Studies on the broodstock production and larval rearing of Coral demoiselle \textit{Neopomacentrus nemurus} (Bleeker 1857)

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Abstract
Steps taken for broodstock development of \textit{Neopomacentrus nemurus}, Coral demoiselle (Bleeker, 1857) in a hatchery, its spawning and development of egg to the juvenile stage are described. Among the three treatments tried, only the trial using \textit{Parvocalanus crassirostris} nauplii as the first diet helped in the development of larvae. Larval and post-larval growth was studied for a period of 65 days post hatch. The larva measured 2–2.4 mm in total length at the time of hatching. The mouth size of larva at the time of hatching measured 237.92 µm. By the 5th day, the width of the larval body had significantly increased. All the fins were conjoined, except for the caudal fin which had begun its appearance on the 2nd day itself. The digestive system was functional by the 5th day and the copepod remains were seen in the digestive tract. The larva metamorphosed by the 15th day with fully developed fins and fin rays. On the 20th day, the larval body had begun to display signs of fin pigmentation. The larva assumed adult pigmentation by the 35th day. By the 65th day, the larva grew to a total length of about 27 mm and weighed about 0.56 g.

Keywords
broodstock development, Damselfish, larval rearing, \textit{Neopomacentrus nemurus}, Pomacentridae

1 | INTRODUCTION

The marine ornamental fish industry depends mostly on the organisms harvested from the wild (Wabnitz, Taylor, Green, & Razak, 2003). The only solution to this problem is aquaculture, which could provide a growing proportion of marine ornamental fish in the near future (Molina & Segade, 2012). Out of the 1,000 species of coral reef fish traded (Green, 2003), only 51 have been cultured in captivity for the aquarium trade (Arvedlund, McCormick, & Ainsworth, 2000) and a very few in commercial quantities. Approximately 30 million marine reef fish belonging to roughly 1,800 different species are commercialized every year worldwide (Rhine et al., 2012; Thornhill, 2012; Wabnitz et al., 2003). In 2013, Singapore had exported around US$ 56 million worth of ornamental fish to over 80 countries (FAO, 2014). Over the years since 1985, its international trade has shown an increasing trend (Moorhead & Zeng, 2010), with an average growth rate of approximately 14% per year with an estimated wholesale trade of nearly US$ 1 billion and retail trade of about US$ 3 billion (Olivotto et al., 2005). Large-scale destruction of coral reefs by anthropogenic activities has taken its toll on the fragile coral reef ecosystem that houses many of these fish. In view of this, many captive breeding programmes have been initiated for minimizing harvesting from the wild and maximizing ex-situ production of these fish. It can be said that the marine ornamental fish industry is still in its infancy and many hazards have yet to be faced before we could safely say that we have reached a significant milestone in this field. The constraints faced by this industry are so many that we cannot pinpoint a single factor, many fish need large tanks to spawn while others have to be provided with suitable photoperiods and temperatures (Holt & Riley, 2001) and specific environmental conditions. Some others need hormones to make them spawn (Moe, 1997). But one advantage that Pomacentrids have over most other...
ornamental fish species is that they need not be subjected to the other techniques mentioned before. Pomacentrids are continuous spawners and they lay eggs if provided with the proper water quality conditions and food, the only contradiction to this being members of the genus _Abudefduf_. Even though reports of successful breeding of Pomacentrids of genera _Amphirion_ and _Premnas_ have been published, successful larval rearing of damselfish is still in its infancy. There are only a few reports regarding the successful larval development of many damselfish such as _Abudefduf saxatilis_ (Alshuth, Tucker, & Hatley, 1998; Wittenrich, Turingan, & Cassiano, 2012), _Microspathodon chrysurus_ (Potthoff, Saksena, Moe, & Young, 1987), and _Pomacentrus amboinensis_ (Murphy, Leis, & Kavanagh, 2007). Breeding and larviculture of the sapphire devil damselfish _Chrysiptera cyanea_ were carried out by Gopakumar, Santhosi and Ramamurthy (2009). A comprehensive study on the breeding of _N. nemurus_ (Coral demoiselle) is yet unavailable. This study is an attempt to throw light on the broodstock development, larval rearing, and larval development of _N. nemurus_.

The Coral demoiselle, _N. nemurus_ is distributed throughout the Indian Ocean and Western Central Pacific. Adults inhabit lagoon and inshore coral reefs, and they are found in aggregations of coral reefs from where they feed on zooplankton (Myers, 1991). They occur at a depth of about 1–10 m (Allen, 1991). The fish has a greyish body with transparent pectoral and pelvic fins. The spiny dorsal is grey in colour and the soft dorsal is yellow, same is the case of the anal and caudal fins.

2 | MATERIALS AND METHODS

2.1 | Broodstock development

Eight sexually mature fish of _N. nemurus_ measuring 6.5–10 cm were collected from the wild.

The collection was made from Vizhinjam along the south-west coast of India in September, 1998 by skin diving using scoop net. They were transported to the hatchery of Vizhinjam Research Centre of ICAR-Central Marine Fisheries Research Institute, in jerry cans of 20 L capacity in aerated seawater using battery operated aerators.

2.2 | Broodstock tank

The fish were stocked in a rectangular fibre-reinforced plastic (FRP) tank of 500 L capacity and water was continuously filtered using an in-situ biological filter of volume 21 L. Coral rubble is used as the biological filter material. Coral containing numerous pores provide a large surface area for nitrifying bacteria to colonize which helps in the effective removal of ammonia produced by the fish. The area being near the equator, the photoperiod during the rearing period was of 12 hr L:12 hr D and no photoperiod manipulation was done. The tank is kept in a place where there is minimum disturbance. Physico-chemical parameters, such as salinity, pH, dissolved oxygen, water temperature, and _NH_3 were checked twice in a week using standard methods. For maintaining water quality in the rearing tanks, about 10% of rearing water along with excretory matter and excess feed was siphoned off daily and the water was replaced with fresh seawater. The tank was provided with earthen pots and polyvinyl chloride (PVC) pipes as a substratum for laying eggs. The fish were fed four times a day as per the schedule: pellet feed at 10.00 hr; boiled mussel meat at 12.00 hr and 14.00 hr, and _Artemia_ at 16.00 hr.

Observations on the courtship, parental care, and response to nest disturbance were carried out twice daily for a period of 4 months. In order to assess the behaviour, F1 generation _N. nemurus_ were kept in three aquarium tanks and observations were made. The observations were carried out during 10–11 a.m. and 4–5 p.m. for the presence of any new egg clutches. Observations were made for more than 500 hr. Three months of observation for a period of 2 hr a day were made for making possible conclusions. To monitor the embryonic development, fertilized eggs from three different clutches were taken out three times. It was difficult to observe the frequency as two fish were laying eggs. The characters of newly spawned and fertilized egg were noted for a period of 3 days. The larval development was monitored by sampling five numbers of fish larvae every day until the 5th day and on 10, 15, 20, 30, 40, 50, and 65 dph for growth measurements. The embryonic development was monitored using a Leica DMCS research microscope. Water was taken in a beaker and the number of larvae, their behaviour, and feed availability were observed. The availability of adult copepods, number of rotifers, and number of _Artemia_ were counted by taking subsamples and the feeding was adjusted accordingly during the different stages of larval rearing according to the schedule given in the table. The larval development was monitored using a Leica S8APO stereozoom microscope. From the measurements obtained, the mean and standard deviation were calculated.

2.3 | Live feed culture

Live feeds, such as phytoplankton, rotifer, marine zooplankton including copepods, and _Artemia_, were used for the larval rearing experiments. For feeding copepod culture and for maintaining green water in the rearing tanks, the stock culture of algae viz., _Nannochloropsis oculata_ and _Isochrysis galbana_ was maintained in a stock culture room at 24°C in 100 ml to 3-L flasks and then the cultures were up-scaled to 20-L carboys for feeding. The copepod _Parvocalanus crassirostris_ was cultured using _N. oculata_ and _I. galbana_. Rotifer _Brachionus plicatilis_ was cultured using algae _N. oculata_. Rotifers were enriched using Algamac 2000 (Aquafauna Bio-Marine, USA).

2.4 | Larviculture tank

Three treatments were tried for rearing the larvae and about 400 damselfish larvae were used for each treatment in a rectangular 500-L FRP tank with light blue colour (100 cm length × 80 cm breadth × 60 cm depth). Larvae were stocked at a rate of 10 per ml. Seawater used was filtered using a series of 10-, 5-, and 2-µm cartridge filters and mild aeration was given throughout the rearing period.
Three feeding trials were conducted with copepod (T1), rotifer with and without enrichment (T2 and T3) and in all three treatments, the green water technique was employed. Algal cell density used for the experiment ranged from $1.4 \times 10^6$ to $9 \times 10^6$ per ml. All the three tanks were supplied with the algae *N. oculata* and *I. galbana* (3:1) up to the 20th day. In (T1), the tanks contained adults of *P. crassirostris*. The number of adult copepods was maintained at about $0.7–0.9$ per ml respectively. The presence of adult copepods will ensure the smallest naupliar stage just hatched out from egg available to the altricial fish larvae. The tanks of (T2) contained enriched rotifer *B. plicatilus*, size $170.532–213.565$ µm ($3–6$ per ml). The tanks of (T3) contained rotifer *B. plicatilus* ($3–6$ per ml).

For feeding experiments, the larvae were divided and from day 0 to 8 the first group, T1 was fed with the copepod *P. crassirostris* nauplii and from the 6 to 18th day on rotifer *B. plicatilus* ($3–6$ per ml) and then from 16th day on Artemia nauplii. Boiled mussel meat was given to the larvae from the 35th day onwards. In T2, the larvae were fed with enriched *B. plicatilus* from 1 to 5 dph. In T3, the larvae were fed from day 1 to 5 on *B. plicatilus* (Table 1). Treatments were repeated three times. The data were analysed and the result was expressed as mean ± SD of the data (Table 2).

### Table 1. Feeding schedule of hatchery-reared *Neopomacentrus nemurus*

<table>
<thead>
<tr>
<th>Days</th>
<th>Treatment 1</th>
<th>Treatment 2</th>
<th>Treatment 3</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Nanochloropsis and Isochrysis</td>
<td>Nanochloropsis and Isochrysis</td>
<td>Nanochloropsis and Isochrysis</td>
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<tr>
<td></td>
<td>(0.7–9 per ml)</td>
<td>Brachionous plicatilus (3–6 per ml)</td>
<td>Brachionous plicatilus (3–6 per ml)</td>
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<tr>
<td></td>
<td>Parvocalanus nauplii</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Brachionous plicatilus (3–6 per ml)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Artemia nauplii</td>
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<tr>
<td></td>
<td>Boiled mussel meat</td>
<td></td>
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3.1 | Reproductive behaviour

The fish exhibited territorial behaviour and were observed to attack other fish if they came close to their eggs. The fish preferred to lay the eggs on the sides of the tank other than the earthen pots or PVC pipes (Figure 1). *N. nemurus* is sexually monomorphic and the male parent exhibited a high degree of parental care for the eggs, such as fanning the eggs with the pectoral and caudal fins and removing the infected eggs with the mouth (Figure 2). The fish also tend to consume the egg clutch if it was disturbed. The fertilized eggs were transparent, capsule shaped, demersal, and contained oil globules. The eggs measured $1.168–1.156$ µm in length and $380–400$ µm in width. The yolk length was found to be $725$ µm on the first day after spawning. The egg clutches contained $1,500–1,800$ eggs with an average of $1,653.36 ± 133.55$ per spawn.

The fish exhibited a high spawning frequency with an average of 7–8 per month, indicating that it spawns on every 3/5th day. In the beginning, the spawning frequency was observed to be 2–3 times per month with no specific interval. The highest spawning frequency was observed during the month of March. Hatching took place in the early evening at about 19.00–19.30 hr. The larvae were collected from the broodstock tank and transferred to the larval rearing tanks with the help of a small trough by concentrating the larvae to one area using a torch as the larvae are photopositive in nature. The hatching rate was found out to be 98%.

3.2 | Larval development

Hatching usually occurred on the 3rd day of incubation. Eggs were encased in a transparent capsule, the colour of which changed from transparent to yellowish in the case of spoilage and the spoiled eggs

<table>
<thead>
<tr>
<th>Days</th>
<th>Feeding schedule of hatchery-reared <em>Neopomacentrus nemurus</em></th>
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<tr>
<td>0 1 2 3 4 5 6 7 8 9 10 12 14 16 18 20 25 30 35 40 45 50 55 60 65</td>
<td></td>
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</table>
also exhibited a cloudy appearance. The egg capsule did not exhibit any external morphological alterations during development. In addition to the yolk, the eggs were provided with oil globules which equipped them with the capacity for buoyancy and enabled them to stand upright while attached to the substratum. The number of oil droplets varied from 1 to several. On the first day as a consequence of meroblastic discoidal cleavage, the yolky part of the egg remained uncleaved while the cleavage and micromere formation occurred in the blastodisc where the cytoplasm was confined (Figure 3). On the second day, rudiments of the head fold and tail fold were visible in the embryo. The larval body showed clear signs of development with the appearance of melanophores over the tail region. The yolk had shrunk to half of its normal size. On the 3rd day at the time of hatching, the larval body was almost fully formed. The optic vesicles had completed its development followed by the appearance of the retina. The eyes showed a bluish-silver colour, the tail was totally curved and extended to the head, these two signs indicated that the larvae were about to hatch. The larva had developed an efficient circulatory system, which was evident by the heartbeat. The larval body showed signs of segmentation which was evident by the appearance of paired myomeres. The yolk had shrunken and now occupied one third of the egg, the larval body was transparent except for the presence of a yellow colouration in the gut and melanophores (Figure 4). The larval body occupies the whole of the

<table>
<thead>
<tr>
<th>Days post hatch (dph)</th>
<th>Total length (mm)</th>
<th>Standard length (mm)</th>
<th>Head length (mm)</th>
<th>Head width (mm)</th>
<th>Body width (mm)</th>
<th>Eye diameter (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.18 ± 0.15</td>
<td>1.84 ± 0.11</td>
<td>0.67 ± 0.04</td>
<td>0.47 ± 0.04</td>
<td>0.23 ± 0.04</td>
<td>0.17 ± 0.04</td>
</tr>
<tr>
<td>2-2.4</td>
<td>1.8-2</td>
<td>0.6-0.7</td>
<td>0.4-0.5</td>
<td>0.2-0.3</td>
<td>0.17-0.2</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>3.08 ± 0.26</td>
<td>2.72 ± 0.32</td>
<td>0.86 ± 0.09</td>
<td>0.69 ± 0.12</td>
<td>0.56 ± 0.11</td>
<td>0.26 ± 0.08</td>
</tr>
<tr>
<td>10</td>
<td>3.74 ± 0.50</td>
<td>3.3-3.1</td>
<td>0.8-0.86</td>
<td>0.6-0.9</td>
<td>0.5-0.7</td>
<td>0.15-0.35</td>
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<tr>
<td>10</td>
<td>4.94 ± 0.30</td>
<td>4.32 ± 0.49</td>
<td>1.36 ± 0.15</td>
<td>1.33 ± 0.18</td>
<td>1.20 ± 0.28</td>
<td>0.54 ± 0.11</td>
</tr>
<tr>
<td>4-5-5.2</td>
<td>3.5-4.7</td>
<td>1.2-1.5</td>
<td>1.12-1.52</td>
<td>0.79-1.45</td>
<td>0.4-0.65</td>
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</tr>
<tr>
<td>15</td>
<td>7.24 ± 0.50</td>
<td>5.88 ± 0.4</td>
<td>2 ± 0.11</td>
<td>1.76 ± 0.21</td>
<td>1.76 ± 0.48</td>
<td>0.78 ± 0.15</td>
</tr>
<tr>
<td>20</td>
<td>8.2 ± 0.67</td>
<td>8.1 ± 0.99</td>
<td>3.5 ± 0.34</td>
<td>3.2 ± 0.33</td>
<td>3 ± 0.41</td>
<td>1.1 ± 0.71</td>
</tr>
<tr>
<td>9-11.2</td>
<td>7.2-8.8</td>
<td>2.8-3.5</td>
<td>2.6-3.4</td>
<td>2.7-3.7</td>
<td>1-1.5</td>
<td></td>
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<tr>
<td>30</td>
<td>13.06 ± 1.65</td>
<td>10.3 ± 1.64</td>
<td>4.14 ± 0.34</td>
<td>4.32 ± 0.37</td>
<td>3.96 ± 0.69</td>
<td>1.44 ± 0.34</td>
</tr>
<tr>
<td>10-13.7</td>
<td>8.1-12.6</td>
<td>3.8-4.5</td>
<td>3.9-4.8</td>
<td>2.8-4.5</td>
<td>1.2-2.0</td>
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</tr>
<tr>
<td>40</td>
<td>17.2 ± 1.92</td>
<td>14.1 ± 1.88</td>
<td>4.88 ± 0.37</td>
<td>4.96 ± 0.50</td>
<td>4.68 ± 0.63</td>
<td>1.90 ± 0.29</td>
</tr>
<tr>
<td>15-20</td>
<td>14-17</td>
<td>4.4-5.3</td>
<td>4.5-5.8</td>
<td>4-5.6</td>
<td>1.5-2.18</td>
<td></td>
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<tr>
<td>50</td>
<td>20.3 ± 4.0</td>
<td>17.6 ± 2.1</td>
<td>5.12 ± 0.5</td>
<td>5.49 ± 0.4</td>
<td>5.6 ± 0.7</td>
<td>2.1 ± 0.2</td>
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<td>15-30</td>
<td>15-20</td>
<td>4.4-5.6</td>
<td>5-6.04</td>
<td>4.7-6.3</td>
<td>1.9-2.3</td>
<td></td>
</tr>
<tr>
<td>65</td>
<td>27 ± 2.5</td>
<td>18 ± 1.4</td>
<td>6 ± 0.67</td>
<td>5.9 ± 0.34</td>
<td>6.18 ± 0.57</td>
<td>2.3 ± 0.16</td>
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<tr>
<td>24-30</td>
<td>16-20</td>
<td>5.2-6.8</td>
<td>5.5-6.4</td>
<td>5.6-7</td>
<td>2.1-2.5</td>
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capsule and then breaks it off by vigorous wriggling movements at the end of incubation period (5 days).

3.3 | Feed management

In T1, the presence of *P. crassirostris* adults ensures that the smallest possible live stages of copepod nauplii are available to the newly hatched larvae. In this treatment, the larvae survived. In T2 and T3, the larvae that survived with the available yolk reserve until the 4–5th day perished. The larvae in T1 showed active swimming movements, and the larvae also appeared to favour the sides of the tank where the copepods were concentrated. The microscopic observation of the dissected stomach on the 4th day revealed the presence of copepod nauplii. In T2, when the larvae were given the enriched rotifer *B. plicatilis* of size range (170.532–213.565 µm), they did not feed on rotifers and this resulted in the total mortality of all larvae in the tank by the 4th–5th day. Similar observations were made in T3 fed with *B. plicatilis* without enrichment.

The larva was transparent and measured 2–2.4 mm in total length at the time of hatching. Since the larva was atrial and with a mouth gap of about 237.92 µm, it was provided with a small but regressed yolk sac to meet the requirements of its early development. Although immediately after hatching, the larvae did not feed on any copepod nauplii, it consumed the algae provided in the tank and the dissection of the larval body helped to confirm the same. The first nauplii of copepods are in the range of 50–100 µm, which makes it suitable food for the damsel fish larvae. The eye is well developed in the larvae and is found protruding from the two sides of the head, which is an indication of its carnivorous and voracious habit (Figure 5). The melanophores, which were making their appearance when the fish was in the embryonic stage, have established their presence now. The larva follows a developmental pattern characterized by 4–5 melanophores along the head and with melanophores below the ventral midline of the body which are not much developed as that of the head. They were about 13 in number. All the fins are fused to form a single fin fold which surrounds the body (Figure 6).

By the 5th day, the width of the larval body had drastically increased. The pigmentation on the larval body was well evident in the opercular region which was transparent at first and then steadily attained a yellow colour. The drastic increase in pigmentation was also evident in the ventral midline. All the fins were conjoined except for the caudal fin which had begun its appearance on the 2nd day itself. The larval body did not exhibit any colouration and remained transparent. The dissection of the digestive tract on the 5th day revealed the presence of a well-developed digestive system, which was supported by the fact that copepod nauplii were obtained from the larval stomach. By the 7th and 8th day, one of the notable changes, which occurred in the larval body, was the infiltration of the yellow pigmentation which was restricted to the gut into the orbital region (Figure 7).
On the 10th day, the fin separation was complete during which the larval body showed the appearance of fins and fin rays. The soft dorsal had separated from the spinous dorsal fin. Many of the specimens had undergone notochord flexion. Pigmentation of the deep pre-anal body was characterized by extensive scattered stellate melanophores. Pigmentation of the stomach, operculum, and pelvic fins did not show any notable difference. There was an increase in the number of caudal fin rays, the larva showed increased pigmentation and decreased transparency. A slightly yellowish colour was observed in the region of the body where the melanophores were present (Figure 8).

The larvae developed into subadult fish by the 15th day with fully developed fins and fin rays. They showed vigorous swimming movements and voracious feeding. The melanophores present over the body varied in size and distribution. Stellate and stellate reticulate granules were present above the eyes, the snout and upper jaw, but punctate melanophores were predominant in the gut and anal fin (Figure 9).

On the 20th day, the larval body had begun to display signs of fin pigmentation. Black pigmentation had begun to develop in the centre of the spinous dorsal fin. The rest of the fins were transparent. The depth of the larval body increased conspicuously. Punctate melanophores were present on the ventral midline (Figure 10).

The larva assumed adult pigmentation on the 35th day and by then, the larva resembled the adult in all other aspects except size. Now the fish had attained the characteristic greyish colour of the adult on the anterior region of the body, whereas the posterior region exhibited a yellowish colouration which extended to the two sides of the caudal fin, the middle of the caudal fin was more or less transparent. The pectoral and anal fins were transparent with some yellow pigment granules. The spinous dorsal possessed black
pigment granules whereas the soft dorsal had yellowish granules (Figure 11).

On the 50th day, the pigmentation of the body showed some change in which the anterior part of the body exhibited a greyish colouration and the posterior region exhibited a bluish-grey colour, the fin pigmentation which was restricted to broken pigments changed into one with continuous colouration. The soft dorsal fins exhibited a continuous yellow colour, while yellow colour was also prominent on the caudal peduncle and the caudal fins (Figure 12).

On the 65th day, the pigmentation pattern was unchanged, but the difference in colouration between the anterior and posterior sides decreased notably. From now on, the larva did not undergo any change in skin and fin pigmentation but only increased in body mass (Figure 13). By the 65th day, the larva grew to a total length of about 27 mm and weighed about 0.56 g (Figure 14). The growth in terms of the total length showed a steady increase, whereas the weight increase was slow until 20 days thereafter it increased at a higher rate. The average growth of *N. nemurus* in length and weight is depicted in the table. The average survival rate was found to be 4.9 ± 2.0 from three trials (Figure 15).

In order to assess the sex ratio, 10 fish of F1 generation (9-month-old specimens), which started laying eggs from 177 days which were kept in the rearing tank, were sacrificed and it was observed that two mature males, two mature females, and two maturing females were present in the tank. There were two active females and one active male in the tank. The active male was guarding the eggs and two active females were depositing the eggs.

4 | DISCUSSION

The number of marine ornamental fish species, which are bred on a commercial scale, is very much limited in number. The hatchery production of marine ornamental fish is subjected to many constraints
such as difficulty in producing good quality gametes and producing a large number of larvae which will undergo metamorphosis into good quality juveniles (Holt, 2003). It is now evident that even before the commencement of captive spawning, different factors can affect the quality juveniles (Holt, 2003). It is now evident that even before the commencement of captive spawning, different factors can affect the quality of the larvae, the notable being broodstock genetics (Green 2004), and environmental stress (Schreck, Contreras-Sanzhez, & Fitzpatrick, 2001).

The water quality parameters in broodstock tank, such as temperature, salinity, pH, ammonia, nitrite, and nitrate, were maintained to the optimum level. Nitrate of less than 18 ppm, nitrite and ammonia levels of less than 0.01 ppm were always maintained in the tanks. Temperature, salinity, and pH in the breeding tanks were maintained at 27 ± 02°C, 30–32 ppt, and 8–8.2 respectively. Temperature in the broodstock tanks must be maintained at an optimum as the temperature can accelerate or retard the development rate of the embryo. Bermudes & Ritar, 1999; Das et al., 2006; Kazuyuki, Hisashi, & Shogoro, 1988; Miranda, Cal, & Iglesias, 1990; Moran, Smith, Gara, & Poortenaar, 2007). Epinephelus coioides, which have very small larvae similar to those of Neopomacentrus nemurus, was successfully reared in a hatchery having water quality parameters: water temperature 26.0 ± 0.8°C, ammonia concentration 0.18 ± 0.05 mg/L, nitrite concentration 0.021 ± 0.003 mg/L, and pH 7.8 ± 0.4 (Yousif, Kumar, Balamurugan, Hozifa, & Sagir, 2016).

Brood fish were fed four times a day with pellet feed, boiled mussel meat, and Artemia nauplii. This feeding schedule can be followed as such or can be subjected to slight variations. It is always recommended to feed the broodstock with a well-balanced compounded feed (Olivotto & Geffroy, 2017). In Sunrise dottyback Pseudochromis flavivertex fish were fed twice a day using frozen adult Artemia, frozen plankton, and chopped fish and shrimps (Olivotto et al., 2006). In honeycomb grouper, Epinephelus merra the spawners were provided with fresh sardines on alternate days and squid meat (Jagadis, Ignatius, Kandasami, & Khan, 2006).

Neopomacentrus nemurus performed spawning during the morning, while a similar behaviour was reported in the damselfish Neopomacentrus cyanomos (Rohini Krishna, Anil, Neethu Raj, & Santhosh, 2016). But in Amblyglyphidodon leucogaster, the spawning occurred throughout the day (Goulet, 1995). The eggs were demersal and were deposited on the earthen pots, tiles, PVC pipes, filters, or sides of the tank. In natural habitat, they always deposit eggs over a certain substratum and the choice of the substratum varies with different species. In the case of N. nemurus, they always prefer to deposit eggs on the sea-floor or among coral masses. Whereas in the case of brown chromis, Chromis multilineata eggs were deposited in large masses on Sargassum (Myrberg, Brahy, & Emery, 1967). Beaugregory damsel, Stegastes leucostictus deposited their eggs on empty conch shells or on the under surface of rocks, sometimes they also preferred Sea fans shells, cans, bottles, and other hard substrates (Brinley, 1939). From this, we can conclude that the availability of a suitable substrate is indispensable for spawning.

The spawning frequency in N. nemurus was found to be 7–8 per month, which is much higher when compared with other fish such as Premnas biaculeatus or Amphiprion nigripes (Anil, Santhosh, Prasad, & George, 2012; Madhu, Madhu, & Retheesh, 2012).

The incubation period of N. nemurus is 3 days, which is characteristic of damsel fish having small egg size, here they are about 1,168–1,156 µm long and 380–400 µm wide. Similar results were observed in C. cyanea by Gopakumar, Madhu, et al. (2009) in which the egg size measured from 1.3 mm in length and 0.6 mm in width. Species such as A. saxatilis have larger eggs (1,100–1,250 mm long and 650–670 mm wide) and have an incubation period of about 5–7 days (Alshuth et al., 1998; Thresher, 1984). Kavanagh (1996) recorded bigger eggs in Acanthochromis polyacanthus, which measured 3.7–4.3 mm in length and 1.4–1.5 mm in width and they have an incubation period of about 16 days.

The larval body started forming from the 2nd day of incubation onwards. The myomeres were visible from the 3rd day onwards. This development is noteworthy as muscle development can be taken as a measure of the morphological development and swimming ability (Fisher, Bellwood, & Job, 2000).

For larval rearing, a rectangular FRP tank with light blue colour holding 400 L of seawater was used. Tank colour in the rearing tank may affect the feeding of marine fish larvae. In groupers, the tank colour preference is species dependent (Ma, Guo, Zhang, & Bai,
There is paucity of literature to support the fact that colour of the larval rearing tank affects the larval survival in damsel fish. When *A. saxatilis* larvae were stocked in 120 L and 60 L tanks at a stocking rate of 8 L\(^{-1}\), the larvae survived in the former whereas complete mortality was observed in the smaller tank though appropriate food organisms were present to promote growth and survival in both the tanks (Wittenrich et al., 2012). The above observation and several studies indicated that in addition to other factors mentioned, tank size and stocking density are two crucial factors which affect the survival of damsel fish larvae.

The larval body showed notable development from the 5th day onwards, which included an increase in the body width, the appearance of pigmentation, and the functioning of a full-fledged digestive system. Similar observations on pigmentation were made in *A. saxatilis* larvae, which showed the appearance of pigmentation on the 5th to 7th day (Alshuth et al., 1998). The yolk was also exhausted by the 3–4th day in *A. saxatilis* along with the appearance of a well-developed digestive system.

In *N. nemurus* by the 10th day, fin separation was evident but in *A. saxatilis*, the caudal rays had just begun to make an appearance and in *N. nemurus*, most of the larvae had undergone notochord flexion, similar observations were made in the *A. saxatilis* larvae. In *N. nemurus*, the larval body was characterized by stellate melanophores, a similar pattern of pigmentation was observed in *A. saxatilis*. Pomatocentrids also lack elongate fin spines and the fin elements or scales develop precociously (Murphy et al., 2007). On the 15th day, fin separation has just been completed in *N. nemurus* whereas in *A. saxatilis* fin separation can be witnessed on the 17th day. In *N. nemurus*, the fin pigmentation developed on spinous dorsal fin by the 21st day whereas in *N. nemurus* it took about 21–27 days to complete the pigmentation in the spinous dorsal fin (Wellington & Victor, 1989). The stripping pattern was evident in *A. saxatilis* from the 21st day onwards whereas in *N. nemurus* it took about 35 days to attain the adult colouration.

In sergeant major, *A. saxatilis* the feeding schedule included *B. plicatilis*, 35–90 µm size-sorted zooplankton, 90–250 µm size-sorted zooplankton, 12 hr *Artemia* sp., and 48 hr *Artemia* sp. (Wittenrich et al., 2012). *Neopomacentrus cyanomos* was given zooplankton segregated by passing through a sieve of mesh size 500 µm and then retained in the sieve of 20-µm mesh size, plankton could be replaced successfully by *P. serricaudatus* from the 10th day onwards and from the 15th day onwards freshly hatched *Artemia* nauplii were given to the larvae. No mortality was reported after the 15th day. Pellet feed of size 300 µm and boiled mussel meat were given to the larvae from the 30th day onwards (Rohini Krishna et al., 2016). In the present study, the larvae were divided into three groups and in T1 from day 0 to 8 the larva was fed with the copepod *Parvocalanus* nauplii and from 6 to 18th day on rotifer and then from 16th day on *Artemia* nauplii. Mussel meat was given from 35th day onwards.

The composition of live feed in the rearing tank affects the larval survival and performance. This was demonstrated by the splitting of the larvae into three experimental groups. From day 1 to 5, fish were fed with copepods in T1, enriched rotifer (*B. plicatilis*) in T2, and non-enriched rotifer in T3. From this, we can understand the importance of the starter feed in the larval survival and growth. Even though rotifers are not suitable to be given as a starter feed, they can be successfully employed to feed the larvae from the 6th or 7th day onwards, which is illustrated by the survival of the larvae in the first tank. Although proper prey items are indispensable for larval survival, many other factors influence larval survival such as photoperiod (Arvedlund et al., 2000; Olivotto, Cardinale, Barbareis, Maradonna, & Carnevali, 2003), temperature and salinity (Hart, Hutchison, & Purser, 1996), development of appropriate digestive enzymes (Kolkovski, Tandler, Kissel, & Gartler, 1993), development of the feeding mechanism (Wittenrich, 2007), rearing tank design (Büke, Özden, & Arslan, 2005), and flow field (Sakakura, Shiotani, Chuda, & Hagiwara, 2007).

The survival rate was found to be 4.9 ± 2.0. Similar (3%–5%) survival rates were obtained in dusky grouper *Epinephelus marginatus*, which was fed on copepods (Kerber, Azevedo Silva, Antonio dos Santos, & Sanches, 2012). Copepod nauplii can be used as a starter feed for larvae of marine fish because of their small size and high nutritional value (McKinnon et al., 2003). Lower survival (2%) was obtained in giant grouper *Epinephelus lanceolatus* in which a mix of calanoid copepods (*P. crassirostris*) (0.5–3.0 per ml) and enriched S-type rotifers (10 per ml) was given to the fish larvae at 3 days per hatch (Armando, Adam, & Kevin, 2014). Other pomacentrids such as clown fish have larger eggs and can be reared using rotifers. *Amphiprion nigripes* was fed on rotifer from 0 to 15 dph (Anil et al., 2012). The super small (ss) strain of rotifers has been used successfully as a starter feed in larval culture of altricial fish larvae like groupers, so attempts can be made to use such ss strains of rotifers as starter feed for damsel fish larvae. Damsel fish of genus *Abudedefduf* have larger eggs, longer incubation time and bigger larvae so they could be reared using rotifers during the initial days of hatching. In Sergeant major, *A. saxatilis* 6% survival was obtained by a diet of rotifers and size-sorted copepod dominated zooplankton (Wittenrich et al., 2012). Adding adult copepods in the present study ensured the availability of first naupliar stages (40–45 µm) for the fish larvae to feed upon.

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