

# *Arktocara yakataga*, a new fossil odontocete (Mammalia, Cetacea) from the Oligocene of Alaska and the antiquity of Platanistoidea

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## ABSTRACT

The diversification of crown cetacean lineages (i.e., crown Odontoceti and crown Mysticeti) occurred throughout the Oligocene, but it remains an ongoing challenge to resolve the phylogenetic pattern of their origins, especially with respect to stem lineages. One extant monotypic lineage, *Platanista gangetica* (the Ganges and Indus river dolphin), is the sole surviving member of the broader group Platanistoidea, with many fossil relatives that range from Oligocene to Miocene in age. Curiously, the highly threatened *Platanista* is restricted today to freshwater river systems of South Asia, yet nearly all fossil platanistoids are known globally from marine rocks, suggesting a marine ancestry for this group. In recent years, studies on the phylogenetic relationships in Platanistoidea have reached a general consensus about the membership of different sub-clades and putative extinct groups, although the position of some platanistoid groups (e.g., Waipatiidae) has been contested. Here we describe a new genus and species of fossil platanistoid, *Arktocara yakataga*, gen. et sp. nov. from the Oligocene of Alaska, USA. The type and only known specimen was collected from the marine Poul Creek Formation, a unit known to include Oligocene strata, exposed in the Yakutat City and Borough of Southeast Alaska. In our phylogenetic analysis of stem and node-based Platanistoidea, *Arktocara* falls within the node-based sub-clade Allodelphinidae as the sister taxon to *Allodelphis pratti*. With a geochronologic age between ~29–24 million years old, *Arktocara* is among the oldest crown Odontoceti, reinforcing the long-standing view that the diversification for crown lineages must have occurred no later than the early Oligocene.

**Subjects** Ecology, Evolutionary Studies, Marine Biology, Paleontology, Zoology

**Keywords** Cetacea, River dolphins, Fossil record, Neogene, Evolution, Platanistoidea, Oligocene, Pan-Platanista

## INTRODUCTION

Multiple lines of evidence point to the Oligocene epoch as an important period for the origin and early evolutionary history of crown group Cetacea (*Fordyce, 2003*). This interval, from about ~34 to ~23 million years ago, represents the origin of all extant lineages of cetaceans,

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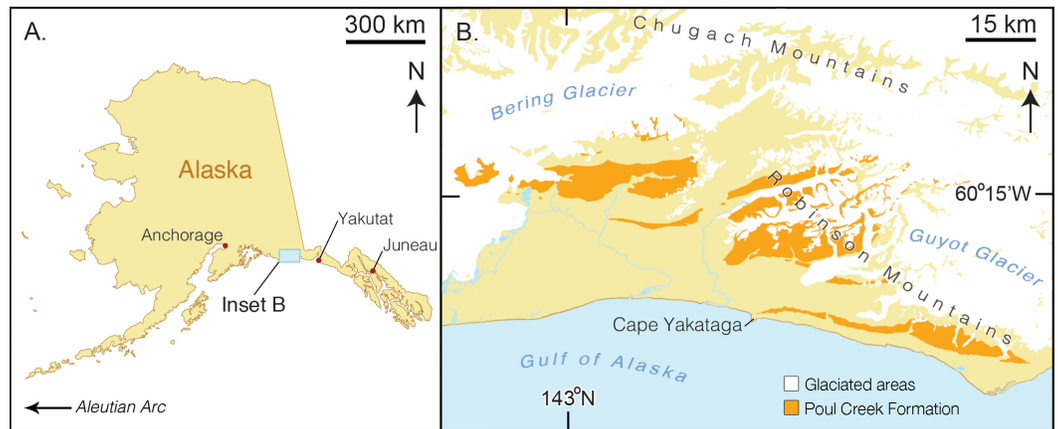
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including crown members of Mysticeti and Odontoceti, as inferred from molecular clock divergence estimates (McGowen, Spaulding & Gatesy, 2009), and fossil data (Geisler et al., 2011; Marx & Fordyce, 2015). Fossil cetaceans from this time are relatively less well known than Neogene ones for several reasons, including a lack of available rock outcrops, and insufficient study and publication on material housed in museum and personal collections (Uhen & Pyenson, 2007). The description of new cetacean taxa from the Oligocene can therefore be significant in resolving phylogenetic patterns of divergences among crown and stem groups, especially within Odontoceti.

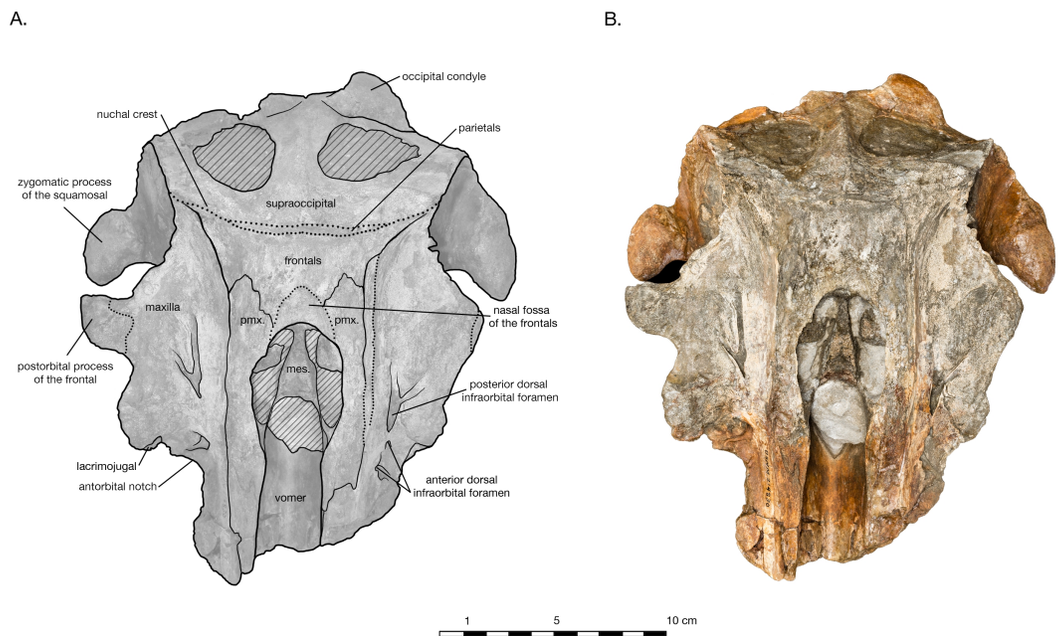
Oligocene fossil cetaceans have played an important role in understanding the evolutionary history of Platanistoidea, once a large group of cosmopolitan marine odontocetes, but now represented by only one freshwater river species: *Platanista gangetica* (Lebeck, 1801), found in the Indus, Ganges-Brahmaputra-Megna and Karnaphuli-Sangu river systems of Southeast Asia. The concept of Platanistoidea has changed drastically since it was first defined by Simpson (1945) to include only one nominal family, Platanistidae, consisting of the four extant river dolphin lineages (*Platanista* Wagler, 1830, *Inia* D'Orbigny, 1834, *Lipotes* Miller, 1918, and *Pontoporia* Gray, 1846) and their closest fossil relatives. Muizon (Muizon, 1984; Muizon, 1987; Muizon, 1988a) later suggested a polyphyletic interpretation of the river dolphin lineages, modifying Simpson's (1945) concept of Platanistoidea to include *Platanista* as the only crown taxon, with *Inia*, *Lipotes* and *Pontoporia* more closely related to Delphinoidea. This concept is consistent with the results from more recent phylogenetic analyses, especially those using molecular datasets (see Geisler et al., 2011 for a review).

Currently, phylogenetic studies of Platanistoidea have reached a general consensus about the inclusion three groups (two of which are completely extinct): Allodelphinidae, Squalodelphinidae and Platanistidae (including extant *Platanista*). The inclusion of Squalodontidae and Waipatiidae in Platanistoidea, as suggested by Muizon (1984), Muizon (1987), Muizon (1988a), Muizon (1994) and Fordyce (1994), have been more heavily debated (Lambert, Bianucci & Urbina, 2014; Tanaka & Fordyce, 2015a). Until now, no comprehensive phylogenetic analysis has used a taxonomic sampling that includes all putative platanistoid lineages, along with appropriate outgroups, such as Delphinida and stem Odontoceti (Tanaka & Fordyce, 2015a; Kimura & Barnes, 2016; Lambert, Bianucci & Urbina, 2014; Geisler & Sanders, 2003; Geisler et al., 2011).

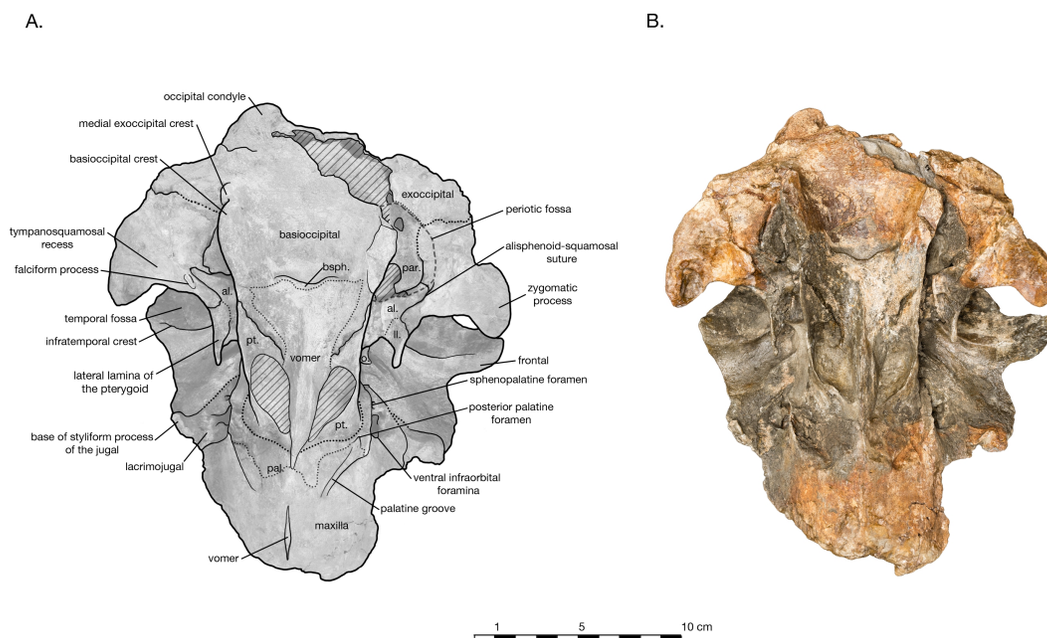
Here we describe *Arktocara yakataga*, a new genus and species of Allodelphinidae, collected in 1951 from the Poul Creek Formation in the Yakutat City and Borough of Southeast Alaska (Figs. 1–6). *Arktocara yakataga* is the most northern platanistoid yet reported, and with an estimated mid-Oligocene age (possibly Rupelian to Chattian), it is both the oldest allodelphinid and among the oldest crown Odontoceti known. The results of our phylogenetic analysis continue to support a monophyletic Platanistoidea, along with traditional sub-clades (including a monophyletic Allodelphinidae), which underscores the importance of Oligocene cetaceans in documenting the early diversification of crown Cetacea.



**Figure 1** Map of type locality for *Arktocara yakataga* (USNM 214830). (A) a map of the state of Alaska, showing major Alaskan cities. (B) simplified geologic map of the Yakutat City and Borough based on the USGS 1971 map by Don J. Miller (available at [http://ngmdb.usgs.gov/Prodesc/proddesc\\_9402.htm](http://ngmdb.usgs.gov/Prodesc/proddesc_9402.htm)). All exposures of the Poul Creek Formation in the Yakutat City and Borough (orange), are potential type localities for *Arktocara yakataga* (USNM 214830). Yellow represents exposures from all other lithological units, not mapped here.



**Figure 2** Skull of *Arktocara yakataga* (USNM 214830) in dorsal view. (A) Illustrated skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. (B) photograph of skull in dorsal view, photography by James Di Loreto, Smithsonian Institution.



**Figure 3** Skull of *Arktocara yakataga* (USNM 214830) in ventral view. (A) Illustrated skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures, dashed lines indicate broad morphological features, and hatched lines indicate sediment obscuring the fossil. (B) photograph of skull in ventral view, photography by James Di Loreto, Smithsonian Institution.

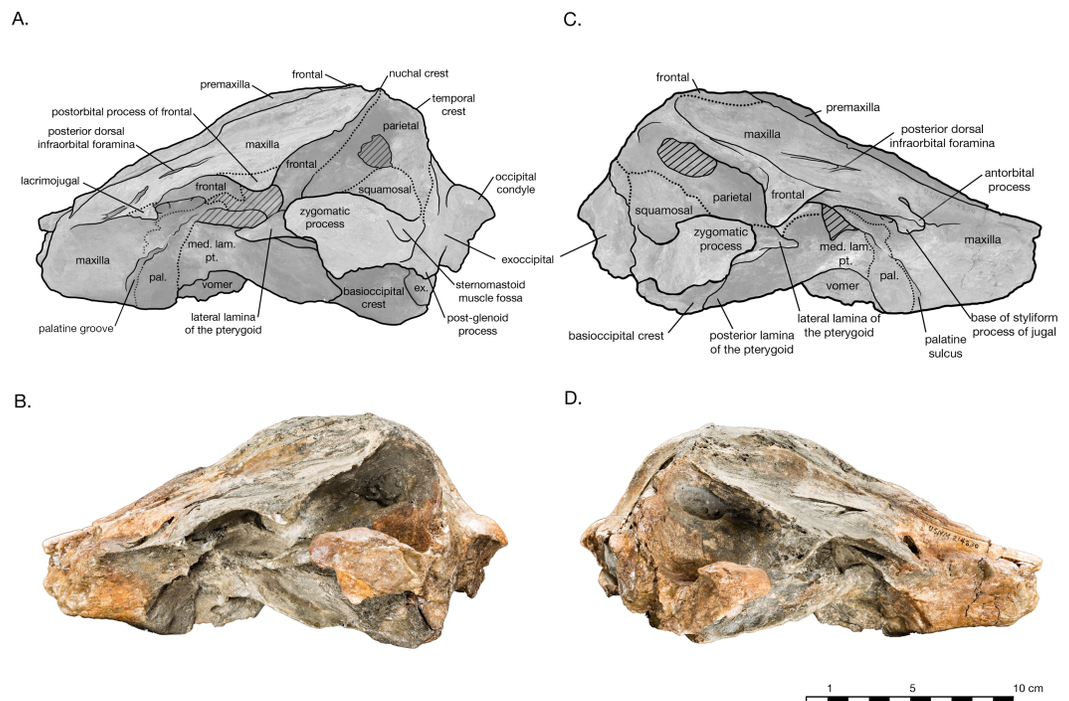
## MATERIALS AND METHODS

### Digital methods

The holotype of *Arktocara yakataga* was scanned using Nikon Metrology's combined 225/450 kV microfocus X-ray and computed tomography (CT) walk-in vault system at Chesapeake Testing in Belcamp, Maryland, USA. Using this vault CT scanner system, we collected CT slices at 0.63 mm, resulting in three-dimensional reconstruction increments of 0.30 mm. We mounted the holotype skull vertically in the vault CT scanner system, with the posterior side down to minimize scanning width. Also, we collected CT scan data for the right periotic (YPM 13408) of *Allodelphis pratti* Wilson, 1935 using their Nikon Metrology's 225 kV microfocus X-ray CT cabinet system. The DICOM files that this produced were processed in Mimics (Materialise NV, Leuven, Belgium) to create a 3D model of the *Arktocara* cranium that will be available for viewing and download on the Smithsonian X 3D website (<http://3d.si.edu>). These 3D files, along with the original DICOM files, are also archived at Zenodo (<http://zenodo.org>) at the following DOI: [10.5281/zenodo.51363](https://doi.org/10.5281/zenodo.51363). While the CT data were useful for making 3D models of the holotype, the density of the fossil material prevented the CT scanner from clearly determining internal morphology of the skull, failing to clarify indistinct sutures between skull elements.

### Phylogenetic analysis

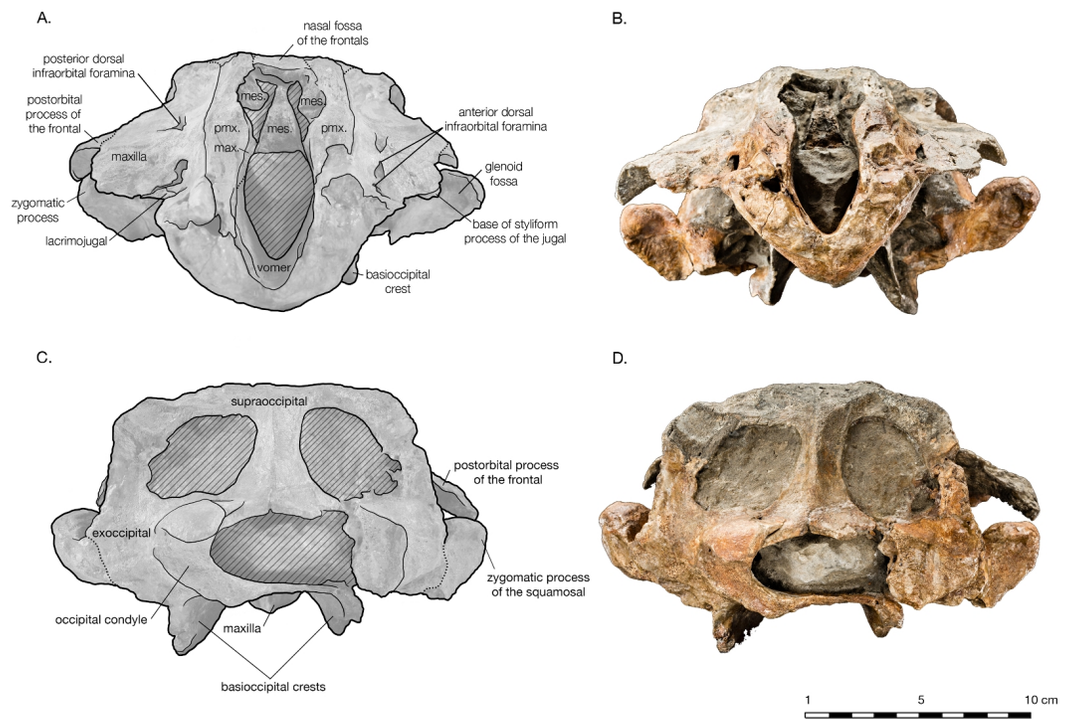
We tested the phylogenetic placement of *Arktocara yakataga* using Tanaka & Fordyce (2015a)'s Odontoceti matrix, adapted from Murakami et al.'s (2012) original version.



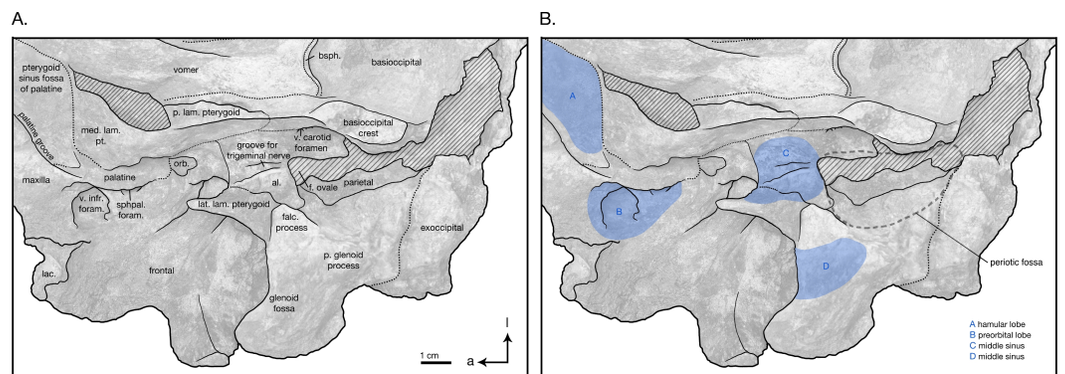
**Figure 4** Skull of *Arktocara yakataga* (USNM 214830) in left (A, B) and right (C, D) lateral views. (A) Illustrated left lateral view of skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. (B) photograph of skull in left lateral view, photography by James Di Loreto, Smithsonian Institution. (C) Illustrated right lateral view of skull with low opacity mask, interpretive line art, and labels for skull elements. (D) photograph of skull in right lateral view, photography by James Di Loreto, Smithsonian Institution.

*Tanaka & Fordyce (2015a)*'s version of this matrix consisted of 292 morphological characters and 83 operational taxonomic units (OTUs), including the fossil platanistoids *Notocetus vanbenedeni* *Moreno, 1892*, *Phocageneus venustus* *Leidy, 1869*, *Squalodon calvertensis* *Kellogg, 1923*, *Waipatia maerewhenua* *Fordyce, 1994*, *Zarhachis flagellator* *Cope, 1868*, and the extant *Platanista gangetica*. The stem cetacean taxon *Georgiacetus vogtlensis* was used as the outgroup. We removed an undescribed specimen (OU 22125), and added 4 allodelphinid taxa (*Zarhinocetus errabundus* (*Barnes, 2010*) *Goedertius oregonensis* *Kimura & Barnes, 2016*, *Allodelphis pratti* and *Arktocara yakataga*), raising the number of OTUs used in the analysis to 86. We also added codings for four periotic characters (288–291) for *Pomatodelphis inaequalis* *Allen, 1921* and *Zarhachis flagellator*, based on material available at the Smithsonian Institution (see 'Specimens observed'). We did not add codings for taxa that we could not directly observe, and therefore some platanistoid taxa were excluded from the analysis, including *Huaridelphis raimondii* *Lambert, Bianucci & Urbina, 2014*, *Ninjadelpheis ujiharai* *Kimura & Barnes, 2016*, and *Allodelphis woodburnei* *Barnes & Reynolds, 2009*.

We performed a cladistic search in TNT\* (Tree analysis using New Technology) using all characters as unordered and equally weighted. We then conducted subsequent statistical support analyses by searching for successively longer trees to calculate bremer decay indices



**Figure 5** Skull of *Arktocara yakataga* (USNM 214830) in anterior (A, B) and posterior (C, D) views. (A) Illustrated skull in anterior view with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. (B) photograph of skull in anterior view, photography by James Di Loreto, Smithsonian Institution. (C) Illustrated skull in posterior with low opacity mask, interpretive line art, and labels for skull elements. (D) photograph of skull in posterior view, photography by James Di Loreto, Smithsonian Institution.



**Figure 6** Skull details & sinus system of *Arktocara yakataga* (USNM 214830). (A) Illustrated detail of right ventrolateral skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. Arrows indicate anatomical direction, with a, anterior and l, left lateral. (B) Illustrated detail of right ventrolateral skull with low opacity mask, interpretive line art, and blue highlighted areas indicating fossa for lobes of the sinus system.

and 100 bootstrap replicates. The complete matrix in .txt format, as well as a description of character states (Table S1 and Appendix S1) are available in Supplemental Information.

### Phylogenetic nomenclature

Following Pyenson *et al.* (2015), we converted several existing, ranked cetacean taxonomic names into clade names that conform to the principles articulated first by Joyce, Parham & Gauthier (2004). We referred traditional names to more inclusive clades (e.g., in the case of extinct families, node-based clade names), where their composition closely resembles current or widely recognized name applications. For these purposes, we used the abbreviation CCN for Converted Clade Name. Below, we clarify our precise definitions for these clades (see PhyloCode, 2014, Article 9.3; Cantino & De Queiroz, 2014), and we also provide full citations for the names of specifier species.

### Nomenclature acts

The electronic edition of this article conforms to the requirements of the amended International Code of Zoological Nomenclature, and hence the new names contained herein are available under that Code from the electronic edition of this article. This published work and the nomenclatural acts it contains have been registered in ZooBank, the online registration system for the ICZN. The ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed through any standard web browser by appending the LSID to the prefix “<http://zoobank.org/>”. The LSID for this publication is: urn:lsid:zoobank.org:pub:0194A593-DBE0-47CA-A41F-04A37931BA2F. The electronic edition of this work was published in a journal with an ISSN, and has been archived and is available from the following digital repositories: PubMed Central, and LOCKSS.

### Specimens observed

*Allodelphis pratti* (YPM 13408); *Allodelphis* sp. (USNM 266608, 256609, 256610); *Goedertius oregonensis* (LACM 123887); *Goedertius* sp. (USNM 335406, 335765, 13673, 314421); *Notocetus* sp. (USNM 206286); *Phocageneus venustus* (USNM 21039, 475496); *Phocageneus* sp. (USNM 182939, 362125); *Platanista gangetica* (USNM 23456); *Pomatodelphis bobengi* Case, 1934 (299775); *Pomatodelphis* sp. (USNM 360054); *Squalodon calvertensis* (USNM 10949, 529246); cast of *Waipatia maerewhenua* (USNM 508061); *Zarhachis flagellator* (USNM 299945, 10911, 13768); *Zarhachis* sp. (USNM 214759, 24868); cast of *Zarhinocetus errabundus* (USNM 526600); *Zarhinocetus errabundus* (USNM 11573, 25425).

## RESULTS

### Systematic paleontology

Cetacea *Brisson, 1762*

Odontoceti *Flower, 1867 sensu Fordyce & Muizon, 2001*

Platanistoidea (CCN) (node-based version of *Fordyce, 1994*)

Allodelphinidae (CCN) (node-based version of *Barnes, 2006*)

*Arktocara*, gen. nov. LSID: urn:lsid:zoobank.org:act:EE11B95B-8338-496B-97F4-1673ED90E709

**Definitions.** Crown group *Platanista* refers to the crown clade arising from the last common ancestor of all lineages descending from *Platanista*, including two subspecies of *Platanista gangetica* (*P. g. gangetica* (Lebeck, 1801) and *P. g. minor* Owen, 1853), as recognized by *The Society for Marine Mammology' Committee on Taxonomy* (2015).

Platanistoidea is a converted clade name for a clade that includes *Platanista gangetica* and fossil taxa that support familial level taxonomic concepts such as: *Allodelphis pratti*; *Squalodelphis fabianii* Dal Piaz, 1917; and *Waipatia maerewhenua*. We do not formally recognize node-based versions of Squalodelphinidae and Waipatiidae at this time (except for in the Diagnosis section) because these familial level groupings are not the explicit focus of this study, and we defer to future work that can better substantiate their taxonomic scope and better test their monophyly (see, for example, Tanaka & Fordyce, 2014; Tanaka & Fordyce, 2015a). This node-based converted clade of Platanistoidea corresponds to the Fordyce (1994)'s concept of Platanistoidea, but differs from Muizon (1987), and Muizon (1991)'s concept, in its exclusion of Squalodontidae. Following Lambert, Bianucci & Urbina (2014), we exclude Squalodontidae and stem Platanistoidea, such as *Prosqualodon davidis* Flynn, 1923, and *Papahu taitapu* Aguirre-Fernández & Fordyce, 2014, from our node-based concept of Platanistoidea. Our concept is more inclusive than Geisler et al. (2011)'s Platanistoidea, which included only *Platanista*, *Zarhachis* and *Squalodelphis*, while excluding *Waipatia* to outside of crown Odontoceti. Moreover, our concept of Platanistoidea shares very little with Simpson (1945)'s articulation, which included all 'river dolphin' lineages, including *Inia*, *Pontoporia*, and *Lipotes*. Subjective synonymies of the converted clade name of Platanistoidea include, among others: Platanistoidea Fordyce, 1994; Platanistoidea Barnes, 2006; Platanistoidea Tanaka & Fordyce, 2014; Platanistoidea Tanaka & Fordyce, 2015a; Platanistoidea Kimura & Barnes, 2016.

Here, we also propose the converted clade name Platanistidae as a node-based clade defined by *Platanista*, *Zarhachis* and *Pomatodelphis*. This node-based converted clade of Platanistidae corresponds to the most recent concepts of the familial level grouping of closest fossil relatives of *Platanista*, such as Platanistidae Barnes, 2006; Platanistidae Barnes, Kimura & Godfrey, 2010; Platanistidae Geisler et al., 2011; and Platanistidae Bianucci et al., 2013.

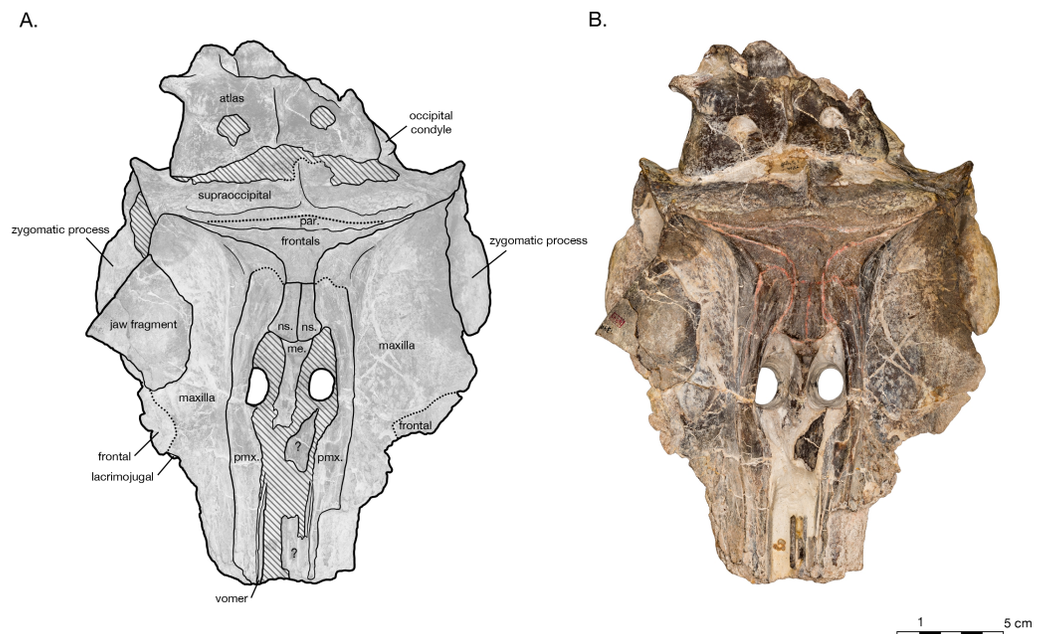
Lastly, Allodelphinidae is the converted clade name for the clade that includes the following fossil odontocete genera: *Allodelphis*, *Arktocara*, *Goedertius*, *Ninjadelpis*, and *Zarhinocetus*. Subjective synonymies of the converted clade name include: Allodelphinidae Barnes, 2006; Allodelphinidae Lambert, Muizon & Bianucci, 2015; Allodelphinidae Kimura & Barnes, 2016. All previous studies have indicated that Allodelphinidae belongs as a sub-clade within a node-based Platanistoidea.

**Type and only included species:** *Arktocara yakataga*, sp. nov.

**Etymology.** The name *Arktocara* derives from the combination of *arktos* from Greek and *cara* from Latin, which together signify "the face of the North." The only preserved material of the type specimen, USNM 214830 consists of the cranium, or its face, and its type locality is the furthest north that a platanistoid has ever been found.

**Age.** Same as that of the species.

**Diagnosis.** Same as that of the species.



**Figure 7** Skull of the holotype of *Allodelphis pratti* (YPM 13408) in dorsal view. (A) Illustrated skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. The symbol “?” denotes a displaced skull fragment of unknown origin. (B) photograph of skull in dorsal view, photography by James Di Loreto, Smithsonian Institution. Courtesy of the Division of Vertebrate Paleontology; YPM 13408, Peabody Museum of Natural History, Yale University, New Haven, Connecticut, USA; [peabody.yale.edu](http://peabody.yale.edu).

*Arktocara yakataga*, sp. nov. (Figs. 2–10 and Table 1)

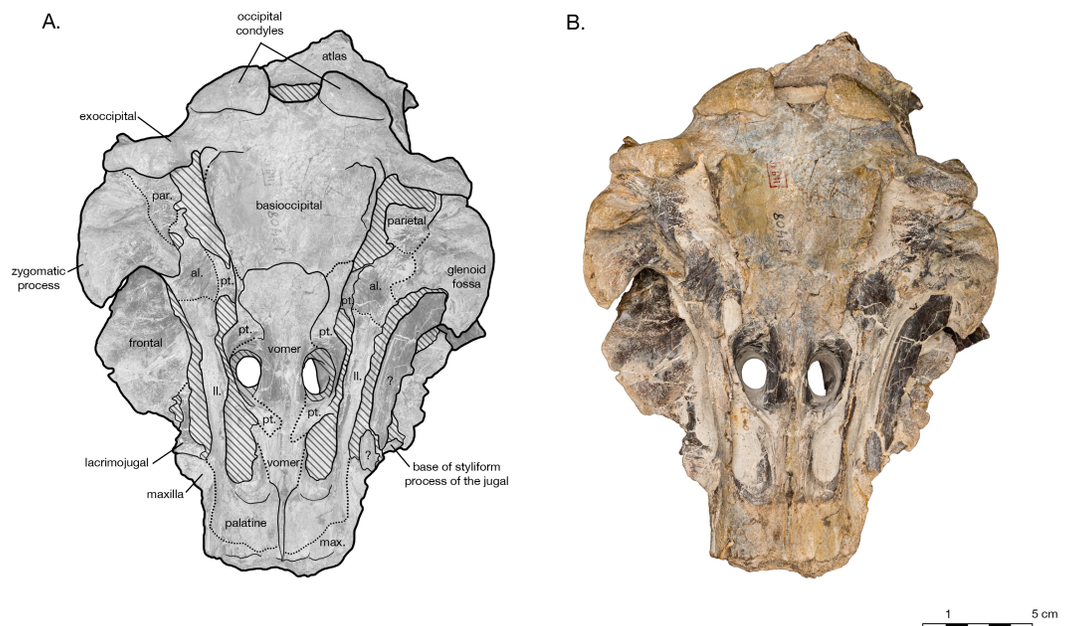
LSID: urn:lsid:zoobank.org:act:FBCF0EAA-7BBB-4EF0-8186-7548993098D1

**Holotype.** USNM 214830, consisting of an incomplete skull lacking the rostrum anterior of the antorbital notches, tympanoperiotics, dentition and mandibles (see Fig. 2).

**Type locality.** The precise geographic coordinates for the type locality of *Arktocara yakataga* are unknown. The type specimen (USNM 214830) was discovered and collected in 1951 by United States Geological Survey (USGS) geologist Donald J. Miller (1919–1961), who was mapping what was then the Yakataga District of Alaska (now the Yakutat City and Borough) between 1944 and 1963. Archival notes housed with the specimen at USNM state that Miller found the specimen in the Poul Creek Formation within the then-Yakataga District (see Age, below). Therefore, we delimit the area for the type’s provenance to exposures of the Poul Creek Formation in the Yakutat City and Borough, Alaska, USA, in a grid ranging approximately from 60°22’N, 142°30’W to 60°00’N, 143°22’W (see Fig. 1). While the formation has been named from its exposures along Poul Creek, it has been suggested that the most abundant macrofossils from this unit have been collected from outcrops along Hamilton Creek, White River, and Big River near Reare Glacier (*Taliaferro, 1932*). It is possible that Miller collected USNM 214830 from one of these exposures.

**Formation.** Poul Creek Formation.

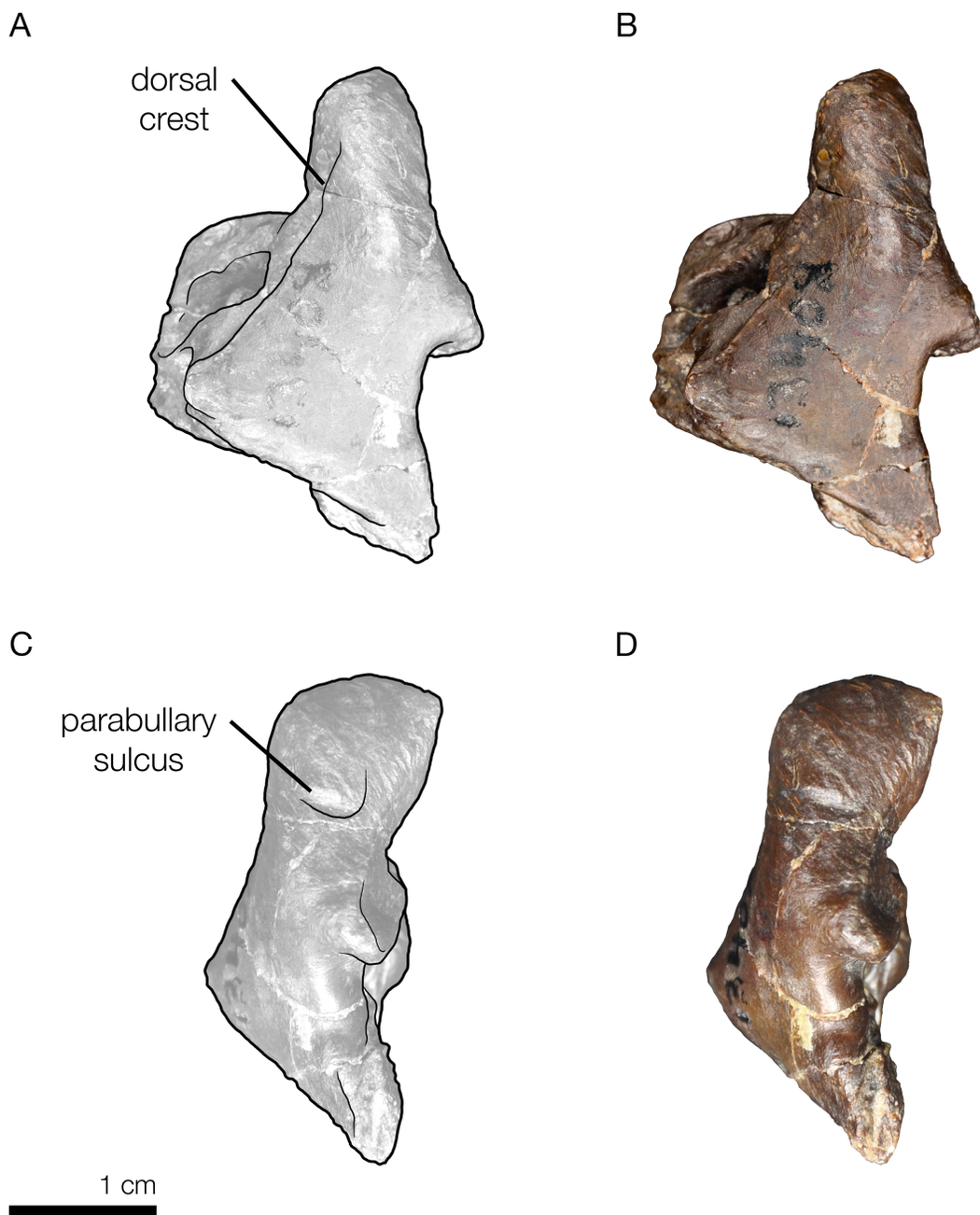
**Age.** Archival documentation accessioned in the Department of Paleobiology with USNM 214830 indicate that the type specimen was collected from an unknown locality exposed



**Figure 8** Skull of the holotype of *Allodelphis pratti* (YPM 13408) in ventral view. (A) Illustrated skull with low opacity mask, interpretive line art, and labels for skull elements. Dotted lines indicate uncertainty of sutures and hatched lines indicate sediment obscuring the fossil. The symbol “?” denotes a displaced skull fragment of unknown origin. (B) photograph of skull in ventral view, photography by James Di Loreto, Smithsonian Institution. Courtesy of the Division of Vertebrate Paleontology; YPM 13408, Peabody Museum of Natural History, Yale University, New Haven, Connecticut, USA; [peabody.yale.edu](http://peabody.yale.edu).

about 400–500 m below the top of the Poul Creek Formation, which has a total stratigraphic thickness of around 1.9 km (Plafker, 1987). The Yakutat terrane of Southeast Alaska consists of the Kulthieth, Poul Creek, and Yakataga Formations (Perry, Garver & Ridgway, 2009; Plafker, Moore & Winkler, 1994; Miller, 1971). The Kulthieth Formation consists of mostly organic-rich sandstones deposited in nonmarine alluvial, deltaic, barrier beach and shallow marine environments, and is Early Eocene to Early Oligocene (~54–33 Ma) in age based on the fossil assemblages present (Perry, Garver & Ridgway, 2009). The Upper Eocene to possibly Lower Miocene (~40–20 Ma) Poul Creek Formation conformably overlies the Kulthieth Formation (Plafker, 1987; Miller, 1971). It is estimated to be approximately 1.9 km thick, and is composed of siltstones and organic-rich sandstones, in part glauconitic recording a marine transgression, interrupted by deposits of the Cenotaph Volcanics (Plafker, 1987). Finally, unconformably overlying the Poul Creek Formation is the Miocene to Pliocene Yakataga Formation (Miller, 1971). It is composed mainly of tillite and marine strata (Perry, Garver & Ridgway, 2009).

The Poul Creek Formation itself is broadly constrained to approximately 40–20 million years in age, from the latest Eocene to possibly early Miocene in age (Plafker, 1987; Miller, 1971). The depositional age of the unit has been further constrained to ~24 to ~29 Ma, or a mid to late Oligocene age, based on detrital zircon fission-track analyses of young grain-age populations (Perry, Garver & Ridgway, 2009). Using the broadest time duration for the formation (~20 million years) and the coarse stratigraphic thickness of the sediments



**Figure 9** Right periotic of the holotype of *Allodelphis pratti* (YPM 13408). Right periotic of *Allodelphis pratti* in dorsal (A, B), and lateral (C, D) views. (A, C), Illustrated periotic with low opacity mask and interpretive line art. The two periotic synapomorphies for the Platanistoidea are labelled: the parabullary sulcus, and the dorsal crest. (B, D), photography by James Di Loreto, Smithsonian Institution. Courtesy of the Division of Vertebrate Paleontology; YPM 13408, Peabody Museum of Natural History, Yale University, New Haven, Connecticut, USA; [peabody.yale.edu](http://peabody.yale.edu).



**Table 1** Measurements for type specimen *Arktocara yakataga* (USNM 214830). Measurements of holotype skull of *Arktocara yakataga* (USNM 214830), in cm (modified after Perrin, 1975 and Tanaka & Fordyce, 2014).

Dimension	Measurement (in cm)
Total preserved length of skull from furthest anterior point to furthest posterior point	23
Cranial length from antorbital notches to occipital condyle	17.5
Distance from preserved rostrum tip to external nares (to mesial end of anterior transverse margin of right naris)	8.5
Distance between upper margin of foramen magnum and nuchal crest	6.8
Height of foramen magnum	2.9
Height of occipital condyle	4.1
Height of temporal fossa	5.9
Height of rostrum at base	6.9
Length of temporal fossa	5.8
Orbit length	4.9
Maximum length of nasal fossa of the frontal	2.6
Length of vertex (nuchal crest to anterior transverse margin of nasal fossa of the frontal)	4
Depth of rostrum at base	6.5
Width of rostrum between antorbital notches	8.8
Width of premaxillae at rostrum base	5.5
Maximum width of premaxillae on cranium	6.2
Width of external bony nares	3.6
Postorbital width of skull	17.7
Bizygomatic width of skull	19.1
Width between temporal crests	11.1
Width of foramen magnum	5.7
Width of occipital condyles	9.8

on the relationship of Allodelphinidae within the Platanistoidea). We also note that, for the purposes of this diagnosis, we use a broad definition of Waipatiidae that includes *Otekaikea* spp. (see Tanaka & Fordyce (2015a)), and Squalodelphinidae *sensu* Lambert, Bianucci & Urbina (2014). See ‘Discussion’ for further comments on systematics of these groups.

*Arktocara* can be differentiated from all other platanistoids by the following combination of character states. *Arktocara* differs from Waipatiidae and Squalodelphinidae in having: a postglenoid process of squamosal greatly reduced (character 114[1]); an occipital shield bearing a distinct sagittal crest (character 118[1]); length of the zygomatic process as percent of the greatest width of the maxillae across the postorbital processes <30% (character 152[1]); lacking a dorsal condyloid fossa (character 119[0]); and lacking any asymmetry in the vertex (character 98[0]).

*Arktocara* also differs from Waipatiidae and Platanistidae in having: frontals posterior to the nasals and between the premaxillae wider than the maximum transverse width across the nasals (character 95[0]); and lacking an anterior transverse ridge and large

tympanosquamosal recess, with middle sinus still inferred to be present (character 144[1]). *Arktocara* further differs from Waipatiidae in having: a lacrimal that wraps around the anterior edge of the supraorbital process of frontal and slightly overlies its anterior end (character 37[0]); maxilla forming the dorsolateral edge of the internal opening of the infraorbital foramen (character 43[0]); ventromedial edge of the internal opening of infraorbital foramen formed by maxilla and palatine (character 44[1]); a nuchal crest weakly convex anteriorly in dorsoposterior view (character 117[1]); a lateral end of the groove for the mandibular branch of the trigeminal nerve wrapping laterally around posterior end of pterygoid sinus fossa and opening primarily anteriorly (character 148[0]); the angle formed by the basioccipital crests in ventral view between 15–40° (character 157[1]); and in lacking a premaxillary crest or posterior maxillary crest adjacent to the nasal (character 72[0]).

*Arktocara* differs from all Platanistidae and Squalodelphinidae in having the anterolateral corner of the maxilla overlying the supraorbital process of frontal being thin and of even thickness to parts posteromedial (character 64[0]). *Arktocara* also differs from all Platanistidae in having: the apex of the postorbital process of frontal projected posterolaterally and slightly ventrally (character 46[0]); the ratio of the greatest width of the premaxillae to the greatest width of maxillae at the level of postorbital processes between 0.49–0.38 (character 76[1]); a shallow emargination of the posterior edge of zygomatic process by the sternomastoid muscle fossa in lateral view (character 111[1]); the width of the squamosal lateral to the exoccipital in posterior view as a percent of the greatest width of the exoccipitals <15% (character 112[0]); fossae for the preorbital lobe of the pterygoid sinus present in the orbit (character 134[1]); shallow posterior portion of the periotic fossa of the squamosal (character 151[1]); posteroventral-most point on the basioccipital crest forming a closely appressed flange with a narrow crease separating exoccipital dorsally from the rest of crest (character 156[1]); and lacking a pneumatic maxillary crest overhanging medially (character 65[0]). Finally, *Arktocara* differs from all Squalodelphinidae in having: a narrower width of the premaxillae at the antorbital notches as a percent width of the rostrum at the antorbital notch (50–64%) (character 51[1]).

*Arktocara* differs from all other Allodelphinidae in having: a reduced postglenoid process of the squamosal (character 114[1]); and the posteroventral-most point of the basioccipital crest forming a closely appressed flange separating the exoccipital dorsally from the rest of the crest by a narrow crease (character 156[1]). *Arktocara* differs from *Allodelphis pratti* and *Goedertius oregonensis* in having: both premaxillae extending posterior to the nasals (character 58[1]); and the ratio of the greatest width of the premaxillae to greatest width of the maxillae at the level of the postorbital processes between 0.49–0.38 (character 76[1]).

*Arktocara* also differs from *Goedertius oregonensis* and *Zarhinocetus errabundus* in having: the width of the premaxillae between 50–64% of the width of the maxillae at the level of the antorbital notches (character 51[1]); a uniformly thin anterolateral corner of the maxilla overlying the supraorbital process of the frontal (character 64[0]); length of the zygomatic process of the squamosal  $\leq 30\%$  of the width of the maxillae at the postorbital processes (character 152[1]); lacking a rostral basin (character 50[0]), lacking a posterior dorsal infraorbital foramen placed posteromedially near the posterior extremity of the

premaxilla (character 60[0]); lacking a dorsal condyloid fossa (character 119[0]) and having a symmetrical cranial vertex (character 98[0]).

*Arktocara* also differs from *Allodelphis pratti* and *Zarhinocetus errabundus* in having: two anterior dorsal infraorbital foramina (character 49[1]); and a shallow emargination of the posterior edge of the zygomatic process by the sternomastoid muscle fossa in lateral view (character 111[1]). *Arktocara* further differs from *Goedertius oregonensis* having: a fused lacrimal and jugal (character 39[1]); the apex of the postorbital process of the frontal directed posterolaterally and slightly ventrally (character 46[0]); a triangular or anteroposteriorly widened falciform shaped postorbital process of the frontal (character 47[2]); one posterior dorsal infraorbital foramina of the maxilla (character 59[1]); nuchal crest weakly convex anteriorly in dorsoposterior view (character 117[1]); and the posterior edge of the vomer terminating on the basisphenoid (character 154[0]).

*Arktocara* further differs from *Zarhinocetus errabundus* in having: the transverse width of the nasal >70% the length of the nasal (character 91[2]); a distinct sagittal crest on the occipital shield (character 118[1]); and in lacking an anterior transverse ridge and large tympanosquamosal recess (character 144[1]).

Lastly, *Arktocara* displays the following apomorphies: a weakly developed antorbital notch (character 10[0]); straight lateral margin of the right premaxilla posterior to premaxillary foramen (character 56[1]); a U-shaped nasal frontal suture (character 94[2]); and a greatly reduced postglenoid process of the squamosal (character 114[1]).

**Etymology.** The species epithet ‘yakataga’ derives from the Tlingit name for the point of land along the southeast coast of Alaska between modern day Kayak Island and Ice Bay. This point, currently called Cape Yakataga, is located directly southwest of Watson Peak and represents the southeastern boundary of a floodplain drained by the Bering Glacier. The name Yakataga was first published by *Tebenkov* (1852: map 7), who was a cartographer and hydrographer of the Imperial Russian Navy, as “M[ys] Yaktaga” on an 1849 map of Alaska. The geographic place name has been alternatively spelled Cape Iaktag, Cape Yakaio, Cape Yakatag, and Yokataga Reef (*Orth, 1967*). According to the Geographic Names Information System (*GNIS, 2016*), developed by USGS in cooperation with the United States Board of Geographic Names (BGN), the name “Yakataga” means “canoe road,” referring to two reefs that form a canoe passage to the shore of the village.

## Description

Anatomical terminology follows *Mead & Fordyce (2009)*. In most cases, description of individual elements derives from the most informative side of the skull, in terms of preservation; we note any morphological asymmetry if present. Besides overall erosion of the bone surfaces and loss of some skull elements, there appears to be no significant burial-related distortion of the skull.

## Skull

The holotype of *Arktocara yakataga* (USNM 214830) consists of an incomplete skull, measuring 23 cm in preserved length. The majority of the rostrum is missing, with an asymmetric transverse break approximately 2–5 cm anterior to the antorbital notch. The

skull also lacks both nasals, jugals, tympanoperiotics, and the right occipital condyle (see Figs. 2 and 3). Most of the paroccipital processes of the exoccipitals are gone, large portions of the palatines and pterygoids are missing, small fragments along the lateral margins of the frontals and maxillae are incomplete, and the general condition of many osteological elements in the skull are poorly preserved. The skull may have been both mechanically and chemically prepared in the past (with no known documentation), including acid preparation, which may have contributed to the poor state of preservation for the osteological surfaces of many elements. Portions of the skull are obscured by a fine-grained grey matrix of siltstone, especially infilling the mesorostral canal, the bony nares, the recesses of tympanoperiotic region, and the braincase (which is exposed via the foramen magnum and fenestrae in the supraoccipital).

In dorsal view, the preserved skull is roughly hexagonal in overall shape (Fig. 2). The external nares are vertically oriented, and positioned at a level between the antorbital and postorbital processes. The vertex is particularly table-like and square, composed of frontals, premaxillae, and nasals (missing). The parietals appear to be narrowly exposed on the vertex immediately anterior to the nuchal crest, between the frontals and supraoccipital—whether this is natural, or an ontogenetic characteristic, is unclear. The vertex is bordered anteriorly by the external nares, laterally by the maxillae and posteriorly by the nuchal crest of the supraoccipital. The nuchal crest is straight along the posterior edge of the vertex, but begins to curve posterolaterally as it approaches the temporal crest. The temporal fossae are visible in dorsal view due to an intertemporal constriction just anterior to the level of the nuchal crest, and the fossa is floored by a narrow valley (the squamosal fossa) between the squamosal plate and supramastoid crest.

In lateral view, the profile of the skull gradually slopes upwards from the level of the antorbital notch to posterior of the nares, where it levels out on the vertex (Fig. 4). The nuchal crest is well defined, and about the same height as the frontals on the vertex. The orbit is shallowly rounded dorsally (4.9 cm in length), with the maxilla completely overlying the frontal on the thin supraorbital process, except on the postorbital process, where the frontal is exposed laterally. It is unclear whether the antorbital process of the frontal is completely covered by maxilla or not, but most of the medial antorbital process is composed of the lacrimojugal. The temporal fossa is trapezoidal in shape, with the temporal crest forming a right angle with the dorsal margin of the zygomatic process of the squamosal. The dorsal margin of the temporal fossa is roofed over by the frontal.

The preserved posterior portion of rostrum anterior of the antorbital notch is wide (8.8 cm anterior to the antorbital notch) and deep (6.5 cm at the level of the antorbital notch), with a broadly open and deep mesorostral canal (2.4 cm wide and 4.6 cm deep at the level of the antorbital notch). While there is material missing around the antorbital notch, the posterior margin of the right antorbital notch appears to be real, demonstrating it to be weakly developed, forming an L-shape with the lateral margin of the rostrum (Fig. 2). Though some of the anteromedial antorbital process is missing, not enough is missing to have drastically changed the shape of the antorbital notch region. In anterior view, the maxilla abruptly slopes upwards medially to the distinct premaxilla-maxilla suture, and underlies the premaxilla along the entire preserved length of rostrum and likely the cranium

(Fig. 5). The premaxilla therefore forms an anteroposteriorly elongated rectangular plateau surrounding the external bony nares, elevated in relation to the plane of the maxillae, appearing similar in transverse cross-section to a horst and graben system.

In ventral view, there is clearly a fossa for the preorbital lobe of the pterygoid sinus (a shallow depression surrounding the ventral infraorbital and sphenopalatine foramina), as well as for the hamular lobe of the pterygoid sinus (fossa anterior to the choanae). Also present are an extremely shallow tympanosquamosal recess for the middle sinus, and a middle pterygoid sinus fossa anterior to the periotic fossa (Fig. 6). There is not enough rostral material or paroccipital process to determine the presence of an anterior or posterior sinus respectively, and the presence of a peribullary sinus is difficult to ascertain.

### **Premaxilla**

Both of the premaxillae are symmetrical, and overlie either the maxillae or the frontals for their entire preserved length (Fig. 2). In lateral view, the premaxilla thins slightly as it passes around the external nares, otherwise maintaining a relatively even thickness on the cranium (Fig. 4). The premaxilla also appears to thin anterior of the antorbital notch (especially in right lateral view), lowering to the same level as the maxilla instead of rising dorsally above it. However, in anterior view, it is evident that the ventral margin of the left premaxilla sinks ventrally into a medial trough created by the maxilla while the dorsal margin remains level, accounting for the apparent reduction in thickness (Fig. 5). Both premaxillae are broken anteriorly, the right further anterior than the left and missing some of its anterolateral edge. The premaxilla-maxilla suture is clear in dorsal view along the entire lateral length of the premaxilla, as well as in anterior view at the transverse cross-section of the rostrum. In dorsal view, the lateral margin of the premaxilla is mostly rectilinear, widening only 0.5 cm from the rostral break to a level anterior to the nares. As it passes laterally around the nares, the premaxilla gently bows out laterally, with the medial edge retreating more than the lateral edge so that the total width is reduced (0.8 cm on the right premaxilla). Posterior to the nares, the lateral edge remains straight posteriorly, but the medial edge expands slightly medially, once again widening the premaxilla. On the left side of the skull, lateral and posterior of the external nares, a narrow ledge of the medial margin of the maxilla laterally borders the premaxilla, where the premaxilla is separated from the maxilla (possibly diagenetically).

On the vertex, the posterior termination of the premaxilla lies on the frontal as an asymmetrical, spatulate lobe, tapering posterolaterally and bordered medially by frontal and the nasal fossa of the frontal, and laterally the maxilla (Fig. 2). There is no division of the premaxilla into a posteromedial splint and posterolateral plate (a widespread characteristic amongst odontocetes). The premaxillary sac fossa, roughly at the level of the antorbital notch, is shallow with a rough surface. No premaxillary foramina or associated premaxillary sulci are visible on the preserved length of the skull.

### **Maxilla**

Only a small portion of rostral maxilla remains, including part of the maxillary flange on the left side, and just over 5 cm of the body of this element on the right side. In anterior view, the maxilla preserved on the rostrum slopes dorsally from the lateral edge to the premaxillary

suture (Fig. 5). The premaxillary suture is distinct and unfused. The preserved maxillary flange on the left side is thin and flat. In lateral view, the maxilla gradually increases in depth posteriorly until anterior of the nares, where it reduces in dorsoventral depth to a thin plate forming the facial portion and ascending process of the maxilla (Fig. 4). In dorsal view, the maxilla flares out laterally at the level of the antorbital notch, which is shallow and L-shaped. Posterior to the antorbital process is broad and relatively flat. The right side bears two infraorbital foramina: one immediately posteromedial to the antorbital notch, and one at the level of the nares. The left maxilla has three infraorbital foramina, all in a sagittal plane from immediately posteromedial of the antorbital notch to a level anterior of the nares. The posterior-most dorsal infraorbital foramen on either side is the largest foramen, with two sulci each—one directed posterolaterally and the other posteromedially—giving the foramen a v-shaped appearance in dorsal view. In the facial region, the maxilla gradually curves dorsomedially from the supraorbital process to the premaxillary suture, and the facial fossa is essentially flat. The maxilla does not extend to the postorbital process, though the suture with the frontal on the postorbital process is unclear. The ascending process suddenly curves dorsomedially towards the lateral edge of the vertex, and the posteromedial margin of the maxilla curves dorsolaterally to terminate in a sharp triple-point junction with the nuchal, temporal, and orbitotemporal crests.

In ventral view, the hard palate of maxilla curves dorsolaterally from the midline to the lateral edge, where it flattens out on the maxillary flange (Fig. 3). No alveoli are present in the preserved palatal surface of the maxillae. A small gap between the maxillae along the midline of the hard palate reveals a thin ridge of the vomer, approximately 2.6 cm long and a maximum of 1 mm wide. Just anterior to the level of the antorbital notch, a narrow palatine groove of the maxilla for vessels and/or nerves begins approximately 1.5 cm lateral of the midline, and curves posterolaterally to terminate ventral of the sphenopalatine foramen. Midway along the palatine groove is the posterior palatine foramen, and the posterior portion of the groove is partially roofed over by fragments of the palatine. The distal maxillopalatine suture would have been at some point anterior and lateral of the palatine groove, though the palatine is incomplete and is missing this portion. Between the palatine groove and the medial lamina of the pterygoid, the maxilla is overlain by fragments of the palatine, though the sutures between the two are extremely unclear (Figs. 2, 4 and 6). Lateral of the palatine groove, the maxilla overlies the medial process of the lacrimal, and encircles both the ventral infraorbital foramen and the dorsal margin of the sphenopalatine foramen. Posterior of the foramina, the maxilla terminates in an abutment with frontal.

### **Frontal**

In dorsal view, the frontals are mostly covered by the maxillae, with the exception of exposures on the postorbital processes and the vertex (Fig. 2). The postorbital processes are asymmetrical in lateral view, with a larger and more robust right postorbital process pointing ventroposteriorly and reaching within 0.7 cm of the tip of the zygomatic process (Fig. 4). The left postorbital process is shorter and more smoothly curved ventroposteriorly. Though more of the length appears to be preserved in the right postorbital process than the

left, it is difficult to determine whether the asymmetry is real or preservational. Moreover, the dorsal rim of the right orbit is missing and heavily eroded into the supraorbital process, yielding an incomplete view of the orbit on this side of the skull. The frontal-maxillary suture is indistinct as it passes in an anteroposterior direction across the postorbital processes.

The frontal-maxillary suture is also indistinct along the lateral edge of the vertex, where the maxilla rises dorsomedially to the edge of the vertex's tabular surface (Fig. 2). The sutures are posteromedially convex in dorsal view, on either side of the vertex, curving from the temporal crest to the posterior premaxilla-maxilla suture. On the vertex, the frontals are exposed as a wide, flat surface. They are separated from supraoccipital posteriorly by a narrow exposure of the parietals which may be an artifact of ontogeny. As such, it is possible that the frontals did contact the supraoccipital in older individuals of this taxon. The frontal exposure is bordered by the maxillae laterally, and the premaxillae and mesethmoid anteriorly. Between the premaxillae, the frontals bear a shallow fossa for the missing nasal bones (see "Nasal" for further description).

### **Lacrimojugal**

Most of the medial margin of the antorbital process is composed of the lacrimal. In dorsal view, a narrow margin of the lacrimal emerges from beneath the maxilla as a thin plate along the lateral and anterior edges of the antorbital process (Fig. 2). In ventral view, the medial process of the lacrimal extends posteromedially towards the ventral infraorbital foramen, but is overlapped by the maxilla (Fig. 3). The styliiform portion of jugal is mostly missing, save a small piece of the jugal base where it is fused to the lacrimal. The jugal base is transversely wide and anteroposteriorly narrow. The lacrimal is covered posteriorly by the frontal.

### **Nasal**

Though both nasal bones are missing, the frontal bones bear a distinct fossa between the premaxillae on the vertex that indicates where the bones would have been (Fig. 2). The fossa is bordered anteriorly by the mesethmoid and laterally by the premaxillae. Based on the extent of the nasal fossa, the nasals were wider anteriorly than posteriorly (from 2.1 to 1.6 cm), and minimally 1.9 cm in length, but could have been longer if they overhung the bony nares, as seen in the holotype of *Allodelphis pratti* (Fig. 7). The height of the nasals cannot be estimated, considering that they rise above the frontal to varying degrees in similar taxa such as *Allodelphis pratti*. All codings for the nasals of *Arktocara yakataga* in the phylogenetic analysis were made from measurements of this nasal fossa of the frontals, considered the minimum extent of the nasals.

### **Vomer**

At the broken rostral tip in anterior view, the mesorostral canal is deeply v-shaped (Fig. 5). The cross-section reveals the damaged vomer to be extremely thin (<1 mm), and lining all sides of the mesorostral canal. Anterior to the nares, the maxilla is exposed dorsal of the vomer, so that it forms the dorsal edge of the mesorostral canal and the anterior wall of the external bony nares, similar to *Tursiops truncatus* (Montagu, 1821) (based on USNM

504560). Anterior to the nares, the vomer also curves medially to form the medial wall of the external bony nares, and the lateral walls of the nasal septum. Posterior of the nares, the vomer is obscured in dorsal view by unprepared matrix.

In ventral view, the vomer is visible as a long, thin crest running down the midline of the hard palate (Fig. 3). Anterior to the antorbital notch, the vomer is mostly obscured by the maxillae, but emerges at the level of the antorbital notch as a vertical wall separating the choanae, slightly wider at its base and thinning to a sharp crest ventrally. Posterior of the nares, the crest flattens and the vomer flares out laterally, adhering to the basisphenoid. Its suture with the basisphenoid is indistinct, roughly straight transversely, between the posterior lamina of the pterygoid and immediately anterior to the basisphenoid-basioccipital suture.

### ***Mesethmoid***

The mesethmoid composes the bulk of the nasal septum, narrowing dorsally and flanked on either side by the thin lamina of the vomer. Behind each choana is a rounded bony protuberance, likely composed of the lateral wings of the mesethmoid (referred to as the ectethmoid by some authors, e.g., *Mead & Fordyce, 2009*) (Fig. 2).

### ***Parietal***

The parietals are visible in dorsal and lateral view in the temporal fossa, where they are in contact with the frontals under the temporal crest, the supraoccipital along the occipital border, and partially underlie the squamosal plate with a semi-circular suture across the temporal fossa. All of the sutures are indistinct. The parietal forms the lateral wall of the braincase as a thin, laterally convex plate. As with the supraoccipital, both parietals in the temporal fossa are broken, with a small rounded window revealing the matrix-filled braincase (Fig. 4). As previously mentioned, the parietals in dorsal view are exposed as an anteroposteriorly narrow strip along the posterior margin of the vertex. In ventral view, the parietal is again visible in the periotic fossa; having passed under the squamosal to form the lateral wall of the braincase, it emerges medial to the squamosal in ventral view as small, slightly concave surface, just posterior to the foramen ovale (Figs. 3 and 6).

### ***Supraoccipital***

The supraoccipital is broadly visible in dorsal view, contributing to the entire length of the nuchal crest. The supraoccipital and frontals are very slightly separated along the vertex by an exposure of the parietals sutured to the supraoccipitals. The supraoccipital is also in contact with the parietals along the occipital border (Fig. 2). Lateral of the vertex along the nuchal crest, the supraoccipital is in direct contact with the maxilla. The nuchal crest is straight medially, but begins to curve posteriorly as it approaches the temporal crest. Along the parietal margin, the supraoccipital is a thin plate, with the edge oriented posterolaterally as it curves around the posterior edge of the temporal fossa. In posterior view, the supraoccipital is rectangular in shape (Fig. 5). A prominent external occipital crest divides supraoccipital sagittally, from the midpoint of the nuchal crest to the opisthion. On either side of the crest, the supraoccipital is very thin and slightly concave. Both these surfaces are broken into fenestrae, with rounded margins that reveal openings

filled with matrix in the braincase. There is no evidence of a dorsal condyloid fossa of any significance. The contact of the supraoccipital with the exoccipital is indistinct, except around the foramen magnum, where the nuchal tubercle of the supraoccipital clearly tucks underneath the dorsal portion of the occipital condyle.

### ***Exoccipital***

Both exoccipitals are incomplete, missing all or part of the occipital condyle, and most of their ventral portions (Fig. 5). The supraoccipital suture is indistinct, but the contact with the squamosal is clear, along the posteroventral temporal crest, and on the ventral side of the skull. The exoccipital is thin along the lateral margin posterior of the temporal crest, thickening ventrally. The broken remains of the occipital condyles are sufficient to observe their robust size and width, composing approximately 70% of the total combined width of the exoccipitals. Only the dorsal portion of the left condyle remains. Its surface is smooth, posteriorly curved and laterally broad. The foramen magnum is elliptical in shape, almost twice as wide as it is tall (2.9 cm wide, 5.8 cm tall). Both ventral and dorsal condyloid fossa are very shallow and ill-defined. Though both exoccipitals are missing most of their ventral portions, including the jugular notches and paroccipital processes, the left exoccipital does bear a small foramen that may represent the hypoglossal foramen, immediately lateral to the posterior end of the basioccipital crest. The exoccipital also forms a small knob separated from the posterior end of the basioccipital crest by a narrow cleft, possibly a vestigial medial crest of the exoccipital.

### ***Basioccipital***

In ventral view, the basioccipital widens posteriorly from 6.2 cm wide at its suture with the basisphenoid, to 8 cm at the posterior end of the basioccipital crest (Fig. 3). The element is ventrally concave, with basioccipital crests oriented laterally from a sagittal plane, and at an angle of approximately 12 degrees from the midline (opening posteriorly). The tympanic plates are thin where they overlies the basisphenoid, increasing in width posteriorly before tapering slightly and rounding off at their posterior ends. The right side of the posterior basioccipital crest is missing, though the left side is complete. The posterior end of the basioccipital crest is interrupted by a narrow cleft that separates a small knob from the rest of the tympanic plate. This small knob is immediately medial to the hypoglossal foramen (on the left side), and presumably the jugular notch. Though consistent with the rest of the basioccipital crest, the knob is actually formed by the exoccipital—a condition described in *Fordyce's* (1994) description of *Waipatia maerewhenua*, and also seen in a number of other platanistoids, including *Allodelphis pratti* (Figs. 7–9). The suture with the basisphenoid, along the anterior margin of the basioccipital, is represented by a wavy margin near the midline. The suture is increasingly less distinct laterally, where the basioccipital extends anteriorly, overlapping the lateral margins of the basisphenoid and bordering the posterior lamina of the pterygoid along its medial edge. There is no strong evidence of a muscular tubercle for the insertion of the ventral rectus capitis muscle.

### **Sphenoid**

The basisphenoid is visible on the ventral side of the skull, though it is mostly obscured by the basioccipital and vomer (Fig. 3). The basioccipital crests extend anteriorly to cover the lateral portions of the basisphenoid, completely obscuring any view of the contact between the basisphenoid and alisphenoid. The posteroventral plate of the vomer obscures the anterior margin of the basisphenoid, and spreads over the basisphenoid's medial section to terminate immediately anterior to the wavy basioccipital suture, which appears partially fused. In ventrolateral view, the sphenoid re-emerges from beneath the basioccipital, with the ventral carotid foramen tucked under the dorsolateral margin of the basioccipital crest (Fig. 6). A small portion of the basisphenoid is visible, wrapped laterally around the ventral carotid foramen. Anterolateral of the foramen, the alisphenoid extends laterally anterior to the periotic fossa as a thin plate. The alisphenoid passes anterior to the foramen ovale, and bears a long, thin groove for the mandibular nerve, extending anteriorly from the foramen ovale to the anterior margin of the alisphenoid, medial of the subtemporal crest of the squamosal. The alisphenoid continues to extend lateral of this groove, posteriorly contacting the medial edge of the falciform process, and anteriorly fusing to a remnant of the lateral lamina of the pterygoid (see 'Pterygoid' below). A very small portion of the orbitosphenoid is visible on the right ventral side of the skull, emerging ventral to the posterior lamina of the pterygoid on the medial wall of the orbit and dorsal of the alisphenoid (only visible on the right side of the skull).

### **Pterygoid**

Both pterygoids are incomplete, missing the pterygoid hamulus and almost all of the lateral lamina. In ventral view, the medial lamina is an extremely thin sheet, meeting the vomer/palatine anterior to the nares and curving posterolaterally to form the anterior and lateral walls of the external bony nares, slightly overlying the palatine (Fig. 3). The posterior lamina rises ventrolaterally as a thin plate, ventrally concave, and forming the posterior wall of the external bony nares. The posterior lamina is bordered posteromedially by the anterior basioccipital crest, separately by a widely open suture. This open suture is unusual among fossil and living odontocetes, and may represent either an ontogenetic feature or diagnostic feature for *Arktocara*. The posterior lamina is bordered posterolaterally by the alisphenoid.

Platanistoids bear a bony structure on the ventral side of their skulls: a thin, bony lamina that extends from the ventral surface of the hard palate and runs parallel to the posterior lateral lamina of the pterygoid to finally attach to the alisphenoid medial of the squamosal in the ear region. The holotype of *Arktocara yakataga* is mainly missing this feature, except for a process protruding from the alisphenoid medial of the subtemporal crest. We interpret this feature as an eroded remnant of the lateral lamina of the pterygoid, that is fused to the alisphenoid (there is no detectable suture line between the two elements in *Arktocara*). For further discussion of the lateral lamina of pterygoid as a platanistoid feature, see discussion in "Platanistoid systematics."

### **Palatine**

Both of the palatine bones are missing large portions, including the palatine surface, maxillary process, palatal crest, and lateral lamina. However, in ventral view a prominent

pterygoid sinus fossa for the hamular lobe is formed by the palatine, anterior to the medial pterygoid lamina and posterior to the palatine groove of the maxilla (Fig. 3). These grooves represent the minimum extent of the palatine extending anteriorly over the maxilla, since the maxillopalatine suture is impossible to detect. A very small fragment of the horizontal portion of the palatine remains immediately ventral to the ventral infraorbital foramen, partially covered by the medial lamina of the pterygoid.

### **Squamosal**

In dorsal view, the short, wide, and rounded zygomatic process of the squamosal points anterolaterally (Fig. 2). The temporal fossa is floored by the squamosal fossa, forming narrow valley between the supramastoid crest and the squamosal plate. In lateral view, the squamosal plate is a thin sheet, slightly convex laterally, and overlaps the parietal at an indistinct, rounded suture traversing the temporal fossa (Fig. 4). The zygomatic process is rounded off. The postglenoid process is greatly reduced and missing its ventral edge on the right side, the postglenoid notch is either absent or too greatly reduced to determine, and the external auditory meatus is difficult to determine. In posterior view, the squamosal is widely visible lateral to the exoccipital, and the temporal crest where these later two elements meet is well developed (Fig. 5). In ventral view, glenoid fossa on the zygomatic process is broad and shallow (Fig. 2). The incomplete postglenoid process is square in cross section. The contribution of the squamosal to the periotic fossa is wide and shallow, sloping medially from the anterior meatal crest, and bordered medially by the parietal exposure in the periotic fossa. The falciform process is transversely thin and flat, and projects ventromedially from the shallow tympanosquamosal recess. Anteromedial of the falciform process, the anterior margin of the squamosal plate and the anterodorsal margin of the falciform process are extended and join to form an anterior protrusion, which overlies the lateral margin of the alisphenoid. This anterior protrusion is observed in other platanistoids, including *Platanista* and an undescribed platanistoid USNM 214911 (Fig. S1), where it articulates with the alisphenoid and the lateral lamina of the pterygoid. Further study will determine if this feature is present in more platanistoid taxa.

### **Body size estimate**

Total body length (TL) was estimated using the formula created by [Pyenson & Sponberg \(2011\)](#) for calculating body size in stem Platanistoidea (*sensu* [Pyenson & Sponberg, 2011](#)) based on a bizygomatic width (BIZYG):

$$\text{Log}(L) = 0.92 * (\text{log}(\text{BIZYG}) - 1.51) + 2.49.$$

The bizygomatic width of USNM 214930 was measured as 19.1 cm, and using the formula produced a reconstructed body length of 2.28 m. Based on this estimate, *Arktocara* would have been similar to the adult size of *Platanista*, which averages a length of 2.4 m and at least 85 kg in weight ([Jefferson, Webber & Pitman, 2008](#)). It is likely that, in life, *Arktocara* possessed a rostrum that was relatively elongate, based on its near relatives *Zarhinocetus* and *Goedertius*; the rostrum of *Allodelphis* is poorly known, based on several incomplete fragments belonging to the type specimen YPM 13408. In addition, all known allodelphinids

have elongated necks, indicated by unfused, proportionally large cervical vertebrae with anteroposteriorly elongated centra, which is in contrast to the general cetacean trend of shortened, hydrodynamic necks (*Buchholtz, 2001*). Though the type specimen of *Arktocara* lacks any post-cranial material, it most likely also had an elongated neck given its close relationships to other allodelphinids. Such longirostry and neck elongation may add to *Arktocara*'s reconstructed total length, and although *Pyenson & Sponberg's (2011)* equations took such allometry into account, we propose that a TL of 2.28 m for *Arktocara* may be a slight underestimate.

### Ontogeny

We assessed skeletal maturity based on long-established osteological indicators, particularly the fusion of cranial sutures and textural surface of the occipital condyles (*Pyenson & Sponberg, 2011*). Most sutures are clearly distinguishable and closed, with some exception of sutures on the ventral side of the skull that appear un-sutured. Most pronounced are the open sutures between the dorsal lamina of the pterygoids and the basioccipital on the medial ventral surface (*Fig. 3*). It is unclear whether this feature is an ontogenetic trait unique to *Arktocara*, or whether it is more broadly observed in other allodelphinids (for example, *Zarhinocetus*). Also, the missing nasals and palatines suggest that their sutures to adjacent skeletal elements were unfused. *Pyenson & Sponberg (2011)* described the presence of a pitted periosteal surface of the occipital condyles as an indication of immaturity. The preserved occipital condyles of USNM 214830 are smooth, indicated a more advanced ontogenetic age. Based on these combined observations, we suggest that the skull of USNM 214930 belonged to a skeletally mature individual.

### Phylogenetic analysis results

The phylogenetic analysis resulted in 430 most parsimonious trees, all with a score of 1960, consistency index of 0.233 and retention index of 0.631. The strict consensus tree, which was created from the 430 trees, shows a similar topology to the equally weighted analysis of *Tanaka & Fordyce (2015a)*. *Arktocara* is the sister taxon to *Allodelphis*, nested within a broader clade of Allodelphinidae, which includes *Zarhinocetus* and *Goedertius*. This is the first phylogenetic analysis to include these latter two genera, which were not included in *Barnes (2006)*'s original matrix. *Lambert, Bianucci & Urbina (2014)* recovered a monophyletic Allodelphinidae in their 50% majority consensus tree and *Lambert, Muizon & Bianucci (2015)* resolved a polyphyletic Allodelphinidae, although both analyses only included *Zarhinocetus* and *Allodelphis* among their allodelphinid sample of Platanistoidea. Our analysis yields robust support for the monophyly of Allodelphinidae, with higher support values (decay index 5, bootstrap 50) than those recovered for the node-based clade of Platanistoidea (decay index 1, bootstrap value <50) (*Fig. 10*). Like *Tanaka & Fordyce (2014)* and *Tanaka & Fordyce (2015a)*, we failed to recover a monophyletic Squalodelphinidae (*sensu Lambert, Bianucci & Urbina, 2014*), yet in contrast, we did find low support for a monophyletic Waipatiidae, an idea proposed by *Fordyce (1994)*, but not explicitly tested until recently. Our analysis is not the first one to recover a clade that includes both species of *Waipatia* and both species of *Otekaikea*—*Tanaka & Fordyce's*

(2015a; 2015b) results are consistent with ours—however Tanaka and Fordyce excluded the *Otekaikea* species from their definition of Waipatiidae. We recommend defining Waipatiidae more inclusively (both species of *Waipatia* and *Otekaikea*).

Though it is beyond the scope of this study, the incorporation of molecular data into future analyses of odontocete systematics will help to better resolve the relationships among extant cetaceans, and consequently their fossil relatives. See below for further comment on the implications of these results on the systematics of Platanistoidea.

## DISCUSSION

### Platanistoid systematics

The present day concept of Platanistoidea has its origins with *Simpson (1945)*, although by the late 20th century, it became clear that genera such as *Inia*, *Pontoporia*, and *Lipotes* were more closely related to Delphinoidea than to *Platanista* (*Muizon, 1984; Muizon, 1985; Muizon, 1987*), especially with the advent of molecular datasets in the 21st century (see *Geisler et al., 2011* for a comprehensive review). *Muizon (1984)* provided the first modern articulation of Platanistoidea to include the numerous fossil forms that appeared to be most closely related to *Platanista* than any other odontocete, living or extinct, including Platanistidae, Squalodelphinidae (=Squalodelphidae *sensu Muizon, 1984*, an alternative spelling that according to *Rice (1998)* was wrongly formed), and Squalodontidae. Later, *Muizon (1987)* described two synapomorphies for Platanistoidea: a loss or reduction of the coracoid process and suprascapular fossa of the scapula; and the acromion process located on the anterior edge of the scapula. In a review of fossil and extant Delphinida, *Muizon (1988a)* added another extinct family, Dalpiazinidae, to the aggregate of extinct families in Platanistoidea, tentatively placing it as sister group to Squalodontidae within Platanistoidea.

*Muizon (1994)* modified this diagnosis of the Platanistoidea to include three more characteristics: a deep subcircular fossa located dorsal to the spiny process of the squamosal; a hook-like articular process or rim on the periotic; and the migration of the palatines dorsolaterally, surrounded by the maxilla and pterygoid which partly overlap them. The type and only specimen of *Arktocara* does not possess any of the elements required to evaluate these synapomorphies, though the palatine is located dorsolaterally (ventral of the sphenopalatine foramina) and is slightly overlapped by the pterygoid (though not the maxilla) (see ‘Description’, palatine).

*Muizon (1994)* maintained that Dalpiazinidae may be a sister group to Squalodontidae, but admitted that the available material referable to Dalpiazinidae was too fragmentary to evaluate any synapomorphies of Platanistoidea. As a result, *Fordyce (1994)* excluded Dalpiazinidae from his analysis of Platanistoidea. *Dalpiazina ombonii (Muizon, 1988b)* (IGUP 26405), which was originally given the genus name *Champsodelphis* by Longhi (1898), was later reviewed by and given its new genus by *Muizon (1988b)*, and is the only described member of this group. Based on observations by one of us (NDP) of the type specimen, we follow *Fordyce (1994)* in excluding this taxon from consideration as a platanistoid until a more detailed study can resolve the confusing history of associated material that forms the basis for this taxon (and potential membership of other odontocetes).

In his description of *Waipatia maerewhenua*, [Fordyce \(1994\)](#) articulated the current concept of Platanistoidea (and largely the basis for the node-based definition used here), which narrowed Muizon's ([1987, 1991](#)) definition to include only the families Squalodontidae, Squalodelphinidae, and Platanistidae, although [Fordyce \(1994\)](#) hinted at possibly platanistoid affinities of other taxa, such as *Prosqualodon davidis*. [Fordyce \(1994\)](#) also added two synapomorphies: the anterior process of the periotic roughly cylindrical in cross section; and the anterior process smoothly deflected ventrally. [Fordyce \(1994\)](#)'s diagnosis of Platanistoidea also omitted any mention of synapomorphies related to the palatines, and noted that the previous two synapomorphies of the scapula were equivocal, as they are not seen in all platanistoids. The type specimen of *Arktocara* has no associated tympanoperiotics, but the periotics of both *Allodelphis pratti* and *Zarhinocetus errabundus* possess both periotic synapomorphies of the Platanistoidea ([Fig. 9](#)).

More recent revisions of the Platanistoidea have supported the exclusion of Squalodontidae, restructuring Platanistoidea to some combination of the families Platanistidae, Allodelphinidae, Squalodelphinidae and Waipatiidae. [Lambert, Bianucci & Urbina \(2014\)](#)'s description of the squalodelphinid *Huaridelphis* pointed to the inclusion of Platanistidae, Allodelphinidae and Squalodelphinidae in a monophyletic Platanistoidea based on a number of descriptive synapomorphies: deeply grooved rostral suture between the premaxilla and maxilla; elevation of the antorbital region higher than dorsal margin of rostrum base in lateral view; widening of cranium; presence of a deep fossa in orbit roof; vertex distinctly shifted to the left compared with the sagittal plane of the skull; reduction of the ventral exposure of palatine; hamular fossa of the pterygoid sinus extended anteriorly on the palatal surface of rostrum; presence of an articular rim on the periotic; elongation of anterior spine on the tympanic bulla and associated anterolateral convexity; loss of double rooted posterior teeth; and tooth count greater than 25. Of these synapomorphies, *Arktocara* lacks two: the antorbital region is not higher than the rostrum base, and the vertex is not shifted to the left. Both *Squalodon* and *Waipatia* were excluded from Platanistoidea in the results, though a broader sample size may change the relationship between the heterodont and homodont platanistoids.

In contrast to [Lambert, Bianucci & Urbina \(2014\)](#), [Tanaka & Fordyce's \(2015a\)](#) equally weighted strict consensus recovered a monophyletic Platanistoidea that included both *Waipatia maerewhenua* and *Waipatia hectori* ([Tanaka & Fordyce, 2015b](#)), both *Otekaikea* spp., Platanistidae, *Squalodelphis fabianii*, and *Notocetus vanbenedeni* (i.e., a paraphyletic Squalodelphinidae). However, in their implied weighting strict consensus, *Squalodon* was added to Platanistoidea. Allodelphinidae was not included in their analysis. [Tanaka & Fordyce \(2015a\)](#) diagnosed Platanistoidea *sensu stricto* (i.e., with *Squalodon* excluded) based on 6 synapomorphies: presence of the posterior dorsal infraorbital foramina of the maxilla (character 59); C-shaped or weakly curved parabullary sulcus (character 169); presence of the articular rim on the periotic (character 186); presence of the anterior spine of the tympanic bulla (character 195); presence of the anterolateral convexity of the tympanic bulla with anterolateral notch (character 196); and presence of the ventral groove (median furrow) of bulla anteriorly (character 212). [Tanaka & Fordyce \(2015a\)](#) also mentioned that

character 59 was seen in other odontocete lineages besides the Platanistoidea, and it is the only character that is preserved in *Arktocara*.

In a broad review of Allodelphinidae, *Kimura & Barnes (2016)* described three new allodelphinids from the Miocene of western North America and revised the definition of Platanistoidea to include Waipatiidae, Squalodelphinidae, Allodelphinidae, Squalodontidae, and Platanistidae. *Kimura & Barnes (2016)*, however, did not provide a computer-assisted phylogenetic analysis to support their claim about the familial level relationships among platanistoids, pointing instead to a matrix and an analysis in *Barnes (2006)* that included only two outgroups in a taxon list that exclusively contained presumed platanistoids. More crucially, *Kimura & Barnes (2016)* did not perform a phylogenetic analysis nor code the character states for the three novel allodelphinid taxa that they described (i.e., *Goedertius oregonensis*, *Ninjadelpheis ujiharai*, and *Zarhinocetus donnamatsonae* *Kimura & Barnes, 2016*).

Our phylogenetic analysis herein addresses some of the shortfalls of previous studies by including type genera belonging to all potential platanistoid families that have been presented in recent phylogenetic analyses (i.e., *Lambert, Bianucci & Urbina, 2014*; *Tanaka & Fordyce, 2015a*; *Kimura & Barnes, 2016*). We resolved a monophyletic Platanistoidea that included Platanistidae, Waipatiidae (*Waipatia maerewhenua* + *Waipatia hectori* + *Otekaikea marplei* + *Otekaikea huata*), Allodelphinidae and a polyphyletic Squalodelphinidae. We note that, for *Phocageneus venustus*, we followed *Tanaka & Fordyce (2015a)*'s coding, which is primarily based on USNM 21039 (*Kellogg, 1957*). *Lambert, Bianucci & Urbina (2014)* provide a valuable discussion of material that has been referred to this taxon. Our analysis departs most sharply from *Tanaka & Fordyce (2015a)* with the addition of the four allodelphinid genera. Though our recovery of a monophyletic Waipatiidae consisting of all described species of *Waipatia* and *Otekaikea* is consistent with *Tanaka & Fordyce (2015b)*, the authors chose to limit their definition of Waipatiidae to both species of *Waipatia*, differing from the results in *Tanaka & Fordyce (2014)* where the authors defined Waipatiidae as including *Waipatia maerewhenua* and *Otekaikea marplei*. Our results are consistent with *Tanaka & Fordyce (2015a)*'s findings with the resolution of a polyphyletic Squalodelphinidae, with *Squalodelphis fabianii* as a basal member of Platanistoidea and an unnamed clade of *Notocetus vanbenedeni* + *Phocageneus venustus* as the sister group to Platanistidae. A more detailed coding of Squalodelphinidae in future work, especially one that includes *Huaridelphis raimondii*, will provide more insight into the relationships among this group.

We diagnose a node-based Platanistoidea by the following synapomorphies: moderately elevated coronoid process (character 33\*); premaxillae >65% of width of rostrum at antorbital notches (character 51\*); alisphenoidal-squamosal suture coursing along groove for mandibular branch of trigeminal nerve in ventral view (character 147[1]); lateral groove or depression with profile of periotic becoming slightly to markedly sigmoidal in dorsal view (character 166[1]); anteroposterior ridge on dorsal side anterior process and body of periotic (character 167[1]); parabullary sulcus on the periotic weakly to strongly curved and c-shaped (character 169[1,2]); and ventral surface of the posterior process of the periotic not flat along a straight path perpendicular to its long axis (character 191[1,2]) Of

these synapomorphies, the two marked by an asterisk (\*) are equivocal across the group, demonstrating character state reversals or independent origins (characters 33, 51). Two characters are ambiguous and show independent origins (characters 169 and character 191), but we argue remain useful for characterizing this group.

Only one of the six synapomorphies presented by *Tanaka & Fordyce (2015a)* is consistent with ours (character 169). The other 5 characters are all equivocal across the Platanistoidea, but some are still useful for diagnosing members of certain sub-clades. For example, the presence of the articular rim or on the periotic (character 186) is seen in all platanistoids except *Allodelphis pratti*, where there is no distinguishable rim lateral to the posterior process and separated by a sulcus (Fig. 9). In *Zarhinocetus errabundus*, this trait is present as an extremely reduced rim. *Kimura & Barnes (2016)* make no mention of an articular rim or process on the periotic of *Ninjadelphus ujiharai*, and there is no evidence of it from the published photos of the type specimen. The presence of the anterior spine of the tympanic bulla (character 195), the anterolateral convexity of the tympanic bulla with anterolateral notch (character 196), and the ventral groove (median furrow) of bulla anteriorly (character 212) are all ambiguous characters, represented by two states each across Platanistoidea. All of the latter traits are present in *Allodelphis pratti* and *Zarhinocetus errabundus*, with perhaps the exception of the ventral groove of the anterior surface of the bulla in *Allodelphis pratti*, which could not be determined from the photos of the referred specimen (UCMP 83791) provided by *Kimura & Barnes (2016)*, nor was not mentioned in their description of this taxon.

### Systematics of Allodelphinidae

Our analysis recovered Allodelphinidae as a well-supported sub-clade within a node-based Platanistoidea, rooted in a polytomy with *Squalodelphis fabianii* and an unnamed sub-clade that includes *Notocetus vanbenedeni* + *Phocageneus venustus* + Platanistidae. Allodelphinidae in our study is supported by the following synapomorphies: rostral constriction anterior to the antorbital notch (character 9[1]); premaxillae in dorsal view contacting along midline for approximately half of the entire length of the rostrum and partially fused (character 14[3]); buccal teeth entocingulum absent (character 24[1]); greatest diameter of largest functional tooth <3% of greatest width of maxillae at postorbital processes (character 25[2]); angle of anterior edge of supraorbital process and the median line oriented anteromedially (character 35[1]); dorsolateral edge of internal opening of infraorbital foramen formed by maxilla (character 43[0]); posterolateral sulcus shallow or absent (character 57[1]); lack of premaxillary crest or posterior maxillary crest adjacent to nasals (character 72[0]); temporal fossa roofed over by lateral expansion of the maxillae (character 101[1]); palatines partially covered by pterygoid dividing it into medial and lateral exposures (character 121[1]); lateral lamina of palatine (character 122[1]); lateral end of groove for mandibular branch of trigeminal nerve wrapping laterally around posterior end of pterygoid sinus fossa and opening anteriorly (character 148[0]); lack of anterior bullar facet (character 172[1]); elevated caudal tympanic process of periotic with ventral and posterior edges forming a right angle in medial view (character 178[1]); tubular fundus of internal acoustic meatus (character 182[1]) angle between posterior process of periotic and

long axis of pars cochlearis  $\leq 135^\circ$  from dorsal or ventral view (character 189[1]); and ventral surface of posterior process of periotic convex along a straight path perpendicular to its long axis (character 191[2]). Based on the published descriptions and illustrations provided by [Kimura & Barnes \(2016\)](#), the three allodelphinid taxa not included in our phylogenetic analysis (*Allodelphis woodburnei*, *Ninjadelphis ujiharai*, and *Zarhinocetus donnamatsonae*) each possess all of the allodelphinid synapomorphies presented by our analysis.

In their review of Allodelphinidae, [Kimura & Barnes \(2016\)](#) based their diagnosis of this group on comparative characters rather than phylogenetic synapomorphies. Many of these comparative characters can be readily observed in all platanistoids, such as the posteriorly extended lateral lamina of the pterygoid and palatine (except in species of *Waipatia* and *Otekaikea* where the palatine is poorly preserved or missing), and a tympanic bulla with elongated and pointed anterior process, among others. Nevertheless, our diagnosis is consistent with [Kimura & Barnes \(2016\)](#)'s concept of Allodelphinidae with only two exceptions. First, [Kimura & Barnes \(2016\)](#) report that, in allodelphinids, the posterior ends of the premaxillae are separated from the lateral sides of the corresponding nasal bones, beginning with a more "primitive" state in *Allodelphis pratti* where only one premaxilla is separated from the corresponding nasal by a tiny exposure of maxilla, to further "derived" states in *Ninjadelphis* and *Zarhinocetus* where the premaxillae are further retracted anteriorly onto the facial region and away from the nasals. However, it is unclear in the more "primitive" state of *Allodelphis* whether the lack of contact between the premaxilla and nasal could be a result of diagenetic breakage, or individual variation. Furthermore, speculations on the more "derived states" in taxa such as *Ninjadelphis*, are based on specimens with incomplete premaxillae. In *Goedertius oregonensis*, the premaxillae are not separated from the nasals. This condition is likely also true for *Arktocara yakataga*: although the nasals are missing, the premaxillae directly abut the nasal fossa of the frontal, and therefore would most likely have been in direct contact with the nasals. Further extensive comparative work on allodelphinid taxa (including the multiple specimens housed at USNM that can readily be referred to *Goedertius* sp. ([Fig. S2](#))) will help to clarify the distribution and diagnostic utility of these traits.

Second, [Kimura & Barnes \(2016\)](#) diagnosed Allodelphinidae by an absence of both the preorbital and postorbital lobe of the pterygoid sinus. Both fossae for the pre- and postorbital lobe of the pterygoid sinus are unclear in the type specimen of *Allodelphis pratti*, in part due to obstruction by unprepared matrix. However, in *Arktocara yakataga*, though there is no obvious indication of a postorbital lobe of the pterygoid sinus, the deep and broad fossa surrounding the ventral infraorbital foramen and the sphenopalatine foramen anteromedial of the orbit suggests the presence of a preorbital lobe.

Originally assigned to Platanistidae by [Wilson \(1935\)](#), *Allodelphis pratti* was referred to the Platanistidae by [Barnes \(1977\)](#), and later [Barnes \(2006\)](#) erected a new group, Allodelphinidae, for it. However, in both instances, [Barnes \(1977\)](#) and [Barnes \(2006\)](#) did not provide an explanation for why the Allodelphinidae belong to the Platanistoidea. Of the 7 synapomorphies for Platanistoidea identified by our phylogenetic analysis, the Allodelphinidae possessed 4 of the 5 unequivocal characters: lateral groove or depression with profile of periotic becoming slightly to markedly sigmoidal in dorsal view (character

166[1]); anteroposterior ridge on dorsal side anterior process and body of periotic (character 167[1]); parabullary sulcus on the periotic weakly to strongly curved and c-shaped (character 169[1,2]); and ventral surface of the posterior process of the periotic not flat along a straight path perpendicular to its long axis (character 191[1,2]). The fifth unequivocal character (147), could not be observed in any of the Allodelphinidae specimens. In addition, the type specimen of *Ninjadelphis* is the only allodelphinid specimen with an associated scapula, and it is missing the coracoid process. This agreement with *Muizon's* (1987) platanistoid synapomorphy—the loss or reduction of the coracoid process of the scapula—suggests that, though the process is still present in some putative platanistoids (i.e., *Otekaikea huata*), the character may still be relevant for diagnosing Platanistoidea (*Kimura & Barnes, 2016*). We urge future studies on Allodelphinidae to not only include all available genera (if not putative species), but also to explicitly test phylogenetic hypotheses in a repeatable analytical framework.

### Morphological comparisons

Of the 7 supporting synapomorphies for Platanistoidea in our study, none of the unequivocal synapomorphies are preserved and demonstrated on the skull of *Arktocara*. However, one equivocal synapomorphy is preserved in *Arktocara*: width of the premaxillae >50% of the width of the rostrum at the antorbital notch (character 51[1]). Though the type specimen of *Arktocara* lacks tympanoperiotics, it is closely allied with *Allodelphis pratti*, whose periotic shares three more of the platanistoid synapomorphies: presence of lateral groove or depression with the profile of the periotic becoming slightly to markedly sigmoidal in dorsal view (character 166[1]); anteroposterior ridge developed on anterior process and body of periotic in dorsal view (character 167[1]); and a curved C-shaped parabullary sulcus (character 169[2]; see [Fig. 9](#) for illustration of the periotic synapomorphies on the type specimen of *Allodelphis pratti*). Therefore, in the absence of tympanoperiotics associated with new cranial material of *Arktocara*, we are confident that these elements would share many features with *Allodelphis pratti*, the sister taxon of *Arktocara*.

Overall, the allodelphinid that most resembles *Arktocara* in morphology is *Allodelphis pratti* ([Figs. 7–9](#)), originally described by *Wilson (1935)* from the Jewett Sand in Kern County, California, USA. Having examined the holotype—consisting of a skull with associated rostrum and jaw fragments, other skull fragments, and a right periotic—we note that the material included in the holotype actually belongs to more than one individual. For example, the holotype includes an isolated left postglenoid process, even though the holotype skull still has this feature intact. However, having examined the right periotic in relation to the periotic fossa, we are confident in the association of this element to the holotype skull. Any further study including the holotype of *Allodelphis* should take this caveat into consideration. *Allodelphis* is similar in size and shape to the type of *Arktocara*, with wide, hexagonally shaped craniums and postorbital widths within 2 cm of one another. In dorsal view, the two genera are alike in having their premaxillae rise above the maxillae for the entire length of the cranium from the level of the antorbital notch to the cranial vertex, forming an anteroposteriorly elongated and dorsally elevated plateau in relation to the broad, flat maxilla across the facial region. In both genera, this premaxillary

plateau continues posteriorly to a tabular vertex, posterior to the external bony nares. The exposures of the frontals and nasals are symmetrical on the vertex, and there is no evident leftward skew or other facial asymmetry. The nasals are also transversely widened anteriorly, setting these two genera apart from all other allodelphinids. *Kimura & Barnes (2016)* showed photos of the holotype of *Allodelphis pratti* with a feature on the skull labeled “posterior dorsal infraorbital foramen.” However, having examined the holotype, we do not see any evidence of a posterior infraorbital foramina—the featured labelled in the photo is a small break in the maxilla overlying the frontal. While it is possible that a posterior dorsal infraorbital foramen is hidden under the jaw fragment that is adhered to the right maxilla, we refrain from definitively stating any foramen exists. Both *Arktocara yakataga* and *Allodelphis pratti* have a nuchal crest weakly convex anteriorly, a widely open mesorostral canal anterior to the bony nares, the maxilla covering almost all of the frontal along the supraorbital process, and the posterior end of the basioccipital crest separated from the rest of the crest by a narrow crease.

The coded character state differences between *Arktocara yakataga* and *Allodelphis pratti* are listed in the Diagnosis section, above, although we provide more descriptive differences between these two taxa, as follows. First, *Arktocara* differs from *Allodelphis* in dorsal view by having: a deeper mesorostral canal anterior to the external nares; straight lateral margins of the premaxillae lateral and posterior of the external nares; no exposure of the maxillae on the vertex; a greater transverse constriction of the lateral margins of the maxilla/frontal anterior to the nuchal crest; a less extreme flaring of the posterior temporal crest along the occipital border; and more prominent dorsal infraorbital foramina, with posteriorly directed sulci. In lateral view, *Arktocara* shows a markedly reduced postglenoid process and zygomatic process of the squamosal, and a more posterolaterally directed postorbital process as opposed to a ventrally oriented process in *Allodelphis*. In ventral view, *Arktocara* has a more elevated vomerine keel. We argue that these differences, along with those coded in the phylogenetic analysis, provide the basis for *Arktocara yakataga*'s status as a new genus of allodelphinid.

*Arktocara* also differs in clear ways from three allodelphinids (*sensu Kimura & Barnes, 2016*) that were not included in the phylogenetic analysis: *Ninjadelphus ujiharai*, *Allodelphis woodburnei*, and *Zarhinocetus donnamatsonae*. *Arktocara* differs from both *Ninjadelphus ujiharai* and *Zarhinocetus donnamatsonae* in having: a wider opening of the mesorostral canal, anterior to the external nares in dorsal view; anteroposteriorly straight lateral margins of the premaxillae both lateral and posterior of the external bony nares, in dorsal view; the posterior ends of the premaxillae extending posterior of the nasals; nasals expanding in width anteriorly rather than narrowing anteriorly; a reduced post-glenoid process; and a broader extent of the maxilla above the supraorbital process of the frontal. *Arktocara* further differs from both *Ninjadelphus ujiharai* and *Zarhinocetus donnamatsonae* in lacking a dorsal depression on the base of the rostrum formed by ventromedially sloping of the premaxillae and maxillae, and lacking an asymmetrical skew to the vertex or nuchal crest.

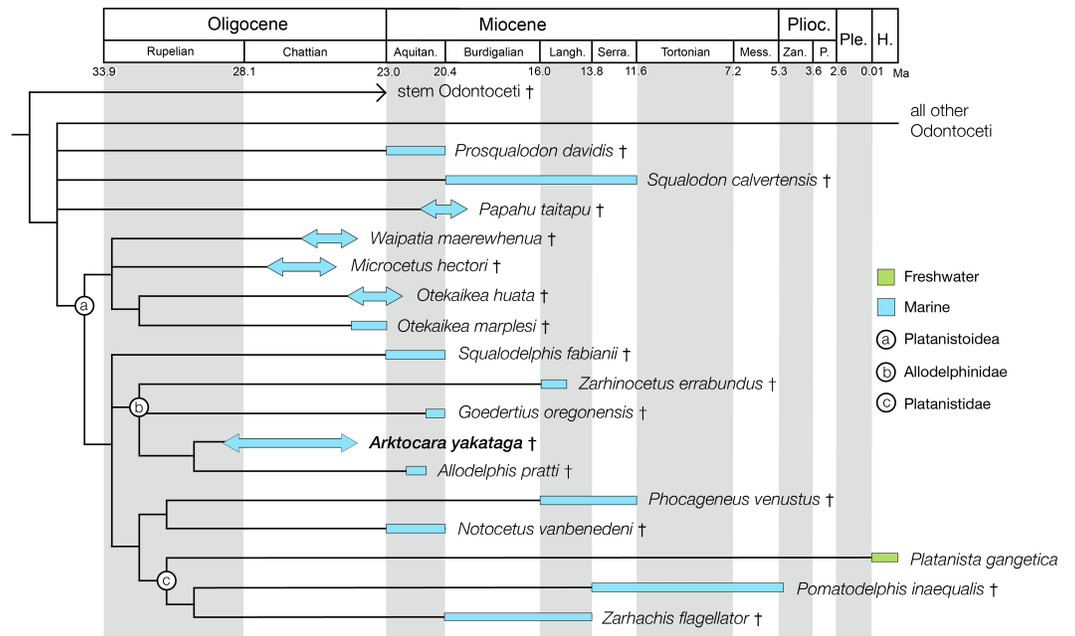
*Arktocara* further differs from *Ninjadelphus ujiharai* in: lacking exposures of the maxillae on the vertex; lacking a glenoid fossa facing anteriorly as opposed to anteromedially; lacking widely diverging basioccipital crests; and lacking a depressed pit of the posterior

end of the maxilla with an overhanging lip of the nuchal crest. *Arktocara* also differs from *Zarhinocetus donnamatsonae* in having: a more prominent and flaring temporal crest; a zygomatic process more tapered anteriorly in lateral view; the absence of a maxillary tuberosity on the lateral edge of the maxillary flange immediately anterior to the antorbital notch; no reduction of the maxilla on the supraorbital process to expose a thick band of frontal; and lacking a maxillary crest on the supraorbital process in dorsal view. *Arktocara* differs from *Allodelphis woodburnei* in having: a smaller and more anteriorly tapered zygomatic process; a reduced postglenoid process; the absence of a prominent fossa on each side of the sagittal crest on the supraoccipital; the premaxillae sloping medially towards the mesorostral canal on the posterior rostrum; and a glenoid fossa directed anteriorly rather than anteroventrally.

### Geological & geographic significance

Today, *Platanista gangetica* is distributed in two subspecies across the Indus, Ganges-Brahmaputra-Megna and Karnaphuli-Sangu river systems of Southeast Asia, and remains highly threatened by human activities, including by-catch, fishing, and habitat modification (e.g., [Braulik et al., 2014a](#)). The fossil record of all other Platanistoidea demonstrates that the immediate relatives of *Platanista gangetica* comprise a morphologically diverse group of small to medium sized odontocetes that are distributed globally in marine sediments of Oligocene and Miocene age (see [Bianucci et al. \(2013\)](#) and [Hulbert & Whitmore Jr \(2006\)](#) for two exceptional occurrences of platanistid specimens in freshwater sediments of Peru and Alabama, respectively). There is no fossil record for the genus *Platanista*, but recent work on mitochondrial DNA haplotype diversity ([Braulik et al., 2014b](#)) places the divergence between subspecies across at around 550,000 years ago (with 95% posterior probability 0.13–1.05 million years ago). The strong ecological disparity between *Platanista*'s obligate freshwater lifestyle and the presumed marine lifestyle of all other named platanistoids ([Fig. 11](#)) implies some kind of differential evolutionary success for this group, with potentially higher extinction rates in Platanistoidea. [Fordyce & Muizon \(2001\)](#) first proposed that competition between platanistoids and early delphinooids may explain the strong difference in taxonomic richness observed in their fossil records, but this suggestion has never been tested in a rigorous framework ([Fordyce, 2003](#); [Marx, Lambert & Uhen, 2016](#)).

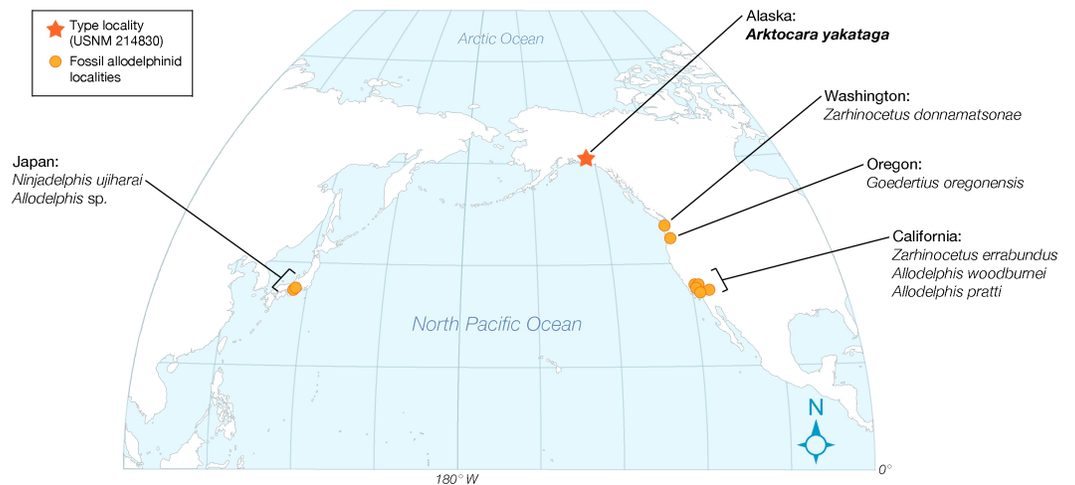
Platanistoids first appear in the fossil record in the late Oligocene, and reach peak richness in the early Miocene ([Kimura & Barnes, 2016](#); [Tanaka & Fordyce, 2015a](#)). The oldest platanistoids with solid age constraints are the waipatiids, all found in the Oligocene-Miocene Otekaike Limestone of New Zealand ([Graham et al., 2000](#); [Benham, 1935](#); [Fordyce, 1994](#); [Tanaka & Fordyce, 2014](#); [Tanaka & Fordyce, 2015a](#)). Based on both the lithology and the presence of age-diagnostic planktic foraminifera and ostracod species, *Waipatia hectori* ([Benham, 1935](#)) is the oldest reported waipatiid, from the uppermost Duntroonian Stage of the Otekaike Limestone, approximately 25.2 Ma ([Tanaka & Fordyce, 2015b](#)). *Arktocara* is possibly very similar in age to *Waipatia hectori*, constrained to the Chattian Stage of the upper Oligocene in the Poul Creek Formation, approximately ~24–29 Ma ([Perry, Garver & Ridgway, 2009](#)). Unfortunately, the lack of robust locality data for either *Waipatia hectori* or *Arktocara* makes impossible to determine which is the oldest.



**Figure 11** Phylogenetic results of Platanistoidea and major odontocete groups, calibrated for geologic time. Time calibrated phylogenetic tree of the Platanistoidea, pruned from the consensus cladogram in Fig. 10. The groups ‘stem Odontoceti’ and ‘all other Odontoceti’ were left as collapsed outgroups. Stratigraphic range data were derived from published accounts for each taxon, including global ranges. Geologic time scale based on *Cohen et al. (2013)*. Stem Odontoceti node depth follows mean divergence date estimates by *McGowen, Spaulding & Gatesy (2009)*; all other nodes (Platanistoidea, Allodelphinidae) should be considered graphical heuristics, and do not reflect divergence dates. Thick bars correspond to the stratigraphic ranges of each taxon, with arrows indicating lower confidence in stratigraphic boundaries. Ecological habitat preference (freshwater vs. marine) is indicated by bar colour, and is based on depositional environment or extant habitat. Labelled circles denote node-based clades. Abbreviations: Aquitan., Aquitanian; H., Holocene; Langh., Langhian; Mess., Messinian; P., Piacenzian; Ple., Pleistocene; Plioc., Pliocene; Serra., Serravallian; Zan., Zanclean.

*Arktocara* is, however, very clearly the oldest known allodelphinid, expanding the previously reported age range of Allodelphinidae by as much as 9 million years (*Kimura & Barnes, 2016*). Other allodelphinids span temporally from the early to middle Miocene, which largely matches the stratigraphic range of other platanistoid lineages (Fig. 11). Interestingly, *Arktocara* is among the oldest crown Odontoceti, reinforcing the long-standing view that the timing for the diversification for crown lineages must have occurred no later than the early Oligocene.

Lastly, Allodelphinidae appear uniquely limited, in terms of geography, to marine rocks of the North Pacific Ocean, with occurrences in Japan, Alaska, Washington State, Oregon, and California (see Fig. 12; *Kimura & Barnes, 2016*). *Arktocara* expands this geographic range to sub-Arctic latitudes. At approximately 60° N in the Yakutat City and Borough, *Arktocara* is the most northern platanistoid yet reported. The next most northern platanistoid reported is an incomplete and unnamed specimen from the late Chattian marine Vejle Fjord Formation in northern Denmark, approximately 56.7°N, 9.0°E (*Hoch, 2000*).



**Figure 12** Distribution map of fossil Allodelphinidae. Mapped of fossil localities of allodelphinids, projected on a truncated Winkel Tripel map and centered on 25°N and 170°W. Occurrences for fossil data derive from location of type and referred localities for each taxon, are listed alphabetically by region, and are represented by orange dots.

### Anatomical Abbreviations

<b>al.</b>	alisphenoid
<b>bsph.</b>	basisphenoid
<b>ex.</b>	exoccipital
<b>falc. process</b>	falciform process
<b>f. ovale</b>	foramen ovale
<b>fr.</b>	frontal
<b>lac.</b>	lacrimojugal
<b>ll./ lat. lam. pterygoid</b>	lateral lamina of the pterygoid
<b>infr. Foram.</b>	infraorbital foramina
<b>Ma.</b>	mega-annum period of 1 million years
<b>max.</b>	maxilla
<b>m. lam. pt.</b>	medial lamina of the pterygoid
<b>mes.</b>	mesethmoid
<b>ns.</b>	nasal
<b>o./ orb.</b>	orbitosphenoid
<b>pal.</b>	palatine
<b>par.</b>	parietal
<b>p. glenoid process</b>	postglenoid process
<b>p. lam. pterygoid</b>	posterior lamina of the pterygoid
<b>pmx.</b>	premaxilla
<b>pt.</b>	pterygoid
<b>sphpal. foram.</b>	sphenopalatine foramen
<b>v.</b>	ventral
<b>?</b>	displaced skull fragment of unknown origin

### Institutional Abbreviations

IGUP	Geological Institute of Padua University, Padua, Italy.
LACM	Departments of Mammalogy and Vertebrate Paleontology, Natural History Museum of Los Angeles County, Los Angeles, California, USA.
OU	Geology Museum, University of Otago, Dunedin, New Zealand.
UCMP	University of California Museum of Paleontology, Berkeley, California, USA.
USNM	Departments of Paleobiology and Vertebrate Zoology (Division of Mammals), National Museum of Natural History, Smithsonian Institution, Washington, District of Columbia, USA.
YPM	Division of Vertebrate Paleontology, Yale Peabody Museum, New Haven, Connecticut, USA.

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Nicholas D. Pyenson is an Academic Editor for PeerJ. This does not alter the authors' adherence to PeerJ policies on sharing data and materials.

### Author Contributions

- Alexandra T. Boersma and Nicholas D. Pyenson conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, wrote the paper, prepared figures and/or tables, reviewed drafts of the paper.

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The following information was supplied regarding the registration of a newly described species:

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Publication LSID: urn:lsid:zoobank.org:pub:0194A593-DBE0-47CA-A41F-04A37931BA2F

*Arktocara* LSID: urn:lsid:zoobank.org:act:EE11B95B-8338-496B-97F4-1673ED90E709

*Arktocara yakataga* LSID: urn:lsid:zoobank.org:act:FBCF0EAA-7BBB-4EF0-8186-7548993098D1.

### Supplemental Information

Supplemental information for this article can be found online at <http://dx.doi.org/10.7717/peerj.2321#supplemental-information>.

## REFERENCES

- Aguirre-Fernández G, Fordyce RE. 2014.** *Papahu taitapu*, gen. et sp. nov., An Early Miocene Stem Odontocete (Cetacea) from New Zealand. *Journal of Vertebrate Paleontology* **34**(1):195–210 DOI [10.1080/02724634.2013.799069](https://doi.org/10.1080/02724634.2013.799069).
- Allen GM. 1921.** A new fossil cetacean. *Bulletin of the Museum of Comparative Zoology* **65**(1):1–14.
- Barnes LG. 1977.** Outline of eastern North Pacific fossil cetacean assemblages. *Systematic Zoology* **25**:321–343.
- Barnes LG. 2006.** A phylogenetic analysis of the superfamily Platanistoidea (Mammalia, Cetacea, Odontoceti). *Beitrage zur Palaontologie* **30**:25–42.
- Barnes LG, Kimura T, Godfrey SG. 2010.** The evolutionary history and phylogenetic relationships of the superfamily Platanistoidea. In: Ruiz-Garcia M, Shostell JM, eds.

- Biology, evolution and conservation of River Dolphins within South America and Asia*. New York: Nova Publishers, 445–488.
- Barnes LG, Reynolds RE. 2009.** A new species of Early Miocene allodelphinid dolphin (Cetacea, Odontoceti, Platanistoidea) from Cajon Pass, southern California. *Museum of Northern Arizona Bulletin* **65**:483–507.
- Benham WB. 1935.** The teeth of an extinct whale, *Microcetus hectori* n. sp. *Transactions of the Royal Society of New Zealand* **65**:239–243.
- Bianucci G, Lambert L, Salas-Gismondi R, Tejada J, Pujos F, Urbina M, Antoine P-O. 2013.** A Miocene relative of the Ganges River dolphin (Odontoceti, Platanistidae) from the Amazonian Basin. *Journal of Vertebrate Paleontology* **33**(3):741–745  
DOI [10.1080/02724634.2013.734888](https://doi.org/10.1080/02724634.2013.734888).
- Braulik GT, Arshad M, Noureen U, Northridge SP. 2014a.** Habitat fragmentation and species extirpation in freshwater ecosystems; causes of range decline of the Indus River Dolphin (*Platanista gangetica minor*). *PLoS ONE* **9**(7):e101657  
DOI [10.1371/journal.pone.0101657](https://doi.org/10.1371/journal.pone.0101657).
- Braulik GT, Barnett R, Odon V, Islas-Villanueva V, Hoelzel AR, Graves JA. 2014b.** One species or two? Vicariance, lineage divergence and low mtDNA diversity in geographically isolated populations of South Asian River dolphin. *Journal of Mammalian Evolution* **22**(1):111–120.
- Brisson MJ. 1762.** *Regnum animale in Classes IX distributum, sive synopsis methodica sistens generalem animalium distributionem in Classes IX, et duarum primarum Classium, Quadrupedum scilicet & Cetaceorum, particulare divisionem in Ordines, Sectiones, Genera, et Species*. Paris: T. Haak.
- Buchholtz EA. 2001.** Vertebral osteology and swimming style in living and fossil whales (Order: Cetacea). *Journal of Zoology* **253**:175–190 DOI [10.1017/S0952836901000164](https://doi.org/10.1017/S0952836901000164).
- Cantino PD, De Queiroz K. 2014.** *Phylocode: a phylogenetic code of biological nomenclature*. Version 4c. Ohio University. Available at <http://www.ohiou.edu/phylocode> (accessed on 11 May 2016).
- Case EC. 1934.** A specimen of a long-nosed dolphin from the Bone Valley Gravels of Polk County, Florida. *Contributions from the Museum of Paleontology, University of Michigan* **4**(6):105–113.
- Cohen KM, Finney SC, Gibbard PL, Fan J-X. 2013.** The ICS international chronostratigraphic chart. *Episodes* **36**:199–204.
- Cope ED. 1868.** Second contribution to the history of the Vertebrata of the Miocene period of the United States. *Proceedings of the Academy of Natural Sciences of Philadelphia* **1868**:184–194.
- Dal Piaz G. 1917.** Gli Odontoceti del Miocene bellunese, Parte Terza. *Squalodelphis fabianii*. *Memoire dell' Istituto Geologico della R. Università di Padova* **5**(1):1–34.
- D'Orbigny A. 1834.** Notice sur un nouveau genre de cétacé des rivières du centre de l'Amérique méridionale. *Nouvelle Annales du Muséum d'Histoire Naturelle Paris* **3**:28–36.

- Flower WH. 1867.** Description of the skeleton of *Inia geoffrensis* and of the skull of *Pontoporia blainvillii*, with remarks on the systematic position on these animals in the order Cetacea. *Transactions of the Zoological Society of London* **6**:87–116.
- Flynn TT. 1923.** A whale of bygone days. *Australian Museum Magazine* **1**:266–272.
- Fordyce RE. 1994.** *Waipatia maerewhenua*. New genus and new species (Waipatiidae, New Family), an archaic late oligocene dolphin (Cetacea: Odontoceti: Platanistoidea) from New Zealand. In: Berta A, Demere TA, eds. *Contributions in marine mammal Paleontology honoring Frank C. Whitmore, Jr. Proceedings of the San Diego Society of Natural History*, vol. 29. San Diego, 147–176.
- Fordyce RE. 2003.** Cetacea evolution and Eocene-Oligocene ocean revisited. In: Prothero DR, Ivany LC, Nesbitt E, eds. *From greenhouse to icehouse: the marine Eocene-Oligocene transition*. New York: Columbia University Press.
- Fordyce RE, De Muizon C. 2001.** Evolutionary history of whales: a review. In: Mazin JM, De Buffrenil V, eds. *Secondary adaptation of tetrapods to life in water*. München: Pfeil, 169–234.
- Geisler JH, McGowen MR, Yang G, Gatesy J. 2011.** A supermatrix analysis of genomic, morphological and paleontological data from crown Cetacea. *BMC Evolutionary Biology* **11**:112 DOI 10.1186/1471-2148-11-112.
- Geisler JH, Sanders AE. 2003.** Morphological evidence for the phylogeny of Cetacea. *Journal of Mammalian Evolution* **10**(1):23–129 DOI 10.1023/A:1025552007291.
- Geographic Names Information System (GNIS) USGS. 2016.** Feature detail report for: Cape Yakataga. Available at [http://geonames.usgs.gov/apex/f?p=gnispq:3:0::NO::P3\\_FID:1399935](http://geonames.usgs.gov/apex/f?p=gnispq:3:0::NO::P3_FID:1399935) (accessed on 9 May 2016).
- Graham IJ, Morgans H, Waghorn DB, Trotter JA, Whitford DJ. 2000.** Strontium isotope stratigraphy of the Oligocene Miocene Otekaike Limestone (Trig Z section) in southern New Zealand: age of the Duntroonian/Waitakian Stage boundary. *New Zealand Journal of Geology and Geophysics* **43**(3):335–347 DOI 10.1080/00288306.2000.9514891.
- Gray JE. 1846.** On the cetaceous animals. In: *The zoology of the voyage of H M. S. Erebus and Terror, under the command of Capt. Sir J. C. Ross, R. N., F. R. S., during the years 1839 to 1843*. 1 & 2. 1–53.
- Hoch E. 2000.** Olfaction in whales: evidence from a young odontocete of the Late Oligocene North Sea. *Historical Biology* **14**:670–689.
- Hulbert RC, Whitmore Jr FC. 2006.** Late Miocene mammals from the Mauvilla Local Fauna, Alabama. *Bulletin of the Florida Museum of Natural History* **46**(1):1–28.
- Jefferson TA, Webber MA, Pitman RL. 2008.** *Marine mammals of the world: a comprehensive guide to their identification*. Oxford: Academic Press.
- Joyce WG, Parham JF, Gauthier J. 2004.** Developing a protocol for the conversion of rank-based taxon names to phylogenetically defined clade names, as exemplified by turtles. *Journal of Paleontology* **78**:989–1013 DOI 10.1666/0022-3360(2004)078<0989:DAPFTC>2.0.CO;2.

- Kellogg R. 1923.** Description of two squalodonts recently discovered in the Calvert Cliffs, Maryland; and notes on the shark-toothed cetaceans. *Proceedings of the US National Museum* **62**(16):1–69.
- Kellogg R. 1957.** Two additional Miocene porpoises from the Calvert Cliffs, Maryland. *Proceedings of the U.S. National Museum* **107**(3387):279–337.
- Kimura T, Barnes LG. 2016.** New Miocene fossil Allodelphinidae (Cetacea, Odontoceti, Platanistoidea) from the North Pacific Ocean. *Bulletin of the Gunma Museum of Natural History* **20**:1–58.
- Lambert O, Bianucci G, Urbina M. 2014.** *Huaridelphis raimondii*, a new early Miocene Squalodelphinidae (Cetacea, Odontoceti) from the Chilcatay Formation, Peru. *Journal of Vertebrate Paleontology* **34**(5):987–1004 DOI [10.1080/02724634.2014.858050](https://doi.org/10.1080/02724634.2014.858050).
- Lambert O, De Muizon C, Bianucci G. 2015.** A new archaic homodont toothed cetacean (Mammalia, Cetacea, Odontoceti) from the early Miocene of Peru. *Geodiversitas* **37**(1):79–108 DOI [10.5252/g2015n1a4](https://doi.org/10.5252/g2015n1a4).
- Lebeck HJ. 1801.** *Delphinus delphis* beschrieben von Herrn Heinrich Julius Lebeck. *Der Gesellschaft Naturforschender Freunde zu Berlin, Neue Schriften* **2**:280–282.
- Leidy J. 1869.** The extinct mammalian fauna of Dakota and Nebraska, including an account of some allied forms from other localities, together with a synopsis of the mammalian remains of North America. *Journal of the Academy of Natural Sciences, Philadelphia* **2**:1–472.
- Marx FG, Fordyce RE. 2015.** Baleen boom and bust: a synthesis of mysticete phylogeny, diversity and disparity. *Royal Society Open Science* **2**(4):140434 DOI [10.1098/rsos.140434](https://doi.org/10.1098/rsos.140434).
- Marx FG, Lambert O, Uhen MD. 2016.** *Cetacean paleobiology*. Oxford: John Wiley & Sons.
- McGowen MR, Spaulding M, Gatesy J. 2009.** Divergence date estimation and a comprehensive molecular tree of extant cetaceans. *Molecular Phylogenetics and Evolution* **53**(3):891–906 DOI [10.1016/j.ympev.2009.08.018](https://doi.org/10.1016/j.ympev.2009.08.018).
- Mead JG, Fordyce RE. 2009.** *The Therian skull: a lexicon with emphasis on the odontocetes*. Washington, D.C.: Smithsonian Institution Scholarly Press.
- Miller GS. 1918.** A new river-dolphin from China. *Smithsonian Miscellaneous Collections* **68**(2):1–12.
- Miller DJ. 1971.** Geologic map of the Yakataga district, Gulf of Alaska Tertiary Province, Alaska. United States Geological Survey Map I-610, scale 1: 125,000. Available at [http://ngmdb.usgs.gov/Prodesc/proddesc\\_9402.htm](http://ngmdb.usgs.gov/Prodesc/proddesc_9402.htm).
- Montagu G. 1821.** Description of a species of *Delphinus*, which appears to be new. *Memoirs of the Wernerian Natural History Society* **3**:75–82.
- Moreno FP. 1892.** Lijeros apuntes sobre dos generos de cetaceos fosiles de la Republica Argentina. *Revista del Museo de La Plata* **3**:393–400.
- Muizon C de. 1984.** Les vertébrés fossiles de la Formation Pisco (Pérou) Deuxième partie: les Odontocètes (Cetacea, Mammalia) du Pliocène inférieur de Sud-Sacaco. *Travaux de l'Institut Français d'Études Andines* **27**:1–188.

- Muizon C de. 1985.** Nouvelles données sur le diphylétisme des Dauphins de rivière (Odontoceti, Cetacea, Mammalia). *Comptes rendus l'Academie des Sciences series 2* **301**:359–362.
- Muizon C de. 1987.** The affinities of *Notocetus vanbenedeni*, an Early Miocene Platanistoid (Cetacea, Mammalia) from Patagonia, Southern Argentina. *American Museum Novitates* **2904**:1–27.
- Muizon C de. 1988a.** Les relations phylogénétiques des Delphinida (Cetacea, Mammalia). *Annales de Paleontologie* **74**:159–227.
- Muizon C de. 1988b.** Le polyphylétisme des Acrodelphidae, Odontocètes longirostres du Miocène européen. *Bulletin du Muséum National d'Histoire Naturelle, Paris Section C: Sciences de la Terre, Paléontologie, Géologie, Minéralogie* **10**:31–88.
- Muizon C de. 1991.** A New Ziphiidae (Cetacea) from the early Miocene of Washington State (USA) and phylogenetic analysis of the major groups of odontocetes. *Bulletin du Muséum National d'Histoire Naturelle Paris (4)* **12**:279–326.
- Muizon C de. 1994.** Are the squalodonts related to the platanistoids? In: Berta A, Demere TA, eds. *Contributions in marine mammal paleontology honoring Frank C. Whitmore, Jr. Proceedings of the San Diego Society of Natural History*, vol. 29. San Diego, 135–146.
- Murakami M, Shimada C, Hikida Y, Hirano H. 2012.** Two new extinct basal phocoenids (Cetacea, Odontoceti, Delphinoidea), from the upper Miocene Koetoi Formation of Japan and their phylogenetic significance. *Journal of Vertebrate Paleontology* **32**:1172–1185 DOI [10.1080/02724634.2012.694337](https://doi.org/10.1080/02724634.2012.694337).
- Orth DJ. 1967.** *Dictionary of Alaska place names*. Washington, D.C.: United States Geological Survey.
- Owen R. 1853.** *Descriptive catalogue of the osteological series contained in the Museum of the Royal College of Surgeons of England*. London: Taylor & Francis.
- Perrin WF. 1975.** Variation of spotten and spinner porpoise (genus *Stenella*) in the eastern Pacific and Hawaii. *Bulletin of the Scripps Institution of Oceanography of the University of California* **21**:1–206.
- Perry SE, Garver JI, Ridgway KD. 2009.** Transport of the Yakutat Terrane, southern Alaska: evidence from sediment petrology and detrital zircon fission-track and U/Pb double dating. *The Journal of Geology* **117**:156–173 DOI [10.1086/596302](https://doi.org/10.1086/596302).
- Plafker G. 1987.** Regional geology and petroleum potential of the northern Gulf of Alaska continental margin. In: Scholl DW, Grantz A, Vedder JG, eds. *Geology and resource potential of the continental margin of western North America and adjacent ocean basins: Beaufort Sea to Baja California*. Earth Science Series. Houston: Circumpacific Council for Energy and Mineral Resources, 229–268.
- Plafker G, Moore JC, Winkler GR. 1994.** Geology of the southern Alaska margin. In: Plafker G, Berg HC, eds. *The geology of Alaska (Geology of North America, volume G-1)*. Boulder: Geological Society of America, 389–449.
- Pyenson ND, Sponberg SN. 2011.** Reconstructing body size in extinct crown Cetacea (Neoceti) using allometry, phylogenetic methods and tests from the fossil record. *Journal of Mammalian Evolution* **18**(4):269–288 DOI [10.1007/s10914-011-9170-1](https://doi.org/10.1007/s10914-011-9170-1).

- Pyenson ND, Velez-Juarbe J, Gutstein CS, Little H, Vigil D, O’Dea A. 2015.** *Isthminia panamensis*, a new fossil inioid (Mammalia, Cetacea) from the Chagres Formation of Panama and the evolution of ‘river dolphins’ in the Americas. *PeerJ* **3**:e1227 DOI 10.7717/peerj.1227.
- Rice DW. 1998.** Marine mammals of the world, systematics and distribution. *Society for Marine Mammology Special Publication* **4**:1–231.
- Simpson GG. 1945.** The principles of classification, and a classification of mammals. *Bulletin of the American Museum of Natural History* **85**:1–350.
- Taliaferro NL. 1932.** Geology of the Yakataga, Katalla, and Nichawak districts, Alaska. *Bulletin of the Geological Society of America* **43**:749–782 DOI 10.1130/GSAB-43-749.
- Tanaka Y, Fordyce RE. 2014.** Fossil Dolphin *Otekaikea marplesii* (Latest Oligocene, New Zealand) expands the morphological and taxonomic diversity of Oligocene Cetaceans. *PLoS ONE* **9**(9):e107972 DOI 10.1371/journal.pone.0107972.
- Tanaka Y, Fordyce RE. 2015a.** A new Oligo-Miocene dolphin from New Zealand: *Otekaikea huata* expands diversity of the early Platanistoidea. *Palaeontologia Electronica* **18**(2.23A):1–71.
- Tanaka, Fordyce RE. 2015b.** Historically significant late Oligocene dolphin *Microcetus hectori* Benham 1935: a new species of *Waipatia* (Platanistoidea). *Journal of the Royal Society of New Zealand* **45**(3):135–150 DOI 10.1080/03036758.2015.1016046.
- Tebenkov MD. 1852.** *Atlas sieu-zapadeykh beregov Ameriki (Atlas of the N.W. shores of America from Bering Strait to Cape Corientes and the Aleutian Islands, together with some places on the N.E. shore of Asia)*. St. Petersburg: Tipografiia Morskogo Kadetskogo Korpusa.
- The Society for Marine Mammology’ Committee on Taxonomy. 2015.** List of marine mammal species & subspecies. Available at <https://www.marinemammalscience.org/species-information/list-of-marine-mammal-species-subspecies/> (accessed on 11 May 2016).
- Uhen MD, Pyenson ND. 2007.** Diversity estimates, biases, and historiographic effects: resolving cetacean diversity in the Tertiary. *Palaeontologia Electronica* **10**(2):1–22.
- Wagler JG. 1830.** *Natürliches System der Amphibien, mit vorangehender classification der Säugethiere und Vögel: ein Beitrag zur vergleichenden Zoologie*. München: J.G. Cotta’schen Buchhandlung.
- Wilson LE. 1935.** Miocene marine mammals from the Bakersfield region, California. *Peabody Museum of Natural History Bulletin* **4**:13–26.