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Freshwater Bryozoa of Tonle Sap, Cambodia

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We identified a collection of freshwater bryozoa from Tonle Sap (meaning Tonle Lake), Cambodia, a body of water fed by the Mekong River and characterized by extreme fluctuations in water level between the wet and dry seasons. The collection also included specimens from the moat of Angkor Wat, located at the north end of the lake. We found four phylactolaemate species (Plumatella bombayensis, Plumatella casmiana, Plumatella vorstmani, Hyalinella lendenfeldi) and one ctenostome species (Hislopia cambodgiensis) from the lake, and only a single, additional phylactolaemate species (Plumatella javanica) from the moat. We provide brief descriptions of these species, photographs of colonies for some, and photomicrographs by light and scanning electron microscopy (SEM) of statoblasts. None of the species encountered in this study is endemic to Cambodia, and the wide distributions of the species are possibly related to the dispersability of floatoblasts by birds. We briefly discuss some of the taxonomic problems surrounding Hislopia cambodgiensis.

Key words: biodiversity, freshwater Bryozoa, Phylactolaemata, Ctenostomata, Cambodia, Tonle Sap

INTRODUCTION

Bryozoans are benthic colonial animals, mostly sessile, that form colonies of various morphologies by asexual budding. About 6,000 living species have been described from marine environments (Gordon, 1999), and about 65 species from freshwater (Wood and Wood, 2000; Wood and Okamura, 2005). Modern bryozoans are divided into three classes: Gymnolaemata, Stenolaemata, and Phylactolaemata. Most freshwater bryozoans belonging to Class Gymnolaemata, although a few species of the gymnoalaemate order Ctenostomata also occur in freshwater.

Phylactolaemata differ from the other classes in several respects, including a U-shaped rather than a circular feeding lophophore. The colonies of most species grow in a branched tubular morphology or as a compact gelatinous mass. Unlike other bryozoans, phylactolaemata form encapsulated, asexually produced dormant bodies called statoblasts. Statoblasts are generally resistant to cooling and drying, and germinate under favorable conditions following an obligate period of dormancy, in response to a variety of environmental cues including changes in water temperature and light (Rogick, 1938, 1941a; Oda, 1959, 1979; Mukai, 1974; Bushnell and Rao, 1974). There are two types of statoblasts; one is a free-floating buoyant statoblast called a "floatoblast," and the other is a sessile statoblast, or "sessoblast." Many species can form both types. The morphological features of statoblasts are the most important diagnostic characters for the identification of phylactolaemate species.

Most of the approximately 260 described species of living ctenostome bryozoans inhabit the sea, but at least 12 species worldwide in the genera Hislopia, Sineportella, Paludicella, Pottsiella, and Victorella are known from freshwater habitats. Hislopia is the most speciose genus of ctenostome bryozoans occurring in freshwater; seven species have been reported worldwide. However, there have been few detailed studies of Hislopia species, and the existing taxonomical information is generally insufficient. For example, Jullien (1880) originally described Hislopia cambodgiensis from the Mekong River, but this species has not been reported since 1916, and the paucity of information in the original description makes it difficult to identify specimens as this species with any certainty.

Freshwater bryozoans were studied in Southeast Asia mostly from the late nineteenth century to the early twentieth century (Jullien, 1880; Kraepelin, 1906; Annandale, 1908, 1910, 1911, 1916a, b, 1919; Vorstman, 1928a, b). Recently, there has been renewed interest in freshwater bryozoans in the region (Wood and Wood, 2000; Wood et al., 2006). However, only one study (Jullien, 1880) has included freshwater bryozoans from Cambodia.

From 2003 to 2005, Japanese and Cambodian scientists (S. Tsukawaki, A. Ohtaka, H. Katakura, T. Kamiya, Y. Hirabuki, S. Endoh, Y. Okumura, S. Sotham, A. Cheouleu) collaborated in research to evaluate mechanisms for sustaining biodiversity in Tonle Sap, Cambodia (S. Tsukawaki, pers. comm.). These biologists collected several freshwater bryozoan species from the lake and from the moat of Angkor Wat. Here we present brief descriptions of these species and discuss the life cycle and ecology of the freshwater bryozoans in Tonle Sap, to the extent permitted by our limited data.

MATERIALS AND METHODS

Study area

Tonle Sap (= 12°50'N, 104°06'E; surface elevation = 10–20 m asl), located in the mid-western part of Cambodia, is the largest...
natural lake in Indochina (Fig. 1). Among all lakes in the world, this lake shows the greatest change in water level between wet and dry seasons. In the dry season (November to April), it is about 3,000 km² in area and only about 1 m in maximum depth. In the wet season (May to October) (Fig. 2), the lake is fed by a reverse flow of the Mekong River via the small Tonle Sap River. The area of the

![Map of Southeast Asia](image1)

**Fig. 1.** Map of Southeast Asia showing the location of Tonle Sap in Cambodia and the extent of the lake during the dry season. The Tonle Sap River connects the lake with the Mekong River (heavy line).

![Photograph of Tonle Sap](image2)

**Fig. 2.** Photograph of Tonle Sap during the wet season. The bushes shown here are completely emergent during the dry season, when the water level is about 10 m lower than in the wet season.
lake increases to about 16,000 km$^2$, over four times that in the dry season, with a maximum change in depth of about 10 m (CNMC/NEDECO, 1998). As a consequence of the changes in water level, bushy vegetation around the lake that is submerged in the wet season, and provides a substratum for freshwater bryozoans, dries up in the dry season. Situated in the tropics, Tonle Sap shows little annual variation in temperature; the annual range is about 26–30°C (Endoh et al., 2002). At the north end of Tonle Sap is located the ancient temple of Angkor Wat, which is surrounded by a moat about 190 m across.

Collection and treatment of specimens

Prof. A. Ohtaka collected bryozoans from the moat of Angkor Wat in 2002 and at two localities in Tonle Sap from 2003–2005; the only substrates on which he found specimens were submerged branches and the roots of water hyacinths. Specimens were initially fixed in 10% formalin, except for *Hislopia* specimens collected from the lake in 2005, which were fixed in 90% ethanol. Statoblasts were dehydrated with ethanol and HMDS (hexamethyldisilazane), coated with Pd-Pt in a Hitachi E-1030 sputter-coater, and observed with a Hitachi S-2380N scanning electron microscope (SEM) at 15 kV accelerating voltage. Ctenostome bryozoans were dehydrated with butanol, embedded in paraffin, and sectioned at a thickness of 7 μm. Sections were stained with hematoxylin-eosin and observed by light microscopy to measure the size of the gizzard and count the number of tentacles.

RESULTS

In all, we identified six species of freshwater bryozoans, including four species of phylactolaemates and one ctenostome in Tonle Sap and an additional phylactolaemate species in the moat of Angkor Wat.

Taxonomy

**Class PHYLACTOLAEMATA** Allman, 1856

**Family Plumatellidae** Allman, 1856

**Genus Plumatella** Lamarck, 1816

*Plumatella bombayensis* Annandale, 1908

(Fig. 3)

*Plumatella bombayensis* Annandale, 1908, 169–171, Figs. 1, 2.


*Plumatella longigemmis*: Lacourt, 1968, 73, 75 (in part).

*Plumatella tanganyikae*: Annandale, 1911, 225–227, Fig. 44 (in part).

Material examined. A few statoblasts were collected from Baribour Province in the southern part of Tonle Sap, December 2004.

Description. Colony dark brown, with well-chitinized ectocyst (Wood and Okamura, 2005). Floatoblast elongate, with a large fenestra on ventral valve. Annulus especially broad at both ends. Dorsal valve almost flat; fenestra is smaller than on ventral valve. In young sessoblasts, surface of fenestra is reticulate. With age, tubercles appear in each cell of the reticulation, and in mature sessoblasts (Fig. 3A, B) tuberculation almost entirely obscures the reticulation, which is evident only near the annulus (Fig. 3B).

Remarks. Annandale (1908) originally described *P. bombayensis* from Igatpuri, near Bombay, India. Lacourt (1968) regarded *P. bombayensis* as a synonym of *P. longigemmis*. However, compared to *P. longigemmis*, the floatoblast of *P. bombayensis* has a much wider annulus on the ventral valve, a smaller dorsal fenestra, and a different microstructure on their fenestrae (lightly tuberculate on dorsal valve and strongly reticulate on ventral in *P. bombayensis*; tuberculate on both valves in *P. longigemmis*).

Distribution. *P. bombayensis* has been reported from throughout southern Asia and at a single, artificially heated site in Belarus (Wood and Okamura, 2005).

*Plumatella casmiana* Oka, 1907

(Fig. 4)

*Plumatella casmiana* Oka, 1907, 121–123, Fig. 3.

*Plumatella casmiana*: Behning, 1924, 172; Vorstman, 1928a, 7–8, Fig. 5, Pl. 1, Fig. 8; Rogick, 1941b, 211–219, Pl. 1, Figs. 1–4, Pl. 2, Figs. 5–6; 1943, 265–270, Pl. 1, Figs. 1–8; Lacourt, 1968, 52–56, Pl. 13, Figs. g–l, Pl. 15, Fig. a; Rogick and van der Schalie, 1950, 142–144, Figs. 6–12; Torumi, 1955, 76–77, Figs. 1–3; Sebestyen, 1961, 127–131, Figs. 3–8; Wiebach, 1983a, 581–595, Figs. 1–4; 1963b, 77–78, Abb. 1–5; Bushnell and Wood, 1971, 229–230, Fig. 1; Wood, 1979, 59, 61, 68–69, Pl. 1, Figs.

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Fig. 3. *Plumatella bombayensis* Annandale. Scanning electron micrographs; (A) mature sessoblast; (B) enlargement of a mature sessoblast showing that some reticulation remains evident near the annulus.
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7–8, Pl. 3, Figs. 4–5; 1989, 28–31, Fig. 15; Wood and Wood, 2000, 424, 426; Wood and Okamura, 2005, 46, 78–80, Figs. 16–18, 45–46; Wood et al., 2006, 12, Figs. 11–13.

\textit{Plumatella casmiana} Abрикосов, 1927, 309.

\textit{Plumatella repens} Loppens, 1908, 159 (in part); 1910, 145 (in part).

\textit{Plumatella repens} var. \textit{annulata} Hozawa and Toriumi, 1940, 428–429, Fig. 3 (A–D); 1941, 236–237, Fig. 3, Pl. 14, Fig. 2; Rogick and Brown, 1942, 131–132, Pl. 2, Figs. 10, 14–16, 26, Pl. 3, Fig. 30.

\textit{Plumatella repens} var. \textit{casmiana} Toriumi, 1941a, 203–204, Fig. 7, Pl. 12, Figs. 4–5, Pl. 13, Fig. 15; 1942, 209–211, Figs. 1–3.

\textit{Plumatella repens} var. \textit{flabellum} \textit{type casmiana} Toriumi, 1941b, 418–419, Figs. 5–7.


\textit{Plumatella annulata} Lacourt, 1955, 88, Pl. 5 (bottom center and bottom right).

\textbf{Material examined.} Specimens were collected from the northern part of Tonle Sap in November 2003. These included a few small, brownish, translucent colonies found on roots of the water hyacinth, \textit{Eichhornia crassipes} (Mart.) Solms. Statoblasts were not observed.

\textbf{Description.} Colony light brown and often fouled. Colony varies in form from having zooids in branched uniserial series (Fig. 4) to having zooids crowded in a compact mass; compact colonies sometimes assume a honeycomb-like shape (Hozawa and Toriumi, 1940; Bushnell and Wood, 1971). Tentacle number about 24. \textit{Plumatella casmiana} produces two types of free floatoblasts: the usual type of plumatellid floatoblast (capsuled floatoblast) with well a developed periblast, and a type (leptoblast) lacking the internal capsule. The leptoblast has a very narrow annulus and germinates immediately after release from the colony. The capsuled floatoblast has weak tubercles on surface of fenestra; these are sometimes slight, with the surface almost smooth. The sessoblast has a wide annulus and the surface of the capsule is smooth.

\textbf{Remarks.} Oka (1907) originally described \textit{P. casmiana} from Kasumigaura, Japan, and Rogick (1941b, 1943) showed detailed drawings of colony morphology and statoblasts from North America. Lacourt (1948) regarded \textit{P. casmiana} as a synonym of \textit{P. philippinensis} on the basis of floatoblast morphology. However, the floatoblasts and leptoblasts of these two species differ in features of the periblast and the capsule. The weakly tuberculate surface of the capsulated floatoblast is diagnostic for \textit{P. casmiana}.

\textbf{Distribution.} This is one of the most common and widely distributed species worldwide (Bushnell, 1973; Wood et al., 2006).

\textit{Plumatella javanica} Kraepelin, 1906 (Figs. 5, 6)

\textit{Plumatella javanica} Kraepelin, 1906, 143–146, Figs. 1–3.

\textit{Plumatella javanica}: Annandale, 1910, 42, 50; 1911, 221–223; Toriumi, 1952, 267–269; Wiebach, 1964, 12, Figs. 4–8, Taf. 1, Fig. 4, Taf. 2, Figs. 5–7, Taf. 3, Figs. 8–12, Taf. 4, Fig. 13; 1967, 177–178, Taf. 1, 2, Abbs. 1–8; 1970, 356, Taf. 1, Abbs. 4a, 4b, 5a, 5b; Lacourt, 1968, 72–73, Pl. 15, Fig. j; Smith and Wood, 1995, 362–365.

\begin{figure}[h]
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\caption{\textit{Plumatella casmiana} Oka. Light micrograph showing a portion of a colony attached to the roots of the water hyacinth, \textit{Eichhornia crassipes} (Mart.).}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig5.png}
\caption{\textit{Plumatella javanica} Kraepelin. Light micrograph showing a portion of a colony attached to a branch under the water.}
\end{figure}
Fig. 6. *Plumatella javanica* Krajev. Photomicrographs of floatoblasts taken by light (A, B) and scanning electron (C–G) microscopy; (A) ventral valve; (B) dorsal valve; (C) ventral valve; (D) dorsal valve; (E) enlargement from (C) showing textures of annulus and fenestralula; (F) enlargement from (D) showing textures of annulus and fenestra; (G), higher magnification enlargement from (C) showing tubercles surrounded by rounded-triangular pores in the angles between tubercles.
Material examined. Several light-gray colonies with some floatoblasts were collected from the south side of the moat of Angkor Wat in October 2002.

Description. Colony is light brown or light gray, its surface is heavily fouled and encrusted. Colony almost entirely recumbent, with few branches and with a pronounced keel; ends of the branches often rising from the substratum (Fig. 5). Ectocyst of new buds sometimes transparent. Tentacle number about 45. Floatoblast elliptical; large, about 400 $\mu$m long (Fig. 6A, B). Fenestrae of floatoblast large (Fig. 6A–D); in SEM view, surface entirely covered with minute, rounded tubercles (Fig. 6C–F) rounded-hexagonal in outline, with 5–8 (average, 6) small rounded-triangular pores in the angles (Fig. 6G). Annulus of floatoblast covered with coarser, more irregular tubercles (Fig. 6F). Floatoblast valves asymmetrical in lateral and transverse views, the ventral valve strongly convex and dorsal valve almost flat. Sessoblast also with tuberculate surface.

Remarks. Kraepelin (1906) originally described \textit{P. javanica} from Indonesia. Previous authors have considered statoblast morphology, colony color, and colony form as important diagnostic characters of this species. Annandale (1911) reported a transparent, glassy ectocyst and a strong raphe, and Smith and Wood (1995) also indicated these characters. However, these features vary ecophenotypically, as with other plumatellid species. Annandale (1911), for example, specifically mentioned variability in the form of zoaria attached to dead wood. Like \textit{P. javanica}, \textit{P. vorstmani} (following description) also has a transparent ectocyst, but the tentacle number in \textit{P. javanica} is almost twice that in \textit{P. vorstmani}, and statoblast size and morphology are different.

Distribution. This species occurs in tropical regions (Bushnell, 1973), including Southeast Asia (Lacourt, 1968; Wood \textit{et al.}, 2006), central Africa (Wiebach, 1964), and the Amazon River in South America (Wiebach, 1967, 1970).

\textit{Plumatella vorstmani} (Toriumi, 1952)

(Fig. 7)

\textit{Plumatella vorstmani}: Mukai, 1984, 51–52, Fig. 2; Wood and Wood, 2000, 423, 425, 427, Fig. 5; Wood \textit{et al.}, 2006, 18–19, Figs. 28, 29, 39.
\textit{Plumatella javanica}: Vorstman, 1928a, 6–7, Fig. 4, Pl. 2, Figs. 6–7; 1928b, 163.
\textit{Hyalinella vorstmani}: Lacourt, 1968, 86–87, Pl. 12, Figs. g, i; Wiebach, 1973, 546.

Material examined. Only a few floatoblasts were collected from Baribour Province in the southern part of Tonle Sap in December 2004.

Description. According to Toriumi (1952), Mukai (1984), and Wood \textit{et al.} (2006), the colony is transparent, almost entirely recumbent; the zooids are never fused with one another; and tentacle number is 20–27. Floatoblast small, about 300–350 $\mu$m long by 180–220 $\mu$m wide (Fig. 7A–D); symmetrical in lateral view; with tubercles on fenes-

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Fig. 7. \textit{Plumatella vorstmani} (Toriumi). Photomicrographs of floatoblasts taken by light (A, B) and scanning electron (C, D) microscopy; (A) ventral valve; (B) dorsal valve; (C) ventral valve; (D) dorsal valve.
trae (Fig. 7C, D). Sessoblast with tubercles on surface of capsule and reticulation on annulus (Toriumi, 1952). The sessoblast is shorter than the floatoblast (Toriumi, 1952; Mukai, 1984).

Remarks. Toriumi (1952) recognized *P. vorstmani* as a new species from material that Vorstman (1928a) had reported as *P. javanica*, on the basis of differences from *P. javanica* in tentacle number, floatoblast size, and the surface sculpturing of the sessoblast. Toriumi reported tubercles on the sessoblast. However, Wood and Wood (2000) reexamined specimens that Mukai (1984) had identified as *P. vorstmani* and found the surface of the capsule to be reticulate rather than tuberculate. Wood et al. (2006) reported *P. vorstmani* from many sites in Thailand as having sessoblasts with tuberculate fenestrae, and the features of the floatoblast are completely compatible with those from Tonle Sap. Uncertainties in the diagnosis of this species are due to equivocal specimens from Japan. Toriumi’s and Mukai’s phylocladomate collections contain both tuberculated and reticulated sessoblasts identified as those of *P. vorstmani* (Hirose, unpublished data), and this material possibly includes more than one species.

Lacourt (1968) referred *P. vorstmani* to the genus *Hyalinella*. However, this species fits better in *Plumatella* in having sessoblasts and zooloids that are loosely attached to one another.

Distribution. *Plumatella vorstmani* has been reported from Asia (Bushnell, 1973; Wood et al., 2006), Japan (Toriumi, 1952; Mukai, 1984), India (Lacourt, 1968), and Java, Indonesia (Vorstman, 1928a, b).

*Hyalinella lendenfeldi* (Ridley, 1886) (Figs. 8, 9)


*Plumatella punctata* phase prostrata Vorstman, 1928a, 10, Fig. 7.

Material examined. Three large colonies with many floatoblasts were found on submerged branches of *Persicaria hydropiper* in the northern part of Tonle Sap, November 2003.

Description. Colony tubular, branched, forming extensive thin patches on the substratum; ectocyst transparent, mature colonies thicker than young ones, the branches surrounded by a gelatinous ectocyst about 1–3 mm thick (Fig. 8). *Hyalinella lendenfeldi* produces only floatoblasts (Fig. 9A–D). Floatoblast is oblong, the ends rounded, the margins of the long axis parallel; 670–690 μm long by 410–420 μm wide, much larger than the floatoblasts of other plumatellid species. Annulus very wide relative to fenestra size. Ventral fenestra about 280 μm long by 215 μm wide; dorsal fenestra smaller than ventral, about 162 μm long by 110 μm wide. SEM images show fenestrae of both valves covered with fine, round tubercles (Fig. 9E). Annulus sculptured with closely set tubercles coarser than those of fenestra. Ventral valve convex; dorsal valve flat or concave.

Remarks. Ridley (1886) originally described this species from Australia as *Lophopus lendenfeldi*. Rousselet (1904) referred it to *Hyalinella* because of the tubular colony, the soft, gelatinous ectocyst, and the absence of sessoblasts. Annandale (1910) established the genus *Australella* to which he transferred *H. lendenfeldi* and in which he included another species, *A. indica*. However, this genus is not clearly distinct from *Hyalinella* (Wood, 1998). Well-developed colonies of *H. lendenfeldi* resemble those of *Gelatinella toanensis* (Hozawa and Toriumi, 1940) reported from East Asia, but the floatoblast of *G. toanensis* is quite different from that of *H. lendenfeldi* in size and in having a spine on the ventral fenestra. Moreover, *G. toanensis* produces sessoblasts, whereas *H. lendenfeldi* does not.

Distribution. There is a single record of *Hyalinella lendenfeldi* from Australia (Ridley, 1886); the species has been reported as abundant in Thailand, with records from a lake at Chiang Mai and from Vachiralongkorn Reservoir (Wood et al., 2006).

Class GYMNOCLAEMATA Allman, 1856

Order Ctenostomata Busk, 1852

Family Hisliopidae Jullien, 1885

Genus HISLIOPIA Carter, 1858

*Hislopia cambodgiensis* (Jullien, 1880) (Fig. 10)

*Norodonia cambodgiensis* Jullien, 1880, 77–78, Figs. 1–3.

*Norodonia cambodgiensis*: Jullien, 1885, 181–182, Figs. 244–245; Anonymous, 1888, 102, Figs. 17–18; Loppens, 1908, 176, Figs. 28–29.

*Hislopia cambodgiensis* Annandale, 1916a, 34–35, Pl. 1, Figs. 8; Hondt, 1983, 19, 21, Fig. 9 (a, c).

*Norodonia sinensis* Jullien, 1880, 78–79, Figs. 1–3.
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Norodonia sinensis: Jullien, 1885, 182, Figs. 246–247; Anonymous, 1888, 102, Figs. 19–20; Loppens, 1908, 177, Fig. 30.


Material examined. Eight colonies were collected from the roots of Eichhornia crassipes, northern part of Tonle Sap, November 2003 and December 2005, and from Barbour Province in the southern part, December 2004.

Description. Colony encrusting, spreading broadly on the substratum, the zooids arranged in straight, branching uniserial series; columns of zooids distinct even in crowded colonies (Fig. 10). Zooids light- or dark-yellowish brown, with a transparent ectocyst; oval to almost circular in shape, tapering proximally, about 0.80–0.93 mm long by 0.60–0.68 mm wide, on average 1.18 times as long as wide. Tentacle number about 16. Orifice is round or rounded-quadrate; oral spines lacking. Gizzard about 150 μm long by 150 μm wide, with thick lining. Stomach about 280 μm long by 140 μm wide. The primordium of new autozooidal buds begins from a long, narrow, flattened peduncle that gradually expands before differentiation of the polypide. Ancestrula smaller than astogenetically mature zooids, giving rise to several
Remarks. Jullien (1880) originally described *Norodonia cambodgiensis* from the Mekong River of northeastern Thailand and also interior Cambodia; however, his description was limited. Annandale (1916a) reported what he identified as Jullien’s species from Tai-Hu (Great Lake), China and provided a detailed description; he transferred the species to *Hislopia* because of similarities in colony and zooid form to *Hislopia lacustris*, the type species of *Hislopia*.

Jullien (1880) described a similar species, *Hislopia sinensis* (as *Norodonia sinensis*), from the Ngan-Honi Province, southern China. He indicated the key diagnostic character of *H. sinensis* as being the tapering axial keel on the zooidal frontal wall, arising from the proximal lip of the orifice and extending posteriorly for a third to one-half the length of the zooid. However, Jullien observed only dried specimens. The frontal wall of *Hislopia* is weak and readily shrinks when the colony dries; the central area then appears raised above the gut and other internal organs. The main diagnostic feature of *H. sinensis* is thus perhaps an artifact of drying. *Hislopia sinensis* is very similar to *H. cambodgiensis* in colony form and other respects, and Annandale (1916a) regarded the two as synonymous. The holotype specimen of *H. sinensis* has been lost.

Annandale (1916a) described from Thailand *H. malayensis*, which is quite similar to our *H. cambodgiensis*. Both species have zooids about 0.85–0.90 mm long, and *H. malayensis* has about 16 tentacles. The only difference appears to lie in the budding form of new autozooids. The bud of *H. malayensis* is different from that of all other *Hislopia* species; it has been reported to arise as a slender, blunt-ended, flattened cylinder that reaches almost the full zooid length before expanding at the tip as it gradually assumes the oval shape of the adult zooid (Annandale, 1916a; Hondt, 1983; Wood et al., 2006). However, the form of buds might vary with growth rate or environmental conditions, and the taxonomic reliability of this character is questionable. Considering the original description of *H. malayensis* and the similar distributions of Wood’s *H. malayensis* and our *H. cambodgiensis*, these species are possibly synonymous. If this proves to be the case, the latter name will have priority as the earlier description.

Distribution. *Hislopia cambodgiensis* has been reported from Cambodia; the Mekong River on the border of Cambodia and Siam (Thailand); and Canton, Ngan-Honi, and Tai-Hu (Great Lake), China (Jullien 1880; Annandale, 1916a).

**DISCUSSION**

Tonle Sap undergoes great fluctuations in depth and surface area between the wet and dry seasons. Bryozoan colonies that grow on the branches of bushes sunken during the wet season dry out and die (except for their statoblasts) during the dry season. Colonies were also found attached to the water hyacinth (*Eichhornia crassipes*), plants of which can drift to shore and dry out during the dry season.

In many groups of animals, summarized by Pennak (1989), species living in freshwater can produce resting eggs, hibernacula, or other stages that enable these species to overwinter, or to survive periods of drying, as occur in Tonle Sap. In phylactolaemate bryozoans, statoblasts perform this function. The maximum longevity reported for a phylactolaemate colony (*Lophopodella cartari*) is about 23 weeks (Rogick, 1935), but colonies developed from dried statoblasts last only about two months (Rogick, 1938, 1941a; Mukai, 1997). However, this is sufficient time for colonies arising from dried statoblasts after the onset of the wet season to produce sexually a generation of colonies that can in turn form statoblasts that will survive the next dry season. Many of the phylactolaemate colonies collected in October (the end of the dry season) and from November to December (early in the wet season) were mature, and some of them also contained abundant statoblasts. The statoblasts of some species (e.g., *Hyalinella punctata*, *Lophopodella cartari*, *Asajiarella gelatinosa*) need to dry before they can float (Mukai and Oda, 1980; Mukai, 1982, 1984; Wood and Marsh, 1996).

At temperate latitudes, statoblasts are freed from a dependency on light as a germination stimulus by exposure to low temperature during winter, after which rising water temperatures in spring serve as a germination cue (Oda, 1959, 1979; Mukai, 1974, Bushnell and Rao, 1974). In tropical Tonle Sap, there is little fluctuation in water temperature (annual range 26–30°C), so the environmental cue that triggers the germination of statoblasts may involve drying and

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Fig. 10. *Hislopia cambodgiensis* Jullien. Light micrographs of (A) a colony radiating from an ancestrula; enlargement at upper right shows the ancestrula (dark zooid) and periancestrular zooids; (B) zooids.
rehydration, rather than an increase in temperature. In addition, the lake does not entirely dry out during the dry season, and colonies attached to permanently submerged or floating water plants may be able to continually populate the lake by production of larvae and statoblasts.

*Hislopia cambodgiensis* is a ctenostome bryozoan and does not produce statoblasts. However, species of some other genera of freshwater ctenostomes produce hibernacula, which are masses of cells surrounded by a thick, resistant coat (Pennak, 1989); hibernacula perform essentially the same functions as statoblasts. We did not observe hibernacula in our specimens of *Hislopia*. Nonetheless, numerous colonies of *Hislopia* were found in every year of our survey in Tonle Sap (November 2003, December 2004, December 2005). Wood et al. (2006) reported that older zooids of *Hislopia malayensis* eventually develop thickened walls and form hibernacula. Thus, the production of hibernacula could be one means by which *Hislopia cambodgiensis* repopulates the lake following the dry season. Alternatively, as with the phylactolaemates, colonies of this species occurring on vegetation that had been exposed during the dry season could have settled there from larvae released by colonies deeper in the lake or dispersed from the Mekong River with the reverse current in the wet season.

None of the phylactolaemate bryozoan species we identified in Tonle Sap is endemic to Southeast Asia. *Hyalinella lendenfeldi*, known from Australia, Thailand, and Cambodia, and *Hislopia cambodgiensis*, known from Southeast Asia and China, have relatively limited distributions. The other four species have broader distributions, including two (*Plumatella casmiana* and *Plumatella javanica*) reported from both the paleotropical and the neotropical regions.

These distributions might suggest that the collection we examined contained specimens of only common, widely distributed species and missed individuals of rare or narrowly distributed species. This is not necessarily the case, however. Phylactolaemate bryozoans generally tend to be broadly distributed (Lacourt, 1968; Bushnell, 1973; Wood and Wood, 2000, 2001; Wood and Okamura, 2005), because their floatoblasts provide a mechanism for broad dispersal. Ducks, some amphibians, and some reptiles feeding on the surface of the water are known to ingest floatoblasts, which can remain viable and germinate after passing through the digestive tract (Brown, 1933; Charalambidou et al., 2003). In addition, floatoblasts often attach to feathers of birds (Okamura and Hatton-Ellis, 1995; Freeland, 2001; Wood, 2001). Floatoblasts attached to or ingested by waterfowl can be especially broadly dispersed. The great East Asian–Australian flyway passes through Southeast Asia, and the occurrence of *Hyalinella lendenfeldi* in Cambodia and Australia is at least consistent with dispersal along this flyway. The central Asian and western Pacific flyways pass close to Southeast Asia, potentially facilitating dispersal to other areas. It is noteworthy the ctenostome bryozoan *Hislopia cambodgiensis*, which lacks floatoblasts, also has the narrowest distribution of any of the species we found in Tonle Sap; its known distribution is restricted to southern China and the Mekong region of Southeast Asia.

Most taxonomic studies of freshwater bryozoans in Southeast Asia were conducted in the late 19th and early 20th centuries. Geographical coverage of the region by these early studies was limited; some species have been detected only a few times since their original description, and both specimens and redescriptions are few. There were only a few descriptions of new species in tropical and subtropical Asia in the latter half of the 20th century: *Plumatella vorstmani* from Java (Toriumi, 1952), *Swarupella andamanensis* from Andaman Island (Rao, 1961), *Varunella coronifera* and *Varunella indorana* from India (Wiebach, 1974), and *Plumatella ganapati* and *Hyalinella diwaniensis* from Central India (Rao et al., 1985). Recently, Wood et al. (2006) described several new species, including a new *Hislopia*, after an extensive survey of freshwater bryozoans in Thailand.

Compared to Southeast Asia, freshwater bryozoans have been studied more intensively in Japan (e.g., Oka, 1907; Toriumi, 1941a, 1955; Muki, 1982, 1984; Oda, 1990; Hirose et al., 2006), where 19 species have been reported. In contrast, only about 30 species have been reported from the much larger area of Southeast Asia (Oriental) (Lacourt, 1968; Bushnell, 1973; Wood and Wood, 2000). This suggests that much remains to be learned about the diversity and distributions of freshwater bryozoans in Southeast Asia, and that more species are likely to be discovered in the region. Reliable taxonomy of both previously described and new species will best be accomplished by both the detailed examination of archival specimens and the use of modern technologies, such as molecular data and SEM observation of the statoblasts.

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