Chapter One

BIOLOGY AND ECOLOGY

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For the culture of any species to evolve from tradition or art to science, basic information on the biology of the species is required.

This paper reviews current information on Penaeus monodon including taxonomy, morphology, distribution, and bionomics and life history. The last covers reproduction, development of embryo, larva up to adult, spawning, food and feeding, and physiology.

Problems that have cropped up with the intensification of prawn culture, e.g. discharge of pesticides from grow-out ponds, are highlighted. Other conflicts such as the conversion of mangroves and other estuaries, considered nursery grounds of various marine fauna including P. monodon, into fishponds; overexploitation of wild spawners with no stock assessment data; and indiscriminate throwing away of other prawn and finfish fry from wild collections in favor of P. monodon fry could adversely affect the ecology of mangroves and other marine ecosystems.

TAXONOMY

The genus Penaeus Fabricius (1798) was placed on the Official List of Generic Names in Zoology as Name No. 498 upon the discovery and description of Penaeus monodon by John Christ Fabricius in 1798 (Mohamed 1970). With the revision of the specific name monodon by Holthuis, the two species have become stabilized and the name P. monodon is generally accepted for the present species (Hall 1961, Mohamed 1970, Motoh 1981). No subspecies are currently recognized for this species and P. monodon manillensis (Villaluz and Arriola 1938) proved to be based on an abnormal specimen of P. semisulcatus (Mohamed 1970, Motoh 1981).

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Definition

The taxonomic definition of the giant tiger prawn is as follows:

Phylum Arthropoda
Class Crustacea
Subclass Malacostraca
Order Decapoda
Suborder Natantia
Infraorder Penaeidea
Superfamily Penaeoidea
Family Penaeidae Rafinesque, 1815
Genus Penaeus Fabricius, 1798
Subgenus Penaeus
Species monodon

Scientific name: *Penaeus (Penaeus) monodon* Fabricius, 1798.

It has four synonyms:

*Penaeus carinatus* Dana, 1852
*P. caeruleus* Stebbings, 1905
*P. monodon* var. *manillensis* Villaluz
and Arriola, 1938
*P. bubulus* Kubo, 1949

The FAO names are giant tiger prawn (English), crevette géante tigree (French), and camaron tigre gigante (Spanish).

The term shrimps and prawns are common English names used synonymously due to the absence of systematic basis to mark a distinction (Wickins 1976, Holthuis 1980). In an attempt to clarify the issue, Holthuis (1980) traced the origin of the names shrimps and prawns and its usage in various countries. In general, shrimps refer to the smaller animals and prawns to the larger ones, while according to Food and Agriculture Organization (FAO) Convention, shrimps refer to marine
penaeids while prawns refer to freshwater palaemonids. In the present view, the term prawn will be used following the accepted usage at SEAFDEC Aquaculture Department.

Considerable taxonomic works on the Penaeidae have been published throughout the world, many providing detailed information to interested workers. Motoh (1977) also compiled various common and vernacular names of commercially important penaeid prawns and shrimps.

For the identification of *P. monodon* postlarvae, Motoh and Buri (1981) published a key for penaeid postlarvae from Panay Island, Philippines; so did Prawirodihardjo et al (1975) in Indonesia, while Rao and Gopalakrishnan (1969) identified *P. monodon* and *P. indicus* juveniles in India.

**MORPHOLOGY**

The morphological features of *P. monodon* have been described in detail by workers from various countries, among whom are Bate 1888; Blanco and Arriola 1937; Villaluz and Arriola 1938; Kubo 1949; Holthuis 1949; Racek 1955, 1957, 1972; Hall 1956, 1961, 1962; Dall 1957; Cheung 1960; Racek and Yaldwin 1971; Motoh 1981; and Motoh and Buri 1984.

The following description includes important features sufficient for the identification of this species. The shell is smooth, polished, and glabrous. The rostrum extends beyond the tip of the antennular peduncle, is sigmoidal in shape, and possesses 6-8 dorsal and 2-4 ventral teeth, mostly 7 and 3, respectively. The carapace is carinated with the adrostral carina almost reaching the posterior margin of the carapace. The gastro-orbital carina occupies the posterior one-third to one-half distance between the post-orbital margin of the carapace and the hepatic spine. The hepatic carina is prominent and almost horizontal. The antennular flagellum is subequal to or slightly longer than the peduncle. Exopods are present on the first four pereopods but absent in the fifth. The abdomen is carinated dorsally from the anterior one-third of the fourth, to the posterior end of the sixth, somites. The telson has a median groove but without dorso-lateral spines. Figure 1 shows the various parts of *P. monodon* and the technical terms with taxonomic importance.
Fig. 1. Lateral view of *P. monodon* showing important parts (Motoh 1981)
A live giant tiger prawn has the following characteristic coloration: carapace and abdomen are transversely banded with red and white, the antennae are greyish brown, and the pereopods and pleopods are brown with crimson fringing setae. In shallow brackish waters or when cultured in ponds, the color changes to dark and, often, to blackish brown (Motoh 1981).

**DISTRIBUTION**

The giant tiger prawn is widely distributed throughout the greater part of the Indo-Pacific region, ranging northward to Japan and Taiwan, eastward to Tahiti, southward to Australia, and westward to Africa (Racek 1955; Holthuis and Rosa 1965; Motoh 1981, 1985).

In general, *P. monodon* is distributed from 30°E to 155°E in longitude and from 35°N to 35°S in latitude with the main fishing grounds located in tropical countries, particularly Indonesia, Malaysia, and the Philippines (Motoh 1985, Figure 2).

The fry, juvenile, and adolescent inhabit shore areas and mangrove estuaries, while most of the adults inhabit deeper waters down to 162 m (Motoh 1985). Distribution is sparse as evidenced by a few prawns collected at any one time.

![Fig. 2. Geographic distribution of Penaeus monodon (Motoh 1981)](http://repository.seafdec.org.ph)
Reproduction

*P. monodon* is heterosexual. The female attains a relatively larger size than the male. The sexually mature prawn can be distinguished by the presence of the external genital organs: a joined petasma, a pair of appendix masculina on the exopods of the second pleopods, and a genital opening on the coxa of the fifth of pereopod for the male. In females, the thelycum is situated between the fourth and fifth pereopod with the genital opening on the coxa of the third pereopod.

The reproductive system of male and female *P. monodon* is shown in Figure 3. The following description is based on the studies of Motoh (1981, 1985).

**Male genital organ.** The internal reproductive organ of the male consists of paired testes, vasa deferentia, and terminal ampoules located in the cardiac region dorsal to the hepatopancreas. The testis is translucent and composed of six lobes, each connected in the inner margins leading to the vas deferens. The vas deferens consists of four portions, namely: the short narrow proximal vas deferens, a thickened larger median portion or the medial vas deferens, the relatively long narrow tube as the distal vas deferens, and the muscular portion referred to as terminal ampoule. The terminal ampoule contains the spermatophore and opens at the base of the coxopod of the fifth pereopods.

The spermatozoa of *P. monodon* are minute globular bodies composed of the head of about 3 microns in diameter and a short spike.

The petasma is a pair of endopods of the first pleopods formed by the interlocking hook-like structures. The appendix masculina is oval and is located on the endopod of the second pleopod.

**Female genital organ.** The internal reproductive organ of the female consists of paired ovaries and oviducts. The ovaries are bilaterally symmetrical, partly fused, and extend almost the entire length of the mature female. It is composed of the anterior lobe located close to the esophagus and the cardiac region of the stomach; the lateral lobes located dorsal to the
hepatopancreas; and the abdominal lobe which lies dorso-lateral to the intestine and ventro-lateral to the dorsal abdominal artery. The oviducts originate at the tips of the sixth lateral lobe and lead to the external genital opening at the coxopods of the third pair of pereopods.

The thelycum, located between the fifth pair of pleopods, consists of an anterior and a pair of lateral plates. It receives the spermatophores during mating. In penaeids, the thelycum may be classified as closed or open type, and *P. monodon* belongs to the closed type.

Motoh (1981) compared the detailed internal reproductive organs of *P. monodon* with those of *P. setiferus* and *P. indicus*.

**Sexual maturity.** Motoh (1981) defined sexual maturity as the minimum size at which spermatozoa are found inside the terminal ampoule of the males and inside the thelycum in the females. The later indicates that copulation or the transfer of spermatophores from the male to the thelycum of the female has taken place. On this basis, Motoh (1981) reported that wild *P. monodon* males possess spermatozoa at 37 mm carapace length (CL) (about 35 g body weight or BW) and females at 47 mm CL (about 67.7 mm BW) although pond-reared prawns were mature only at 31 mm CL (about 20 g BW) and 39 mm CL (about 41.3 g BW), respectively. Primavera (1980) reported the presence of spermatozoa in both pond-reared and wild *P. monodon* males of 40 g body weight (38.5 mm CL), a minimum of 63 weight (about 46 mm CL) for wild females, and about 40 g body weight (41 mm CL) for pond-reared prawns.

From the viewpoint of reproduction, Primavera (1985) emphasized the importance of gonadal maturation and the presence of fully developed spermatozoa with tail or spike. Motoh (1981) reported that sperms without tail were observed in wild *P. monodon* males of smaller size or about 37 mm CL, while Primavera (unpubl.) recently made mention of 10-month old pond-reared *P. monodon* with immature (spikeless) sperm.

**Ovarian maturation stages.** The maturation of the ovary has been categorized into five stages, the classification of which is based on ovum size, gonad expansion, and coloration (Villaluz et al 1969, Primavera 1980, Motoh 1981, Tan-Fermin and Pudadera, in press). Figure 4 illustrates the stages of ovarian development in *P. monodon*. 
Fig. 3. Reproductive system of *Penaeus monodon* (Motoh 1981)
ANTERIOR LOBE
LATERAL LOBE
OVIDUCT
ABDOMINAL LOBE

MATURE SPERMATOZOA
( x 200 magnification)

SPIKE

FEMALE (INTERNAL)

5 MICRONS

20 mm
Stage I and V (undeveloped and spent stages). Ovaries are thin, transparent, and not visible through the dorsal exoskeleton. Histological studies show that the ova averaging 36 microns are covered with a layer of follicle cells and the larger ones have nucleus and yolk granules (Motoh 1981). Tan-Fermin and Pudadera (in press) described Stage I as the perinuclear stage composed of perinuclear oocytes (46-72 microns) negatively stained with AB-PAS and Sudan Black. Oocytes bigger than 55 microns are enveloped by a single layer of follicle cells.
Similar features are observed in the spent stage which also contains some yolky oocytes, thicker follicle layer, or irregularly shaped perinucleolar oocytes (Tan-Fermin and Pudadera in press).

*Stage II (developing stage).* Referred to as early maturing stage, the ovaries are flaccid and white to olive green in color, and discernible as a linear band through the exoskeleton. The developing ova averaging 177 microns in diameter have yolk granules and cells believed to be nutritive bodies (Motoh 1981). The cells referred by Tan-Fermin and Pudadera (in press) as cystoplasmic inclusions are composed of small granules of glycoproteins, medium-sized globules of lipoglycoproteins, and few large lipid droplets.

*Stage III (nearly ripe stage).* Ovaries have glaucous color with the anterior portion thick and expanded. They are very visible through the exoskeleton, particularly at the first abdominal segment, when viewed against the light (Motoh 1981, Tan-Fermin and Pudadera, in press). The ova average 215 microns in diameter.

*Stage IV (ripe stage).* The ovary classified as ripe (mature) stage is diamond-shaped, expanding through the exoskeleton of the first abdominal segment. The isolated ovary appears dark olive green, filling up all the available space in the body cavity (Primavera 1980). Motoh (1981) reported the presence of a characteristic margin of peripheral rod-like bodies, the apexes of which radiate from the center of the egg. The ova average 235 microns in diameter. Tan-Fermin and Pudadera (in press) characterized this stage to consist mostly of yolky oocytes (288-408 microns) with additional rod-like bodies which contain acid and basic mucopolysaccharides but without lipids.

In some cases, ovaries are observed to be discontinuous, i.e., white in color in either the anterior or posterior portions with olive green color in the opposite ends. This condition is referred to as partially spent ovaries.

At present, these categories are used in the selection of wild spawners and prove to be generally effective. Prawns of Stage IV are used in hatchery operations. In the field, handling of the prawn for visual observation of the ovary color, size, and shape can not be avoided and can be stressful to the animal.
Fecundity. The number of eggs spawned varies according to the condition of the spawning female. Estimate of fecundity is mostly undertaken in the laboratory by counting the eggs from aerated spawning tanks.

For wild spawners of _P. monodon_, Motoh (1981) reported 248,000 to 811,000 eggs/spawn. Primavera (1985) mentioned that several researchers have observed lower fecundity. For captive and ablated females, fecundity ranges from 60,000 to 600,000 eggs/spawn because of small body size and uneven development of ovaries. Further detailed discussion on the subject is referred to Primavera (this volume).

In general, larger females produce more eggs than smaller females (Motoh 1981, Primavera 1985, Villegas et al 1986).

Morphological Development

Embryo. Eggs are spherical, yellowish green, and very minute, having a diameter ranging from 0.27 to 0.31 mm with an average of 0.29 mm. Eggs tend to sink slowly in still waters. Cleavage to 2-celled, 4-celled, morula, and embryonic nauplius stages occur approximately 0.5, 1, 1.8, and 11 hours, respectively, after spawning (Figure 5). The nauplius in each egg is observed to move intermittently before hatching (Villaluz et al 1969; Kunvankij 1976; Motoh 1979, 1981, 1985).

With significance in hatchery operations, Primavera and Posadas (1981) classified the eggs of _P. monodon_ based on morphological criteria and hatching rates.

Larva. The larval stage consists of 6 nauplius, 3 protozoea, 3 mysis, and 3 or 4 megalopa substages, requiring about 1.5 days, 5 days, 4-5 days, and 6-15 days, respectively, for development (Villaluz et al 1969; Kunvankij 1976; Motoh 1979, 1981, 1985). Figure 6 illustrates the larval stages of _P. monodon_. Larvae exhibit planktonic behavior offshore with antennal propulsion for swimming in nauplius, antennal and thoracic propulsion in mysis, and abdominal propulsion in megalopa. While the nauplii utilize yolk granules within their body, feeding starts in protozoea and mysis (collectively called zoea) substages. The megalopa with the earlier juvenile stage (traditionally called postlarva or "fry" for commercial purpose) is transparent with, dark brown streak on the ventral side tip of
the antennular flagellum to the tip of the telson. Under laboratory conditions, postlarvae become benthic on the sixth day of the post-larval stage. In natural conditions, the megalopa enters the nursery ground. The carapace length of megalopa varies between 1.2 and 2.3 mm.

**Juvenile.** The earlier juvenile stage has transparent body with dark brown streak on the ventral side as in the megalopa. Motoh (1985) described the earlier juvenile stages as follows: (1) relatively shorter sixth abdominal segment compared to the carapace length, (2) greater body size, (3) complete rostral spine formula, (4) complete gill system, and (5) benthic behavior.
Naupliar Substages
(Scales represent 0.2 mm.)

First nauplius, lateral (A) and ventral (B) views. A1, first antenna; A2, second antenna; En, endopod; Ex, exopod; Md, mandible; O, ocellus.

First nauplius, lateral (A) and ventral (B) views. A1, first antenna; A2, second antenna; En, endopod; Ex, exopod; Md, mandible; O, ocellus.

First nauplius, lateral (A) and ventral (B) views. A1, first antenna; A2, second antenna; En, endopod; Ex, exopod; Md, mandible; O, ocellus.

First nauplius, lateral (A) and ventral (B) views. A1, first antenna; A2, second antenna; En, endopod; Ex, exopod; Md, mandible; O, ocellus.
Fig. 6. Larval stages of *P. monodon* (Motoh 1981, 1985)
In the later stage, the body becomes blackish in color and bulky, and the rostrum has 7 dorsal and 3 ventral spines. The juvenile crawls using the pereopods and swims using the pleopods as in adults. The carapace length varies from 2.2 to 11.0 mm.

Motoh (1981, 1985) and Motoh and Buri (1980, 1981) have described the early postmysis stages of the giant tiger prawn.

**Adolescent.** This stage resembles the adult prawn. Sexes are now distinct beginning at 11 mm CL. The minimum size of males possessing a jointed petasma is about 30 mm CL and the minimum size of females possessing adultlike thelycum is about 37 mm CL. The carapace length of the adolescent varies between 11 and 34 mm.

**Subadult.** This stage is the onset of sexual maturity. The male possesses spermatozoa in its terminal ampoules. The thelycum of the female now contains spermatozoa. At this stage (30 mm CL), females grow faster and migration from nursery to spawning grounds begins. In the course of migration, first copulation takes place between males and females having a minimum of 37 mm and 47 mm CL respectively.

**Adult.** This stage has appendages very similar to the subadult and is characterized by the completion of sexual maturity. It differs only with the subadult in size and habitat. Males possess spermatozoa, and females start to spawn offshore although a few spawn in shallow water. A second or more copulations may occur in majority of the species. Major habitat is the offshore area up to about 160 m depth.

The maximum total length recorded was 336 mm (Holthuis 1980), while a mature female of 307 mm from Madagascar was reported by Crosnier (1965) as cited by Mohamed (1970) and 330 mm total length by Racek (1972). In the Philippines, the largest male ever found was 71 mm CL while the female was 81 mm CL with 270 mm body length or 240 g weight (Motoh and Buri 1980). Carapace lengths of adults vary between 37 and 71 mm in males and 47 and 81 mm in females.

The life history phases of the giant tiger prawn are summarized in Table 1, and the diagram of the life history is shown in Figure 7.
Longevity

There is no reliable method developed to determine the age of an individual prawn. Villaluz et al (1969) believed that the life span of *P. monodon* is one to two years; Motoh (1981) estimated it to be about one and a half years for males and about two years for females. Mohamed (1970) cited Srivatsa (1953) who reported that the life span of prawns (including *P. monodon*) in the Gulf of Kutch is 12-14 months.

Spawning

Spawning is the release of eggs and spermatozoa by the female prawn into the water for fertilization. The spermatophore which contains the spermatozoa is deposited in the female thelycum during copulation long before spawning. Although there is no report on the actual process observed in the natural condition, the spawning behavior of *P. monodon* has been documented based on laboratory observations (Villaluz et al 1969, Aquacop 1977, Primavera 1980, Motoh 1981). Discussion on spawning behavior is described in detail by Primavera (this volume).

In the Philippines, Villaluz et al (1969) reported that no *P. monodon* spawners below 50 mm CL have been collected in the Panguil Bay area, and concluded that first spawning occurs at 56 mm CL. However, Motoh (1981) reported that spawning females ranged from 47 to 81 mm CL and came in four size groups, namely: 48-50; 60-62, 66, and 72 mm CL. This finding indicates that *P. monodon* spawns four times in its life span and probably has multiple spawnings in a single season (Primavera 1980). In Orissa, India, Rajyalakshmi et al (1985) reported gravid *P. monodon* with size range of 100-250 g (about 54-76 mm CL) off the Paradip Coast.

Specific location of spawning area depends greatly on secondary evidence like the presence of abundant spawners and larval forms. In the Philippines, *P. monodon* spawns in the sea close to the coast (Delmendo and Rabanal 1956) or in the mouth of the bays with water depth of about 20 m but mostly spawns in offshore water to about 70 m (Motoh 1981). Hall (1962) calculated a more specific spawning area of *P. indicus* with *P. monodon* at about 18-36 m deep. In the Paradip Coast, Orissa *P. monodon* spawns at 30-40 m
Table 1. Life history phases of the giant tiger prawn, *P. monodon* (Motoh 1981)

<table>
<thead>
<tr>
<th>Phase</th>
<th>Begins at</th>
<th>Duration</th>
<th>Carapace length (mm)</th>
<th>Mode of Life</th>
<th>Habitat</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Male</td>
<td>Female</td>
<td></td>
</tr>
<tr>
<td>Embryo</td>
<td>Fertilization</td>
<td>12 hours</td>
<td>0.29*1</td>
<td></td>
<td>Planktonic Outer littoral area</td>
</tr>
<tr>
<td>Larvae</td>
<td>Hatching</td>
<td>20 days</td>
<td>0.5-2.2</td>
<td></td>
<td>Planktonic Outer/inner littoral area</td>
</tr>
<tr>
<td>Juvenile</td>
<td>Completion of gill system</td>
<td>15 days</td>
<td>2.2-11.0</td>
<td>Benthic</td>
<td>Estuarine area</td>
</tr>
<tr>
<td>Adolescent</td>
<td>Stability of body proportion,</td>
<td>4 months</td>
<td>11-30<em>2, 11-37</em>3</td>
<td>Benthic</td>
<td>Estuarine area</td>
</tr>
<tr>
<td></td>
<td>development of outer genitalia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Subadult</td>
<td>Start of sexual maturity, first</td>
<td>4 months</td>
<td>30-37<em>4, 37-47</em>5</td>
<td>Benthic</td>
<td>Inner/outer littoral area</td>
</tr>
<tr>
<td></td>
<td>copulation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adult</td>
<td>Completion of sexual maturity</td>
<td>10 months</td>
<td>37-71<em>6, 47-81</em>6</td>
<td>Benthic</td>
<td>Outer littoral area</td>
</tr>
</tbody>
</table>

*1 Egg diameter.
*2 Minimum size with jointed petasma.
*3 Minimum size with adult-like thelycum.
*4 Minimum size with spermatozoa in terminal ampoules.
*5 Minimum size with spermatozoa in thelycum.
*6 Maximum size ever found.
Fig. 7. Diagram of the life history of the giant tiger prawn, *P. monodon* (Motoh 1981)
In the Philippines, spawning of *P. monodon* is year-round but there seems to be two peak spawning seasons in a year: February-March or July and October-November although these vary from year to year (Motoh 1981). Hall (1962) reported February to April in Singapore; Rajyalakshmi et al (1985) in October through April corresponding to the post monsoon stability in the water movement and the increasing salinity in Orissa Coast, India; and Su and Liao (1986) from June to December in Taiwan.

Food and Feeding Habit

Hall (1962) generally considered penaeids to be omnivores with *P. monodon* in particular preferring crustaceans, vegetable matter, polychaetes, molluscs, fish, and insects. Thomas (1972) supported this finding and explained that mud and sand found in the gut were accidentally ingested. Villadolid and Villaluz (1951) reported that the fry stage of *sugpo* relishes plankton (*lablab*) food. Marte (1980) reported that *P. monodon* food also consisted of crustacea (small crabs and shrimps) and molluscs, making up 85% of ingested food. The remaining 15% consisted of fish, polychaetes, ophiuroids, debris, sand, and silt. This indicates that the giant tiger prawn is more of a predator of slow-moving benthic macroinvertebrates rather than a scavenger or detritus feeder. Kuttyama (1973) observed that debris composed of mud and organic matter constituted the main portion of the stomach content while crustaceans ranked next in quantity. Similar food items were also observed by Su and Liao (1986). All these findings suggest that *P. monodon* is more of a carnivore with preference for crustaceans particularly when in the natural environment, but it also feeds on other available organisms including algae.

*P. monodon* seems to have increased feeding activity during ebb tide (Marte 1980) and shows some food preferences during seasonal variations of food (Kuttyama 1973). This species feeds by seizing the food with its pinchers and pushing food to the mouth to nibble (Villadolid and Villaluz 1951). Undigested food is defecated four hours after ingestion (Marte 1980).
Physiology

*Molting.* Growth and the increase in size in crustacea are generally a function of the frequency of molting. Molting can occur anytime but more often at nighttime. Cited in the review by Wickins (1976), some conditions bring about a reduced increment during ecdysis so that the prawn may continue to molt but not grow.

During molting, the cuticle splits between the carapace and intercalary sclerite, paving the way for the cephalothorax and anterior appendage to withdraw, followed by the abdomen and posterior appendage emerging from the old shell with a forceful body flexure. The new cuticle takes about a few hours to harden in small prawns, to one or two days in larger animals (Villadolid and Villaluz 1951, Wickins 1976). Molting prawns characterized by soft shell are sensitive to stress and are good indicators of adverse environmental or nutritional conditions in the culture population (Wickins 1976).

Detailed reviews of the endocrine control of molting and reproduction in prawns were made by Adiyodi and Adiyodi (1970), Mantel and Farmer (1983), Adiyodi (1985), Kleinholz (1985), Skinner (1985), and Truchot (1983). Molting in crustaceans is believed to be controlled by two different hormones, namely: (1) molt-inhibiting hormone (MIH) secreted by the X-organ-sinus gland complex of the eyestalk (Kleinholz 1985, Skinner 1985). Molt-inhibiting hormone inhibits release of ecdysone by the molt gland. Closely associated with molting is reproductive maturation controlled by the gonad-inhibiting hormone (GIH) produced by the neurosecretory cells of the X-organ and transported to the sinus gland for storage and release. Induced gonadal maturation of penaeids through endocrine manipulation, such as eyestalk ablation, influences the molting cycle. Pudadera et al (1985) found marked changes in the internal structure of setae and cuticle throughout the molt cycle so that it is possible to properly time eyestalk ablation for induced ovarian maturation.

The physiological aspect of molting of *P. monodon* has received little attention. Although some works have been done on external factors, e.g., light, temperature, photoperiod, and salinity that may affect molting in penaeid prawns and shrimps (Bishop and Herrnkind 1976, Wickins 1976), no studies have been conducted on *P. monodon*. Meanwhile, Ferraris et al (1986)
reported a high degree of interaction between molting and salinity on osmotic and ionic regulation in *P. monodon*.

**Osmoregulation.** Crustaceans when subjected to change in water salinity have built-in mechanisms to adapt themselves to such change. It can be a mechanical response, such as burrowing, but the more efficient mechanisms are their physiological processes like osmoregulation by the organism. This process has been thoroughly reviewed by Mantel and Farmer (1983), Truchot (1983), Kleinholz (1985), and Skinner (1985). On the other hand, osmoregulation of *P. monodon* has been studied only by a few workers, e.g. Ferraris et al (1986) and Cheng and Liao (1986).

In the life history of *P. monodon*, spawning occurs in offshore waters where the larval stages are subsequently found. Post-larval, juvenile, and adolescent stages inhabit the nursery areas which are subjected to wide variations in salinity, temperature, and other environmental conditions. With proper acclimatization procedure, *P. monodon* postlarvae can also survive in fresh water (Pantastico and Oliveros 1980, Motoh 1981). The subadults and adults migrate to offshore areas where conditions are more or less stable. Behavior and survival are better understood, among others, in terms of the prawn's osmoregulation abilities. These physiological responses can be monitored by the changes in osmolality (or the number of particles such as ions, amino acids in solution) and the ion concentration of the blood (hemolymph) in relation to those of the medium (Ferraris et al 1986).

In the event when there is no osmotic gradient between the medium and the hemolymph, isosmotic point is attained. *P. monodon* in low salinity sea-water responds to osmotic gradient between the blood and external medium by gaining water, losing ions or both (hyperosmotic regulation) (Mantel and Farmer 1983). *P. monodon* juveniles are highly efficient regulators between 103 and 1480 mOsm/kg (30-50 ppt) and adults in over 444 mOsm/kg (about 15 ppt) (Cheng and Liao 1986). Isosmotic point for *P. monodon* juveniles of about 10 g is 730 mOsm/kg (Cheng and Liao 1986) or 676-720 mOsm/kg (26-28.5 ppt) (Ferraris et al 1986); for subadult (about 30 g weight or about 35 mm CL) 724-792 mOsm/kg (26-28.5 ppt) (Ferraris et al 1986); and for adults, 750 mOsm/kg (Cheng and Liao 1986). In general, *P. monodon* have isosmotic concentration at 20-30 ppt.
The isoionic points for _P. monodon_ are 352 mEq/l for sodium and 320 mEq/l for potassium (Cheng and Liao 1986); with Ferraris et al (1986) finding chloride between 324 to 339 mM in 10 g prawns.

Relating these findings to the distribution of _P. monodon_ in their natural habitat, Cheng and Liao (1986) attributed the abundant distribution of postlarvae to low salinity with due regard to other factors such as being genetically highly euryhaline species.

**Identification of Postlarvae**

Wild fry still support the seed requirements of the extensive culture operation. Inasmuch as there are many other _Penaeus_ fry during collection, fry gatherers and users often mistakenly identify _P. monodon_ fry with that of other penaeids. At present, scanty information has been found on the identification of _P. monodon_ fry. Some published articles on the subject are those by Rao and Gopalakrishnan (1969) for the juveniles of prawns in India, Prawirodihardjo et al (1975) on fry of _Penaeus_ in Indonesia and Motoh and Buri (1981) on fry along the shoreline of Panay Island in the Philippines.

Morphological characters and color (chromatophore or pigment patterns) are often used in the identification of penaeid postlarvae. The following morphological characters were used by Motoh and Buri (1981) to identify _Penaeus_ postlarvae along the coast of Panay Island, Philippines: (1) shape of rostrum, (2) number of rostral spines, (3) relative length of the antennular flagellum, (4) presence of antennal spine, and (5) presence of spinules on the sixth abdominal segment (6th AS). Chromatophore patterns on the 6th AS, telson, and uropods were also utilized, especially with fresh materials.

Compared with larvae and postlarvae of known parentage and reared in the laboratory, Motoh and Buri (1981) distinctly separated the postlarvae of _P. monodon_ from _P. semisulcatus_, whereas with the absence of distinct differences in morphological characters for species identification, _P. indicus_ and _P. merguiensis_ were combined as a group, and _P. japonicus_, _P. latisulcatus_, and _P. longistylus_ as another group.
P. monodon. The postlarvae of P. monodon are the largest among the species or groups. The body is slender and the modal CL is 2.6 mm. The rostrum is either straight or slightly bent upward at the tip, usually having five dorsal spines but devoid of ventral spines. The inner (lower) antennular flagellum is twice lower than the outer (upper) flagellum. The 6th AS does not have spinules (Motoh and Buri 1981).

Distinct with fresh postlarvae, chromatophores of dark brownish red extend from the tip of the inner antennular flagellum to the tip of the telson. When viewed microscopically, thirteen chromatophores align ventrally along the 6th AS or are densely distributed to form an almost continuous pattern (Prawirodihardjo et al 1975, Motoh and Buri 1981). The chromatophore on the antero-lateral margin of the 6th AS is absent.

P. semisulcatus. The postlarvae of this species are relatively small. The rostrum is usually bent upward and has six dorsal spines and one or no ventral spine. The inner antennular flagellum is about one and a half to two times longer than the other flagellum. The antennal spine is very small or absent (Motoh and Buri 1981).

Chromatophores are not so dense, numbering from six to twelve along the 6th AS. Only the base and the tip portions of the telson and uropods are pigmented (Prawirodihardjo et al 1975, Motoh and Buri 1981). One chromatophore is present at the antero-lateral margin of the 6th AS.

Motoh and Buri (1981) confirmed the similarities between the postlarvae of P. monodon and P. semisulcatus in the Philippines and that of the specimens from Indonesia.

P. merguiensis group. The postlarvae of this group, composed of P. merguiensis and P. indicus are the smallest among the other species or groups. The rostrum is long or one and one-half times larger than the CL. About two-thirds of the anterior portion of the rostrum is toothless both dorsally and ventrally, but the remaining portion has three to four dorsal teeth and none below. Antennal spine is absent.

In older postlarvae, the rostrum becomes more bent upward having six to seven dorsal and three to five ventral teeth. As in its adult form, the postlarvae are poorly pigmented,
hence, the species are referred to as white prawns. One prominent chromatophore at the antero-lateral margin of the 6th AS is present (Motoh and Buri 1981).

*P. japonicus* group. The postlarvae of this group represents *P. japonicus*, *P. latisulcatus*, and probably *P. longistylus*. The body is short and bulky. The rostrum is short and does not exceed the tip of the eye. There are five to seven dorsal but no ventral teeth. The inner flagellum is about one and one half longer than the length of the outer flagellum. The carapace has a modal length of 2.0 mm. These postlarvae are often mistaken as smaller post-larvae of *P. monodon* because they have similar longitudinal streaks of dark brown chromatophores. The chromatophores at the 6th AS are more than eight or sometimes countless and are usually absent at the antero-lateral margin (Motoh and Buri 1981).

Chromatophore patterns of different *Penaeus* species shown in Figure 8 and the chromatophore patterns on the 6th AS, telson, and uropods shown in Figure 9 can aid in quick identification of each species or group.

![Chromatophore patterns for quick identification](image)

*Fig. 8. Dorsal view of postlarval Penaeus showing chromatophore patterns for quick identification. (A) P. monodon (B) P. semisulcatus (C) P. merguiensis group, (D) P. japonicus group. Scales represent 2.5 mm (Motoh 1981)*
Fig. 9. Dorsal view of the sixth abdominal segment, telson, and uropods of postlarval *Penaeus* showing chromatophore patterns. (A) *P. monodon* (B) *P. semisulcatus* (C) *P. merguiensis* group, (D) *P. japonicus* group. Scale represents 1 mm (Motoh 1981)

The following is the key prepared by Motoh and Buri (1981) to identify post-larval *Penaeus* appearing at the shoreline of Panay Island, Philippines.

**Key to postlarval *Penaeus* appearing at shore waters, based on morphological features**

1) Rostrum stout and inferior to tip of eye, spinules on the sixth abdominal segment present*, antennal spine prominently present, carapace slightly longer than sixth abdominal segment. ........................................... *P. japonicus* group.

Rostrum slender and exceeding tip of eye, spinules on the sixth abdominal segment absent, antennal spine absent or minute, carapace slightly or distinctly shorter than the abdominal segment. ........................................... 2.

2) Inner (lower) antennular flagellum nearly 1.6 times the outer (upper), exceeding the latter by its distal one segment. ........................................... *P. merguiensis* group.

Inner antennular flagellum 1.6 to 2.0 times the outer (upper) exceeding the latter by its distal two segments. ........................................... *P. semisulcatus*.

Inner antennular flagellum more than 2.0 times the outer, exceeding the latter by its distal three segments. ........................................... *P. monodon*.

*When the number of rostral teeth is less than four, the spinules are sometimes poorly present or absent. In this case, other criteria are useful.
Key to the postlarval *Penaeus* appearing at shore waters, based on chromatophore patterns

1) Number of chromatophores on the sixth abdominal segment less than seven. Antero-lateral chromatophore of the sixth abdominal segment present

*P. merguiensis* group.

Number of chromatophores on the sixth abdominal segment more than seven.

Antero-lateral chromatophore of the sixth abdominal segment present or absent

2) Number of chromatophores on the sixth abdominal segment less than 12. Antero-lateral chromatophore of the sixth abdominal segment present, chromatophores on the middle portion of telson and inner uropods absent

*P. semisulcatus*

Number of chromatophores on the sixth abdominal segment more than 12, antero-lateral chromatophore of the sixth abdominal segment absent. Chromatophores on the middle portion of the telson and inner uropods present

3) Chromatophores on the sixth abdominal segment dense and thickly continuous

*P. monodon*.

Chromatophores on the sixth abdominal segment discontinuous or confluent

*P. japonicus*.

Common to all workers is the use of morphological characters and color patterns for identification, with Motoh and Buri (1981) also using morphometric measurements in identifying *P. monodon* from *P. semisulcatus* and two other groups, *P. merguiensis* and *P. japonicus*.

Motoh and Buri (1981) confirmed the similarities between the post-larvae of *P. monodon* and *P. semisulcatus* in the Philippines and Indonesia.

For detailed descriptions, Motoh and Buri (1981) provided a key with illustrations while Prawirodihardjo et al (1975) tabulated the differences between *P. monodon* and *P. semisulcatus*.
PROBLEMS AND PROSPECTS

Findings on biology and ecology of *P. monodon* have contributed significantly to its aquaculture. What was before an extrinsic species in the brackishwater pond culture of milkfish is now a cash crop of great demand in intensive monoculture. Fry collection from the wild has intensified to meet the demand for seeds usually for extensive culture while improvement in hatchery techniques has resulted in the proliferation of commercial hatcheries supplying the seeds for intensive culture. Despite these advances, a myriad of problems remain.

First, there is a conflict in the use and management of natural resources. The estuarine areas, rivers, and mangroves which are considered nursery grounds of wild *P. monodon*, are often identified for fishpond development or reclamation for social, commercial, or industrial purposes. This is a complex problem requiring concerted efforts of intergovernmental agencies to formulate and to religiously implement policies on the conservation of natural resources.

Second, there is no conservation effort during collection from the wild of *P. monodon* fry where finfish fry and larvae, other penaeid fry, and crustaceans of potentially high economic value are often discarded indiscriminately. Measures should be taken to return the live fry or larvae of other organisms to the shorewater. Policies on proper management and collection of wild fry should be adopted.

Third, overexploitation of wild stock, especially the spawners of *P. monodon*, is apparent. Stock assessment studies are thus necessary to pave the way for sea ranching open-water stocking in certain protected areas to replenish wild stocks.

For aquaculture scientists, the physiological aspects of the biology of *P. monodon* remain a major area for future research. More baseline information on molting is needed. The manipulation of hormonal and environmental conditions that control frequency and time of molting could improve growth and survival in culture, reproductive maturation for broodstock, and lastly, artificial fertilization techniques beneficial for genetic stock improvement and hybridization studies.
LITERATURE CITED


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