Absonifibula estuarina sp. n. (Monogenea: Diclidophoridae) parasite of juvenile Cynoscion guatucupa (Osteichthyes) from southwestern Atlantic Ocean

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Absonifibula estuarina sp. n. (Diclidophoridae, Absonifibulinae), is described from the gills of juvenile striped weakfish, Cynoscion guatucupa (Cuvier), from the southwestern Atlantic, Argentinean coast. This marine fish migrates to estuarine areas to spawn where exclusively juveniles are found parasitized; adult fish in marine water were never found to be parasitized by this monogenean. A. estuarina sp. n. is characterized mainly by the pedunculate clamps dissimilar in size, the shape of anterior jaw with sclerite ‘a’ attached to a sub-trapezoidal lamellate extension and fused to sclerites ‘c’ and ‘d’. It differs from Absonifibula bychowskyi Lawler & Overstreet, 1976, the only known species of the genus, in the shape and arrangement of the genital corona, which is armed with six similar hooks disposed in circle and the sub-trapezoidal shape of lamellate extension (‘b’). The restriction to juvenile sciaenids is a shared feature among the Absonifibulinae indicating an estuary-dependent life cycle.

Key words: Absonifibula estuarina sp. n. - Monogenea - Cynoscion guatucupa - Argentina - estuaries

The striped weakfish, Cynoscion guatucupa (Cuvier) (= Cynoscion striatus) (Sciaenidae), is a demersal fish distributed in South American Atlantic waters, from Rio de Janeiro, Brazil, to north Patagonia, Argentina (43°S) (Cousseau & Perrotta 1998, Froese & Pauly 2009); like other sciaenids, the striped weakfish inhabits coastal waters, including bays and estuaries during the spawning season (Cassia 1986, Cordo 1986, López Cazorla 2000).

The parasite fauna of this species is relatively well-known throughout its distributional range (Martorelli 1992, Timi & Etcheogin 1996, Timi et al. 1997, 2005, Santos et al. 2002, Sabas & Luque 2003, Sardella et al. 2005), comprising at present four Monogenea species: Cynoscionica julia (Rudolphi) from Brazilian waters, an unknown species of Diclidophoridae found in fish from Argentina and Diplectanum squamatum Santos, Timi & Gibson registered in both zones; all of them reported in fishes larger than 23 cm in total length (TL) (Santos et al. 2002, Sabas & Luque 2003, Timi et al. 2005).

During a survey on the helminth parasites of juvenile C. guatucupa from coastal waters of Buenos Aires province, Argentina, a different diclidophorid monogenean was found exclusively on the gills of the smallest sciaenid fish (TL< 16 cm). Their detailed study has shown that they represent a new species, which is described herein.

MATERIALS AND METHODS

A total of 500 juvenile specimens of striped weakfish, C. guatucupa were examined for monogeneans. Samples were obtained at irregular intervals from 1990-1994 as accidental catches of prawn fishery at Mar del Plata Port (38°08’S 57°32’W) following sampling schedules shown in Table.

Fish were examined immediately after capture or deep frozen in plastic bags at -18°C, until examination. After defrosting, each fish was measured for TL (cm). Parasites were recovered from the gills after microscopic examination.

The worms recovered were fixed in formaldehyde solution 4% and stored in 70% ethanol, stained in Giemsa’s trichromic and Meyer’s acetocarmine, dehydrated in ethanol series, cleared in cresote or clove oil and mounted in Canada balsam. Illustrations were made with the aid of a drawing attachment on a Leica DM LS2 microscope. Measurements are presented in micrometers as the range, with the mean in parentheses, unless otherwise stated. The terminology used in the description of clamp sclerites follows Llewellyn (1958).

Confocal laser scanning microscopy studies were carried out using an Olympus BX51 with Fluvioview version 3.2, the specimens analyzed having previously been stained with alcoholic chloride carmine. Type-specimens are deposited in the Helminthological Collection of the Museo de La Plata (CHMLP), La Plata, Argentina, and in the Helminthological Collection of Oswaldo Cruz Institute (CHIOC). Voucher specimens of Absonifibula bychowskyi Lawler and Overstreet, 1976 (CHIOC 37204 a-c) were also studied.

Host TL was compared among samples by mean of Kruskal-Wallis tests followed by *a posteriori* Tukey tests for unequal samples (Zar 1999). Prevalence and abun-
dance of infection were calculated following Bush et al. (1997). The effect of fish size on parasite abundance was analysed by mean of Spearman rank correlations after pooling samples from the same season to ensure a broader range of fish size.

_Absonifibula estuarina_ sp. n. (Figs 1-3)

**Description** - Diclidophoridae, Absonifibulinae. Body symmetrical, elongate, tapering at anterior end and enlarging to haptor level, TL 1.3-2.5 (2.0) mm, 370-650 (491) (n = 14) wide at ovary level. Haptor not distinctly set off from body proper with four pairs of pedunculated clamps which slightly decrease in size towards posterior end (Fig. 1A). Clamps of closed type, asymmetrical. Anterior jaw supported by fused sclerites: ‘a’ (a1, a2, a3, a4) with sub-trapezoidal lamellate extension (‘b’), fused to sclerites ‘c’ and ‘d’ (c1, c2, c3, d1, d2, d3); sclerites ‘a’, ‘b’ and proximal part of ‘c’ form a ring with elongate central space. Posterior jaw supported by five sclerites: two pairs of symmetrical sclerites (g1, g2, i, k) and a single median sclerite T-shaped (f) (Fig. 2A-E). First pair of clamps inverted, 125 - 200 × 112 - 180 (161 × 140) (n = 13); second clamp 125 - 180 × 105 - 160 (150 × 133) (n = 13); third clamp 100 - 160 × 102 - 150 (128 × 120) (n = 7); fourth clamp 60 - 100 × 55 - 110 (76 × 74) (n = 11). Terminal lappet 20 - 45 × 32 (n = 8) long with two pairs of larval hooks 13 - 18 (16) (n = 9) long (Fig. 1B). Eyespots absent. Buccal suckers muscular, aseptate, 32 - 50 × 32 - 50 (42 × 37) (n = 14). Mouth terminal. Pharynx ovoid 60 - 82 × 40 - 62 (72 × 50) (n = 13). Intestinal bifurcation at level of male copulatory organ. Intestinal ceca with lateral branches, extending into the haptor, confluent posteriorly. Testes 5 - 10 (8) (n = 9) in number, post-ovarian, intercelcal. Vas deferens runs anteriorly dorsal to uterus. Male copulatory organ with genital corona 27 - 37 × 25 - 37 (32 × 32) (n = 14) in diameter, armed with six similar hooks 13 - 15 (14) (n = 14) long, curved inwards (Fig. 1C). Ovary tubular, folded, pre-testicular. Oviduct connects to genitor-intestinal canal. Uterus intercelcal run medially to genital atrium. Vagina absent. Vitelline follicles extend from level of genital sac to haptor; transverse vitelline duct Y-shaped in ovarian region. Eggs fusiform with two polar filaments were damaged.

_Type-host - Cynoscion guatucupa_ (Cuvier) (Sciaenidae) (body length 3-16 cm).

*Site of infection* - gills.

*Host-locality* - Mar del Plata (38°08’S 57°32’W), Buenos Aires Province, Argentina.

*Type data and depository* - Holotype CHMLP (5939), paratypes CHML (5940), paratypes CHIOC 37205 a-d.

*Etymology* - The specific name relates to the host-parasite interaction as parasites exclusively occur in estuarine sciaenid fish.

Host-parasite data - Fish size varied significantly among samples (H5 = 351.3, p < 0.01), being host from autumn 1994 larger than those from other samples (p < 0.01); weakfish caught in both spring and summer, 1994 were similar in size (p > 0.01) and larger than those from previous years (p < 0.01); finally samples from 1990, 1992 and 1993 were homogeneous in size (all p > 0.01). Parasites occurred seasonally, being present during spring and summer only. Samples composed by smaller fish showed higher prevalence than those containing larger ones (Table). _A. estuarina_ sp. n. was found exclusively on the gills of the smallest sciaenid fish (TL < 16 cm). A significant decrease of abundance with increasing host size was observed for both spring (Rs = -0.55, p < 0.01) and summer (Rs = -0.39, p < 0.01) samples.

**DISCUSSION**

The evolutionary significance of the clamp structure and the shape of the genital corona were used in the revision of Diclidophoridae proposed by Mamaev (1976) to discriminate the genera in an evolutionary tree. In this tree, _Absonifibula_ Lawler & Overstreet was included among the higher genera (least primitive) on account of the morphology of the clamps, with fused sclerites in the posterior dorsal jaw (‘a’, ‘b’, ‘c’, ‘d’), a T-shaped median sclerite (Fig. 2E-F) and a genital corona with six dissimilar hooks.

The type-species of the monotypic _Absonifibula, A. bychowskyi_ Lawler & Overstreet was described from the Atlantic croaker _Micropogonias undulatus_ (L.) from the Gulf of Mexico (Lawler & Overstreet 1976). It was later reported from this same host in Chesapeake Bay.
and Pamlico Sound (Thoney 1991, Hendrix 1994). More recently it was found in the whitemouth croaker *Micropogonias furnieri* (Desmarest) from mixohaline areas of the Buenos Aires Province, Argentina, and neighbouring marine environments, such as the coastal waters off Mar del Plata (Martorelli et al. 2007), where it is sympatric with the new species described above. Apparently, absonifibuline genera have an estuary-dependent life cycle, where low salinity provides a suitable condition for their transmission and survival, whereas the establishment of viable populations in marine waters seems to be impeded (Lawler & Overstreet 1976).

*A. estuarina* sp. n. and *A. bychowskyi* show affinities in terms of their host group, which interestingly includes estuarine juvenile sciaenid fish. The main difference between these species is the shape and arrangement of the genital corona, which has six similar hooks disposed in a circle in the new species, whereas *A. bychowskyi* has six dissimilar hooks disposed in two rows, with the posterior pair being larger and directed anteriorly. Furthermore, in *A. estuarina* sp. n., sclerites ‘a’, ‘b’ and the proximal part of ‘c’ form a ring with an elongate central space and the shape of lamellate extension (‘b’) is more sub-trapezoidal with well marked angles. On the other hand, in *A. bychowskyi*, the central space of the ring (‘a’, ‘b’, ‘c’) is more open and elliptical and the lamellate extension is almost triangular.

A related genus to *Absonifibula* is *Mamaevicotyle* Lamothe-Argumedo, which Lamothe-Argumedo (1984) described from the serranid fish *Paralabrax maculo-
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Fig. 2: *Absonifibula estuarina* sp. n. (A-E). A: clamp, postero-lateral view; B: clamp in postero-lateral view [median sclerite (‘a’), lamellate extension (‘b’), fused sclerites (‘c’ and ‘d’)], posterior jaw [median sclerite T-shaped (f), sclerites g1, g2, i and k]; C: clamp, lateral view; D: clamp, anterior view; E: schematic design of clamp with anterior jaw [median sclerite (‘a’), lamellate extension (‘b’), fused sclerites ‘c’ and ‘d’], posterior jaw [median sclerite T-shaped (f), sclerites g1, g2, i and k]; F: *Absonifibula bychowskyi* (schematic design of clamp sclerites); G: *Mamaevicotyle* (schematic design of clamp sclerites). Scale bars: A-D = 100 μm.

Fig. 3: *Absonifibula estuarina* sp. n. (confocal laser scanning microscopy). A: clamp of closed type with anterior jaw with lamellate extension and membranous part; B: larger and left smaller clamps with posterior jaw. Right smaller clamp with detail of anterior jaw. Scale bars: A, B = 50 μm.
fasciatus (Steindachner) from off the Pacific coast of Mexico. *Mamaevicotyle*, although placed in the Diclidophoridae, resembles *A. estuarina* sp. n. in the more sub-trapezoidal shape of the lamellate extension (‘b’) (Fig. 2C) and the number and disposition of the genital corona spines, but sclerites ‘c’ and ‘d’ are not fused and the haptor is not pedunculate.

Specimens of *A. estuarina* sp. n. were not evenly distributed in the host population, but showed a marked seasonality and preference for younger fishes. In fact, the host size range parasitized by these monogeneans corresponds to an age of 0+ (Lopez Cazorla 2000, Sardiña & Lopez Cazorla 2005). Both variables affecting the distribution of the new species in the host population, namely season and host size, are correlated with each other.

*C. guatucupa* is a coastal species inhabiting offshore waters, which migrates to estuarine and other protected areas to spawn (Sardiña & López Cazorla 2005); juveniles remain in these areas during their first year (Lopez Cazorla 2000). Environmental factors in nursery areas, mainly salinity and temperature, are determinants of age-class abundance along the estuarine-marine gradient, with the youngest age-classes decreasing proportionally from brackish to marine waters (Jaureguizar et al. 2006).

This fish is a typical species of the inner coastal shelf waters of the mouth of the Río de la Plata, which is characterized by low salinity and a high bottom temperature (Jaureguizar et al. 2006). The Río de la Plata estuary dominates the coastal oceanographic conditions on the northern Argentinean sea (Acha et al. 2008), where the surface salinity distribution is controlled by the balance between onshore and offshore winds, the river discharge and the Coriolis force. As a result of the combined effects of these forces, two periods can be identified as affecting the salinity distribution. Autumn-winter is characterized by a balance between onshore and offshore winds and a maximum in the continental drainage, generating a main NNE drift of the estuarine waters along the Uruguayan coast. During spring-summer, onshore winds become dominant, resulting in a surface drift introducing freshwater southwards along the Argentine coast (Guerrero et al. 1997). This seasonal northeast/southwest oscillation implies that the coastal area around Mar del Plata falls under the influence of estuarine waters during warmer seasons (Lucas et al. 2005). This could explain the absence of monogeneans during autumn-winter, when the sampling zone is occupied by saline and colder continental shelf waters penetrating from the southwest (Jaureguizar et al. 2006).

On the other hand, a clear effect of host size on parasite burdens was observed. In fact, this parasite was never recorded in adult weakfish from either Argentina or Brazil (Sabas & Luque 2003, Timi et al. 2005). This restriction to juvenile sciaenids is a shared feature with *A. bychowskyi* (Lawler & Overstreet 1976, Martorelli et al. 2007). Several processes, such as changes in biochemistry and decreasing population density as fish growth and an increasing water salinity when fish migrate from estuarine to marine environments, have been pointed as causes of the lower parasite burden in older croakers (Lawler & Overstreet 1976, Martorelli et al. 2007). A similar combination of factors could be responsible for the observed host size-related patterns of distribution of parasite populations.

*A. estuarina* sp. n. is the second species of the genus to be described from the coastal areas of the Atlantic, confirming the predilection of absonifibulines for juvenile sciaenids in estuarine areas.

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