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***Koristocetus pescei* gen. et sp. nov., a diminutive sperm whale (Cetacea: Odontoceti: Kogiidae) from the late Miocene of Peru**

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Abstract. Among odontocetes, members of the family Kogiidae (pygmy and dwarf sperm whales) are known as small-sized and in many respects enigmatic relatives of the great sperm whale *Physeter macrocephalus*. Most of the still scanty fossil record of Kogiidae is represented by isolated skulls and ear bones from Neogene deposits of the Northern Hemisphere, with the significant exception of *Scaphokogia*, a highly autapomorphic genus from late Miocene deposits of the Pisco Formation exposed along the southern coast of Peru. Here we report on a new fossil kogiid from Aguada de Lomas, a site where the late Miocene beds of the Pisco Formation are exposed. This specimen consists of an almost complete cranium representing a new taxon of Kogiidae: *Koristocetus pescei* gen. et sp. nov. *Koristocetus* mainly differs from extant *Kogia* spp. by displaying a larger temporal fossa and well-individualized dental alveoli on the upper jaws. Coupled with a relatively elongated rostrum, these characters suggest that *Koristocetus* retained some degree of raptorial feeding abilities, contrasting with the strong suction feeding specialization seen in Recent kogiids. Our phylogenetic analysis recognizes *Koristocetus* as the earliest branching member of the subfamily Kogiinae. Interestingly, *Koristocetus* shared the southern coast of present-day Peru with members of the genus *Scaphokogia*, whose unique convex rostrum and unusual neurocranial morphology seemingly indicate a peculiar foraging specialization that has still to be understood. In conclusion, *Koristocetus* evokes a long history of high diversity, morphological disparity, and sympatric

habits in fossil kogiids, thus suggesting that our comprehension of the evolutionary history of pygmy and dwarf sperm whales is still far from being exhaustive.

1 Introduction

Among extant odontocetes, members of the currently monotypic family Kogiidae (*Kogia breviceps* and *K. sima*, pygmy and dwarf sperm whales) are known as small-sized and in many respects enigmatic relatives of the great sperm whale, *Physeter macrocephalus*. In spite of their wide geographic distribution, the natural history of pygmy and dwarf sperm whales remains largely speculative, and most of our knowledge about them comes from the study of stranded individuals (e.g., Nagorsen, 1985; McAlpine, 2002; Bloodworth and Odell, 2008; Taylor et al., 2012a, b, and references therein). The still scanty fossil record of Kogiidae is mostly represented by isolated skulls and ear bones from Neogene deposits of the Northern Hemisphere (e.g., Barnes, 1973, 1998; Pilleri, 1987; Whitmore, 1994; Bianucci, 1996, 1997; Cigala Fulgosi, 1996; Luo and Marsh, 1996; Bianucci et al., 1998, 2011; Bianucci and Landini, 1999; Lambert, 2008; Whitmore and Kaltenbach, 2008; Velez-Juarbe et al., 2015, 2016), with the significant exception of *Scaphokogia*, a highly autapomorphic genus from the late Miocene marine deposits of the Pisco Formation exposed along the southern coast of Peru (de Muizon, 1988). The aim of this paper is the

description of another kogiid cranium from the Pisco Formation referred to a new genus and species. The specimen was discovered in late Miocene beds exposed at the locality of Aguada de Lomas in the Sacaco Basin. We then undertake a phylogenetic analysis of the relationships of this new taxon and briefly discuss its paleobiological meaning.

(ZooBank LSID for *Koristocetus* gen. nov.: urn:lsid:zoobank.org:act:790FADA0-A6C5-4345-95AE-7CD5CF00149D, record created by Alberto Collareta on 30 October 2017)

(ZooBank LSID for *Koristocetus pescei* sp. nov.: urn:lsid:zoobank.org:act:805D6750-10CF-4FB0-ABDB-30160400A8D6, record created by Alberto Collareta on 30 October 2017)

2 Material and methods

2.1 Institutional abbreviations

IRSNB: Institut Royal des Sciences Naturelles de Belgique, Brussels, Belgium; MBMS, Museo Balseros del Mar del Sur, Salango, Ecuador; MGPUF: Museo di Storia Naturale, Sezione di Geologia e Paleontologia, Università degli Studi di Firenze, Florence, Italy; MNHN: Muséum National d'Histoire Naturelle, Paris, France; MSNC: Museo Civico di Storia Naturale di Comiso, Comiso, Italy; MUSM: Museo de Historia Natural de la Universidad Nacional Mayor de San Marcos, Lima, Peru; MUSNAF: Museo di Storia Naturale dell'Accademia dei Fisiocritici, Siena, Italy; UCR: University of California, Riverside, California, USA; USNM: National Museum of Natural History, Smithsonian Institution, Washington DC, USA; ZMA: Zoological Museum Amsterdam, Amsterdam, the Netherlands.

2.2 Kogiid specimens analyzed for comparison

In addition to MUSM 888 we have directly examined for comparison the following fossil and extant Kogiidae: *Aprixokogia kelloggi* Whitmore and Kaltenbach, 2008 (USNM 187015); *Kogia breviceps* (de Blainville, 1838) (MBMS 4000; MNHN 1976-37; USNM 22015, 22559, 22893, 270979, 283625, 302041, 395699, 504314; ZMA 14.817, 14.818); *Kogia pusilla* (Pilleri, 1987) (MGPUF 1540V); *Kogia sima* (Owen, 1866) (MSNC 3450; MUSNAF Mam410; USNM 550482, 550487, 573734; ZMA 14.765); *Praekogia cedrosensis* (UCR 15229); *Scaphokogia cochlearis* de Muizon, 1988 (MNHN PPI 229, MUSM 971, MUSM 1998); *Scaphokogia* sp. (MUSM 972); *Scaphokogiinae* sp. (MUSM 3291, MUSM 3405); *Thalassocetus antwerpiensis* Abel, 1905 (IRSNB M.525).

2.3 Phylogeny and taxonomy

The phylogenetic analysis was undertaken with PAUP (Phylogenetic Analysis Using Parsimony; Swofford, 2001) using a modified version of the character–taxon matrix of Lambert et al. (2017) (see appendices A and B for the list of characters and the character–taxon matrix, respectively). We rescored some characters for *Scaphokogia cochlearis* based on personal observations of two specimens kept at the MUSM and slightly modified characters 29 and 30 in order to better account for the variability recognized in Kogiidae. In addition to MUSM 888, we inserted in our matrix an undescribed taxon of Kogiidae, sharing similarities with *Scaphokogia cochlearis* and represented by two partial skulls (MUSM 3291 and MUSM 3405) from the highly fossiliferous sites of Cerro Los Quesos (Lambert et al., 2009; Gariboldi et al., 2015, 2017; Bianucci et al., 2016a, c; Di Celma et al., 2016; Gioncada et al., 2018) and Cerro Blanco (Brand et al., 2004, 2011; Esperante et al., 2008, 2015), respectively (late Miocene, Pisco Formation, Peru).

We used the tree-bisection–reconnection algorithm and the heuristic search option, considering all characters as unordered and unweighted. We used the phylogenetic definitions proposed by Bianucci and Landini (2006) and Lambert (2008) for Physeteroidea, Physeteridae, and Kogiidae. Embracing the subdivision of Kogiidae in subfamilies proposed by de Muizon (1988) and then reappraised by Bianucci and Landini (1999), Lambert (2008), and Whitmore and Kaltenbach (2008), here we redefine Kogiinae Gill, 1871 as the most inclusive clade including *Kogia* but not *Scaphokogia*, whereas we redefine *Scaphokogiinae* de Muizon, 1988 as the most inclusive clade including *Scaphokogia* but not *Kogia*.

3 Geographical and stratigraphical framework

Exposed along the southern coast of Peru, the Pisco Formation is a late Neogene shallow-marine sedimentary unit, known as one of the most impressive Cenozoic marine Fossil Lagerstätten worldwide. Indeed, the fossil record of vertebrates from Pisco Formation stands out for the exceptional abundance of specimens, high diversity and disparity, and remarkable quality of preservation (e.g., Hoffstetter, 1968; de Muizon, 1981, 1984, 1988, 1993; de Muizon and DeVries, 1985; Pilleri, 1989–1990; de Muizon and McDonald, 1995; de Muizon et al., 1999, 2002, 2003, 2004a, b; McDonald and de Muizon, 2002; de Muizon and Domning, 2002; Brand et al., 2004, 2011; Esperante et al., 2008, 2015; Lambert et al., 2008, 2009, 2010a, b, 2015, 2017; Ehret et al., 2009, 2012; Bianucci et al., 2010, 2016a, b, c; Bisconti, 2012; Lambert and de Muizon, 2013; Amson et al., 2014, 2015a, b, 2016; Collareta et al., 2015, 2017b; Esperante and Poma, 2015; Gariboldi et al., 2015; Stucchi et al., 2015, 2016; Gioncada et al., 2016; Marx and Kohno, 2016; Di Celma et al., 2017;

Landini et al., 2017a, b; Marx et al., 2017; Ramassamy et al., 2017). One of the most significant paleontological sites of the Pisco Formation is Aguada de Lomas (Fig. 1; indicative geographic coordinates: 15°30' S; 74°48' W), about 8 km NNE of the village of Puerto Lomas, within the Neogene Sacaco Basin (de Muizon and DeVries, 1985). According to Lambert and de Muizon (2013), the thick sedimentary succession exposed at Aguada de Lomas includes four vertebrate-bearing levels; namely, ELJ (El Jahuay), AGL (Aguada de Lomas), MTM (Montemar), and SAS (Sud-Sacaco) as defined by de Muizon and DeVries (1985) and de Muizon (1988). The cetacean skull studied here (MUSM 888) was collected about 20 years ago by two of us (Christian de Muizon and Mario Urbina) from the Aguada de Lomas area; it comes from the late Miocene beds of the Pisco Formation located above the AGL vertebrate level of de Muizon and DeVries (1985) (personal observation by Mario Urbina). Therefore, MUSM 888 should be referred to either the MTM or SAS vertebrate levels. These two levels were originally dated to 6.0–5.5 Ma and ca. 5 Ma, respectively, based on the composition of the molluscan assemblages and radiometric dating of tuff layers (de Muizon and DeVries, 1985; de Muizon and Bellon, 1986; de Muizon, 1988). More recently, U–Pb dating on zircon grains from tuff layers and Sr-isotope analyses on marine mollusc shells provided new insights into the age of several fossiliferous exposures of the Pisco Formation in the Sacaco Basin (Ehret et al., 2012). In particular, for the SAS level, these geochemical methods yielded ages of 7 ± 1 Ma (U–Pb) and ca. 6.59 Ma (strontium ratio); such values are supported by Sr-isotope analyses on shells, collected at the site of Sud-Sacaco (West) from two mollusc-rich intervals located above the vertebrate-bearing tuff layer discussed by de Muizon and DeVries (1985), resulting in ages of 5.93 and 5.89 Ma, respectively (Ehret et al., 2012). Furthermore, Sr-isotope analyses yielded an age of ca. 7.30 Ma for the MTM vertebrate level, whereas the AGL vertebrate level was younger than 7.46 Ma, which in turn represents the estimated age of the beds just above the ELJ vertebrate level (Ehret et al., 2012) (see also Lambert and de Muizon (2013), Collareta et al. (2017a), and Lambert et al. (2017) for a discussion about the chronostratigraphic framework of the Pisco Formation cropping out in the Aguada de Lomas valley). Therefore, embracing these recent age estimates, and accounting also for the rather wide confidence intervals associated with the strontium-ratio analyses on shells provided by Ehret et al. (2012), it appears that MUSM 888 should be latest Tortonian or early to mid-Messinian in age.

4 Systematic paleontology

Cetacea Brisson, 1762

Pelagiceti Uhen, 2008

Neoceti Fordyce and Muizon, 2001



Figure 1. Geographical position (star) of the upper Miocene site of Aguada de Lomas (Pisco Formation, Sacaco Basin, southern coast of Peru, where the holotype of *Koristocetus pescei* gen. et sp. nov. MUSM 888 was discovered). Modified after Collareta et al. (2017a).

Odontoceti Flower, 1867

Physeteroidea Gray, 1821

Kogiidae Gill, 1871

Kogiinae Gill, 1871

Koristocetus gen. nov.

Type and only known species: *Koristocetus pescei* gen. et sp. nov.

Etymology: The generic name originates from the combination of the Greek word *koristos*, meaning “with helmet” or “crested”, for the presence of a well-developed sagittal facial crest, and the Latin word *cetus*, meaning “whale”. Gender masculine.

Known range: Late Miocene (latest Tortonian or early to mid-Messinian, following the chronostratigraphic framework provided by Ehret et al. (2012); see above for more details) of southern Peru.

Differential diagnosis: Same as for the type species until other species are described.

Koristocetus pescei sp. nov.

Figs. 2–5

Holotype and only known specimen: MUSM 888, a partial cranium.

Etymology: The specific name honors the memory and heritage of Hugo Pesce Pescetto (1900–1969), Peruvian physician, politician, writer, and “hero of public health in Peru” (Burstein, 2003), who occupied a position of professor at the Universidad Nacional Mayor de San Marcos, Lima.

Type locality: The holotype is from the locality of Aguada de Lomas, about 8 km NNE of the village of Puerto Lomas, Arequipa Province, southern coast of Peru (Fig. 1). Indicative geographic coordinates: 15°30' S, 74°48' W.

Type horizon: Pisco Formation, late Miocene (latest Tortonian or early to mid-Messinian; see above).

Differential diagnosis: *Koristocetus* is a small-sized physeteroid, similar in size to *Kogia* spp. and *Nanokogia*. It is recognized as a kogiid based on the following features: bizygomatic width smaller than 40 cm, presence of a sagittal facial crest, external nares greatly asymmetric, and nasals absent. Differs from *Aprixokogia*, *Kogia*, and *Thalassocetus* by the presence of a more delicate postorbital process of the frontal. Differs from *Aprixokogia* and *Kogia* by the smaller angle between the frontal–maxilla suture line and the horizontal plane. Differs from *Aprixokogia*, *Kogia breviceps*, and *K. sima* by having a supracranial basin that does not significantly extend onto the dorsal face of the rostrum. Differs from *Aprixokogia*, *Nanokogia*, and *Praekogia* by the foramen magnum being proportionally smaller. Differs from *Aprixokogia*, *Scaphokogia*, and *Thalassocetus* by the antorbital notches extending within the supracranial basin and by the presence of a deep and round notch for the posterior process of the tympanic bulla. Differs from *Aprixokogia* and *Thalassocetus* by the slit-like geometry of the antorbital notches. Differs from *Kogia*, *Nanokogia*, *Praekogia*, *Scaphokogia*, and *Thalassocetus* by the presence of a single, not subdivided right posterior dorsal infraorbital foramen. Differs from *Kogia*, *Nanokogia*, and *Praekogia* by presenting no lateral expansion of the postnarial eminence of the right premaxilla. Differs from *Kogia*, *Nanokogia*, and *Thalassocetus* by the larger, more anteroposteriorly elongated temporal fossa. Differs from *Kogia* and *Nanokogia* by presenting well-individualized dental alveoli on the maxilla. Differs from *Kogia* and *Praekogia* by the less posteroven- trally elongated postorbital process of the frontal. Differs from *Kogia* by the proportionally smaller lacrimojugal complex. Differs from *Kogia breviceps* and *K. sima* by the proportionally longer rostrum, the supracranial basin not being laterally expanded, the more obliquely oriented frontal groove, and the thin and uninflated lateral maxillary crests. Differs from *Kogia sima* and *Scaphokogia* by the lack of a left premaxillary foramen. Differs from *Kogia sima* by the presphenoid which does not extend anteriorly within the mesorostral groove. Differs from *Nanokogia* by having a wider mesorostral groove in its posterior portion and a maxilla wider than the premaxilla all along the rostrum. Differs

from *Praekogia* by having the left premaxilla that does not reach the sagittal facial crest and the presphenoid largely exposed on the ventral aspect of the skull. Differs from *Scaphokogia* by the rostrum having a dorsal face that is not semicylindrical, the postnarial eminence that significantly contributes to the sagittal facial crest, the sagittal facial crest that is not significantly dislocated towards the left side of the skull, the supracranial basin that is not spoon-shaped, and the supraoccipital shield that does not dip anteriorly.

Description, comparisons, and remarks: MUSM 888 is an incomplete skull, lacking parts of the ventral and posterodorsal portions of the braincase, most of the auditory bones, both mandibles, all the teeth, and the left zygomatic process (Figs. 2–5). The rostrum is almost complete. The facial aspect of the skull is generally well preserved, although some surfaces are moderately abraded and the posterior part of the supracranial basin (including the vertex of the skull) is lacking. In ventral view, the surfaces of the squamosals and exoccipitals are poorly preserved. Locally, the skull is still encrusted by hardened sediment. An endocranial cast is preserved as a concretion.

The skull of *Koristocetus* is small, distinctly asymmetric, with a rather short rostrum (ratio between length of the rostrum and condylobasal length ca. 0.45). Nevertheless, the skull appears anteroposteriorly elongated and drop-shaped in dorsal view (Fig. 2) with respect to Recent kogiids (ratio between condylobasal length and skull width across the postorbital processes ca. 1.7, versus 1.0–1.2 in extant *Kogia* spp.), thus recalling *Nanokogia* in this regard (ratio ca. 1.8 in *Nanokogia*). Sutures between adjacent cranial bones are fused in MUSM 888, thus indicating that it belonged to a full-grown (i.e., physically adult) individual. Measurements of the cranium of *Koristocetus* are reported in Table 1.

Premaxilla. In dorsal view, the lateral margins of the premaxillae are slightly concave laterally and weakly convergent towards the apex of the rostrum, which in turn presents a significant constriction at mid-length (Fig. 2). Medial to each premaxilla, the mesorostral groove opens starting from the base of the rostrum. It widens progressively towards the anterior tip of the skull as the medial margins of the premaxillae weakly diverge forward, reaching a transverse width of 23 mm at about two-thirds of the rostrum length (measured from its base). The posterior half of the mesorostral groove is wider than in *Nanokogia*, recalling instead the condition observed in *Aprixokogia* and *Kogia breviceps*. In the posterior half of the rostrum, anterior to the bony nares, the dorsal surface of the right premaxilla is convex transversely, whereas the dorsal surface of the left premaxilla is distinctly concave transversely; both premaxillae progressively flatten towards the apex of the rostrum. No premaxillary foramina are observable on the right and left premaxilla; however, the dorsal surface of the right premaxilla is moderately eroded at the base of the rostrum, and the possibility that a tiny right premaxillary foramen was present cannot be completely ruled

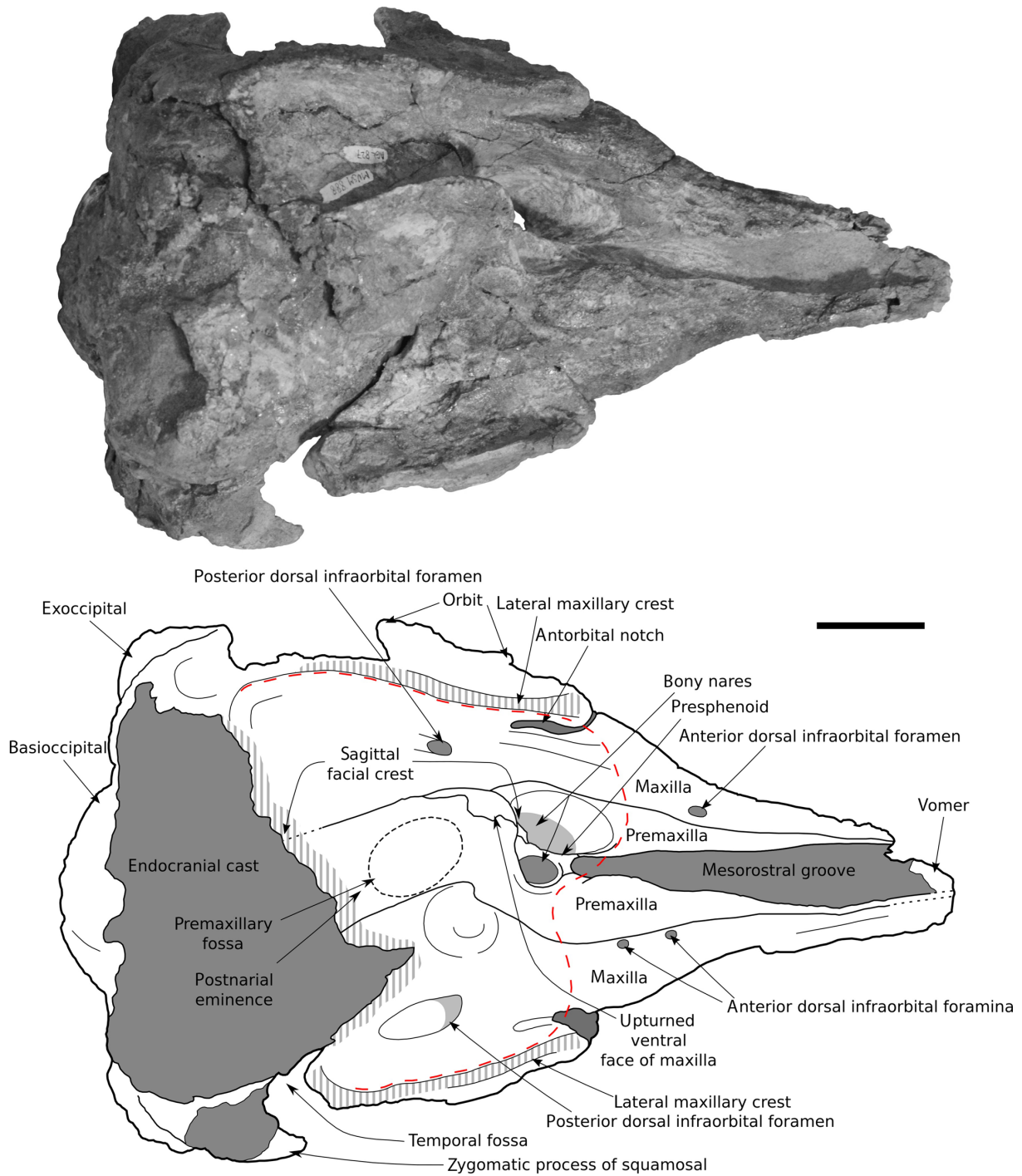


Figure 2. Dorsal view of the skull MUSM 888, holotype of *Koristocetus pescei* gen. et sp. nov., Pisco Formation, late Miocene of Aguada de Lomas, southern coast of Peru. The dark-grey-shaded areas correspond to hardened sediment. The light-grey-shaded areas correspond to openings in the skull. The grey vertical stripes denote damaged areas. The red dashed line denotes the indicative extent of the supracranial basin. Scale bar: 4 cm.

out. By contrast, the unambiguous absence of the left premaxillary foramen is a feature shared with *Aprixokogia*, *Kogia breviceps*, *K. pusilla*, and *Nanokogia*, whereas this foramen is present (although reduced in size) in *Kogia sima* and *Scaphokogia*. The external nares are observed at the base of the rostrum (Fig. 2). They are strongly asymmetrical and displaced towards the left side of the skull (a feature shared by all Physeteroidea). The left naris is large and oval-shaped; it is anteroposteriorly elongated and is situated in a similarly oriented funnel-shaped depression (a character also observed in *Kogia* spp.); its transverse diameter is 24 mm. The right naris is significantly smaller than the left one, having a transverse diameter of 9 mm. The left premaxilla presents a lobe-like posterior end which forms the anterior and lateral walls of the left naris (a feature reminiscent of *Aprixokogia*, *Kogia*, and *Nanokogia*); as also observed in all Kogiidae except presumably *Praekogia*, the posterior end of the left premaxilla does not take part in building the sagittal facial crest. In turn, the right premaxilla forms the anterior, lateral, and posterior walls of the right naris. The right premaxilla is constricted transversely at the level of the right naris; posterior to this level, the right premaxilla is significantly expanded posteromedially as a stripe of bone that joins the upturned ventral face of the medial margin of the left maxilla to form the sagittal facial crest (Fig. 2). Such a posterior expansion of the right premaxilla (defined by Whitmore and Kaltenbach, 2008, as the postnarial eminence) is present, more or less developed, in all Kogiidae. The postnarial eminence significantly contributes to build the sagittal facial crest in all Kogiidae except *Scaphokogia*, in which the right premaxilla constitutes only a small fraction of the supracranial basin and sagittal facial crest, being exposed only at the anterior end of the latter feature. In contrast with *Kogia*, *Nanokogia* and, to a lesser extent, *Praekogia*, no lateral expansion of the postnarial eminence posterior to the right naris is observed in *Koristocetus* (a feature somewhat reminiscent of *Aprixokogia*). On this portion of the right premaxilla, which dips anterolaterally, a very shallow depression (premaxillary fossa of Barnes, 1973) occupies the area posterior to the right naris (Fig. 2). This premaxillary fossa is oval-shaped and roughly anteroposteriorly elongated; it measures 37 mm in length and 27 mm in width. Several other Kogiidae exhibit a more or less shallow depression located in the right part of the supracranial basin and affecting the right premaxilla (*Kogia* spp. and *Thalassocetus*) or both the right premaxilla and maxilla (*Praekogia*). In extant kogiids, this depression corresponds to the spermaceti chamber hosting the case in which the spermaceti organ is located (Thornton et al., 2015). The posterior half of the facial sagittal crest is not preserved in MUSM 888. In ventral view, the premaxillae are not exposed at the apex of the rostrum (a character shared with *Kogia*, *Nanokogia*, and *Scaphokogia*; for the other fossil kogiids the anterior portion of the rostrum is not known); as such, they bear no teeth.

Maxilla. In dorsal view, the maxilla reaches the anterior end of the skull (Fig. 2). As in *Nanokogia*, the dorsal expo-

Table 1. Measurements (in mm) of the skull of *Koristocetus pescei* gen. et sp. nov. (list of measurements modified after Velez-Juarbe et al., 2015).

Condylbasal length	321
Length of rostrum	144
Width of rostrum at base	107
Height of rostrum at base	54
Max width of mesorostral groove	23
Width of rostrum at mid-length	54
Height of rostrum at mid-length	34
Width across premaxillae at mid-length of rostrum	34
Width of mesorostral groove at mid-length of rostrum	19
Width of rostrum at three-quarters the length, measured from posterior end	42
Width of mesorostral groove at three-quarters the length of rostrum	22
Distance from tip of rostrum to right external naris	145
Greatest postorbital width	185e
Width of left naris	24
Width of right naris	9
Maximum width of external nares	36
Greatest width across premaxillae	56
Greatest length of temporal fossa	96
Greatest width of internal nares	38
Maximum width across occipital condyles	71
Height of foramen magnum	20
Width of foramen magnum	17

sure of the maxilla tapers transversely towards the apex of the rostrum. However, in contrast to *Nanokogia*, the maxilla is transversely wider than the premaxilla all along the rostrum. Medial and anteromedial to the antorbital notches, the dorsal surface of the maxilla is transversely convex (almost crest-like on the left side of the skull); then it flattens progressively forwards. Both antorbital notches are very narrow, slit-like (as in *Kogia*, *Nanokogia*, and *Scaphokogia*); they enter the supracranial basin (thus resembling *Kogia*, *Nanokogia*, and, at least for the right side, *Praekogia*). Lateral to the antorbital notch, the lateral maxillary crest is sharp and not transversely inflated on both sides (thus differing from those of *Kogia breviceps* and *K. sima*, resembling instead those of *Aprixokogia*, *K. pusilla*, and *Scaphokogia*). The right and left lateral maxillary crests slightly converge anteriorly and form part of the lateral borders of the supracranial basin. When viewed dorsally, the supracranial basin appears roughly trapezoidal (Fig. 2) as in *Kogia pusilla*, *Nanokogia*, and *Praekogia*, whereas it appears as more laterally and posteriorly expanded (thus appearing roughly semicircular) in *K. breviceps*, *K. sima*, and *Scaphokogia* (the outline of the supracranial basin is poorly preserved in *Praekogia* and *Thalassocetus*). As in *Kogia pusilla* and *Scaphokogia*, the supracranial basin does not significantly extend onto the dorsal surface of the rostrum (Fig. 2); in turn, in *Aprixokogia*, *Kogia breviceps*, and *K. sima*, the anterior part of the supracranial basin is open into the region above the rostrum

at both sides of the bony nares. Posterior to the antorbital notches, the maxillae form the major part of the floor of the preserved part of the supracranial basin, whose deepest region is located on the right maxilla, posterolateral to the right external naris (a condition reminiscent of what is observed in *Aprixokogia* and *Nanokogia*). In extant kogiids, the posterodorsal surface of the right maxilla hosts a wide fossa that accommodates the vocal chamber, in which the vocal cap and the phonic lips (two soft-tissue structures comprising part of the kogiid echolocation system) are situated (Thornton et al., 2015). The upturned median margin of the left maxilla contributes to build the sagittal facial crest, being exposed on the right side of this crest along its dorsal (subhorizontal) edge. The sagittal facial crest is bent leftwards, thus partially hiding the left part of the supracranial basin in dorsal view (a condition reminiscent of *Aprixokogia* and *Kogia pusilla*). Two dorsal infraorbital foramina are present on the left maxilla (Fig. 2). The posterior foramen is located posterolateral to the left external naris, roughly at mid-length between the level of the preorbital process and the level of the postorbital process of the frontal; it continues posterolaterally as a short and shallow groove. The anteriormost left dorsal infraorbital foramen is located at about one-third of the length of the rostrum near the maxilla–premaxilla suture; it is anteroposteriorly elongated, with no groove extension. Three dorsal infraorbital foramina are observed on the right maxilla (Fig. 2). The posterior foramen is situated roughly at the same level of the left posterior maxillary foramen; it is large and is followed posterolaterally for about 25 mm by an elliptical groove. The two anterior right dorsal infraorbital foramina are very small-sized and oval-shaped, located slightly posterior to the level of the left dorsal infraorbital foramen along the maxilla–premaxilla suture. In *Kogia*, the posterior dorsal infraorbital foramina are commonly subdivided (e.g., duplicated), and a similar condition is observed in *Nanokogia*, *Praekogia*, *Scaphokogia*, and *Thalassocetus*; in turn, in *Aprixokogia*, a single posterior dorsal infraorbital foramen is present on each maxilla at the level of the external nares. The palatal exposure of the maxilla is significantly transversely convex in its rostral portion (Fig. 3). Both maxillae bear circular dental alveoli starting from the posterior third of the rostrum. They are shallow, but clearly individualized where the bone surface is nicely preserved; their mean diameter is about 6 mm. Six dental alveoli have been detected on each maxilla; due to poor preservation of the ventral aspect of the anterior half of the rostrum, however, a few more alveoli could have been present. The presence of discrete dental alveoli on the maxillae of MUSM 888 is shared with *Aprixokogia* and *Scaphokogia*, whereas just a faint alveolar groove is present in *Kogia* and *Nanokogia* (condition unknown in *Praekogia* and *Thalassocetus*). We regard the dental alveoli on the maxillae of *Koristocetus* as presumably bearing functional teeth. The maxillae contact each other medially for about 15 mm posterior to the posteriormost termination of the left alveolar row. Anterior and posterior to this

area, the vomer is exposed between the maxillae. No palatine foramina have been detected on the ventral surface of the maxilla near the contact with the palatine. Posterior to the base of the rostrum, each maxilla projects between the palatine and the lacrimal as a lobe-shaped extension that forms the medial walls of the ventral infraorbital foramen (Fig. 3). This foramen is rather large and oval-shaped; it is elongated along a posteromedial to anterolateral axis, posterolateral to the choanae (i.e., anteromedial to the frontal groove). Anterior to the ventral infraorbital foramen, two shallow depressions are present on each maxilla (Fig. 3); similar features have also been observed in *Kogia* and *Nanokogia* and could be related to the anteriormost portion of the pterygoid sinus complex. This interpretation would imply a development of the pterygoid sinus beyond the main fossa outlined in *Kogia breviceps* by Fraser and Purves (1960: plate 16).

Nasal. As in all Kogiidae known to date, the nasals are absent.

Palatine. Because of the loss of the palatal part of the pterygoid, the ventral surface of the left palatine is clearly observable in MUSM 888; in turn, the right palatine is largely missing, but its outline is nonetheless inferable from the observation of the corresponding suture with the overlying right maxilla. The palatines are located posteromedially to the maxillae and anterolaterally to the choanae; they are anterolaterally bent (i.e., roughly kidney-shaped), and the left palatine is distinctly transversely convex (Fig. 3). A cusp-like projection of the maxillae is present between the palatines (a condition reminiscent of that observed in *Scaphokogia*). The left palatine contributes to the anterolateral border of the left choana.

Lacrimal + Jugal. As in all kogiids and, more generally, in all Physeteroidea, the boundary between the lacrimal and the jugal is not discernible due to complete fusion. In lateral view, the lacrimojugal complex is roughly triangular (Fig. 4a) and smaller than in *Kogia*. As observed in *Kogia*, *Nanokogia*, and *Scaphokogia*, the lacrimal contacts the maxilla forming a sigmoidal suture line. The posterodorsal corner of the lacrimal is wedged between the frontal and the maxilla. This portion of the lacrimal is posterodorsally projected as in *Kogia*, *Nanokogia*, and *Praekogia*, whereas it is distinctly posteriorly projected in *Scaphokogia*; it is shorter than in *Kogia* and *Praekogia* (where it terminates just anterior to the apex of the orbit) but longer than in *Nanokogia* and *Scaphokogia*. The ventralmost tip of the lacrimojugal complex is missing on both sides of the skull. When viewed ventrally, the lacrimojugal complex is posteromedially to anterolaterally elongated and tapers caudally (Fig. 3), thus recalling the condition observed in *Nanokogia*.

Frontal. The frontal is mostly exposed in lateral and ventral views, the dorsal face of this bone being largely covered by the maxilla. In lateral view, the suture line between the frontal and the maxilla is inclined at about 30° with respect to the horizontal plane (Fig. 4a), an angle smaller than in *Aprixokogia* and *Kogia* but similar to that observed in

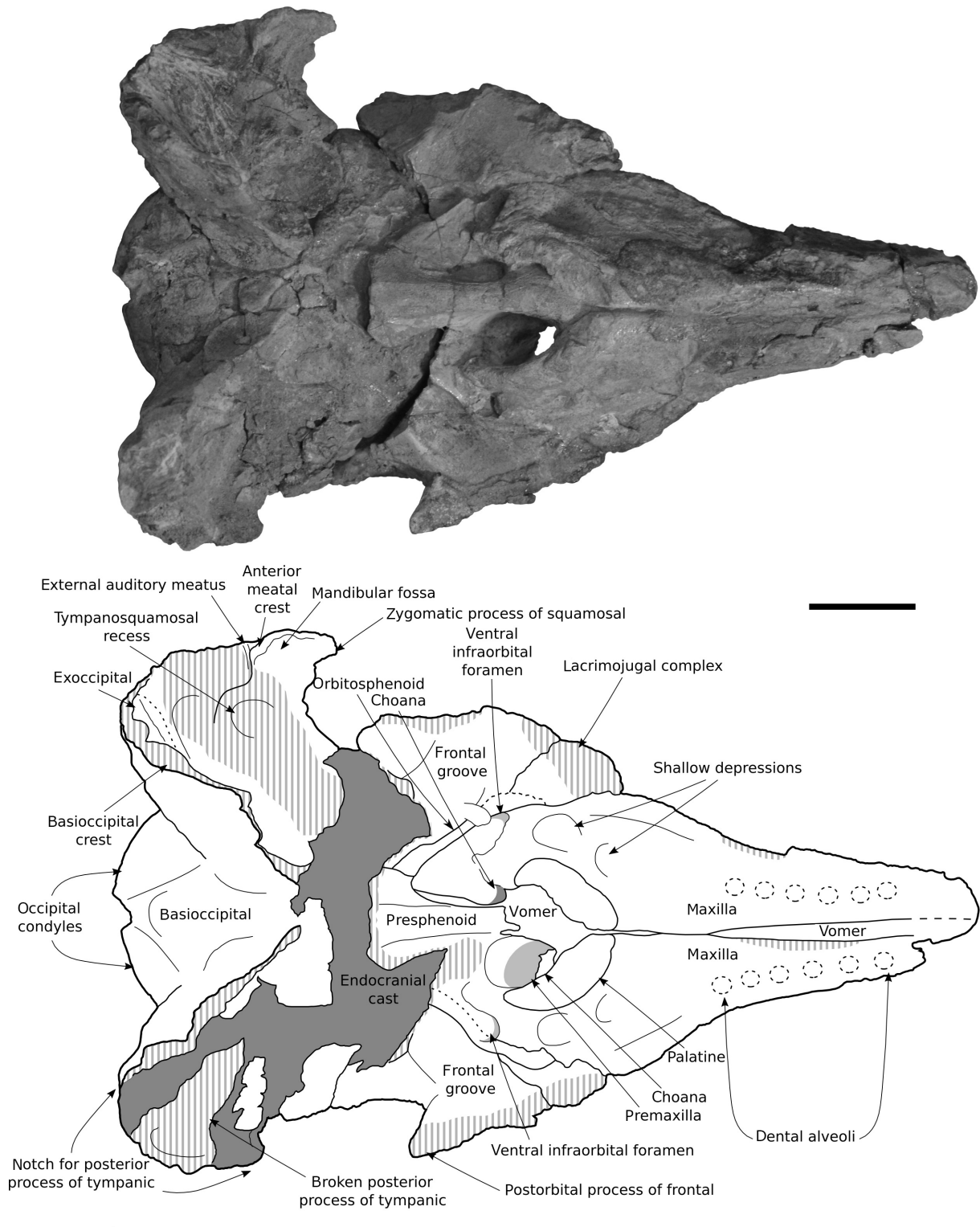


Figure 3. Ventral view of the skull MUSM 888, holotype of *Koristocetus pescei* gen. et sp. nov., Pisco Formation, late Miocene of Aguada de Lomas, southern coast of Peru. The dark-grey-shaded areas correspond to hardened sediment. The light-grey-shaded areas correspond to openings in the skull. The grey vertical stripes denote damaged areas. Scale bar: 4 cm.

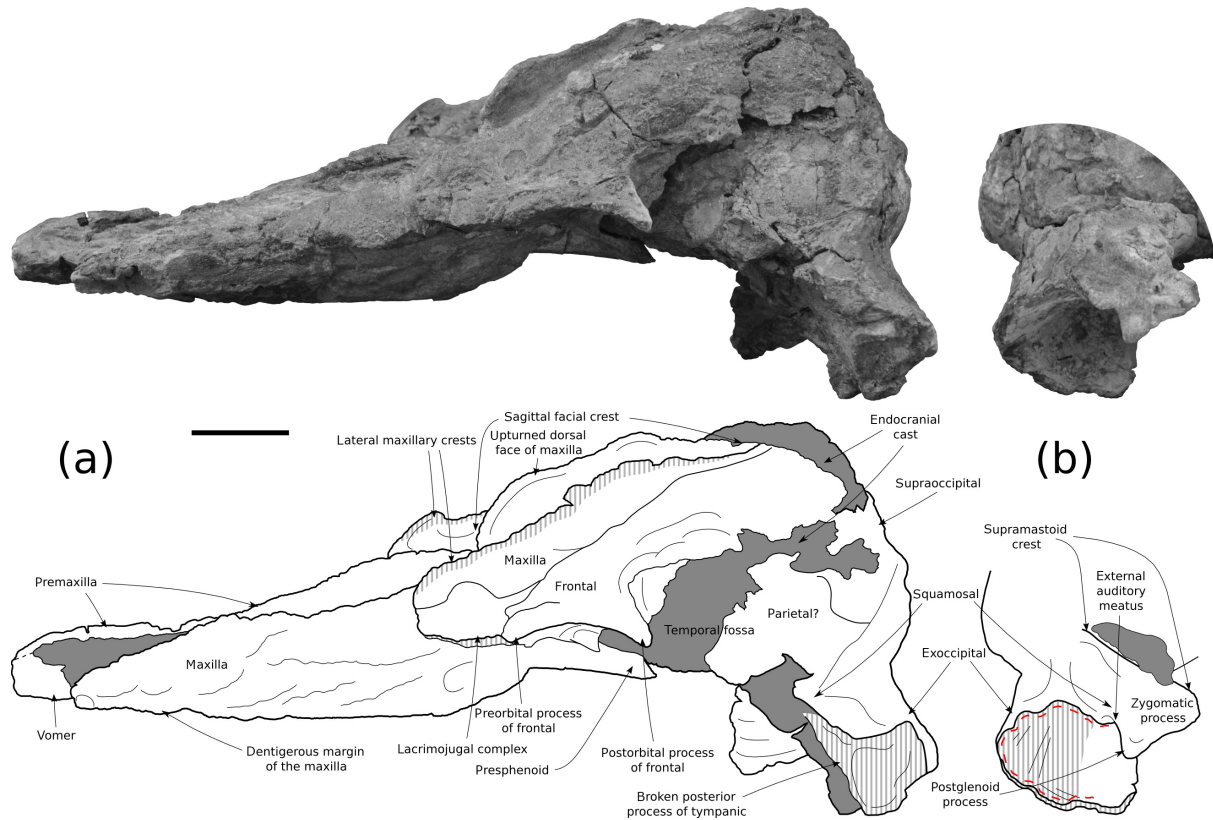


Figure 4. Lateral views of the skull MUSM 888, holotype of *Koristocetus pescei* gen. et sp. nov., Pisco Formation, late Miocene of Aguada de Lomas, southern coast of Peru. (a) MUSM 888 in left lateral view; (b) MUSM 888 in right lateral view (only a detail of the exoccipital region is shown). The dark-grey-shaded areas correspond to hardened sediment. The grey vertical stripes denote damaged areas. The red dashed line denotes the extent of the notch for the enlarged posterior process of the tympanic bulla. Scale bar: 4 cm.

Nanokogia. When viewed dorsally, the lateral edge of the supraorbital process is slightly S-shaped, being bent medially in its anterior portion and laterally in its posterior portion (Fig. 2). In lateral view, the roof of the orbit is less concave downwards than in *Kogia* and *Nanokogia*. The postorbital process is short and slender; it projects posteroventrolaterally. The postorbital process is less massive than in *Aprixokogia*, *Kogia*, and *Thalassocetus*; furthermore, it is less posteroventrally elongated than in *Kogia* and *Praekogia*. The frontal constitutes the upper part of the roof of the large temporal fossa. This fossa is large (Figs. 4a, 5a), being distinctly anteroposteriorly longer than the distance between the preorbital process of the maxilla and the anteriormost wall of the temporal fossa; with a ratio between the greatest length of the temporal fossa and the condylobasal length being ca. 0.30, MUSM 888 has a fossa longer than in *Nanokogia* (ratio in *Nanokogia* ca. 0.23), contrasting even more with the anteroposteriorly shortened temporal fossa of *Kogia* and *Thalassocetus* and resembling instead that of *Aprixokogia* and *Praekogia*. The dorsalmost end of the fossa was most likely located near the lost vertex of the skull. The ventral surface of the frontal is only partly preserved in MUSM 888. It displays a wide frontal groove following anterolaterally the

optic canal (Fig. 3). The frontal groove widens and flattens anterolaterally; it is more obliquely oriented with respect to the transverse plane than in extant *Kogia*, thus resembling *Nanokogia* and other fossil Kogiidae.

Vomer. The vomer reaches the anterior tip of the skull (Figs. 2, 3, 4a, 5a, c). In dorsal view, it forms the floor of the mesorostral groove (Fig. 2). In ventral view, the vomer is exposed along the rostrum as a narrow slit, running anteriorly between the maxillae from the point where they contact each other anterior to the palatines (Fig. 3). Posterior to this level, the vomer covers ventrally the anterior portion of the presphenoid, thus forming the anteromedial segment of the border of the inner nares; it also covers the lateral sides of this bone projecting posteriorly as a pair of slice-like bony projections. This condition recalls that observed in most Kogiidae (Velez-Juarbe et al., 2015: fig. 12) except for *Praekogia*, in which the vomer partially drapes the ventral surface of the presphenoid.

Sphenoid. In dorsal view, as in all Kogiidae, the presphenoid forms the septum that separates the external nares medially (Fig. 2). It is transversely thin, and its dorsal edge is sharp. The presphenoid does not extend anteriorly within the mesorostral groove, thus differing from many adult speci-

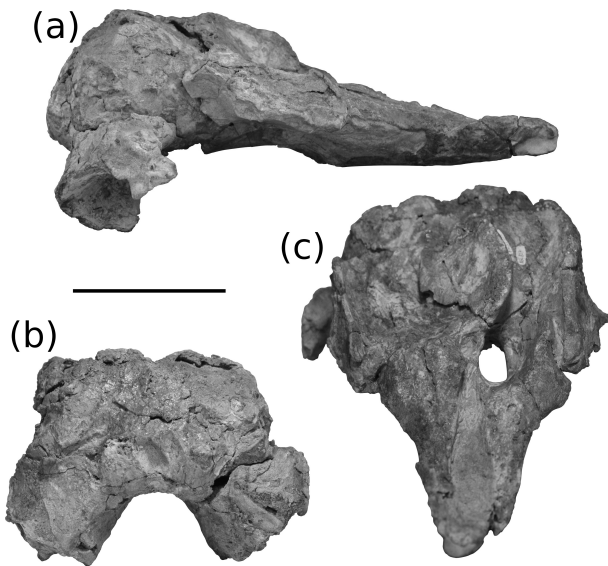


Figure 5. Additional views of the skull MUSM 888, holotype of *Koristocetus pescei* gen. et sp. nov., Pisco Formation, late Miocene of Aguada de Lomas, southern coast of Peru. (a) Skull in right lateral view; (b) skull in posterior view; (c) skull in anterodorsal view. Scale bar: 10 cm.

mens of *Kogia sima* and some specimens of *Kogia breviceps*. In ventral view, the presphenoid is partially exposed, being bordered laterally and anteroventrally by the vomer. Posterior to the inner nares, the presphenoid is robust and expands posterolaterally, thus recalling the shape of an upturned T (Fig. 3). The orbitosphenoid is probably present in MUSM 888 as a thin, elongated bony slice lying anterior to the anteromedial margin of the optic canal/frontal groove. The orbitosphenoid runs posteromedially–anterolaterally, from the posterolateral corner of the presphenoid towards the posteromedial termination of the lacrimal (Fig. 3). Among Kogiidae, the orbitosphenoid is visible in *Aprixokogia*, *Kogia*, and possibly *Scaphokogia*; apparently, it has not been observed in *Nanokogia* and *Praekogia*. Only shreds of the posteriormost portion of the alisphenoid are preserved on both sides of the skull. The basisphenoid is not preserved in MUSM 888.

Pterygoid. Most of the pterygoid is lost on both sides of the skull and only a shred of that bone is tentatively detectable in MUSM 888. It consists of a thin slice of bone located posterolateral to the right bony naris and lateral to the posterior projection of the vomer; it also contacts the frontal (anteriorly), the orbitosphenoid (posterolaterally), and the presphenoid (posteriorly).

Parietal. The parietal is not clearly identifiable but still present in MUSM 888. It likely constitutes most of the posterodorsal portion of the medial wall of the temporal fossa (see description of the frontal bone above).

Squamosal. The zygomatic process is incompletely preserved on the right side of the skull (Figs. 2, 3, 4b, 5a, c). Its proximal portion is triangular in lateral view, and

its transverse section is roughly triangular. The supramastoid crest, which represents the anterolateral continuation of the temporal crest, appears as well defined and rather salient. The mandibular fossa is shallow, anteroposteriorly elongated, and facing anteroventrally (Fig. 3). Posteromedial to the mandibular fossa the tympanosquamosal recess is a shallow fossa in which the middle sinus probably resided (Fraser and Purves, 1960: plate 16; see also Barnes, 1973); this depression is observed in *Nanokogia* and *Praekogia* and is seemingly less well defined in *Kogia*. In lateral view, the postglenoid process is robust and triangular in outline (Figs. 4b, 5a), with a long, anteroventrally directed posterior wall; it is followed medially by the sharp, spine-like anterior meatal crest. This crest is gently sigmoidal and anterolaterally–posteromedially elongated in ventral view, whereas it is generally more transversely oriented in other Kogiidae for which this feature is known. A narrow groove is observed posterior and parallel to the anterior meatal crest; it is here interpreted as the vestigial external auditory meatus (Fig. 3). Posterior and medial to this level the ventral surface of the squamosal is poorly preserved, and features such as the falciform process and the sutures between the squamosal, the exoccipital, the parietal, and the basioccipital cannot be defined. The posteroventral portion of the squamosal and the anteroventral portion of the paroccipital process of the exoccipital (see below) are shaped as a wide and deep round notch (Figs. 4b, 5a), as in *Kogia breviceps*, *K. sima*, *Nanokogia*, and *Praekogia*. A similar feature is present but distinctly shallower in *Aprixokogia* and *Scaphokogia*, whereas it is not developed in *Thalassocetus*; as in extant Kogiidae, this notch hosts the inflated posterior process of the tympanic bulla (see below).

Occipital. The basioccipital crests are known only for their posterior portions; they are salient and markedly posterolaterally diverging (Fig. 3), as observed in other Kogiidae. The foramen magnum is roughly oval-shaped (Fig. 5b), higher than wider, and proportionally smaller than in *Aprixokogia*, *Nanokogia*, and *Praekogia*. The occipital shield is only partially preserved: as observed in most Kogiidae, it forms an angle of about 60–70° with the horizontal plane. The occipital condyles are dorsoventrally elongated and separated ventrally by a shallow furrow. Dorsolateral to the condyles, the shallow dorsal condyloid fossae appear to be present. Lateral and ventrolateral to the condyles, the exoccipitals are broad and flare ventrolaterally to form the paroccipital processes. A reduced jugular notch separates the paraoccipital process of the exoccipital from the basioccipital crest. The preservation of an endocranial cast (Figs. 2, 3, 4a) allows for the reconstruction of the general shape of the braincase of MUSM 888. The two cerebral hemispheres appear as separated dorsally by a wide but shallow longitudinal sulcus, strongly reminiscent of the interhemispheric cleft.

Tympanic bulla. Part of the posterior process of the left tympanic bulla is preserved in MUSM 888 (Figs. 3, 4a). It is broken and its original shape cannot be ascertained. As in extant Kogiidae, the posterior process of the tympanic bulla is

enlarged and occupies the wide and deep notch present on the posteroventral portion of the squamosal and the anteroventral portion of the paroccipital process of the exoccipital by firmly attaching to the neurocranium.

5 Phylogeny and paleoecology

Our phylogenetic analysis resulted in six most parsimonious trees having a consistency index (CI) of 0.5379 and a retention index (RI) of 0.7185. The strict-consensus tree is shown in Fig. 6. The base of Physeteroidea is marked by a polytomy involving (i) *Eudelphis*, (ii) a clade embracing the macroraptorial physeteroid genera (i.e., *Acrophyseter*, *Brygmophyseter*, *Livyatan*, and *Zygophyseter*), and (iii) a clade formed by “*Aulophyseter*” *rionegrensis* plus the crown Physeteroidea. Differing from previous phylogenetic analyses of physeteroids (e.g., Lambert et al., 2010, 2017; Boersma and Pyenson, 2015; Velez-Juarbe et al., 2015), our consensus tree recovers all the middle to late Miocene macroraptorial sperm whales (as defined by Lambert et al., 2017) as a monophyletic group, with *Livyatan* as a sister group of *Acrophyseter* spp. Crown physeteroids consist of two clades corresponding to the families Physeteridae and Kogiidae. In agreement with Lambert et al. (2017), but differing from other recent analyses (Boersma and Pyenson, 2015; Velez-Juarbe et al., 2015), *Orycterocetus* and *Placoziphius* are here recognized as Physeteridae.

All the phylogenetic analyses of Physeteroidea including all extant and fossil kogiids published to date (Bianucci and Landini, 2006; Lambert, 2008; Lambert et al., 2008, 2010, 2017; Boersma and Pyenson, 2015; Velez-Juarbe et al., 2015) agreed on the existence of three groups among Kogiidae: (i) a paraphyletic group including early branching forms such as *Aprixokogia* and *Thalassocetus*; (ii) a lineage (subfamily Scaphokogiinae) consisting of the sole bizarre genus *Scaphokogia*; (iii) a clade (subfamily Kogiinae), sister group to *Scaphokogia*, whose members are *Kogia* spp., *Nanokogia*, and *Praekogia*. Our cladistic analysis confirms this phylogenetic structure for Kogiidae but identifies a sister group to *Scaphokogia* in MUSM 3291 and MUSM 3405, which are thus recognized as representatives of a new form of Scaphokogiinae. Although the osteoanatomical differences that separate the four genera of Kogiinae known to date are rather minor, our cladistic analysis is able to fully resolve the phylogenetic relationships among *Kogia*, *Nanokogia*, *Praekogia*, and *Koristocetus*. The latter genus is here recognized as the earliest branching member of the subfamily Kogiinae, whereas the late Tortonian *Nanokogia* from Panama, the Messinian *Praekogia* from Baja California, Mexico, and the late Pliocene to Recent, globally distributed *Kogia* represent subsequent branches.

Therefore, the late Miocene genus *Koristocetus* represents the first representative of the subfamily Kogiinae from the Southern Hemisphere, and it retains a series of archaic traits,

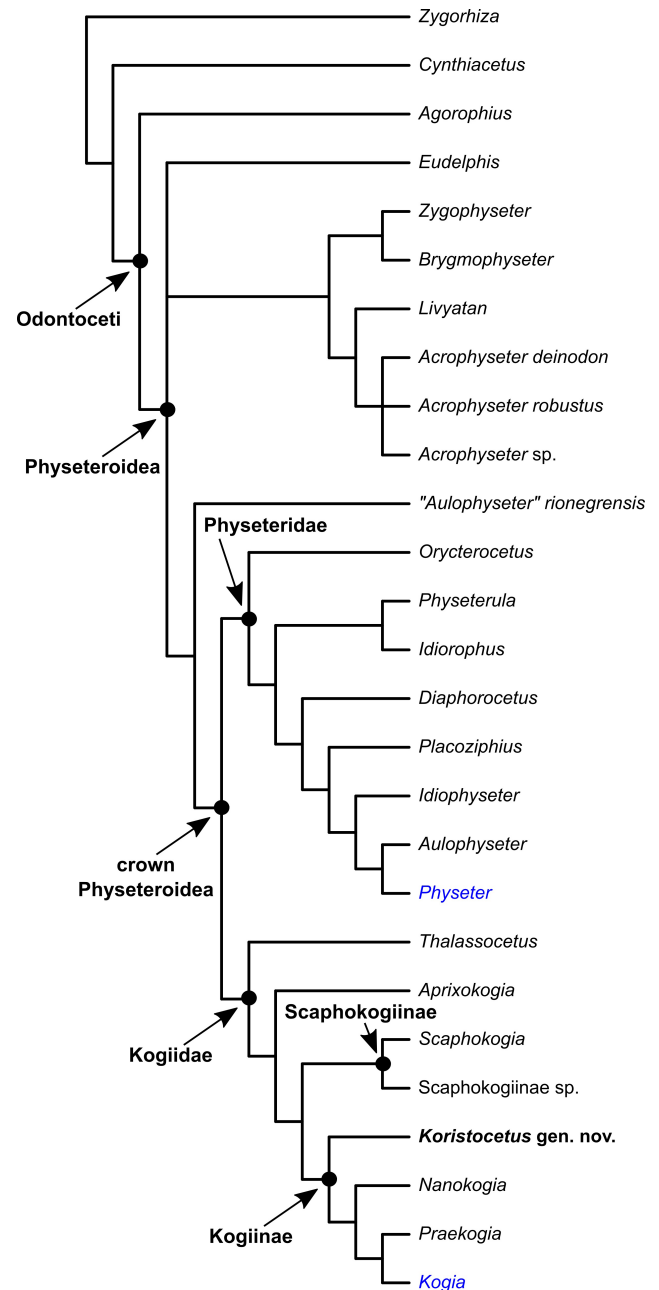


Figure 6. Consensus tree showing the phylogenetic relationships of *Koristocetus pescei* gen. et sp. nov. with other physeteroids. *Acrophyseter* sp. refers to MUSM 2182, a partial skull from the site of Cerro Los Quesos (late Miocene, Pisco Formation). Scaphokogiinae sp. refers to MUSM 3291 and MUSM 3405, two partial skulls from the sites of Cerro Los Quesos and Cerro Blanco, respectively (late Miocene, Pisco Formation), representing a new form of Kogiidae sharing similarities with *Scaphokogia cochlearis*. Extant genera are highlighted in blue.

including the presence of well-individualized dental alveoli on the upper jaws. Extant dwarf and pygmy sperm whales primarily forage on deep-water cephalopods and, secondar-

ily, on fish and crustaceans; these odontocetes exhibit a strong suction feeding specialization that relies on a peculiar craniomandibular architecture (featuring the proportionally shortest rostrum among Recent cetaceans) and on a highly derived hyoid apparatus (e.g., Caldwell and Caldwell, 1989; McAlpine, 2002; Bloodworth and Marshall, 2005; Werth, 2006; Bloodworth and Odell, 2008; Hocking et al., 2017). As reported above, *Koristocetus pescei* mainly differs from extant *Kogia* spp. by displaying a longer rostrum, a larger and more anteroposteriorly elongated temporal fossa, and a presumably functional maxillary dentition. Coupled with a rather robust and not ventrally deflected zygomatic process, these characters suggest that *Koristocetus* retained some degree of raptorial feeding ability, contrasting with the obvious suction feeding specialization observed in extant dwarf and pygmy sperm whales. The discovery of new specimens of *Koristocetus*, and especially of remains of the mandibles and hyoid apparatus, will hopefully contribute to clarify the ecology of this fossil kogiid.

During the late Miocene *Koristocetus* shared the southern coast of present-day Peru with members of the genus *Scaphokogia*, whose unique pachyostotic and downwards-deflected semicylindrical rostrum evokes a peculiar feeding/foraging specialization still to be understood. Whereas at least five skulls of *Scaphokogia* – either collected and deposited in museum collections or still in the field – are known from the late Miocene coastal marine deposits exposed at Aguada de Lomas (Alberto Collareta, personal observation), a single specimen of *Koristocetus* has been found in the same deposits (and, more generally, from the whole Pisco Formation) to date. Such an uneven distribution could reflect an intense ecospace use of the coastal area of Aguada de Lomas by members of the genus *Scaphokogia*; in turn, *Koristocetus* could have exhibited more open-ocean habits, thus recalling the habitat preferences of extant *Kogia* in this aspect (McAlpine, 2002). However, since skulls of *Scaphokogia* are much more robust and possibly less prone to degradation than those of other small-sized odontocetes such as *Koristocetus*, a preservation bias cannot be ruled out.

In conclusion, *Koristocetus* evokes a long history of high diversity, morphological disparity, and sympatric habits in fossil kogiids (as already highlighted elsewhere; Velez-Juarbe et al., 2015, 2016), thus suggesting that our comprehension of the evolutionary history of pygmy and dwarf sperm whales is still far from being exhaustive, especially considering the broad geographical distribution of extant species.

Data availability. No data sets were used in this article.

Appendix A: List of characters used in the cladistic analysis

Characters taken from Lambert et al. (2017). Modifications are highlighted in bold.

1. Rostrum length: **0**, rostrum elongated, ratio between rostrum length and skull width > 1.2 ; **1**, ratio ≤ 1.2 and ≥ 0.95 ; **2**, short rostrum, ratio < 0.95 .
2. Maxillae, premaxillae and vomer, all reaching the tip of the rostrum which is not formed only by the premaxillae: **0**, absent; **1**, present.
3. Supracranial basin of the skull: **0**, absent; **1**, present; **2**, extended onto the whole dorsal surface of the rostrum.
4. Dorsal exposure of the maxilla on the rostrum: **0**, exposure limited to less than half the rostrum length; **1**, maxilla exposed on more than half the length of the rostrum, narrower than the premaxilla at some levels; **2**, wider than the premaxilla all along.
5. Constriction of premaxilla anterior to antorbital notch followed by anterior expansion: **0**, absent, suture maxilla-premaxilla on the rostrum roughly anteriorly directed; **1**, present, suture maxilla-premaxilla distinctly anterolaterally directed.
6. Upper tooth row: **0**, deep alveoli; **1**, alveoli shallow or absent.
7. Premaxillary teeth: **0**, present; **1**, absent. This character cannot be coded for taxa lacking distinct upper alveoli.
8. Maximum width of skull (postorbital or bizygomatic width): **0**, < 40 cm; **1**, ≥ 40 and < 60 cm; **2**, ≥ 60 and < 100 cm; **3**, ≥ 100 cm.
9. Antorbital notch: **0**, absent; **1**, present; **2**, transformed into a very narrow slit.
10. Right antorbital notch: **0**, outside the supracranial basin; **1**, inside the supracranial basin.
11. Number and size of dorsal infraorbital foramina, in the area of the right antorbital notch and posteriorly: **0**, small to moderate size foramina, at least three-four; **1**, three large foramina; **2**, two large foramina; **3**, one large foramen (maxillary incisure).
12. Right premaxilla: **0**, posteriorly extended as the left premaxilla; **1**, more posteriorly extended than the left premaxilla.
13. Right premaxilla: **0**, not widened posteriorly; **1**, posterior extremity of the right premaxilla laterally widened, occupying at least one third of the width of the supracranial basin, mostly on the right side.
14. Presence of a sagittal crest: **0**, absent; **1**, present as a shelf covered by the pointed right premaxilla.
15. Left premaxillary foramen very small or absent: **0**, absent (i.e. foramen present and not reduced); **1**, present.
16. Increase in size of the right premaxillary foramen: **0**, absent, ratio between width of foramen and width of premaxilla at that level ≤ 0.20 ; **1**, present, ratio > 0.20 .
17. Anteroposterior level of right premaxillary foramen: **0**, distinctly anterior to antorbital notch; **1**, slightly anterior to antorbital notch; **2**, same level or posterior to antorbital notch.
18. Asymmetry of the bony nares: **0**, absent or reduced; **1**, strong, left bony naris significantly larger than right naris.
19. Lack of nasals: **0**, both nasals present; **1**, one nasal absent; **2**, both nasals absent.
20. Widening of the supracranial basin on the right side: **0**, absent; **1**, present, basin overhangs the right orbit.
21. Right maxilla reaching the sagittal plane of the skull on the posterior wall of the supracranial basin: **0**, absent; **1**, present.
22. Fusion of lacrimal and jugal: **0**, absent; **1**, present.
23. Projection of the lacrimal-jugal between frontal and maxilla: **0**, short or absent; **1**, long.
24. Dorsoventral level of the antorbital process of the frontal: **0**, higher than the lateral margin of rostrum base; **1**, at approximately the same level; **2**, considerably lower.
25. Frontal-maxilla suture, with skull in lateral view: **0**, forming an angle $< 15^\circ$ from the axis of the rostrum; **1**, $15-35^\circ$; **2**, $> 35^\circ$.
26. Temporal fossa: **0**, anteroposteriorly longer than distance between preorbital process of the maxilla and anterior wall of temporal fossa; **1**, approximately same length; **2**, distinctly shorter.
27. Zygomatic process of squamosal in lateral view: **0**, 'L'-shaped with dorsal margin ventrally bending in its posterior portion; **1**, triangular, with dorsal margin dorsally bending in its posterior portion.
28. Postglenoid process of the squamosal: **0**, significantly ventrally longer than post-tympanic process; **1**, roughly same ventral extent as post-tympanic process.

29. In lateral view of the skull, wide notch posterior to the postglenoid process of the squamosal for the enlarged posterior process of the tympanic: **0**, absent; **1**, present but only partially developed, paraoccipital concavity moderately excavated; **2**, present and well developed, paraoccipital concavity transformed in a wide and deep notch.
30. Occipital shield: **0**, convex and forming an angle of about 40° from the axis of the rostrum; **1**, as state 0 with an angle of about 60°; **2**, flat or concave forming an angle of about 90°; **3**, flat or concave forming an angle distinctly greater than 90°.
31. Falciform process of the squamosal: **0**, contacting the corresponding pterygoid; **1**, forming a thin plate not contacting the pterygoid; **2**, reduced to a simple peg or absent.
32. Anterior bullar facet of the periotic: **0**, very anteroposteriorly elongated; **1**, reduced; **2**, absent or very small.
33. Posterior extension of the posterior process of the periotic parallel to the general plane of the bone and not ventrally orientated: **0**, absent; **1**, present.
34. Accessory ossicle of the tympanic bulla: **0**, absent or small; **1**, enlarged and partially fused with the anterior process of the periotic.
35. Involucrum of the tympanic bulla with an evident central concavity, visible in ventral and medial views, due to the marked pachyostosis of its anterior and posterior portion: **0**, absent; **1**, present.
36. Size of teeth (greatest transverse diameter of root expressed as percentage of the maximum width of skull): **0**, < 5 %; **1**, > 5 %. Considering the strong heterodonty in *Cynthiacetus* and *Zygorhiza* this character is restricted to single-rooted teeth.
37. Loss of dental enamel: **0**, absent; **1**, present.
38. Number of mandibular teeth: **0**, 11; **1**, 12–14; **2**, > 14.
39. Labiolingual compression of the posterior lower teeth (portion out of the alveolus): **0**, strong; **1**, weak or absent.
40. Ventral position of the mandibular condyle: **0**, absent, well developed angular process; **1**, present, angular process low or absent.
41. Anteroposterior level of last upper alveolus or posterior end of vestigial alveolar groove: **0**, posterior to antorbital process; **1**, at level of antorbital notch or slightly anterior; **2**, distinctly anterior to the notch.
42. Lateral margin of the supraorbital process of the maxilla: **0**, dorsoventrally thin; **1**, significantly dorsoventrally thickened, making a subvertical wall.
43. Postorbital process of the frontal: **0**, moderately posteroventrally extended; **1**, much ventrally extended (vertical length of process equal or greater than horizontal length of orbit), with a correspondingly low position of the zygomatic process of the squamosal.
44. Height of temporal fossa: **0**, dorsal margin at top of skull or somewhat lower; **1**, much lower, temporal fossa making less than half the skull height.
45. Contact between jugal and zygomatic process of squamosal: **0**, anteroposteriorly long contact; **1** proportionally short, more rounded contact; **2**, no contact. In specimens with no jugal preserved, the contact surface can sometimes be observed on the zygomatic process (e.g., *Orycterocetus crocodilinus* USNM 22926).
46. Length of the zygomatic process of the squamosal (horizontal length from anterior tip to posterior margin of squamosal): **0**, ratio between length of the process and bizygomatic width of skull > 0.35; **1**, ratio < 0.35.
47. Medial to tympanosquamosal recess, deep and rectilinear narrow groove in ventral surface of squamosal, from spiny process area to temporal fossa: **0**, absent or shallow and poorly delineated; **1**, present.
48. Dorsal process of the periotic: **0**, dorsally extended and anteroposteriorly long; **1**, anteroposteriorly shorter, but dorsally extended beyond the medial margin of the internal acoustic meatus; **2**, dorsally short.
49. Posteromedial outline of the pars cochlearis in dorsal view: **0**, angular; **1**, flattened, barely convex, and roughly continuous with posterior margin of dorsal process.
50. Curvature of the mandible in lateral view: **0**, absent or reduced, ventral margin roughly rectilinear or rising moderately anterodorsally; **1**, conspicuous, ventral margin distinctly convex rising both posterodorsally and anterodorsally; **2**, present, ventral margin concave.
51. Symphyseal angle on the mandibles: **0**, < 35°; **1**, 35–55°; **2**, > 55°.
52. Lateral margin of atlas: **0**, roughly rectilinear or laterally concave; **1**, convex, with laterally pointed transverse process at mid-height of the bone. Not applicable to *Kogia* (single block of cervical vertebrae).
53. Notch in the anterior margin of the basihyal: **0**, wide and shallow notch; **1**, narrow and deep notch; **2**, no notch, rectilinear or convex anterior margin.

Appendix B: Character/taxon matrix

Data matrix of 53 characters for three outgroups (*Agorophius*, *Cynthiacetus*, and *Zygorhiza*) and 24 extinct and extant physeteroid taxa (*Eudelphis*, *Zygodphyseter*, *Brygmophyseter*, *Acrophyseter deinodon*, *Acrophyseter robustus*, *Acrophyseter* sp., *Livyatan*, “*Aulophyseter*” *rionegrensis*, *Orycterocetus*, *Physeterula*, *Idiorophus*, *Diaphorocetus*, *Placoziphius*, *Aulophyseter*, *Physeter*, *Idiophyseter*, *Thalassocetus*, *Aprixokogia*, *Scaphokogia*, *Praekogia*, *Kogia*, *Koristocetus*, *Nanokogia*, and *Scaphokogiinae* sp.). *Acrophyseter* sp. refers to MUSM 2182, a partial skull from the site of Cerro Los Quesos (late Miocene, Pisco Formation). *Scaphokogiinae* sp. refers to MUSM 3291 and MUSM 3405, two partial skulls from the sites of Cerro Los Quesos and Cerro Blanco, respectively (late Miocene, Pisco Formation), representing a new form of Kogiidae sharing similarities with *Scaphokogia cochlearis*. Extant genera are underlined. All characters are unordered. 0, primitive state; 1, 2, 3, derived states; a, variable between 0 and 1; b, variable between 1 and 2; c, variable between 1, 2, and 3; n, character inapplicable; ?, missing character.

	5	10	15	20	25	30	35	40	45	50	53
<i>Zygorhiza</i>	00000	0000n	?000n	nn00n	n000n	00002	00000	00000	0n000	00000	?0?
<i>Cynthiacetus</i>	00000	0010n	?000n	nn00n	n000n	00002	01000	10000	0n000	00?00	000
<i>Agorophius</i>	??010	0?01n	00000	0000n	0?010	00000	?????	?0???	1000?	0????	???
<i>Eudelphis</i>	101?0	0?110	01??0	1?1?0	?????	?100?	1????	00???	?????	?0???	???
<i>Zygodphyseter</i>	10110	01210	21101	12111	?1001	0110?	12?11	10101	100?0	0?2?0	11?
<i>Brygmophyseter</i>	??1??	0?210	21101	??1??	????1	01101	????1	101?1	???0?	????0	??1
<i>Acro. deinodon</i>	20110	001?0	3?100	111?1	0??02	01101	?2011	10101	1??0?	01201	1??
<i>Acro. robustus</i>	??110	0?020	2110?	01101	01002	01101	????1?	10?21	10?00	01??1	?11
<i>Acrophyseter</i> sp.	00110	00120	????0	??1??	?????	0110?	?????	10?01	1?0??	01??1	???
<i>Livyatan</i>	21211	01320	2?1?0	????0	?1?12	01?0?	?????	10011	10?0?	1????	1??
' <i>Aulo.</i> ' <i>rionegr.</i>	?0110	00110	11100	01110	0?011	01101	?????	??2?1	2001?	0????	???
<i>Orycterocetus</i>	10110	00110	11101	01110	1?011	11102	22011	012??	20?00	1011?	???
<i>Physeterula</i>	??10?	0?210	21101	??1?0	0??11	?1102	2????	0121?	?000?	?0??0	1??
<i>Idiorophus</i>	00110	00210	?????	?????	?????	?????	?????	0121?	?0?0?	?????	???
<i>Diaphorocetus</i>	??110	0?120	?1101	11?0?	?????	1110?	?????	??2??	20?1?	10???	???
<i>Placoziphius</i>	?0120	??110	?1101	11?10	??011	?110?	2????	?????	20???	?0???	?0?
<i>Aulophyseter</i>	10100	1n210	21101	02110	01022	21102	?201?	01???	200??	1010?	???
<i>Physeter</i>	a022a	1n310	31101	02110	01022	21102	22011	0121a	20011	10110	102
<i>Idiophyseter</i>	?????	1?1?0	2110?	121?0	?????	1???	?????	?????	?????	?????	???
<i>Thalassocetus</i>	?????	??010	1101?	???	1???	?1101	?????	?????	?011?	?????	???
<i>Aprixokogia</i>	??2?0	0?010	01111	021?0	?1121	01112	2????	?????	1110?	10???	???
<i>Scaphokogia</i>	?1111	1n020	01011	02120	11121	a1013	?211?	?????	1100?	1????	???
<i>Praekogia</i>	?????	??021	0101?	??120	1?1?1	11021	2????	?????	?111?	10???	???
<i>Kogia</i>	21220	1n021	01011	0b120	11122	2102b	22111	01c1a	21112	10212	2?2
<i>Koristocetus</i> gen nov.	21120	01021	31011	??120	?11?1	01121	?????	?????	2100?	1????	???
<i>Nanokogia</i>	21110	1n021	01011	??120	11001	01021	2????	0?11?	11012	10?0?	1??
<i>Scaphokogiinae</i> sp.	?1111	01020	01011	02120	11121	?????	?????	?????	2100?	?????	???

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