2% dilution (3.2 ± 0.9 min) and to 0.3-0.8 min at a 4% dilution (0.55 ± 0.15 min; ANOVA F2,27= 476.7,
Abalone, hatchery and seed production trials in Oman

p < 0.001). Recovery times were 14.7 ± 1.9 min (at 1%), 10.4 ± 2.4 min (at 2%) and 9.0 ± 2.2 min (at 4%) (ANOVA F2, 27= 35.5, p < 0.001). The later result is contrary to the expectation that recovery should take longer at higher concentrations of ethanol (Hahn, 1989). Nevertheless, the juveniles anesthetized with 4% ethanol did not resume feeding for three days, whereas feeding of the 2% group was apparently normal from the first recovery day. The drop-off and recovery times using 2% ethyl alcohol was similar to that found by Hahn (1989), using ethyl carbonic acid, magnesium sulfate and chloral hydrate on juvenile abalone.

After 13 months of rearing, juveniles reached an average size of 52.9 mm shell length (range 42–64.4 mm; Fig. 7). These translate to an average growth of 4.1 mm.mo⁻¹. Stirn and Al-Hashmi (1996) reared abalone (<35 mm) collected from the wild in aquaria under closed water system and different diet regimens for 6 months, and obtained average growth of 4.8 mm.mo⁻¹ with artificial diet, 3.3 mm.mo⁻¹ with kelp. Ogawa (1997) grew juveniles up to 1.5 mm.mo⁻¹ over 2 months of hatchery production. Al-Rashdi (2002) carried out a 1-month feeding experiment at MASPS using 28 mm juveniles fed with different diets, and obtained a growth rate of 6.39 mm.mo⁻¹ with a formulated diet, 4.17 mm.mo⁻¹ with frozen U. fasciata, and 1.17 mm.mo⁻¹ with frozen brown alga N. zanardinii. These studies indicate that cultured H. mariae can grow fast on formulated diets, and also when fed fresh and frozen U. fasciata.

**Conclusion**

Artificial hatching and seed production of H. mariae have been achieved in Oman, but several challenges remain, such as locating wild abalone that are ripe enough to be spawned, improving broodstock conditioning protocols, and improving spawning success particularly for females. The low survival and settlement rates during larval development are the most important impediments to large-scale commercial aquaculture, nevertheless, the preliminary research trials described in this paper confirms that H. mariae can be cultured successfully in Oman. Further studies on the standardization of the culture techniques would help in stock enhancement programmes and commercial farming.

**Acknowledgements**

The authors would like to express their sincere thanks to Joji Ogawa who developed the research proposal of the MASPS, and to Mohamed Bal-Khair and Ali Al-Mishaikhi for technical participation in MASPS activities. Thanks are also due to Johan Groeneveld, Armando Fermin and Fahad Al-Ajmi for reading the manuscript, and to Saud Al-Habsi for his encouragement. This study was financed by the
Oman Agricultural and Fisheries Development Fund, Sultanate of Oman.

References


Grubert, M.A. and A.J. Ritar. 2005. The effect of temperature and conditioning interval on the spawning success of wild-caught blacklip (Haliotis rubra, Leach 1814) and greenlip (H. laevigata,


