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# THE FOLLICLE-SINUS COMPLEX OF THE BOTTLENOSE DOLPHIN (*Tursiops truncatus*). FUNCTIONAL ANATOMY AND POSSIBLE EVOLUTIONAL SIGNIFICANCE OF ITS SOMATO-SENSORY INNERVATION.

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#### 1 Abstract

2 Vibrissae are tactile hairs found mainly on the rostrum of most mammals. The follicle which is 3 surrounded by a large venous sinus is called "follicle sinus complex" (FSC). This complex is highly innervated by somatosensitive fibers and reached by visceromotor fibers that innervate the 4 5 surrounding vessels. The surrounding striate muscles receive somatomotor fibers from the facial 6 nerve. The bottlenose dolphin (*Tursiops truncatus*), a frequently described member of the delphinid 7 family, possesses this organ only in the postnatal period. However, information on the function of the 8 vibrissal complex in this latter species is scarce. Recently, psychophysical experiments on the river-9 living Guiana dolphin (Sotalia guianensis) revealed that the FSC could work as an electroreceptor in 10 murky waters. In the present study we analyzed the morphology and innervation of the FSC of 11 newborn (n. = 8) and adult (n. = 3) bottlenose dolphins. We used Masson's trichrome stain and antibodies against neurofilament 200 kDa (NF 200), protein gene product (PGP 9.5), substance P 12 13 (SP), calcitonin gene related peptide (CGRP) and tyrosine hydroxylase (TH) to characterize the FSC 14 of the two age classes. Masson's trichrome staining revealed a structure almost identical to that of 15 terrestrial mammals except for the fact that the FSC was occupied only by a venous sinus and that the 16 vibrissal shaft lied within the follicle. Immunostaining for PGP 9.5 and NF 200 showed 17 somatosensory fibers finishing high along the follicle with Merkel nerve endings and free nerve 18 endings. We also found SP-positive fibers mostly in the surrounding blood vessels and TH both in 19 the vessels and in the mesenchymal sheath. The FSC of the bottlenose dolphin, therefore, possesses 20 a rich somatomotor innervation and a set of peptidergic visceromotor fibers. This anatomical 21 disposition suggests a mechanoreceptor function in the newborns, possibly finalized to search for the 22 opening of the mother's nipples. In the adult, however, this structure could change into a 23 proprioceptive function in which the vibrissal shaft could provide information on the degree of 24 rotation of the head. In the absence of psychophysical experiments in this species, the hypothesis of 25 electroreception cannot be rejected.

26 (**344 words**)

27

#### 28 Keywords:

Vibrissae; Whiskers; Follicle-Sinus Complex; Innervation; *Tursiops truncatus;* bottlenose dolphin
 30

#### 31 Introduction

32 Vibrissae, also called whiskers, are modified tactile hairs that occur in most mammals except 33 monotremes, anteaters, rhinoceroses, and humans (Cave, 1969; Van Horn, 1970; Chernova, 2006; 34 Muchlinski, 2010). They are mainly located around the muzzle but can be also present in other parts 35 of the head and under the carpus, depending on the species (Fundin et al., 1995; Sarko et al., 2011). 36 Their main function is to convey mechanical (tactile) stimuli to the somatosensory cortex (Woolsey 37 and Van der Loos, 1970). The hair follicle of each vibrissa is surrounded by a large venous sinus, 38 together forming the "Follicle-Sinus Complex" (FSC) (Rice et al., 1986). The presence of vibrissae 39 in Pinnipeds, Odobenids, Sirenids and otters, suggests that their somatosensory function is functional 40 also in the water. However, vibrissae are present only in newborn cetaceans and generally disappear 41 in adults. Therefore the question arises if the vibrissae of very young cetaceans perform a temporary 42 function that is lost within a few weeks after birth, or are just a remnant of a structure that evolution 43 discarded in these mammals.

44

45 The morphology and innervation of the vibrissae have been studied extensively in rodents and cats (Rice et al., 1986, 1993; Ebara et al., 2002; Park et al., 2003), and thus our present knowledge of the 46 47 structure and function of the FSC mostly derives from these species, although efforts have been 48 developed in marsupials (Lyne, 1958; Hollis and Lyne; 1974; Marotte et al., 1992). In general, the 49 FSC of terrestrial mammals consists of epidermal and dermal components. The epidermal parts 50 include the hair bulb, the vibrissal shaft (VS), the inner and outer root sheaths, surrounded by a glassy 51 membrane. The latter separates these components from the dermal parts, that is the mesenchymal 52 sheath (MS) and the venous sinus. The sinus is horizontally divided into a proximal ring sinus 53 (containing the ringwulst and the inner conical body), and a distal cavernous sinus (that contains a 54 large number of trabeculae, filled with venous blood). The last dermal part is the connective tissue 55 capsule that limits the follicle and caps it above the inner conical body with the outer conical body. 56 Finally, the rete ridge collar is a thickening of the epidermis where the VS protrudes (Rice et al., 57 1986, Ebara et al., 2002).

58

As mentioned above, marine mammals also develop vibrissae, and a description of their morphology and dimensions in seals and otter has been recently reported in comparison with several terrestrial species (Dougill et al., 2020). Walruses have the highest number of vibrissae (up to 350 on each side), while pinnipeds possess large and richly innervated FSCs, divided in three parts, with up to 1600 axons reaching it (Hyvärinen, 1989, 1995; Hyvärinen *et al.*, 2010; Ling, 1966, 1977; Marshall *et al.*, 2006). In manatees, extensive studies have described the vibrissae, which are spread out on the 65 muzzle and the body (Reep et al., 1998, 2001; Sarko et al., 2007). Mysticetes have vibrissae in large 66 quantity caudally to the blowhole and on the rostro-lateral sides of the upper and lower jaws with 67 numbers up to 250 in the bowhead whale (Balaena mysticetus) (Slijper, 1962; Yablokov and Klevezal, 1964). On the contrary, most adult toothed whales have no facial hair and show 2-10 68 69 bilateral rows of vibrissae only during fetal life and the early postnatal period (Yablokov et al., 1972; 70 Ling, 1977; Reidenberg and Laitman, 2009). Toothed whales show fully developed vibrissae only in 71 the early phases of their post-natal life (Czech-Damal et al. 2013; Cozzi et al., 2017; Dehnhardt and 72 Hanke, 2017). From morphological comparisons among odontocetes, a classification divided them 73 into four groups based on the development of the FSC (Yablokov et al., 1972). Following this 74 classification, the bottlenose dolphin falls into a group comprising species in which the VS is still 75 present in the early postnatal period but disappears in the majority of adult individuals. This is not the 76 case in river dolphins such as the Guiana dolphin (Sotalia guianensis) of which a recent study 77 described the FSC (Czech-Damal et al., 2012). The FSC of this species was renamed vibrissal crypt 78 because of its different anatomical structure, characterized by the absence of the VS, hair papilla, 79 clear root sheaths, blood sinus and capsule (Czech-Damal et al., 2012). The FSC lumen is filled with 80 desquamated corneocytes and keratinous fibers, that together may be considered a highly electrically 81 conductive biogel (Czech-Damal et al., 2012), part of an electrosensory system that facilitates the 82 hunt of small bottom-living prey in turbid water, where echolocation is not possible or potentially not 83 efficient enough, by detection of their electric field (Czech-Damal et al., 2012).

84 The somatosensory innervation of mystacial vibrissae is provided by three subdivisions of the 85 maxillary branch of the trigeminal nerve. The deep vibrissal nerve, originates directly from the 86 infraorbital nerve, supplies a single FSC, penetrates the capsule and arborizes dorsally at various 87 levels. The superficial vibrissal nerves (SVNs) come from superficial cutaneous nerves and supply 88 several FSCs. Small- to fine-caliber nerve fiber branches reach the FSC from the base and supply the 89 hair papilla and hair bulb (Rice et al., 1986; Ebara et al., 2002). The deep vibrissal nerve ends in 90 mechanoreceptors such as Merkel nerve endings (MNEs), lanceolate endings and free nerve endings 91 (FNEs) along the follicle. The SVNs, instead, provide innervation to lanceolate endings at the level 92 of inner conical body and MNEs at the level of the rete ridge collar (Fundin et al., 1997a; Ebara et 93 al., 2002). The somatomotor innervation is provided by motoneurons placed in the lateral part of the 94 facial nucleus and innervates the extrinsic (mimic) and intrinsic musculature of the mystacial pad 95 (Haidarliu et al., 2010; Herfst and Brecht, 2008). The visceral innervation (sympathetic and 96 parasympathetic) regulates blood flow in the FSC, supplied by the deep vibrissal artery, and 97 consequently regulates blood pressure, which is essential for the activation of receptors that respond 98 to specific stimulation thresholds. (Fundin et al., 1997b, Maklad et al., 2004).

99 Here we describe the FSC in a series of postnatal and adult bottlenose dolphins, aiming at 100 characterizing the changes in the anatomy and morphology of this structure at different life stages by 101 histochemical and immunohistochemical techniques. Special attention was dedicated to the 102 innervation of the FSC, the nature of the nerve fibers, and its functional potential.

#### 104 Material and methods

105

#### 106 a) <u>Animals</u>

107 The samples of vibrissae from 11 bottlenose dolphins (Tursiops truncatus, Montagu 1821) were 108 the obtained from Mediterranean Marine Mammal Tissue Bank (MMMTB, 109 http://www.marinemammals.eu), housed in the Department of Comparative Biomedicine and Food 110 Science (BCA) of the University of Padova. The MMMTB is a CITES recognized institution (IT 020) 111 that collaborates with the Italian Ministry of the Environment. The MMMTB collects, processes, and 112 stores samples of tissues of various cetacean species that stranded along the Italian coastline since 113 2000. Additional samples derived from marine mammals that died at marine theme parks and aquaria, and whose bodies were delivered to BCA for diagnostic post-mortem. More details of the specimens 114 115 used in this study can be seen in Table 1.

#### 116 b) <u>Sample processing</u>

Each sample was obtained by carving out around the VS on both sides of the rostrum in the newbornand around the dimple containing the orifice in the adult (Figure 1a, b).

119 The samples were fixed by immersion in 4% neutral buffered paraformaldehyde and stored at 4 °C. 120 Tissues for Masson's trichrome were then included in paraffin and cut in 5 µm- and 10 µm-thick sections either longitudinal or transversal to the main axis of the FSC by use of a rotatory microtome 121 122 (Leica, Germany). Sections were mounted on gelatinized slides and air dried. Samples bound for 123 immunocytochemistry were washed in standard phosphate buffer solution (PBS) overnight at 4 °C, 124 stored in PBS containing 0.1% Na-azide and sucrose at 30%, immersed in OCT Compound (Tissue 125 Tek, Sakura Finetek Europe, NL) and frozen at -80 °C in isopentane cooled with liquid nitrogen. 25 126 µm-thick sections of the longitudinal and transversal planes were subsequently taken with a cryostat

127 (Leica, Germany).

ID	Species	Sex	Age class	Origin
# 83	T. truncatus	М	Newborn	Died in a marine theme park
# 114	T. truncatus	М	Newborn	Died in a marine theme park
# 123	T. truncatus	F	Newborn	Died in a marine theme park
# 124	T. truncatus	М	Newborn	Died in a marine theme park
# 144	T. truncatus	М	Newborn	Died in a marine theme park
# 145	T. truncatus	М	Newborn	Died in a marine theme park
# 162	T. truncatus	М	Newborn	Wild

128 **Table 1:** Origin of specimens

# 229	T. truncatus	М	Newborn	Died in a marine theme park
# 146	T. truncatus	М	Adult	Died in a marine theme park
# 159	T. truncatus	М	Adult	Died in a marine theme park
# 444	T. truncatus	М	Adult	Wild

130 c) <u>Histological techniques:</u>

The morphology of the FSC was stained using a Masson's trichrome staining protocol. Briefly, the 131 132 sections were immersed in 3 baths of xylene for 5 minutes each and subsequently hydrated with a descending series of graded alcohol solutions (100 %, 95 %, 90 %, 80 %, 70 %, 50 %). Then, they 133 134 were stained with Mayer's Emallume for 5-10 minutes and rinsed with tap water. Later, the sections 135 were colored for 5 minutes in a solution of distilled water (300 ml) containing Ponceau 2R (0,2 g), 136 acid fuchsin (0,1 g) and acetic acid (0,6 ml). After rinsing with a 1% acetic acid solution, the sections 137 were put in a solution of distilled water (100 ml), phosphomolybdic acid (3-5 g) and orange G (2 g), for 5 minutes, and rinsed again in an acetic acid solution. The sections were then colored for 5 minutes 138 139 in light green (0,1-0,2 g in 100 ml distilled water) and acetic acid (0,2 ml). After the last rinsing in a 140 1% acetic acid solution, the slides were dehydrated directly with absolute alcohol, and passed in 141 xylene (3 x 3 minutes) and coverslipped with Entellan (Merck, Damstraldt, Germany).

142 The innervation of the FSC was characterized with immunocytochemistry, either via 143 immunoperoxidase (IP) or immunofluorescence (IF), using the neuronal markers shown in Table 2. 144 For IP staining, contiguous sections were initially immersed in a 0,4% solution of Triton X-100 145 (Merck, Darmstadt, Germany) in PBS at 4 °C for 24 hours. They were then rinsed in PBS baths for 3 146 x 10 minutes Next, sections were treated with 1% H<sub>2</sub>O<sub>2</sub> in PBS for 30 minutes. After three 10-minute 147 washes in PBS, a 3% solution of normal goat serum (NGS, Sigma G-9023, Saint Louis, Missouri, 148 USA) was applied for 2 hours, at room temperature. Thus, sections of each sample were incubated in 149 a wet chamber for 48 hours, at 4 °C with the primary antibodies (Table 2a) in antibody diluent (1,8% 150 NaCl in a 0.01 M sodium phosphate solution containing 0,1% Na-azide). After primary incubation, 151 the sections were washed with PBS and incubated with the specific secondary antibodies (Table 2b) 152 diluted in PBS in a wet chamber for 2 hours at room temperature. After further three washes in PBS, 153 they were transferred for 30 minutes in an avidin-biotin complex solution (ABC Standard, ABC kit 154 Vectastain, Vector Laboratories, Burlingame, CA, USA, PK 6100) and washed again in PBS. Finally, 155 immunoperoxidase was developed using 3.3'-diaminobenzidine (DAB kit Vector Laboratories, Burlingame, CA, USA, BA-9200). The sections were dehydrated in ethanol, passed in xylene and 156 157 covered with a coverslip using Entellan.

The slides of both Masson's trichrome and IP were observed with an optic microscope (Zeiss Axioplan, Carl Zeiss, Oberkochen, Germany), captured with the microscope Nikon Coolscope (Nikon, Japan) and subsequently elaborated with the programs Elipsenet 1.20.0 (Nikon, Japan) and GIMP 2 (GNU Image Manipulation Program 2.10).

For the IF procedure, slides were placed in a wet chamber. A first PBS wash was performed to 162 rehydrate the sections. A Blocking Serum solution (0,5% Triton X-100, 10% Normal Goat serum, 163 NGS, Vector, Burlingame, CA, USA, in PBS) or 10% Normal Donkey Serum (NDS, Jackson, Bar 164 Harbor, Maine, USA) was used at room temperature for 2 hours. Then, the sections of each sample 165 166 were incubated in a wet chamber for 48 hours, at 4 °C, with the primary antibodies (Table 2a) in antibody diluent. After 48 hours, the sections were washed with PBS and either pure NGS or 10% 167 168 NDS, (5 x 10 minutes on a stirrer). Next, the sections were incubated for 3 hours, at room temperature, with specific secondary antibodies (Table 2b), diluted in PBS. After further five 10-minute washes in 169 170 PBS, the slides were air dried and prepared with glycerol buffered with 0,5 M sodium carbonate (pH 171 8,6) to be finally sealed with nail polish. 172 The slides obtained were observed under an epifluorescence optical microscope (Axioplan, Carl

173 Zeiss, Oberkochen, Germany), equipped with a system of filters that allowed the distinction of the 174 fluorescence FITC (given by fluorescein) from Alexa 594 fluorescence. The images were acquired 175 using a digital camera and DMC 2 software (Polaroid Corporation, Cambridge, MA, USA). The 176 images were proceeded using Adoba Photochen (Adoba Suptema Ser Less CA, USA).

- 176 images were processed using Adobe Photoshop (Adobe Systems, San Jose, CA, USA).
- 177

178 **Table 2a**: List of the primary antibodies used for immunoperoxidase (IP) or immunofluorescence

179 (IF).

Primary antibody	Used for	Immunogen /host	Supplier	Dilution	Antibody RRID	Validation
Protein Gene Product 9.5 (PGP 9.5)	IP	Polyclonal rabbit	Millipore, Temecula, CA, USA	1:500	AB_91019	PMID:19296476
	IF			1:1000		
Substance P (SP)	IP	Polyclonal rabbit	Immunostar, Hudson, WI, USA Fitzgerald Industries	1:1000	AB_572266	PMID:10087030 PMID:10196365
	IF	Monoclonal rat	International, North Acton, MA, USA	1:200	AB_2313816	PMID:22740069 PMID:26713509
Calcitonin Gene Related Peptide	IP/IF	Monoclonal mouse	Santa Cruz Biotechnology Inc., CA, USA	1:200	AB_2259462	PMID:30971286 PMID:29943954
(CGRP)	IF	Polyclonal rabbit	Peninsula Laboratories Inc., San Carlos, CA, USA	1:1000	AB_2313775	PMID:18186028 PMID:28680400
Human Tyrosine	IP/IF	Monoclonal mouse	Monosan, Uden, Netherlands	1:50	ID: MONX10786*	PMID:29615733

Hydroxylase						
(TH)						
Neurofilamen	IP/IF	Monoclonal	Sigma, Saint Louis,	1:1000	AB_477272	PMID:18022951
t 200kDa (NF		rabbit	Missouri, USA			PMID:19937712
200kDa)						

- 181 **Table 2b**: List of the secondary antibodies used for immunoperoxidase (IP) or immunofluorescence
- 182 (IF).

Secondary antibody	Used for	Immunogen /host	Supplier	Dilution	Antibody RRID	Validation
Biotinylated Anti-Rabbit	IP	Goat	Vector Laboratories, Burlingame, CA, USA	10 µg/ml	AB_2313606	PMID:19127523 PMID:23766132
Anti-Mouse	IP	Goat	Vector Laboratories, Burlingame, CA, USA	10 μg/ml	AB_2336171	PMID:23766132 PMID:25057794
Anti-Mouse Alexa 594	IF	Goat	Thermo Fisher Scientific, Waltham, MA, USA	1:200	AB_141372	PMID:23913443 PMID:25057190
Anti-Rat Alexa 594	IF	Donkey	Thermo Fisher Scientific, Waltham, MA, USA	1:200	AB_2535795	PMID:25933105 PMID:28089909
Anti- Rabbit- FITC	IF	Goat	Calbiochem, Darmstadt, Germany	1:100	ID: 401314*	PMID:29615733

183 \*Antibody RRID are universally identified codes and were taken from the website the antibody

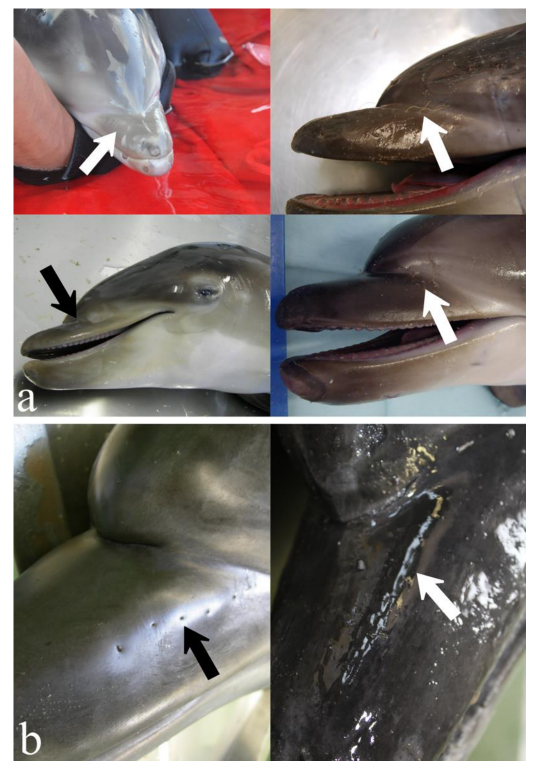
184 registry (https://antibodyregistry.org/) which integrated the antibody database of the Journal of

185 *Comparatve Neurology*. For each antibody, there is at least one publication correlated to a unique

186 PMID (PubMed Identifier). For the antibodies whose lot number are MONX10786 and 401314, there

187 are still no current RRID available, but the validation appears in one publication (Bombardi et al.,

188 2010).



- 190
- 191 Figure 1: Macroscopic images of the rostrum of some specimens of (a, top four) newborn and (b,
- 192 bottom two) adult bottlenose dolphin. The arrows indicate where the vibrissae emerge from the skin
- 193 as can be seen in the newborns (a) or the concavity found in the adults (b).
- 194

195 **Results** 

196

- 197 1. Morphology
- 198 Newborn dolphins

199 In the newborns, the external part of the VS was approx. 10 mm long. The FSC of all specimens 200 consisted of an epidermal and dermal part. The epidermal part comprised the hair with its sheaths, 201 overlying the dermal venous sinus. The VS originated from the bulb and consisted of three concentric 202 layers which, from inside to outside, were identified as the medulla, the cortex, both made of 203 keratinized cells, and the cuticle, which consisted of a simple squamous keratinized epithelium. The 204 MS and the capsule were fused near the follicle outlet. At the base of the FSC, the bulb resembled a 205 highly innervated and vascularized dermal papilla (Figure 2a). The VS was wrapped by the inner root 206 sheath, attached to the cuticle, and the outer root sheath, surrounded by the glassy membrane (Figure 207 2b). The hair shaft was surrounded by a venous sinus and delimited by a connective tissue capsule. 208 The sinus comprised internally the MS, in contact with the glassy membrane and externally by a 209 capsule (Figure 2c).

In the slides analyzed, it was never possible to observe either a ringwulst or a ring sinus. Furthermore,no muscle fiber or gland was present around the follicle.

212

#### 213 Adult dolphins

In the adults, the VS was present but did not reach the skin surface. Apart from this feature, the FSCof the adult dolphins showed the same structure of those of the newborns.

216

217 2. Innervation

Anti-PGP 9.5 immunoreactive (-ir), anti-NF 200-ir, anti-TH-ir and anti-SP-ir nerve fibers were evident in all the samples. No anti-CGRP-ir fibers were observed.

220 Newborn dolphins

PGP 9.5-ir fibers penetrated the FSC at the level of the hair bulb, and yielded an intricate arborized network of ramifications (Figure 3a, b). The nerve fibers protruded at various levels in the mesenchymal sheath, giving rise to button-like terminations characterizing MNEs (Figure 3c). They derived from large and medium-sized fibers that ran to form clusters of button-like endings with a smooth and regular surface, between which fine spiral-like fibers were present (Figure 3d).

226 NF 200-ir fibers were also detected penetrating the bulb (Figure 3e), first running parallel to the VS

and then entering at different levels along the follicle (Figure 3f). Nerve fibers of different calibers

were distributed along the VS, progressing either in a straight line or along a winding path until theyreached the top of the FSC (Figure 3f).

230 Numerous nerve fibers were observed in transversal sections of the FSC, from the hair bulb to the

apex (Figure 4). These fibers innervated the hair bulb (Figure 4b, c) and sent small groups of axons

to surround the follicle (Figure 4d, e), ending into MNEs (Figure 4f). Some of these axons penetrated

the venous sinus, ran along the trabeculae (Figure 4g) and ended at the MS that wrapped the VS with

234 MNEs (Figure 4h). This rich innervation was evident in all transverse sections up to the outlet of the

- vibrissa (Figure 4i). We did not identify other receptors with certainty and, as mentioned above, most
- fibers seemed to end as FNEs.

SP-ir fibers ran either grouped in bundles or alone close to blood vessels (Figure 5a, b). Double immunofluorescence for PGP 9.5 and TH showed that PGP 9.5-ir fibers were qualitatively four-fold the TH-ir fibers (Figure 5c, d). TH-ir fibers were mainly located around the blood vessels, and sometimes presented a tortuous pattern. They contained thin-caliber axons that ran first on the surface of the adventitia and then penetrated the wall (Figure 5e). Few TH-ir fibers were found at the base of the bulb and in the MS (Figure 5f).

243

#### 244 Adult dolphins

Immunohistochemical results in adult dolphins showed the same general pattern of that of newborns, with some notable exceptions. PGP 9.5-ir and NF 200-ir nerve fibers were clear (Figure 6a-c) and the MNEs bound to the mesenchymal sheath were smaller in adults (Figure 6d). SP reactivity was found in large caliber fibers near the dermo-epidermal junction (Figure 7a), where they ran parallel to the skin before bending towards the FSC and ending as FNEs (Figure 7b). TH-ir fibers were rarer. Very thin TH-ir fibers were present in the trabeculae of the venous sinus and the mesenchymal sheath ending with isolated oval corpuscles (Figure 7c-e).

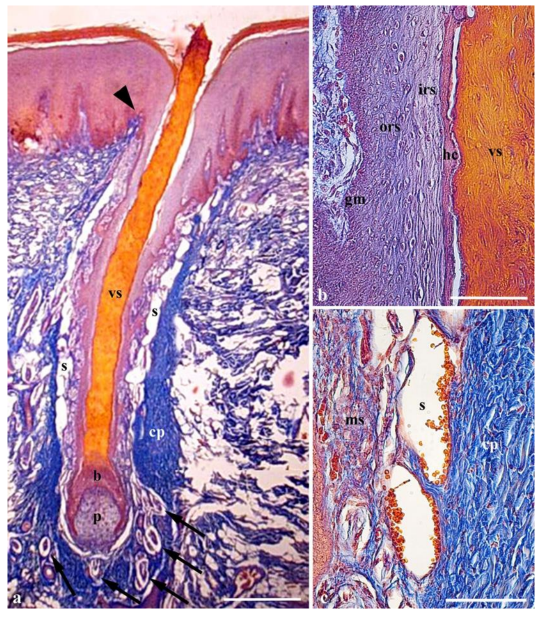
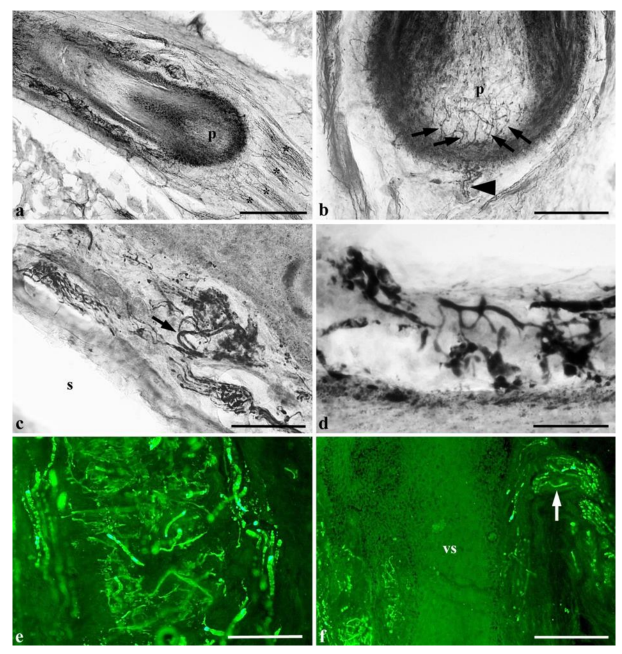
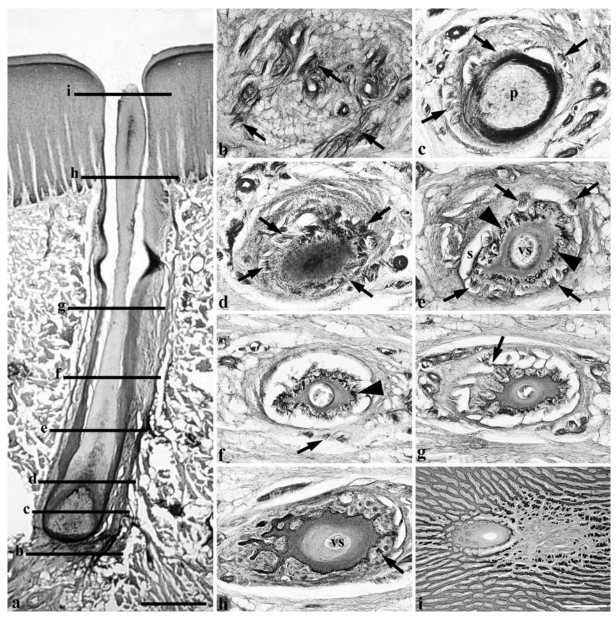


Figure 2 – Longitudinal section of a typical FSC in newborn bottlenose dolphin. a. The vibrissa is surrounded by a venous sinus (s). A capsule (cp) envelops the complex. Several nerves (arrows) reach the root of the vibrissa. The arrowhead indicates the fusion between the capsule and mesenchymal sheat. b. Detail at higher magnification of the epidermal components. c. Detail at higher magnification of the dermal components. b, bulb; cp, capsule; gm, glassy membrane; irs, inner root sheat; ms, mesenchymal sheath; ors, outern root sheat; p, papilla; s, venous sinus; vs, vibrissal shaft. Masson's Trichrome stain. Scale bars: a = 1 mm; b, c = 100 µm.



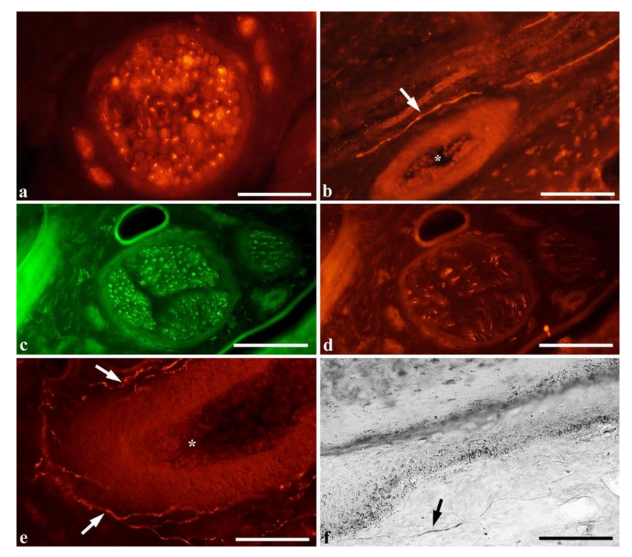
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**Figure 3** – Innervation of the FSC in a newborn bottlenose dolphin. The nerve fibers were immunolabelled for PGP 9.5 (a-d) and NF200kD (e, f). a. Several PGP 9.5-ir nerve bundles (asterisks) reach the root of the vibrissa. p, papilla. b. Few thin-caliber fibers (arrowhead) enter the papilla (p) and terminate as free nerve endings (arrows). c. In the mesenchymal sheath, some nerve fibers (arrow) give rise to MNEs (asterisks). d. High magnification showing MNEs. Note the characteristic buttonlike endings. e. The dense network of nerve fibers around the bulb. f. A nerve bundle penetrate the FSC laterally. vs, vibrissal shaft. Scale bars: a,  $e = 100 \mu m$ ; b, c,  $f = 200 \mu m$ ;  $d = 50 \mu m$ .

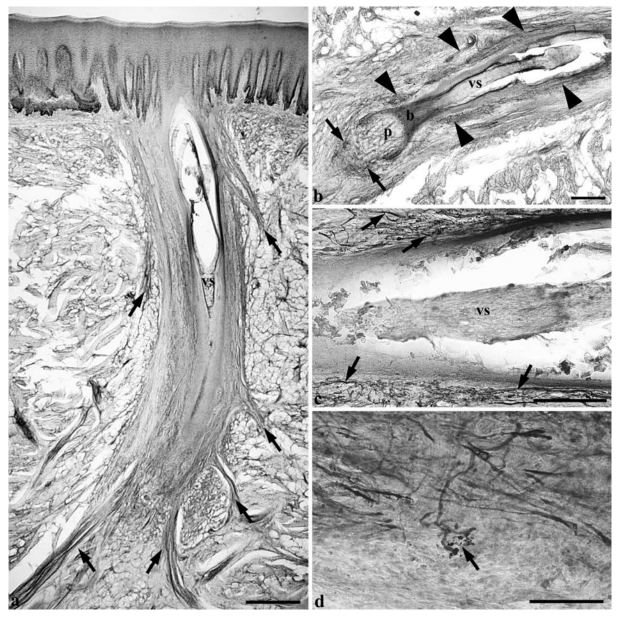


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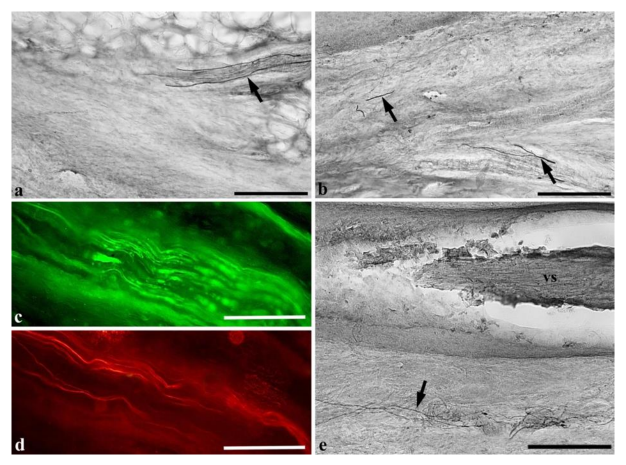
272 Figure 4 – Longitudinal (a) and transverse (b-i) sections (b) of FSC in newborn bottlenose dolphin 273 showing the innervation at different levels from the basal (b) to the apical (i). The nerve fibers were 274 immunolabelled with antibodies to PGP 9.5. b. Some nerves (arrows) reach the root of the vibrissa. 275 c. The nerves break into several fascicles (arrows) that ascend close to the papilla (p). d. The nerve 276 fibers (arrows) surround the follicle. e. Some nerve fibers (arrows) penetrate the venous sinus (s) and 277 branch in the mesenchymal sheath (arrowhead). vs, vibrissal shaft. f. Some fibers terminate on MNEs 278 (arrowhead), while others continue along the FSC (arrow). g. A nerve fiber (arrow) passes through 279 one of numerous trabeculae of the venous sinus. h. At the level of dermo-epidermal border, the nerve 280 fibers disappear but the MNEs are still present (arrow). i. Section through the skin and the dermal 281 papilla. Scale bars: a = 1 mm; b-h = same magnification of i;  $i = 350 \mu \text{m}$ .



**Figure 5** - SP- (a, b) and TH-(c-f) immunoreactive fibers in newborn bottlenose dolphin. a. Transverse section of a nerve bundle showing many immunoreactive fibers. b. A nerve fiber (arrow) runs parallel to a blood vessel (asterisk). c, d. Double immunofluorescence PGP 9.5-FITC (c) / TH-Alexa 594 (d) of a nerve bundle in transverse section. Note the TH immunoreactivity of some fibers. e. Several nerve fibers (arrows) reach the tunica adventitia of a vessel (asterisk). f. A thin fiber run within the ms. Scale bars = 100  $\mu$ m.



**Figure 6** – Longitudinal sections of FSC in adult bottlenose dolphin. The nerve fibers were immunolabelled for PGP 9.5 (a, b) and NF200kD (c, d). a. Note the six nerve bundles (arrows) reaching the FSC. b. A vibrissa is clearly visible inside the follicle. Some nerve fibers (arrows) reach the bulb, others (arrowheads) run within the ms. b, bulb; p, papilla; vs, vibrissal shaft. c. High magnification showing the rich innervation (arrows) of the ms. d. Detail of a MNE (arrow). Scale bars: a, b = 1 mm; c = 200  $\mu$ m; d =100  $\mu$ m.



**Figure 7** – SP- (a, b) and TH- (c-e) immunoreactive fibers in an adult bottlenose dolphin. a. Two positive fibers reach the FSC laterally. b. Few nerve fibers in the ms. c, d. Double immunofluorescence PGP 9.5-FITC (c) / TH-Alexa 594 (d) of a large nerve bundle in longitudinal section. Note the few TH-ir fibers. e. Few fibers (arrow) in the ms. Scale bars: a, b,  $e = 200 \mu m$ ; c, d  $304 = 100 \mu m$ .

#### 306 **Discussion**

307 In the present work, we describe the morphology and the innervation of FSC of newborn and adult 308 bottlenose dolphins. Our findings reveal that newborn specimens possess a complete structure divided

309 in an epidermal and a dermal part, hence the term "follicle-sinus complex" whereas, in adults, the VS

310 lies within the follicle (Figs. 2, 6). The absence of a ring sinus (ringwulst) of the *erector pili* muscle

and of any kind of associated gland constitutes differences with the vibrissae in terrestrial mammals.

312 The blood sinus of the bottlenose dolphin consists of just one cavity forming a trabecular net just like

313 the tammar wallaby (*Macropus eugenii*) (Marotte *et al.*, 1992) and is not divided into two parts as in

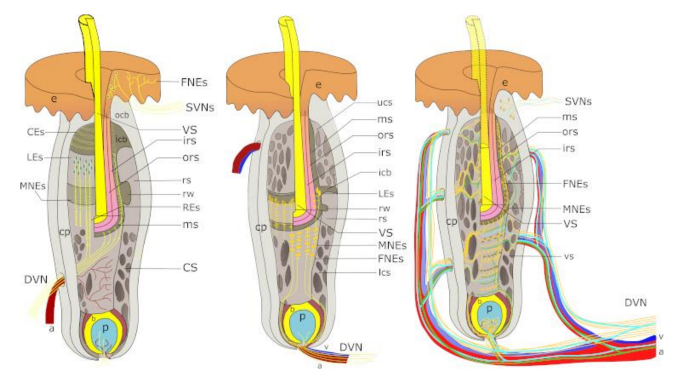
terrestrial mammals (Rice *et al.*, 1986, Ebara *et al.*, 2002) or three parts as in pinnipeds (Marshall *et al.*, 2006) (Figure 8).

316 Anti-neuronal antibodies (anti-PGP 9.5, -NF 200, -TH, and -SP) helped to characterize the 317 innervation of the FSC. Antibodies directed against proteins of the neurofilaments (anti-PGP 9.5, -318 NF 200, Figs. 3, 4, 6) identified several fiber bundles which bifurcated from the deep vibrissal nerve 319 at the bottom of the FSC to reach various heights of the follicle, similarly to what was previously described in manatees (Sarko et al., 2007). Few fibers were also found at the dermo-epidermal 320 321 junction (Figure 4i). Merkel receptors were evenly distributed along the MS of the FSC, although, in 322 the newborn they were also found at the dermo-epidermal junction (Figs. 3, 4). Since no striated 323 muscle fibers and no glands were present in the FSC, the nature of the present innervation is likely 324 somatosensory and derived from the maxillary nerve (V2) of the trigeminal nerve. The V2 runs -325 from its exit of the skull base - rostrally to enter the dolphin equivalent of the maxillary foramen and 326 subsequent canal. The rostralmost fibers of the V2 run as the infraorbital nerve in maxillary canals. 327 Some fibers of the latter penetrate the maxilla in dorsal direction to provide sensorial innervation to 328 the skin of the rostrum (Rauschmann 1992). Dolphins possess no movable lips and have virtually no 329 snout fascia or mimic muscles beyond those, more caudal, that act on the melon, thus implying a 330 virtual absence of a somato-motor component in the facial nerve this far forward on the face. 331 Comparisons with other mammals are difficult. Whisking rodents possess the mystacial pad, a 332 thickening of the snout fascia where mimic muscles, sensory receptors and collagen structures form 333 a highly developed motor-sensory organ (Haidarliu et al., 2020). In the rat and mouse, a column of 334 neocortical neurons in the whisker somatosensory cortex (wS1, or barrel cortex) corresponds to each 335 FSC, with a highly developed layer IV receiving the thalamic afferent (Woolsey and Van der Loos, 336 1970; Van der Loos and Woolsey, 1973; Van der Loos, 1976; Rice and Van Der Loos, 1977; 337 Jeanmonod et al., 1981; Pearson et al., 2006; Bosman et al., 2011; Schröder et al., 2020). However, 338 the barrel cortex is typical of rodents, and is not present in any other species (i.e. mammals of the 339 genus Felis or Panthera), even ones which present whisking behavior such as the short-tailed

opossum (Waite *et al.*, 1991; Ramamurthy and Krubitzer, 2016). Furthermore, the neocortex of dolphins and whales lacks a layer IV and it is currently hypothesized that thalamic projections reach layer II instead of IV (for general description see Cozzi et al., 2017), thus making any comparison with the highly specialized barrel cortex of rodents difficult. In particular the study by van Kann and colleagues (2017) pointed out the main differences in the primary neocortical areas layering between the common dolphin, the wild boar and humans.

The use of antibodies against peptides (SP, CGRP) and against a key enzyme in catecholamine synthesis (TH), allowed further characterization of the innervation. SP is involved in nociception, and a subpopulation of sensory neurons in the mammalian trigeminal ganglion contains SP, colocalized with CGRP (Alvarez *et al.*, 1988, Fundin *et al.*, 1997b; Waite and and Ashwell, 2012). SP-ir neurons have also been described in the dorsal root ganglia of bottlenose dolphins (Bombardi *et al.*, 2010).





#### 352

353 **Figure 8** – Schematic drawn representing the main differences between the FSCs of a terrestrial mammal (left), pinniped (center) and dolphin (right). On the right, the follicle represented is that of 354 355 the adult as the dotted lines and transparent areas (SVNs and MNEs in the dermo-epidermal junction) 356 are of the newborn. In terrestrial mammals the FSC is divided into two halves, the inferior cavernous 357 sinus, and the superior ring sinus. Receptors, coming mainly from the deep vibrissal nerve are 358 positioned at different heights depending on their sensory nature. In pinnipeds it is instead divided into three portions, including an upper cavernous sinus. The fibers, which derive only from the deep 359 360 vibrissal nerve, innervate up to the inner conical body, without reaching the epidermis. Finally, in the dolphin, a trabecular component forms a single venous sinus in which the receptors, also deriving 361 mainly from the deep vibrissal nerve, are distributed along the follicle until they reach the epidermis 362 in the newborn. Also note the TH-ir (green) and SP-ir (light blue) fibers which accompany the blood 363 364 vessels. Abbreviations: VS, vibrissal shaft; e, epidermis; irs, inner root sheat; ors, outern root sheat; ms, mesenchymal sheat; ocb, outern conical body; icb, inner conical body; rw, ringwulst; rs, ring 365 366 sinus; cs, cavernous sinus; lcs, lower cavernous sinus; ucs, upper cavernous sinus; c, capsule; p,

367 papilla; b, bulb; DVN, deep vibrissal nerve; SVNs, superficial vibrissal nerves; MNEs, merkell nerve endings; FNEs, free nerve endings; CEs, circular endings; LEs, lanceolate endings; REs, reticular endings; a, artery; v, vein; vs, venous sinus.

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371 Several SP-ir were located in the dermo-epidermal junction and, deeper lateral to the follicle. Other 372 SP-ir nerves were also found around the sinus' blood vessels. Both SP and CGRP are vasodilators. 373 However, the presence of SP-ir fibers within big nerve bundles suggests a nociceptive function while 374 the presence of SP in thin fibers around blood vessels might indicate a parasympathetic activity on the vessels of the FSC (Fundin et al., 1997b). No CGRP-ir fiber was detected in the vibrissae of the 375 376 dolphins in this study. This absence is difficult to explain from a functional point of view. Indeed, 377 CGRP has been found in the gastrointestinal tract of the striped dolphin (Stenella coeruleoalba) 378 (Domenighini et al., 1997), and has been proposed to be present in CNS of the bottlenose dolphin 379 (Rambaldi et al, 2016). Additionally, the presence of this molecule was demonstrated in the FSC of 380 manatees (Sarko et al., 2007). Therefore, the apparent absence of CGRP in the FSC of the bottlenose 381 dolphin could be due to a loss of nociceptive function or different trigeminal organization.

382 TH-ir fibers consisted of thin branches located around the surface of the blood vessels connecting to 383 the sinus, even though some elements were found also in the MS and trabeculae. Considering their 384 peripheral location, TH-ir fibers are indicative of a noradrenergic sympathetic innervation derived 385 from the cranial [superior] cervical ganglion. In mammals, sympathetic fibers follow both the external 386 and internal carotid arteries. From the former, fibers then run along the infraorbital artery. However, 387 the internal carotid artery obliterates early in postnatal life in dolphins (Boenninghaus, 1903; Cozzi 388 et al., 2017), as in many Cetartiodactyls. Therefore, the precise route of the TH-ir that was observed 389 in the FCS of dolphins remains to be ascertained. It may be possible that the internal carotid artery 390 develops in the odontocete embryo to guide the growing sympathetic fibers from the cervical ganglion 391 since it is known that the axonal outgrowth of mammalian sympathetic precursors proceeds in 392 dependence and thus in parallel to the development of the internal carotid artery (reviewed in Kameda 393 2020). Thus, possibly only after the sympathetic fibers have found their route, the internal carotid 394 artery obliterates in odontocetes. Alternatively, these fibers in cetaceans may rely exclusively on the 395 external carotid path.

396 Contrarily to what was reported by Yablokov et al. (1972), we were able to confirm the presence of 397 the VS in adult specimens, as was previously described by Palmer and Weddell (1964). In all the 398 adult animals analyzed in the present study, the FSC was complete, i.e. the epidermal and dermal 399 components were discernable, and the VS was still present, albeit only in the follicle. In fact, the hair 400 papillae maintained the same morphology found in the newborns. Moreover, the VS did not have the 401 aspect of a formless cluster due to the epithelial regeneration. Since the papilla is responsible for the

402 production of the VS's components, a lack of this structure can justify the absence of the VS in the 403 adult Guiana dolphin, in which it had transformed into an agglomeration of fat cells (Czech-Damal 404 *et al.*, 2012). Interestingly, we found MNEs and FNEs but no other receptor, which is coherent with 405 what was described in the Guiana dolphin (Czech-Damal *et al.*, 2012). In the absence of other markers 406 (e.g. anti-S100 protein), it however remains impossible to rule out the existence of other nerve 407 endings such as Pacini corpuscles.

408 The Guiana dolphin is in fact very similar to the ubiquitous bottlenose dolphin in general body 409 morphology and proportions, although somewhat smaller. Yet, the preferred habitat of the Guiana 410 dolphin is estuarine and coastal, while bottlenose dolphins are distributed worldwide and live both 411 along coastlines and in the high seas. The FSC complex in Guiana dolphins has been linked to the 412 sensitivity to electric fields generated by prey burrowed in sand or hidden by murky waters (Czech-413 Damal et al., 2012, 2013). A progressive morphological adaptation to different specific environments 414 over time may be the explanation to the differences in the structure of the FSC in those two similarly 415 sized species. Whether the evolution was towards a loss or a gain of function remains unclear.

Based on the absence in dolphins of morphological features typical of terrestrial mammals with vibrissae, i.e. the absence of a wide range of receptors, the mystacial pad or the barrel cortex, we can infer that the FSC of dolphins are relatively rigid structures, with a sensitivity potentially reduced which does not allow perception of dynamic changes the way fully formed whiskers do. Yet, since the vibrissae persist in newborn dolphins, contrarily to other structures lost during fetal growth (pelvic limbs, to name the most striking difference with land mammals) it is possible that the vibrissae play a role in the early postnatal life, in the days immediately following parturition.

Pinniped mothers recognize their newborn mostly through olfaction while the newborn, from the first 423 424 minutes of life, vocalizes for most of the first day, a behaviour that diminishes gradually afterwards 425 (Trillmich, 1981). Underwater, this kind of recognition is almost impossible to dolphins since they 426 cannot smell (Cozzi et al., 2017). As proposed by Cozzi et al. (2012, 2015), an evolutionary 427 adaptation in cetaceans to an immediate recognition is the early ossification of the tympanic bulla 428 which would allow the newborn to locate the mother's vocalizations and help it in postpartum period. 429 However, full echolocation capacities are likely not to fully develop until month 1-3 after birth (Harder et al., 2016); and while it is difficult to think that vibrissae may act for this recognition, it is 430 431 more plausible that it could allow the newborn to find the opening of the nipples immediately after 432 birth, thanks to the rich SP innervation and vasomodulation regulated by the TH-ir plexus (Fundin et 433 al., 1997b; Maklad, 2004).

A VS is still present within the FSC in the adult dolphin and the innervation remains relatively
developed. The original tactile function of the FSC may lose value in the growing dolphins that, later

436 in life, may rely on different sensory modalities to interact with their mother and the rest of the pod. 437 This organ could act as a proprioceptor as suggested by Yablokov et al. (1972). The VS could be 438 sensitive to low frequency oscillations and water movements caused by head rotation and would 439 consequently activate the receptors that provide this information to the central nervous system. This 440 would allow the dolphin to always have a perception of the angular position of the head. Nervous 441 fibers could modify the pressure inside the venous system and help maintain thermoregulation and 442 modify the threshold of the receptors (Fundin et al., 1997b). This idea does not preclude that the 443 structure found by Czech-Damal and colleagues (2012) in the Guiana dolphin could function as an 444 electroreceptor. However, to confirm or reject this hypothesis, further psychophysical experiments should be conducted in the bottlenose dolphin. 445

### 446 **Authors contributions :**

- 447 TG, AG, JMG and BC designed the study, TG, AG, CT and MDS acquired and analyzed the data.
- 448 TG, AG, JMG and BC wrote the draft. AG, SdV and SH critically revised the manuscript. All
- 449 authors approved the article.

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#### **Tables:**

Table 1: Origin of specimens 

ID	Species	Sex	Age class	Origin
# 83	T. truncatus	М	Newborn	Died in a marine theme park
# 114	T. truncatus	М	Newborn	Died in a marine theme park
# 123	T. truncatus	F	Newborn	Died in a marine theme park
# 124	T. truncatus	М	Newborn	Died in a marine theme park
# 144	T. truncatus	М	Newborn	Died in a marine theme park
# 145	T. truncatus	М	Newborn	Died in a marine theme park
# 162	T. truncatus	М	Newborn	Wild
# 229	T. truncatus	М	Newborn	Died in a marine theme park
# 146	T. truncatus	М	Adult	Died in a marine theme park
# 159	T. truncatus	М	Adult	Died in a marine theme park
# 444	T. truncatus	М	Adult	Wild

627	Table 2a: List of the primary antibodies used for immunoperoxidase (IP) or immunofluorescence
628	(IF).

Primary antibody	Used for	Immunogen /host	Supplier	Dilution	Antibody RRID	Validation
Protein Gene Product 9.5 (PGP 9.5)	IP	Polyclonal rabbit	Millipore, Temecula, CA, USA	1:500	AB_91019	PMID:19296476
	IF			1:1000		
Substance P (SP)	IP	Polyclonal rabbit	Immunostar, Hudson, WI, USA Fitzgerald Industries	1:1000	AB_572266	PMID:10087030 PMID:10196365
	IF	Monoclonal rat	International, North Acton, MA, USA	1:200	AB_2313816	PMID:22740069 PMID:26713509
Calcitonin Gene Related Peptide	IP/IF	Monoclonal mouse	Santa Cruz Biotechnology Inc., CA, USA	1:200	AB_2259462	PMID:30971286 PMID:29943954
(CGRP)	IF	Polyclonal rabbit	Peninsula Laboratories Inc., San Carlos, CA, USA	1:1000	AB_2313775	PMID:18186028 PMID:28680400
Human Tyrosine Hydroxylase (TH)	IP/IF	Monoclonal mouse	Monosan, Uden, Netherlands	1:50	ID: MONX10786*	PMID:29615733
Neurofilamen t 200kDa (NF 200kDa)	IP/IF	Monoclonal rabbit	Sigma, Saint Louis, Missouri, USA	1:1000	AB_477272	PMID:18022951 PMID:19937712

630 **Table 2b**: List of the secondary antibodies used for immunoperoxidase (IP) or immunofluorescence

631 (IF).

Secondary antibody	Used for	Immunogen /host	Supplier	Dilution	Antibody RRID	Validation
Biotinylated Anti-Rabbit	IP	Goat	Vector Laboratories, Burlingame, CA, USA	10 μg/ml	AB_2313606	PMID:19127523 PMID:23766132
Anti-Mouse	IP	Goat	Vector Laboratories, Burlingame, CA, USA	10 µg/ml	AB_2336171	PMID:23766132 PMID:25057794
Anti-Mouse Alexa 594	IF	Goat	Thermo Fisher Scientific, Waltham, MA, USA	1:200	AB_141372	PMID:23913443 PMID:25057190
Anti-Rat Alexa 594	IF	Donkey	Thermo Fisher Scientific, Waltham, MA, USA	1:200	AB_2535795	PMID:25933105 PMID:28089909
Anti- Rabbit- FITC	IF	Goat	Calbiochem, Darmstadt, Germany	1:100	ID: 401314*	PMID:29615733

\*Antibody RRID are universally identified codes and were taken from the website the antibody
registry (<u>https://antibodyregistry.org/</u>) which integrated the antibody database of the *Journal of Comparatve Neurology*. For each antibody, there is at least one publication correlated to a unique
PMID (PubMed Identifier). For the antibodies whose lot number are MONX10786 and 401314, there
are still no current RRID available but the validation appears in one publication (Bombardi et al.,
2010).

638

#### 640 Figure legends

641

Figure 1 – Rostrum of some specimens of (a) newborn and an (b) adult bottlenose dolphin. The
arrows indicate where the VS emerge from the skin (a) or the concavity (b).

644

**Figure 2** – Longitudinal section of a typical FSC in newborn bottlenose dolphin. a. The vibrissa is surrounded by a venous sinus (s). A capsule (cp) envelops the complex. Several nerves (arrows) reach the root of the vibrissa. The arrowhead indicates the fusion between the capsule and mesenchymal sheat. b. Detail at higher magnification of the epidermal components. c. Detail at higher magnification of the dermal components. b, bulb; cp, capsule; gm, glassy membrane; irs, inner root sheat; ms, mesenchymal sheath; ors, outern root sheat; p, papilla; s, venous sinus; vs, vibrissal shaft. Masson's Trichrome stain. Scale bars: a = 1 mm; b,  $c = 100 \mu \text{m}$ .

652

**Figure 3** – Innervation of the FSC in a newborn bottlenose dolphin. The nerve fibers were immunolabelled for PGP 9.5 (a-d) and NF200kD (e, f). a. Several PGP 9.5-ir nerve bundles (asterisks) reach the root of the vibrissa. p, papilla. b. Few thin-calibre fibers (arrowhead) enter the papilla (p) and terminate as free nerve endings (arrows). c. In the mesenchymal sheath, some nerve fibers (arrow) give rise to MNEs (asterisks). d. High magnification showing MNEs. Note the characteristic buttonlike endings. e. The dense network of nerve fibers around the bulb. f. A nerve bundle penetrate the FSC laterally. vs, vibrissal shaft. Scale bars: a,  $e = 100 \mu m$ ; b, c,  $f = 200 \mu m$ ;  $d = 50 \mu m$ .

660

Figure 4 – Longitudinal (a) and transverse (b-i) sections (b) of FSC in newborn bottlenose dolphin 661 showing the innervation at different levels from the basal (b) to the apical (i). The nerve fibers were 662 663 immunolabelled with antibodies to PGP 9.5. b. Some nerves (arrows) reach the root of the vibrissa. 664 c. The nerves break into several fascicles (arrows) that ascend close to the papilla (p). d. The nerve 665 fibers (arrows) surround the follicle. e. Some nerve fibers (arrows) penetrate the venous sinus (s) and 666 branch in the mesenchymal sheath (arrowhead). vs, vibrissal shaft. f. Some fibers terminate on MNEs 667 (arrowhead), while others continue along the FSC (arrow). g. A nerve fiber (arrow) passes through 668 one of numerous trabeculae of the venous sinus. h. At the level of dermo-epidermal border, the nerve 669 fibers disappear but the MNEs are still present (arrow). i. Section through the skin and the dermal 670 papilla. Scale bars: a = 1 mm; b-h = same magnification of i;  $i = 350 \mu \text{m}$ .

671

Figure 5 – SP- (a, b) and TH-(c-f) immunoreactive fibers in newborn bottlenose dolphin. a.
Transverse section of a nerve bundle showing many immunoreactive fibers. b. A nerve fiber (arrow)

runs parallel to a blood vessel (asterisk). c, d. Double immunofluorescence PGP 9.5-FITC (c) / TH-Alexa 594 (d) of a nerve bundle in transverse section. Note the TH immunoreactivity of some fibres. e. Several nerve fibers (arrows) reach the tunica adventitia of a vessel (asterisk). f. A thin fiber run within the MS. Scale bars =  $100 \mu m$ .

678

**Figure 6** – Longitudinal sections of FSC in adult bottlenose dophin. The nerve fibers were immunolabelled for PGP 9.5 (a, b) and NF200kD (c, d). a. Note the six nerve bundles (arrows) reaching the FSC. b. A vibrissa is clearly visible inside the follicle. Some nerve fibers (arrows) reach the bulb, others (arrowheads) run within the MS. b, bulb; p, papilla; vs, vibrissal shaft. c. High magnification showing the rich innervation (arrows) of the MS. d. Detail of a MNE (arrow). Scale bars: a, b = 1 mm; c = 200  $\mu$ m; d =100  $\mu$ m.

685

**Figure 7** – SP- (a, b) and TH- (c-e) immunoreactive fibers in an adult bottlenose dolphin. a. Two positive fibers reach the FSC laterally. b. Few nerve fibers in the MS. c, d. Double immunofluorescence PGP 9.5-FITC (c) / TH-Alexa 594 (d) of a large nerve bundle in longitudinal section. Note the few TH-ir fibers. e. Few fibers (arrow) in the MS. Scale bars: a, b,  $e = 200 \mu m$ ; c, d  $= 100 \mu m$ .

691

692 Figure 8 – Schematic drawn representing the main differences between the FSCs of a terrestrial 693 mammal (left), pinniped (center) and dolphin (right). On the right, the follicle represented is that of 694 the adult as the dotted lines and transparent areas (SVNs and MNEs in the dermo-epidermal junction) 695 are of the newborn. In terrestrial mammals the FSC is divided into two halves, the inferior cavernous 696 sinus, and the superior ring sinus. The receptors are of various nature, are positioned at various heights 697 depending on their receptor (sensory) nature and come mainly from the deep vibrissal nerve. In the 698 pinniped it is instead divided into three portions, including an upper cavernous sinus. In this case, 699 however, the fibers, which derive only from the deep vibrissal nerve, innervate up to the inner conical 700 body, without reaching the epidermis. Finally, in the dolphin there is a trabecular component that 701 forms a single venous sinus in which the receptors, deriving mainly from the deep vibrissal nerve, are 702 distributed along the follicle until they reach the epidermis in the newborn. Also note the TH-ir 703 (green) and SP-ir (light blue) fibers which accompany the blood vessels. Abbreviations: VS, vibrissal 704 shaft; e, epidermis; irs, inner root sheat; ors, outern root sheat; ms, mesenchymal sheat; ocb, outern 705 conical body; icb, inner conical body; rw, ringwulst; rs, ring sinus; cs, cavernous sinus; lcs, lower 706 cavernous sinus; ucs, upper cavernous sinus; c, capsule; p, papilla; b, bulb; DVN, deep vibrissal 707 nerve; SVNs, superficial vibrissal nerves; MNEs, merkell nerve endings; FNEs, free nerve endings;

708	CEs, circular endings; LEs, lanceolate endings; REs, reticular endings; a, artery; v, vein; vs, venous
709	sinus.

- 710
- 711 **Figure 9** Schematic representation of the course of the maxillary nerve (V2) in adult (left) and
- 712 newborn (right) bottlenose dolphin. The dotted ellipse approximates the area of the FSCs location.
- 713