Feeding in marine mammals: An integration of evolution and ecology through time
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ABSTRACT

Marine mammals are key components of aquatic ecosystems. Feeding strategies identified in extant cetaceans, pinnipeds, sirenians, marine otters, and polar bears are associated with anatomical specializations of the head (rostrum, palate, temporomandibular joint, teeth/baleen, mandible). Genetic and ontogenetic evidence of skull and tooth morphology provide the mechanisms that underlie patterns of feeding diversity. Based on a comprehensive diversity data set derived from the Paleobiology Database, we considered feeding strategies (suction, biting, filter feeding, grazing), prey type (squid, fish, benthic invertebrates, zooplankton, tetrapods, sea grasses), tooth pattern and cusp shape (homodont, heterodont, pointed, rounded, edentulous), and habitat (marine, riverine, estuarine) in fossil and extant marine mammals. These variables were then tested for correlation and their changes through time examined in relation to productivity and climate variables.

We provide an integrated analysis of the evolution of feeding and trophic structure in marine mammals and explore the origin and timing of particular feeding strategies over the last 50 million years. In agreement with earlier reports, updated generic counts reveal that the greatest diversity of pinnipedimorphs and cetaceans occurred during the late Miocene (Tortonian), following the Mid-Miocene Climatic Optimum. These historical data are used as a framework to inform past and present structure and trophic interactions and enable predictions about future marine ecosystems. The drivers of diet and feeding patterns are both environmental (sea level fluctuations, climate change) and biotic (anatomical specializations, competition, predator-prey). The influence of these processes on paleodiversity varies depending on taxonomic group, timing, and geographic scale.

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INTRODUCTION

Marine mammals comprise a diverse assemblage of at least seven distinct evolutionary lineages that independently returned to the sea and spend the majority of their time in water. They occupy aquatic ecosystems ranging from the poles to the equator and riverine, estuarine, and a variety of marine environments from shallow coastal to deep sea. Across this diverse range of habitats they occupy almost every trophic level from top predators to primary consumers. In addition to the three major extant lineages that include pinnipeds (seals, sea lions, and walruses), cetaceans, odontocetes [toothed whales], mysticetes [baleen whales] and sirenians (manatees and dugongs), other marine mammals include sea otters and polar bears plus several entirely extinct lineages: desmostylians, aquatic sloths, and an aquatic bear (Berta, 2017).

Today marine mammals include 130 living species (Committee on Taxonomy, 2017). Marine mammals have been on Earth for a little more than 50 million years, and their past diversity includes more than 450 genera (Paleobiology Database, see Materials and Methods). All of their ancestors had well-developed limbs and feet and lived in terrestrial environments. The transition from land to sea involved the acquisition of many diverse adaptations including the ability to capture and consume prey in the marine environment. Environmental pressures involved in foraging in water, such as increased pressure, density, and viscosity, have resulted in morphological specializations related to locomotion, prey capture, and food processing.

The objective of this study is to provide an integrative analysis of the anatomy, evolution, and ecology of feeding in marine mammals. We begin by examining feeding strategies in extant and extinct marine mammals by associating these strategies with anatomical specializations of the head. We next consider genetic and ontogenetic evidence of skull and tooth morphology, which underlies the evolution of feeding types. Finally, using key morphological features, we performed a comprehensive analysis of the evolution of feeding diversity and trophic structure in marine mammals. Also incorporated in this framework is evidence from diatoms diversity, stable isotopes (e.g., δ18O, δ13C), and sea level changes which track changes in productivity, diet, and the environment over geological time.

Feeding Strategies and Prey

Marine mammals have evolved multiple feeding strategies to capture and consume food or prey underwater. We have adopted the three feeding categories of extant aquatic predators identified by Kienle et al. (2017): biting, suction, and filter feeding with the addition of a fourth category, grazing to include sirenians (Figure 1). It should be noted that, like most animals, marine mammals are opportunistic feeders and capture prey as efficiently as possible, which may require them to switch between prey types depending on their availability, and they also employ combinations of these strategies (Hocking et al., 2013, 2017a, b; Kienle et al., 2017). Furthermore, many species show variation in foraging strategy that can be related to age, sex, season, and prey type (Kienle and Berta, 2016).

**Biting.** Biting (raptorial) feeding, a common marine mammal feeding strategy, can be divided into three

![Figure 1](image-url)
subcategories: crushing, grip-and-tear, and pierce feeding (Kienle et al., 2017). Crushing is performed when prey is fragmented by the teeth during ingestion and intraoral processing. This strategy is exemplified by sea otters (Enhydra lutris) using the molar to break down hard shelled prey (e.g., sea urchins, crustaceans, and molluscs). Grip-and-tear feeding defines animals that hold prey with the jaws/forelimbs, shake prey and/or rip off smaller pieces during ingestion. This strategy has also been documented in some pinnipeds (e.g., leopard seals Hydrurga leptonyx), some odontocetes (e.g., killer whales Orcinus orca), polar bears (Ursus maritimus), and sea otters (Adam and Berta, 2002; Kienle and Berta, 2016). In pierce feeding, animals bite prey during ingestion, often swallowing prey whole with little manipulation or external processing. Suction can be used in combination with both pierce and grip-and-tear feeding to pull prey inside the mouth. Pierce and suction feeding are seen in most pinnipeds (e.g., Weddell seal Leptonychotes weddellii, Hawaiian monk seal Neomonachus schauinslandi), and delphinid odontocetes (Stenella, Delphinus, and Tursiops spp.) (Werth, 2006a; Kienle et al., 2015).

**Suction.** Suction feeding is the most common feeding strategy seen in aquatic vertebrates including fish, sea birds, turtles, and some marine mammals (Werth, 2000; Kienle et al., 2017). Suction feeders generate negative pressure within the oral cavity, which draws water and prey into the mouth (e.g., Marshall et al., 2008; Kane and Marshall, 2009). Most marine mammals, including some pinnipeds (e.g., bearded seal Erignathus barbatus, hooded seal Cystophoca cristata, elephant seal Mirounga spp.) and some odontocetes (e.g., beaked and sperm whales, ziphiids and physeteroids, and globicephaline delphinids) use some form of suction to get prey in the mouth (Kienle and Berta, 2016). Beaked and sperm whales are proficient squid feeders (Heyning and Mead, 1996; Werth, 2006a). Bearded and hooded seals employ suction feeding to prey on shrimp, and the diet of elephant seals is primarily composed of squid (Marshall et al., 2008; Churchill and Clementz, 2015; Kienle and Berta, 2016). Suction feeding is also used by highly specialized feeders. The modern walrus, Odobenus rosmarus, employs this strategy to feed on benthic mollusks (Kastelein et al., 1997).

**Filter.** Two main subcategories of filter feeding have been defined behaviorally: continuous and intermittent (Hocking et al., 2017a; Kienle et al., 2017). Continuous filter feeders (or skim feeders) swim slowly through dense patches of prey with open mouths, and prey passively enter the oral cavity (Werth and Potvin, 2016). This strategy is best exemplified by balaenid whales (bowhead and right whales). Intermittent filter feeders actively engulf a single mouthful of prey and remove water using various filtering structures during a distinct water removal stage. Intermittent filter feeding can be further subdivided into lunge and suction filter feeding based on the method of ingestion (Hocking et al., 2017; Kienle et al., 2017). Lunge filter feeding (also called intermittent ram filter feeding, gulp- ing, and ram gulping) is best exemplified by balaenopterid whales (e.g., minke, fin, humpback, blue, and sei whales also known as rorquals) that swim rapidly at a prey patch while opening their mouths to draw in prey (Goldbogen et al., 2010; Werth and Ito, 2017). In suction filter feeding, animals such as some phocid seals (e.g., crabeater seal Lobodon carcinophaga, and leopard seal Hydrurga leptonyx) use suction to pull prey from the water (e.g., Adam and Berta, 2002; Hocking et al., 2013; Kienle and Berta, 2016; Kienle et al., 2017). Lateral suction filter feeding (also called benthic suction feeding) is a specialized strategy seen in the gray whale (Eschrichtius robustus). It involves gray whales rolling laterally onto their right sides and sucking in prey, primarily amphipods, expelling water and sediment through the baleen (El Adli et al., 2014; Kienle et al., 2017).

**Grazing.** Sireniains, the only extant herbivorous marine mammals, graze on marine grasses and algae. Manatees feed on the blades, leaves, and stems of a variety of aquatic and terrestrial plants, including grasses (e.g., water hyacinth and water-thyme Hydilla), sedges, forbs, herbs, and mangroves. Dugongs graze on offshore and intertidal seagrass beds, plowing through mudflats to dislodge anchored vegetation or stripping leaves from taller stems, targeting the rhizomes of sea grasses (Werth, 2000; Domning and Beatty, 2007).

**Anatomy of Feeding**

Feeding strategies involve a number of specializations of the head including both osteological and soft tissue structures. Major osteological features associated with feeding strategies in various extant and fossil marine mammal lineages include size and shape of the head, palate structure, as well as tooth number, shape, and size. Quantitative examination of morphological features in both extant and fossil marine mammals has been limited (e.g., Jones and Goswami, 2010; Jones et al., 2013; Churchill and Clementz, 2015, 2016; Kienle et al., 2017).
Biting: crushing, grip-and-tear, piercing

1. Crushing

- low-crowned, flat molars

*Enhydra lutris*  
Sea otter

2. Grip-and-tear

- long canines

*Hydrurga leptonyx*  
Leopard seal

3. Pierce

- pointed molars

*Otaria byronia*  
Southern sea lion

*Inia geoffrensis*  
Amazon river dolphin

**FIGURE 2.** Anatomical features associated with biting (pierce, grip-and-tear, and crushing) feeding. 1: Crushing in sea otter (*Enhydra lutris* skull, from Lawlor, 1979), 2: Grip-and-tear in leopard seal (*Hydrurga leptonyx* skull, authors’ work), 3: Pierce in Southern sea lion, (*Otaria byronia* skull, authors’ work) and in Amazon river dolphin (*Inia geoffrensis* skull, authors’ work).

and Berta, 2016, 2019). We describe below anatomical features associated with feeding strategies in both extant and fossil marine mammals.

**Biting.** Pierce, grip-and-tear, and crushing biting strategies are associated with slightly different cranial and dental morphologies (Figure 2). Prior workers have described pierce feeders as having elongated orbits, enlarged pterygoid bones, and unequal but limited tooth spacing (Adam and Berta, 2002; Churchill and Clementz, 2015; Kienle and Berta, 2016). Grip-and-tear feeders have long skulls with wide bases and elongated orbits, auditory bullae, palate, and parietals, and additionally, they sometimes have enlarged canines and long, sharp postcanine tooth cusps (Adam and Berta, 2002; Kienle and Berta, 2016). In contrast to most biting marine mammals, polar bears exhibit sharp, shearing carnassials, characteristic of terrestrial carnivorans.

In addition, marine mammals that employ pierce and grip-and-tear biting strategies typically possess numerous slender pointed teeth to retain and grasp prey (Heyning and Mead, 1996; Werth, 2000; Adam and Berta, 2002; Johnston and Berta, 2011). Prey is swallowed whole if small or torn apart. Polydonty, significant increase in tooth number, and homodonty, presence of simple, conical, tooth crowns, are characteristics of modern cetaceans and pinnipeds and differ from the ancestral mammalian dental pattern of heterodonty (Armfield et al., 2013; Churchill and Clementz, 2015). Extreme cases of polydonty and homodonty are documented in oceanic dolphins, which can have more than 130 conical teeth (Jefferson et al., 2015). Long mandibles and toothrows have been associated with raptorial snap feeding characteristic of river dolphins (e.g., *Inia, Pontoporia*), which developed convergently to gavials (Werth, 2006a; McCurry et al., 2017).

The sea otter is the only extant marine mammal that employs a crushing biting strategy. In the past, the durophagous niche was occupied by the phocid *Hadrokirus* (Amson and de Muizon, 2014) and aquatic bear/pinnipedinform *Kolponomos* (Tseng et al., 2016; Paterson et al., 2020). Marine mammals that employ a crushing biting strategy...
Suction
typically possess massive skulls and broad, low crowned flat molars. The canines are often rounded in cross section and the shovel-like incisors are inclined anteriorly to extract soft invertebrate prey from their shells. In comparison to suction feeders, cetaceans that employ biting strategies rely more heavily on lingual shape changes for prey capture, ingestion, and transport (Werth, 2007).

**Suction Feeding.** Many suction feeders have wide skulls, reduced dentition, and large, arched palates (e.g., beaked and sperm whales and the extinct bizarre cetacean *Odobenocetops* (Adam and Berta, 2002; Churchill and Clementz, 2015; Kienle and Berta, 2016; de Muizon and Domning, 2002) (Figure 3). Experiments, simulations, and biomechanical models have shown that a blunt head and a short, wide rostrum facilitates the formation of a small, circular oral opening characteristic of suction feeders (Werth, 2006a, 2006b). An arched or vaulted palate has been associated with suction feeding as it provides the tongue with an extended space in which it can be rapidly depressed and protracted prior to being retracted to produce suction (Kastelein and Gerrits, 1990). Palatal rugosities (ribbing), which function to hold slippery prey such as cephalopods, have also been associated with suction feeding (Heyning and Mead, 1996; Werth, 2000). Another palatal structure – large pterygoid hamuli – has also been associated with suction feeding in the walrus (Adam and Berta, 2002). This enlargement has been correlated with an expansion of the m. velum palatini in the walrus, which is presumably related to reinforcing the soft palate for large intraoral pressure changes during suction feeding (Kastelein et al., 1991). Similarly, the enlarged pterygoid hamuli on the crania of ziphiids have been associated with suction feeding and tentatively correlated with deep diving (Heyning and Mead, 1996; Ramassamy et al., 2018).

Suction feeders are also characterized by reduced dentition and complete tooth loss in some lineages. The walrus and bearded seal have a reduced number of postcanine teeth lacking cusps (Churchill and Clementz, 2015). Sperm whales and most beaked whales exhibit a reduced tooth count associated with their typical teuthophagous habit. For example, most beaked whale species undergo eruptions of only one or two pairs of teeth in males, and these are likely involved in sexual selection and species recognition rather than feeding (Dalebout et al., 2008).

A fused mandibular symphysis seen in the walrus has been identified as a characteristic feature of suction feeders (Adam and Berta, 2002). By providing a solid foundation, it contributes to an overall increase in mandible robustness, which would allow little or no mandibular deformation accompanying tongue retraction during feeding.
Blunt mandibles have been correlated with dental reduction and associated with suction feeding in many odontocete lineages (e.g., some ziphiids, some physeterids, and globicephaline delphinids) (Werth, 2006a, 2006b; Bloodworth and Marshall, 2007).

Soft tissue structures including the tongue, hyoid apparatus, and associated musculature are specialized in various ways given their different uses in feeding. Enlarged hyoid apparatus and musculature, providing more surface area for tongue muscle attachments, are associated with suction feeding (Reidenberg and Laitman, 1994; Heyning and Mead, 1996; Werth, 2006a, 2006b, 2007; Bloodworth and Marshall, 2007; Johnston and Berta, 2011).

**Filter feeding.** Most pinniped suction filter feeders have morphological adaptations for this strategy that include a long narrow skull, a narrow palate, an elongated toothrow, thick mandibular symphysis, and proportionally large postcanine teeth (Adam and Berta, 2002; Jones et al., 2013; Churchill and Clementz, 2015; Kienle and Berta, 2016, 2019). The crabeater seal uses its highly specialized, multi-cusped teeth as a sieve to filter krill from the water, and prey become trapped by the interdigitating teeth (Adam and Berta, 2002) (Figure 4).

Large heads and mouths have been associated with lunge filter feeding in balaenopterids (e.g., Goldbogen et al., 2010). All modern baleen whales lack teeth and employ baleen, a series of keratin laminae arranged transversely in two racks that erupt from the gums on either side of the upper jaw, as a sieve for filter feeding (Figure 4). The number, length, and texture of baleen plates of modern mysticetes vary according to feeding strategy and diet (e.g., Williamson, 1973; Werth, 2000; Young et al., 2015; Jensen et al., 2017). The high rostral arch of balaenids accommodates their exceptionally long, fine baleen laminae (250 to 350 laminae, some exceeding 3 m in length) that are used to filter tiny copepod prey in skin feeding. The baleen of balaenopterids that feed mostly on fish and krill range from 181 to 402 laminae for engulfment feeding. In contrast to balaenids and balaenopterids, gray whales possess fewer baleen laminae (130-180) that are shorter and coarser, for suction feeding on benthic dwelling amphipods and mysid shrimp.

Several features of the lower jaw facilitate expansion of the oral cavity during bulk filter feeding. The unfused, ligamentous lower jaw symphysis exhibited by modern baleen whales permits independent rotation of the jaws (Lambertsen et al., 2005). The mandibular symphysis of rorquals possesses a gel-like cavity hypothesized as a sensory organ that coordinates lunge feeding (Pyenson et al., 2012). The mandibles of rorquals also exhibit a prominent, laterally deflected coronoid process, which serves as an attachment point for the m. temporalis, an important muscle in lateral rotation of the jaw during engulfment feeding (Brodie, 1993; Lambertsen, 1983). Several other morphological features of the mandible are associated with engulfment feeding in balaenopterids. Laterally bowed mandibles contribute to the expansive gape of balaenopterids (Lambertsen et al., 1995). A fibrocartilaginous temporomandibular joint (TMJ) was found to facilitate alpha (longitudinal), omega (transverse), and delta (vertical) rotation of the mandible (Lambertsen et al., 1995). By contrast, balaenids possess a synovial joint typical of mam-

**FIGURE 4.** Anatomical features associated with filter feeding in crabeater seal (*Lobodon carcinophaga* skull, public domain image), and gray whale (*Eschrichtius robustus* skull, authors’ work).
mals. The TMJ of the gray whale represents an intermediate condition, modified from the balaenopterid condition, with a vestigial joint cavity lacking a fibrous capsule, synovial membrane, and articular disk. In gray whales, this results in an increased importance of alpha rotation of the mandible while limiting the degree of delta rotation (angle of gape) of the lower jaw, an important factor in suction feeding (Johnston et al., 2010; El Adli et al., 2014).

The flaccid and floppy tongue of lunge filter feeding rorquals permits ingested prey and water to extend the throat pouch. This sac-like structure holds the water engulfed during a feeding event (Lambertsen, 1983; Goldbogen et al., 2011; Werth and Ito, 2017). By contrast, the muscular free-moving tongue of the gray whale (Kienle et al., 2015) likely plays an important role in generating the suction forces used in the unique lateral suction feeding employed by this species. Bulk lunge filter feeding rorquals expand the gular cavity by numerous longitudinal throat grooves (70-80) on the ventral throat pouch (cavum ventrale), extending to the umbilicus in some species. Ventral groove blubber consisting of tough ridges separated by deep channels of elastic tissue permits the extraordinary extensibility of the engulfment apparatus (Goldbogen et al., 2010; Pyenson et al., 2012). In the mouths of rorquals, a tendinous part of the m. temporalis, the frontomandibular stay, provides a continuous mechanical linkage between the skull and serves to optimize the gape of the mouth during engulfment filter feeding (Lambertsen et al., 1995). Suction feeding marine mammals typically exhibit smaller throat grooves that aid in tongue retraction. The gray whale possesses two to three throat grooves confined to the gular region (Berta et al., 2015), and both sperm and beaked whales also possess gular throat grooves (Heyning and Mead, 1996).

Grazing. Grazing marine mammals like sirenians possess cranial adaptations allowing them to feed at various depths in the water column. The degree of rostral deflection is correlated with degree of specialization for bottom feeding (Figure 5). Dugongs exhibit extreme rostral deflection (70 degrees) associated with their bottom feeding habit, consuming mostly sea grasses. Manatees exhibit less rostral deflection (26-30 degrees) and feed on aquatic plants throughout the water column (Domning, 2001; Vélez-Juarbe et al., 2012). Similar to this last group, stem sirenians (e.g., pro-rastomids) possessed undeflected rostra, indicating that they likely fed on aquatic plants. Enlarged blade-like, self-sharpening tusks developed in some later diverging dugongines (e.g., Bharatisiren, Rytiodus, Corystosiren, Domniniga, Kutchisiren, Xenosiren, and Diplototherium) were likely employed for digging up rhizomes of larger sea grasses (Domning and Beatty, 2007).

The cheek teeth of most grazing marine mammals are typically bilophodont, arranged in two long ridges, with rounded cusps and thick enamel. The teeth of extinct desmostylians exhibit a distinctive morphology consisting of closely bundled columnar cusps. Both extinct desmostylians and the aquatic sloth Thalassococcus possessed anteriorly extended, wide premaxillae and wide mandibular symphyses, indicating the development of strong lips that presumably facilitated grasping marine vegetation (de Muizon et al., 2004). Although sirenians lack expanded premaxillae, their wide upper lips and horny pads on the maxillae function similarly. In addition to grazing, the vaulted palate of desmostylians has been suggested to enable suction feeding (Beatty, 2004; Uno et al., 2008). The elongate jaws and expanded facial region of some later diverging desmostylians (e.g., Ounalashkastylus) likely facilitated extensive development of hyoid, facial, and adductor musculature, which supports this feeding method (Chiba et al., 2016).

Ontogeny and Growth of the Skull and Teeth

Skull growth. The feeding apparatus of marine mammals is driven by developmental changes in the morphology and growth of the skull and teeth. Skull growth in pinnipeds revealed that ontogenetic shape changes in otariids are greater, due to dimorphism, than in phocids although sample sizes were limited (Jones and Goswami, 2010). Brunner et al. (2004) found that the rostral and palatal regions in otariids required significantly longer periods to complete growth than the neurocranium (Brunner et al., 2004; Tarnawski et al., 2013). The primary importance of the postcanine teeth in feeding is reflected by their more rapid development after weaning compared to growth of the canines in otariids (Brunner et al., 2004). Skull ontogenetic studies in pinnipeds suggest that life history variables (e.g., gestation time, weaning time, growth rate) are more influential than environmental variables (e.g., latitude, temperature, productivity, and seasonality) in phocid cranial evolution (Jones and Goswami, 2010; Jones et al., 2013; Randau et al., 2019). Future studies are needed, comparing feeding components of the skull (e.g., tooth size and shape, postcanine tooth length) across larger sam-
Grazing

**Trichechus senegalensis**
African manatee

rostral deflection

**Paleoparadoxia sp.**
Desmostylia

rounded, columnar molars

**Thalassocnus sp.**
Aquatic sloth

bilocodont teeth

FIGURE 5. Anatomical features associated with grazing in African manatee (*Trichechus senegalensis* skull, from Werth, 2000), Desmostylia (*Paleoparadoxia* skull, public domain), and aquatic sloth (*Thalassocnus* sp. skull, modified from de Muizon et al., 2004).

Little is known of the ontogeny of cetacean cranial morphology during fetal growth (Klima, 1995; Armfield et al., 2011; Lanzetti et al., 2020). Investigations of skull growth changes suggest that, compared to odontocetes, mysticetes begin developing larger mouths in the early fetal stages, presumably related to their filter feeding strategy (Armfield et al., 2011; Lanzetti et al., 2020). However, while the rostrum approximates the adult shape and curvature only at late stages of gestation, the mandible seems to reach its definitive proportions and curvature earlier (Lanzetti et al., 2020).

**Tooth development.** The developmental controls of tooth differentiation and tooth number have been little studied in marine mammals. Four conserved signaling pathways, including the bone morphogenetic proteins (Bmp), fibroblast growth factors (Fgf), sonic hedgehog (Shh), and wingless-related (Wnt) pathways, play important roles in tooth development in mammals. Recent studies show that these signaling pathways interact through positive and negative feedback loops to regulate not only morphogenesis of individual teeth but also tooth number, shape, and spatial pattern (Bei, 2009; Jernvall and Thesleff, 2012; Lan et al., 2014).

Armfield et al. (2013) investigated the genetic pathways that cause characteristic homodont, pointed tooth shape in dolphins. They discovered that, in the early developmental stages, a caudal extension of the expression of a specific growth factor (Bmp4), normally expressed in other mammals only in the anterior portion of the jaw, is responsible for the observed adult tooth shape. Modern mysticetes instead lack teeth postnatally, but they still develop tooth germs that are resorbed before birth (e.g., Ridewood, 1923; Karlsen, 1962; Fudge et al., 2009; Thewissen et al., 2017). Several molecular studies revealed that only some of the putative enamel and tooth genes are inactivated or deleted in mysticetes (Deméré et al., 2008; Meredith et al., 2009, 2011, 2013; Springer et al., 2016). Thewissen et al. (2017) found evidence for the presence of Fgf4, known to be implicated in tooth growth, during baleen growth. This
suggests a partially shared genetic pathway for both teeth and baleen formation.

Among marine mammals and most other mammals, tooth development in modern manatees is unique. The cheek teeth are replaced horizontally; the new tooth erupts from the rear of the tooth row, pushing the anterior tooth out of the tooth row (Domning and Hayek, 1984; Beatty et al., 2012). Tooth replacement is similar in elephants although the latter have a limited number of teeth (Domning and Hayek, 1984). Little is known about the molecular mechanisms behind this mode of tooth replacement. It is possible that in the future a mouse model could be used to study this process, as the formation of supernumerary molars might employ a regulatory pathway similar to the one responsible for the ever-growing incisors of rodents (Jernvall and Thesleff, 2012). A better understanding of tooth development in various marine mammal lineages will provide important data that can then be correlated with major shifts in diet and food processing.

MATERIALS AND METHODS

Data Compilation

Diversity through time. Raw taxon counts for 61 extant and 374 fossil marine mammal genera recorded at the level of geological stage were downloaded from the Paleobiology Database (PBDB, www.paleobiodb.org). The search was performed separately for the following groups: “Archaeoceti, Cetacea”, “Pinnipedimorpha”, “Sirenia”, “Desmostylia”, “Thalassocnus”, “Ursus maritimus”, “Ursus arctos”, “Aonychini, Aonyxini”, “Enhydrini, Enhydriodontini”, “Lontra, Lutra, Pteronura, Siamogale” (sea otters). Occurrences for each group were downloaded for each time bin (geological stage), starting from the Ypresian (early Eocene), using the “major (default)” time rule. Only occurrences with certain generic assignment were considered (“genus certain”). The data include all occurrences entered up to 12/31/2017. Since geological stages in the Pleistocene and Holocene result in time bins that are of much shorter duration than any of the other stages, creating few occurrences in each of these time intervals, we followed Peredo and Uhen (2016) in binning the Pleistocene stages along with the Holocene series in the Quaternary, which at 2.6 Ma, is comparable in length to earlier time bins. Accessory data were also downloaded for these records, such as depositional environment and published sources, to allow verification of each occurrence individually and to be able to use the published references as a starting point to assign genera to the different categories. Diversity plots were obtained by counting the valid genera for each time bin in each group or family after the categorization was completed.

Feeding strategies and ecological adaptations through time. To interpret patterns in the evolution of feeding adaptations through time in marine mammals, we categorized each taxon using four parameters: prey capture, tooth and cusp shape, diet, and habitat. We first defined the categories for each parameter in extant genera and then applied them to the fossils. It is possible that fossil taxa presented completely different adaptations not present in any modern genera. By using broad categories, we aim to characterize trends and possible climatic drivers in the evolutionary history of these lineages that would not be obliterated by the incorrect categorization of single taxa. Categories were assigned to valid taxa only. The categories assigned to each taxon, along with its taxonomic placement, justification for the exclusion of invalid taxa, and all literature sources used are reported in the Supplementary Information that can be found in the Dryad repository (https://doi.org/10.6086/d14671) subdivided by group (Cetacea, Pinnipedimorpha, Sirenia, Desmostylia, and Thalassocoanus, Other Carnivores).

We started by assigning categories to extant marine mammals to then use this information to better categorize extinct taxa. We divided marine mammals into four prey capture categories: biting/crushing, filter, suction, and grazing, as described above. Prey capture types for extant taxa were compiled from primary and secondary sources (e.g., Adam and Berta, 2002). In an effort to further categorize prey selection accurately in the fossils, we examined tooth pattern and cusp shape (tooth presence/absence, homodonty, heterodonty; pointed, rounded, or bilophodont cusps). Type of dentition and cusp shape in extant taxa was determined based on photographs or personal observations of the authors. Diet was characterized using the five distinct dietary categories of Pauly et al. (1998): benthic invertebrates, fish, squid/soft-bodied pelagic invertebrates, zooplankton, and tetrapods. We included an additional dietary category for multicellular primary producers (sea grasses and algae) to characterize herbivorous taxa. Diet of carnivorous living species was categorized based on data from Pauly et al. (1998) and additional literature sources for taxa that were not included in this study. If two categories were indicated by Pauly et al. (1998) for at least one species, both were con-
sidered to categorize the genus if neither made up more than 80% of the diet, but only one was chosen if it accounted for more than 80% of the diet alone. If three or more categories were indicated by Pauly et al. (1998) in a least one species, the first two or three prey items that made up at least 80% of the diet were used. Finally, we considered the environment(s) occupied by marine mammals. Six habitat categories were based on environments used by extant genera: marine pelagic, marine coastal, marine benthic, riverine, estuarine, and lacustrine, and were compiled by relevant secondary literature (e.g., Jefferson et al., 2015).

Based on prey capture categories, tooth shape, diet, and habitat data for extant marine mammals, we inferred relevant groupings for fossil taxa. When any hypotheses about feeding and paleoenvironment for a taxon were reported in the literature, these were used to determine the category assignment. When these data were not available, we based our assessment on a variety of factors. Soft tissue feeding structures (i.e., lips, tongue, and hyoid muscles) typically do not fossilize. However, it is possible to infer feeding strategy in fossil marine mammals from the presence of osteological correlates. For example, the presence of lateral palatal nutrient sulci and foramina in some toothed mysticetes (Aetiocetidae) has been interpreted as bony correlates for the presence of baleen (Deméré and Berta, 2008; but see Peredo et al., 2017). These structures have also been observed in fossil stem baleen whales, although they still retained a full dentition (Deméré and Berta, 2008; but see Peredo et al., 2017). These structures have also been observed in fossil stem baleen whales, although they still retained a full dentition (Deméré and Berta, 2008; but see Peredo et al., 2017). These structures have also been observed in fossil stem baleen whales, although they still retained a full dentition (Deméré and Berta, 2008; but see Peredo et al., 2017). Observations on in situ fossilized stomach contents of associated skeletons and bite marks on bones provide a direct and accurate method for predicting diet (e.g., Boessenecker and Perry, 2011; Collareta et al., 2015). Another method is to catalog potential prey that occur in the same geologic horizon and locality as the predator (e.g., Lambert et al., 2015) although care must be taken to ensure that the carcasses of predator or prey have not been transported prior to fossilization (Adam and Berta, 2002; Lambert et al., 2015). In this case, further confirmation came from fish remains that were actually found in the whale. Diet and habitat can also be assessed using isotope analysis of fossil tooth enamel (e.g., Clementz et al., 2006, 2014). Whereas analyses of tooth microwear (Thewissen et al., 2011; Fahlke et al., 2013) provide clues about prey types that have been consumed, enamel microstructure (Loch et al., 2016) reveal information about tooth behavior (i.e., resistance). By far the most commonly used method for inferring the diet of fossil taxa employs functional morphology and analogy with living forms. By examining the morphology associated with a particular method of capturing and handling prey in living species, both in terms of skull morphology and tooth and cusps shape, inferences of feeding behavior and diet can often be drawn for fossil taxa with similar morphologies (e.g., Kophonomos and sea otters – Tseng et al., 2016). Similarly, habitat preference in fossil was assessed based on modern taxa displaying similar adaptations in skull, limb, and tooth morphology. Depositional environment of the fossil was considered but not used exclusively to determine the habitat of the taxon, as sedimentation, sedimentary outcrop availability, and preservation bias make this parameter unreliable (Uhen and Pyenson, 2007; Marx, 2009).

Statistical Analyses for Environmental Drivers

Using this dataset, we performed statistical analyses to explore the correlations and changes through time of patterns of generic diversity and category distribution. We also examined the correlations between this dataset and productivity and climatic factors that have been used to explain diversity patterns in previous work. Specifically, we used δ18O values that reflect both temperature and global ice volume and δ13C values that serve as a proxy for changes in the global carbon cycle (Zachos et al., 2001). Values of δ18O and δ13C were taken from Prokoph et al. (2008). In particular, data originally published by Zachos et al. (2001) from benthic foraminifera and deep sea locations were used, as they represented the most complete record for the time interval considered. The values are reported using absolute age; we selected the ones belonging to each time bin and averaged them to get a single value representing that geological stage and creating a comparable dataset to the taxa occurrences. In addition, previous studies suggested that an increase in diatom species diversity, which suggests an increase in primary productivity, has driven cetacean diversity (Marx and Uhen, 2010), and therefore we also included this measure in the analyses. The diatom species data were downloaded from the Neptune Database on 03/05/2018 (NSB, www.nsb-mfn-berlin.de). The species present in each time bin were
counted to determine the diversity in that interval. We also considered how changes in sea level might have affected diet and diversity of marine mammals. Estimates of sea level were taken from Miller et al. (2005) and the mean value for each stage was calculated as it was done for the isotope values. Correlation between diversity and category distribution with climatic variables for each group was tested using linear regression analysis. Best model including one to four climatic variables was chosen based on Akaike’s information criterion (AIC) score if a significant correlation was found (p < 0.05). Correlation between Sirenia and Desmostylia diversity and climate variables was tested using Pearson’s correlation test due to their low sample size. All statistical analyses were conducted in XLSTATS (2017). Raw data used for the analyses can be found in the Supplementary Information in the Dryad repository (https://doi.org/10.6086/d14671).

Ancestral State Reconstructions

To expand on the evolutionary significance of these data, pinnipediniform and cetacean phylogenetic trees were assembled using previously published phylogenies and ancestral state reconstructions (ASR) were performed by coding the taxa according to the categories described. The topologies are assembled for the most recent work on each group in order to include the highest number of confidently assigned taxa in the analyses and represent all major groups. The phylogenetic position of some taxa is currently debated in the literature, and different phylogenetic placement might alter the results of the ASR analyses. However, it is beyond the scope of this paper to weigh the literature, and different phylogenetic placement signals will require integrated comparative studies of potential biases of the record (Kelley and Pyenson, 2015).

**Pinnipediniforma.** Pinnipediniforma is a monophyletic group within Carnivora (i.e., dogs, felids, bears, otters, etc.) that originated 30.6-23 Ma (Berta et al., 2018). Pinnipediniforma includes several stem taxa (e.g., *Enaliarctos, Pteronarctos, Pacificotaria,* and *Pinnaarticidion*) in addition to crown pinnipeds composed of the extinct Desmatophocidae and three extant clades: Otariidae (fur seals and sea lions), Odobenidae (walruses), and Phocidae (seals). With the exception of phocids, the pattern of pinnipediniform diversity documented by Boessenecker and Churchill (2018) for the North Pacific is in agreement with our results. Early diverging stem-Pinnipediniforma, also known as 'enaliarctines’, represent the earliest diversification of pinnipeds, originating in the late Chattian, peaking in the Burdigalian, and becoming extinct by the end of the Serravallian (Figure 7). Desmatophocidae appear during the Aquitanian and gradually increase in diversity during the Burdigalian before declining in diversity and apparently becoming extinct in the Messinian (Figure 7). Otariids

**RESULTS AND DISCUSSION**

Diversity through Time

Updated generic plots of marine mammal palaeodiversity reveal that cetaceans and pinnipediniforms achieved greatest diversity during the late Miocene (Tortonian) (Figure 6). Generally, our results confirm earlier reports of diversity changes through time (e.g., Marx and Uhen, 2010; Peredo and Uhen, 2016; Boessenecker and Churchill, 2018). The herbivorous taxa (Desmostylia and Sirenia) appear to maintain a low but constant diversity throughout their distribution, while Cetacea and Pinnipedinorma present wider fluctuations. We confirm previous observations (Peredo and Uhen, 2016) of large drops in marine mammal diversity at the end of the Oligocene, Miocene, and Pliocene (Figure 6). The late Oligocene diversity drop has been explained as the result of poor sampling (Marx and Fordyce, 2015) but a rapidly improving record from this time interval may require revision of this observation (e.g., Fordyce and Marx, 2018; Peredo and Pyenson, 2018; Tsai and Fordyce, 2018). The late Miocene drop in diversity was influenced by the Messinian Salinity Crisis, a geologic event during which the Mediterranean Sea went into a cycle of desiccation resulting in deep, dry basins (Krijgsman et al., 1999). Phocids, cetaceans, and sirenians in the Mediterranean were particularly affected beginning about 6 Ma (Bianucci, 2013). The Pliocene drop in marine mammal diversity has been associated with the onset of Northern Hemisphere glaciation (Marx and Fordyce, 2015) and loss of habitat by sea level fluctuations (Pimiento et al., 2017). Whether these fluctuations in diversity represent real biological signals will require integrated comparative studies of potential biases of the record (Kelley and Pyenson, 2015).
appear in the Burdigalian (but lack a Serravallian record); they diversify in the Messinian and Pliocene and mostly maintain that diversity until the Holocene (Figure 7). Walruses (Odobenidae) were somewhat diverse in the Burdigalian and remained at low diversity through to the Serravallian, gradually increasing until their peak in the Messinian, with their diversity decreasing sharply in the Pliocene and Pleistocene (Figure 7). Phocids appear in the Burdigalian and quickly diversify, reaching peak diversity during the Tortonian, and continue to be the dominant living pinniped group, even after a reduction in diversity in the Pliocene (Figure 7).

Several ecological explanations have been given for these diversity changes. Vélez-Juarbe (2017) suggested possible competitive displacement of desmatophocids by walruses and Boessenecker and Churchill (2018) noted that the extinction of desmatophocids appears to be followed by an increasing diversity of otariids, perhaps as the result of niche filling. The suggestion by these latter authors that several Tortonian odobenids (e.g., *Imagotaria*, *Pontolis*, and *Gomphotaria*) lacking benthic feeding specializations may have exploited similar prey (i.e., fish) by employing a generalized biting strategy like desmatophocids is in agreement with results of our study.

The effects of climate and environmental changes on pinnipedinorph diversity allow the following observations. The peak in phocid diversity and increase in odobenids and otariids during the Tortonian (11 Ma) is probably a result of the mid Miocene Climatic Optimum (MMCO) that occurred in the Langhian (~15 Ma) and is followed by decreases in diversity along with colder temperatures (Figure 7). Previous work suggested that sea level changes could affect pinniped haul-out areas, which provide breeding areas. A rise in sea level may have been involved in the Mio-Pliocene extinction of phocids in South America, resulting in the decrease of sandy beach haul-out areas favored by phocids. Beginning in the Plio-Pleistocene and continuing at present, rocky shore habitats predominated in South America and elsewhere in the Southern Hemisphere (e.g., Africa, Oceania), areas exclusively occupied by otariids (Valenzuela-Toro et al., 2013, 2016). This hypothesis of ecological replacement of phocids by otariids is difficult to confirm at the present time since marine mammal fossils in South America are relatively poorly known and not well sampled during this time. Examination of habitat preferences in several extant otariids and phocids off the coast of Baja California, Mexico, revealed overlap in habitat although phocids (harbor and elephant seals) preferred sandy beaches and cobblestone beaches, which may be associated with their postcranial morphology (i.e., inability of rotating hind limbs forward) and terrestrial locomotion (Arias-del Raza and Heckel, 2016).

*Cetacea*. Cetacea are a monophyletic group nested within Artiodactyla (i.e., pigs, deer, camels, hippos, etc.). Stem cetaceans, or archaeocetes, the paraphyletic group ancestral to all other cetaceans, evolved during the Eocene (Ypresian) approximately 53.5 Ma. Stem cetaceans evolved in
fresh water and later in the Eocene transitioned to marine environments, which is supported by dental stable isotope values (Roe et al., 1998; Clementz et al., 2006). Crown Cetacea (or Neoceti), defined by the common ancestor of all living mysticete and odontocete cetaceans, originated approximately 36.4 Ma at the end of the Eocene (Gatesy et al., 2013; Pyenson, 2017). Absolute divergence dates calculated using molecular data are mostly in line with evidence from the fossil record (Figure 8).

The origin and diversification of crown cetaceans near the Eocene-Oligocene boundary has long been explained as the result of establishment of the Antarctic Circumpolar Current and large-scale changes that accompanied it. Global cooling fed Antarctic ice expansion and Southern Ocean upwelling that resulted in an increase in productivity (e.g., Fordyce, 1980; Steeman et al., 2009). The role of this process in the evolution of Cetacea has been debated given that the origin of crown cetaceans in the late Eocene, recovered both in the fossil record and in phylogenetic analysis using molecular data, actually precedes the estimated onset of the current, which likely occurred right at the epoch boundary at 33.5 Ma (Hassold et al., 2009). As previously noted, testable models that explain the ecological context of cetacean evolution are lacking (Pyenson, 2017).

Late Miocene diversity of mysticetes (Figure 8) appears to be driven dominantly by an increase in diversity of extinct Cetotheriidae (Figure 9 – Marx and Uhen, 2010; Marx and Fordyce, 2015; Slater et al., 2017). Balaenopteridae and Balaenidae instead reach their peak diversity in the Zanclean before sharply declining (Figure 9). Odontoceti also increase in diversity in the late Miocene (Figure 8). This distribution is driven by a simultaneous peak in diversity of four of the five extant groups of toothed whales (Figure 10). The four living or recently extinct river dolphin taxa, Platanistidae (Asian river dolphins), Pontoporiidae (La Plata river/coastal dolphins), Lipotidae (Yangtze river dolphin), and Iniidae (Amazon river dolphins), belong to two distinct groups that appear to have transitioned from marine to freshwater independently at least twice, in the Aquitanian (Platanistidae) and Serravallian (Iniidae/Pontoporiidae/Lipotidae) (Figure 10 – Cassens et al., 2000; Geisler et al., 2011). The explosive radiation of delphinids during the last 3.5 Ma (Figure 10) has been explained as a response to global climate cooling and the beginning of the Northern Hemisphere glaciation and related changes in sea level (e.g., Whit-
FIGURE 8. Cetacean stem and Neoceti taxa diversity through time. “Archaeoceti” includes all stem taxa that are not included in the two extant groups. Dashed vertical lines: black, epoch boundaries; gray, age boundaries.

FIGURE 9. Mysticeti stem and familial level diversity through time. “Cetotheriidae s.l.” is a paraphyletic group including all taxa that belong to crown Mysticeti but are not grouped with any of the four living families. Dashed vertical lines: black, epoch boundaries; gray, age boundaries.
more, 1994; Fordyce and de Muizon, 2001; McGowen et al., 2009; Steeman et al., 2009; Amaral et al., 2018). This increased diversity of the late Pliocene Mediterranean marine mammal assemblages (i.e., cetaceans, pinnipeds, and sireni ans) was hypothesized as possibly due to rapid colonization of this region after the Messinian salinity crisis (Bianucci, 2005). More specifically, the diversification of delphinine dolphins (e.g., *Stenella* spp., *Delphinus* spp., *Lagenorynchus* spp., and *Tursiops* spp.) in the tropical Atlantic and Indian Oceans has been related to pulses of diatom productivity during the late Pliocene and early Pleistocene (Mannocci et al., 2015).

**Sirenia and Desmostylia.** Sireni ans are members of Afrotheria (i.e., elephants, hyraxes, tenrecs, etc.) and include several stem taxa (i.e., Prorastomidae and Protosirenidae) and one crown clade encompassing Trichechidae (manatees) and Dugongidae (dugongs). The stem sireni ans, known from the early to middle Eocene (late Ypresian to early Lutetian) of Africa and Central America (Benoit et al., 2013), reached their peak in diversity during the Lutetian (Figure 11). Appearing at about this same time were dugongids, the most successful and diverse sireni ans, that typically inhabited warm, nearshore waters, with a worldwide distribution. A cold-water adapted lineage, first known in the middle Oligocene in the eastern Pacific, culminated in Steller’s sea cow (*Hydrodamalis gigas*) and evolved large body size as an adaptation to cool climates beginning in the Miocene (Pyenson and Vermeij, 2016). Dugongid diversity dramatically decreased in the late Pliocene and at present they are represented by a single species found in the Indo-Pacific region (Figure 11). The evolution of manatees (Trichechidae) began in South America during the Miocene and is tied to Caribbean paleobiogeography, notably uplift of the Andes, which created erosion and runoff into river systems that

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**FIGURE 10.** Odontoceti stem and familial level diversity through time. The group "Delphinoidea" includes Delphinidae, Monodontidae, and Phocoenidae, as well as extinct members of this crown group of uncertain phylogenetic placement. The "river dolphins" closely related to these taxa (Inioidea and Lipotidae) are plotted separately to emphasize their diversity in the fossil record compared to the present. "Ziphioidea" includes Ziphiidae and closely related stem taxa as identified by Bianucci et al. (2016). Dashed vertical lines: black, epoch boundaries; gray, age boundaries.
resulted in an abundance of abrasive grasses (Domning, 1982). Trichechids adapted to this new food resource by evolving thickened tooth enamel and smaller, more numerous teeth that were continuously replaced. The particular set of adaptations that characterize this lineage might have contributed to their low but stable diversity (Figure 11).

Sirenians appear to have replaced desmostylians during the late Miocene (Tortonian) as the result of competition for the same food resource: sea grasses (Vélez-Juarbe et al., 2012). Desmostylians are a completely extinct, enigmatic group of herbivorous aquatic mammals, exclusively distributed in the North Pacific, likely related to either Afrotheria, Perissodactyla, or Paenungulatomorpha (Matsui and Tsuihiji, 2019). Evidence for their replacement by sirenians is provided by the decrease in diversity of desmostylians after the Tortonian followed by their disappearance in the Messinian, which coincides with the appearance of Hydrodamalis and other dugongids in the same localities and habitats they occupied (Figure 12). Additional data on the paleoecology of these lineages and climatic changes of the region need to be conducted in order to test this hypothesis.

**Other marine mammals.** In the most recent combined phylogeny of otters, the extant sea otter *Enhydra lutris* and its sister taxon *Enhydriodon* spp. are embedded within the river otter (Lutrinae) clade (Wang et al., 2018). Although this same phylogenetic analysis placed *Enhydridtherium* with other lutrine otters and distant from the sea otter we included it in our study since recent fossil evidence suggested that *Enhydridtherium* occupied freshwater environments in addition to estuarine and coastal environments (Tseng et al., 2017). The oldest record of *Enhydra* sp. from the Pacific is Middle Pleistocene in age although an earlier representative of this lineage (*Enhydra reevei*) arose in the North Atlantic during the early Pleistocene about 2.2-1.7 Ma and presumably dispersed into the Pacific though the Bering Strait (Boessenecker, 2016). The extinct sea mink *Neovison macrodon* known from the early to Late Pleistocene in marine and estuarine habitats along the coastal islands of the Gulf of Maine was hunted to extinction, a victim of the fur trade in the late 1800s (Mead et al., 2000).

The polar bear *Ursus maritimus* diverged from brown bears probably less than 500,000 years ago (Liu et al., 2014). Polar bears have a fossil record limited to the Late Pleistocene. The oldest remains of a polar bear are dated at 110-13 ka (Ingólfssson and Wiig, 2008). Stable isotope analyses of fossil polar bear teeth indicate that by at least 110 ka they had acquired a feeding strategy quite different than that of brown bears, having adapted to feeding on fish and marine mammals (Lindquist et al., 2010).

Aquatic sloths (*Thalassocnus* spp.) evolved in the Southeastern Pacific during the Miocene (e.g., de Muizon et al., 2004; Amson et al., 2015). There is some evidence to suggest that they occupied the

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**FIGURE 11.** Sirenia stem and familial level diversity through time. “Protosirenidae / Prorastomidae” includes all taxa that are not included in the two extant groups. Dashed vertical lines: black, epoch boundaries; gray, age boundaries.
FIGURE 12. Hypothesized ecological replacement of desmostylians (circles) by sirenians (squares) in the North Pacific Ocean. On the top, map of the localities of Tortonian age (11 Ma) where desmostylian and sirenian were recorded. On the bottom, map of the localities of Zanclean age (5 Ma) where sirenians were recorded. Desmostylia disappear from the fossil record by the end of the Tortonian (7.2 Ma) when sirenians, particularly *Hydrodamalis* spp., start colonizing this region, likely feeding on the same resources. Only one occurrence per genus is reported in each locality. Locality data for each occurrence of Desmostylia and Sirenia in the Tortonian and Zanclean were downloaded from PBDB (https://paleobiodb.org) using the search parameters described in Materials and Methods and then plotted on the map.
The similarity of these marine sloths to extinct desmostylians in the North Pacific in terms of diet raises the possibility that they may be the ecological homologues of desmostylians in the South Pacific, a hypothesis that requires more rigorous testing.

Biotic and Environmental Drivers of Feeding Diversification

**Pinnipedimorphs.** Pinnipedimorph diversity was most strongly associated with all four climatic variables considered – δ¹⁸O, δ¹³C, diatom species diversity, and sea level changes (Table 1, Appendix 2). The link between specific climatic indicators and pinnipedimorph diversity has not been previously indicated, although research on modern ecosystems has associated these variables (Hirons et al., 2001; Fréon et al., 2009). In particular, when the three modern families of Pinnipedimorpha are examined separately, Otariidae (fur seals and sea lions) appear to be driving the correlation, with the diversity of this group being strongly associated with all four climatic variables (Table 1; Figure 13). This might be the result of habitat and ecological differences. For example, extant otariids are confined to regions of high upwelling and cool water temperature usually less than 24°C (Deméré et al., 2003), which confirms the importance of climate and temperature on their distribution. By contrast, pinnipeds are found in a wider variety of environments including tropical, oligotrophic waters (Chur- chill et al., 2014). Odobenidae (walrus) diversity was also found to be correlated with three variables: δ¹⁸O, δ¹³C, and sea level changes. This fits with a Messinian peak diversity of fossil walruses and the evolution of more specialized feeding systems (i.e., tusks and molluscivory) (Magallanes et al., 2018). The drop in walrus diversity beginning at the end of the Pliocene is associated with the onset of Northern Hemisphere glaciation, large-scale sea level oscillations, and other oceanographic alterations (e.g., changes in productivity and ocean circulation). By the end of the Pleistocene, the loss of shallow foraging areas utilized by molluscivorous odobenines was an important driver of extinction for walruses and other megafauna (Pimiento et al., 2017).

We also examined whether habitat selection and prey type changes in this group were independently associated with climate and primary productivity variables. The distribution through time of the three predominant marine habitats for pinnipedimorphs (pelagic, coastal, and benthic) as well as the three main prey types they feed on (fish, squid, and benthic invertebrates) were each correlated with at least two of the climatic variables (Table 2, Table 3). This confirms the role of climate change and primary productivity on habitat and prey choice in pinnipedimorph evolution and suggests that the diversification of this group should be considered in the context of global environmental changes.

**Cetaceans.** Diversity of cetaceans as a whole, and both odontocetes and mysticetes separately, is correlated with diatom species diversity and δ¹⁸O, a proxy for global ocean temperature (Table 4; Figure 14). These results for Cetacea confirm the findings of some previous studies that linked cetacean and mysticete evolution to climate change (Marx and Uhen, 2010; Marx and Fordyce, 2015). Recent fossil discoveries show that the timing of divergence of both mysticetes and odontocetes is earlier than previously reported, thus raising questions on the role of the establishment of the Antarctic Circumpolar Current in the evolution of these two groups (Pyenson, 2017). For Cetacea, we analyzed the possible correlation between habitat and diet changes through time, as well as prey capture strategy given the high variability in this group (Table 5, Table 6, Table 7). For the habitats (Table 5), riverine taxa diversity appears to be correlated

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**TABLE 1.** Linear regression analyses for Pinnipedimorpha and extant families’ diversity through time against climate and productivity variables: δ¹⁸O (variable A), δ¹³C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Diversity of each group is considered from the time bin of first occurrence ( Chattian for Pinnipedimorpha, Burdigalian for extant families). Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each group are reported, for complete results see Appendix 2, Table S1.

|                | Pinnipedimorpha | Phocidae | Otariidae | Odobenidae |
|----------------|-----------------|----------|-----------|------------|
| R²             | 0.909           | 0.775    | 0.993     | 0.906      |
| AIC            | 29.199          | 22.052   | -14.018   | -1.180     |
| p-value        | 0.008           | 0.087    | 0.001     | 0.016      |
| Best model     | A+B+C+D         | A+B+D    | A+B+C+D   | A+B+D      |
FIGURE 13. Comparison of Otariidae (fur seals and sea lions) diversity with diatom species diversity (productivity), δ¹⁸O values from benthic foraminifers (mean ocean temperature), and sea level changes. Diatom species data: downloaded from Neptune database (https://nsb-mfn-berlin.de) on 03-05-2018; δ¹⁸O values: from Prokoph et al. (2008) (raw data from Zachos et al., 2001, "benthic foraminifera" and "deep sea" series); sea level data: from Miller et al. (2005).
with δ¹⁸O and sea level changes, while we did not find any significant correlations between estuarine habitat preference and changes in climate or primary productivity. These correlations might be influenced by the low counts of fossil occurrences with adaptations compatible with these habitats, but more research needs to be conducted on the causes of re-invasion of freshwater and estuarine habitats by river dolphins. All marine habitats (pelagic, coastal, benthic) instead were correlated with δ¹⁸O and diatom diversity, with marine benthic being correlated also with δ¹³C. This is expected since benthic invertebrates like mollusks are reliant on dissolved carbon for their shells (Byrne, 2011).

The correlations of the habitat preference to climate are confirmed by the strong associations between these same variables and prey type. The most common prey types (fish, soft-bodied free-swimming invertebrates/squid, and zooplankton) are correlated with primary productivity (diatom diversity) and ocean temperature changes (δ¹⁸O), with free-swimming invertebrates again showing significant correlation with carbon isotopes fluctuations (Table 6). Benthic invertebrates were most strongly correlated with δ¹³C and diatom diversity, again showing the importance of carbon fluctuations and primary productivity in the distribution of this prey type. Tetrapods are the only prey type that does not present a significant correlation with climatic variables. This might be due to low counts, but also by the possible opportunistic nature of this prey type choice and other constraints on this prey type (size, dentition). Finally, the three main prey capture strategies (biting/crushing, suction, and filter) present throughout cetacean evolution appear to be associated as well with diatom diversity and δ¹⁸O, with suction also being associated with δ¹³C (Table 7). This is due to the fact that the preferred prey type by suction feeding extant taxa are free-swimming invertebrates (e.g., squid), and this prey type is in turn associated with all three climatic variables.

Sirenia and Desmostylia. When Sirenia and Desmostylia taxa are analyzed together, to account for the overall low taxonomic diversity of these groups, we found that their diversity is correlated with changes in global ocean temperature (δ¹⁸O) and sea level changes (Table 8; Figure 15). This also implies that their prey type (sea grasses and algae) and grazing habits are correlated with these vari-

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**TABLE 2.** Linear regression analyses for Pinnipedimorpha habitat preference through time against climate and productivity variables: δ¹⁸O (variable A), δ¹³C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Only prevalent categories (with at least one time bin with a count > 5) were considered. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S2.

| Category       | Marine coastal | Marine benthic | Marine pelagic |
|----------------|----------------|----------------|---------------|
| R²             | 0.914          | 0.847          | 0.885         |
| AIC            | 28.844         | 0.387          | -7.244        |
| p-value        | 0.004          | 0.004          | 0.036         |
| Best model     | A+C+D          | A+C            | A+B+C+D       |

**TABLE 3.** Linear regression analyses for Pinnipedimorpha prey type preference through time against climate and productivity variables: δ¹⁸O (variable A), δ¹³C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Only prevalent categories (with at least one time bin with a count > 5) were considered. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S3.

| Prey Type          | Fish            | Squid/soft bodied invertebrates | Benthic invertebrates |
|--------------------|-----------------|-------------------------------|-----------------------|
| R²                 | 0.896           | 0.982                         | 0.969                 |
| AIC                | 27.725          | -9.734                        | -4.775                |
| p-value            | 0.011           | < 0.0001                      | < 0.0001              |
| Best model         | A+B+C+D         | A+D                          | A+B+D                |

**TABLE 4.** Linear regression analyses for Cetacea and Odontoceti and Mysticeti diversity through time against climate and productivity variables: δ¹⁸O (variable A), δ¹³C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S4.

| Taxa       | Cetacea | Odontoceti | Mysticeti |
|------------|---------|------------|-----------|
| R²         | 0.674   | 0.667      | 0.706     |
| AIC        | 90.967  | 84.175     | 58.829    |
| p-value    | 0.001   | 0.001      | 0.001     |
| Best model | A+C     | A+C        | A+C       |
FIGURE 14. Comparison of cetacean, mysticete, and odontocete diversity with diatom species diversity (productivity) and δ¹⁸O values from benthic foraminifers (mean ocean temperature). Diatom species data and δ¹⁸O values are from the same sources as in Figure 13.
ables. This is consistent with previous work showing that sirenians (with the exception of cold adapted hydrodamalines) and desmostylians are primarily subtropical and tropical and associated with seagrasses (Vélez-Juarbe, 2014). When performing a Pearson’s correlation test to account for the low counts and check intergroup relationships, we found that, while no single climatic variable is correlated with the diversity of these taxa, the temporal distributions of these two groups are correlated with each other (Table 9). These correlations support the hypotheses of the extinction of Desmostylia being mostly driven by the colonization of the North Pacific by Sirenia. By looking at distribution maps of the main genera for these groups in this area, it appears clear that the appearance of the genus *Hydrodamalis* in the Tortonian (~11 Ma) coincides with the decline of desmostylians, which went completely extinct by the end of the Tortonian (7.2 Ma) (Figure 12). While other variables connected with changes in climate and primary productivity (e.g., δ15N) might be correlated with the diversity of herbivorous groups, their relatively low and stable numbers point to the fact that they occupy a specific ecological niche and are particularly vulnerable to competition.

### Ancestral Reconstructions of Feeding Strategies and Trophic Structure

**Pinnipedimorphs.** Extant and fossil pinnipeds use multiple strategies to capture and consume prey underwater. The ancestral feeding strategy of pinnipeds was a biting strategy (Appendix 3) employing a heterodont dentition in agreement with previous studies based on skull

### TABLE 5. Linear regression analyses for Cetacea habitat preference through time against climate and productivity variables: δ18O (variable A), δ13C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S5.

|                | Riverine | Estuarine | Marine coastal | Marine benthic | Marine pelagic |
|----------------|----------|-----------|----------------|----------------|---------------|
| R²             | 0.601    | 0.240     | 0.676          | 0.906          | 0.710         |
| AIC            | 7.083    | 40.692    | 79.988         | -14.579        | 89.199        |
| p-value        | 0.004    | 0.193     | 0.001          | < 0.0001       | 0.001         |
| Best model     | A+D      | A+D       | A+C            | A+B+C          | A+C           |

### TABLE 6. Linear regression analyses for Cetacea prey type preference through time against climate and productivity variables: δ18O (variable A), δ13C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S6.

|                | Fish      | Squid/soft-bodied inv. | Zooplankton | Benthic inv. | Tetrapods |
|----------------|-----------|------------------------|-------------|--------------|-----------|
| R²             | 0.663     | 0.743                  | 0.698       | 0.837        | 0.036     |
| AIC            | 89.536    | 73.845                 | 59.565      | 6.252        | 27.313    |
| p-value        | 0.001     | 0.001                  | 0.001       | < 0.0001     | 0.500     |
| Best model     | A+C       | A+B+C                  | A+C         | B+C          | C         |

### TABLE 7. Linear regression analyses for Cetacea prey capture strategies through time against climate and productivity variables: δ18O (variable A), δ13C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each category are reported, for complete results see Appendix 2, Table S7.

|                | Biting/ crushing | Suction | Filter |
|----------------|------------------|---------|--------|
| R²             | 0.490            | 0.762   | 0.703  |
| AIC            | 72.148           | 74.033  | 59.146 |
| p-value        | 0.018            | 0.001   | 0.001  |
| Best model     | A+C              | A+B+C   | A+C    |
TABLE 8. Linear regression analyses for Sirenia and Desmostyla diversity through time against climate and productivity variables: Δδ^{18}O (variable A), δ^{13}C (variable B), diatom species diversity (variable C), sea level changes (variable D). Best model, including one, two, three, or all four variables, is selected based on its AIC (Akaike’s Information Criterion) score. Diversity of Desmostyla is considered from the time bin of first occurrence to the last (from Rupelian to Tortonian). To account for their low diversity, these groups were also considered together (“Sirenia and Desmostyla combined”). Significant values (p < 0.05) are in bold. Only data (R², p-value, AIC) for the best model for each group are reported, for complete results see Appendix 2, Table S8.

| Variables | Sirenia | Desmostyla | Sirenia and Desmostyla combined |
|-----------|---------|------------|--------------------------------|
| R²        | 0.243   | 0.603      | 0.460                          |
| AIC       | 27.978  | 5.200      | 37.158                         |
| p-value   | 0.189   | 0.158      | 0.025                          |
| Best model| A+C     | B+D        | A+D                            |

and dental morphologies (e.g., Adam and Berta, 2002; Churchill and Clementz, 2015). Phocoids (phocids + desmatophocids) are biters with several taxa having evolved specialist strategies (Appendix 3). The extant phocid *Hydrurga* (leopard seal) employs biting and filter strategies, and this is predicted for the fossil seal (*Homiphoca*). *Lobodon* (crabeater seal) is the only extant pinnipedimorph that is a specialized filter feeder. *Mirounga* and *Eriphragmus* (bearded seal) employ a suction feeding strategy. Although we reconstructed the extinct phocid *Hadrokiirus* as having a biting/crushing diet largely on the basis of robust, well-worn teeth, Amson and de Muizon (2014) described this taxon as having a more exclusive durophagous habit. Otariids uniformly employ a biting strategy with the exception of some species of *Arctocephalus* that are more generally classified as biting/suction feeders. Odobenids have been hypothesized as exhibiting a biting strategy ancestrally (Boessenecker and Churchill, 2013), and this pattern is also seen in our analysis (Figure 16). The characteristic suction feeding strategy of the modern walrus apparently evolved twice, once among dusignathine walruses (e.g., *Dusignathus*) and again among odobenine walruses (e.g., *Protodobenus*).

Reconstructions of tooth shape indicate that ancestrally the teeth of stem pinnipeds are heterodont with pointed cusps (Appendix 4). Crown pinnipeds are characterized by the development of homodonty with stem taxa in all three major lin-
FIGURE 15. Comparison of Sirenia (manatees and dugongs) and Desmostylia diversity with $\delta^{18}$O values from benthic foraminifers (mean ocean temperature) and sea level changes. $\delta^{18}$O values and sea level change data are from the same sources as in Figure 13.
generally reported (Geisler et al., 2017), although this reconstruction has been challenged (Hocking et al., 2017a; Fordyce and Marx, 2018; Peredo and Pyenson, 2018). Other stem toothed mysticetes (e.g., *Mammalodon*, *Mystacodon*) are reconstructed as having been suction feeders and still other stem taxa (e.g., *aetiocetids*) employed a filter/suction feeding strategy (Deméré and Berta, 2008, but see Marx et al., 2016 and Peredo et al., 2018 for alternate interpretations). Beginning 30 Ma, stem edentulous mysticetes (e.g., *eomysticetids*) exhibited bulk filter feeding that was retained by all crown mysticetes (Gatesy et al., 2013). Our analysis recovers this transition and also highlights that the switch to bulk filter feeding was likely mediated by intermediate forms that were capturing prey using a combination of filter and suction feeding, in line with previous hypotheses described above (Figure 18). The split between skim feeding balaenoids and more diverse generalized and specialized filter feeders, the thalassotherians, was completed by early Miocene (20-18 Ma) (Berta et al., 2016). Among thalassotherians, some cetotheriids (e.g., *Herpetocetus* - Gol’din and Startsev, 2014; El Adli et al., 2014) display a continuous suction feeding mode as early as 11-10 Ma (Berta et al., 2016). The specialized intermittent suction feeding that characterizes Eschrichtiidae, may have occurred as early as 5 Ma. The balaenopterid record extends into the late Miocene (12-10 Ma), which likely marks the origin of engulfment feeding, although the advanced lunge feeding behavior seen in living rorquals (i.e., characterized by

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**TABLE 9.** Correlation test (Pearson’s coefficient) for Sirenia, Desmostylia, and “Sirenia and Desmostylia combined” diversity through time with climate and productivity variables: δ¹⁸O, δ¹³C, diatom species diversity, sea level changes. Significant values (p < 0.05) are in bold. Cells for auto-correlated variables are grayed out. P-values for the correlations are reported in Appendix 2, Table S9.

|                | Sirenia | Desmostylia | Sirenia and Desmostylia combined |
|----------------|---------|-------------|----------------------------------|
| Sirenia        |         | 0.040       |                                  |
| Desmostylia    | 0.040   |             |                                  |
| Sirenia and Desmostylia |       |             | 0.168                           |
| δ¹⁸O           | 0.252   | 0.207       | 0.168                           |
| δ¹³C           | 0.353   | 0.068       | 0.128                           |
| Diatom diversity | 0.870 | 0.660       | 0.742                           |
| Sea level      | 0.823   | 0.379       | 0.554                           |

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**FIGURE 16.** Ancestral state reconstruction of prey capture strategies and tooth pattern and cusp shape in Odobenidae. Topology follows Berta et al. (2018). Details on the tree can be found in Appendix 1. All data matrices and complete trees with branch lengths can be found in the Supplementary Information uploaded to Dryad repository (https://doi.org/10.6086/d14671).
FIGURE 17. Ancestral state reconstruction of prey type preference in stem and crown Pinnipedimorpha. Topology follows Rybczynski et al. (2009), Dewaele et al. (2017), and Berta et al. (2018). Details on the tree can be found in Appendix 1. All data matrices and complete trees with branch can be found in the Supplementary Information (https://doi.org/10.6086/d14671).
greater degree of mandibular alpha rotation and lateral expansion of the mouth) developed later (Berta et al., 2016). The evolution of specialized filter feeding modes allowed mysticetes to feed on different prey types, as highlighted later in the analysis of prey preference.

Odontocetes exhibit more diverse prey capture strategies. Ancestrally, stem odontocetes were similar to stem cetaceans in their employment of a biting/crushing strategy (Figure 19). Although some stem taxa (e.g., Archaeodelphis, Xenorophus, Cotylocara) are reconstructed as using a combination of biting/crushing and suction feeding. Crown odontocetes exhibit a greater diversity of feeding strategies and cranial morphologies (Figure 20). The ancestor of crown odontocetes was likely employing the ancestral biting/crushing prey capture mode. Living sperm whales (Physeteridae and Kogiidae) and beaked whales (Ziphiidae) have been suction feeders since their first appearance in the fossil record, although the teeth and skull morphology of some extinct genera closely related to the modern taxa (e.g., Acrophyseter, Livyatan, and Ninoziphius) support the hypothesis that ancestrally they were able to feed via biting and/or suction on different kinds of prey (Bianucci et al., 2016; Lambert et al., 2016). As characteristic of opportunistic feeding, ancestral Delphinoidea (Delphinidae, Monodontidae, and Phocoenidae) are reconstructed as having employed a biting/crushing/suction feeding strategy.

Fossil stem mysticetes and odontocetes are similar to archaeocetes in exhibiting heterodonty (Figure 19; Appendix 7). Beginning 30 Ma, edentulous stem mysticetes (e.g., eomysticetids) are characterized by their lack of functional teeth and

![Diagram of prey capture strategies and prey type preference in stem and crown Mysticeti.](https://doi.org/10.6086/d14671)
presumably possessed baleen (Appendix 7). A homodont dentition evolved among Physeteroidea (Kogiidae and Physeteridae) and is retained among all later odontocetes with the exception of ziphiiids which feature a reduced dentition (Figure 20).

Inferred diets are a direct result of these feeding adaptations and tooth morphologies. Ancestrally, cetaceans fed on fish (Appendix 8). Some archaeocetes (e.g., Basilosaurus) fed on fish and tetrapods. Most stem mysticetes (e.g., Coronodon and Llanocetus) likely fed on fish and zooplankton whereas several others (e.g., Mystacodon and Mammalodon) consumed benthic invertebrates. Stem taxa at the base of odontocete divergence are reconstructed as predominately having fed on fish. Ancestrally, a diet composed of fish/squid evolved among some stem odontocetes (e.g., Cotylocara, Archaeodelphis) and is characteristic of all crown odontocetes. A diet composed of fish and tetrapods evolved only a few times among crown odontocetes: in Orcinus and in some Physeteridae (e.g., Livyatan – not represented on the tree). In support of this result, diversity of delphinine dolphins (e.g., spinner dolphins, bottlenose dolphins, common dolphins, etc.) in the early Pliocene has been related to phytoplankton groups most abundant in the tropics, which in turn supports high fish abundances (Amaral et al., 2018).

With the exception of Pakicetus which occupied a riverine/estuarine habitat, the ancestral habitat for cetaceans is reconstructed as mixed estuarine/coastal marine environments (Appendix 9). There is isotopic evidence to suggest that the earliest cetaceans inhabited freshwater (Clementz et al., 2006). The protocetid-basilosaurid transition corresponds to the rise of the first fully aquatic cetaceans. Some of these later evolving stem cetaceans occupied more variable marine habitats including both pelagic and coastal environments. Most stem and crown mysticetes are characterized as having occupied these same environments. The extant gray whale Eschrichtius and its close fossil relative Eschrichtoides are reconstructed to inhabit marine coastal and pelagic as well as benthic habitats; the latter is a consequence of their suction feeding strategy. Additional isotopic analyses and other methods of investigation are needed to more narrowly define the habitat of extinct cetaceans, as the depositional environment of the fossils is not always a reliable indicator.

**Other marine mammals.** We did not perform ancestral state feeding reconstructions for sireniens, desmostylians, marine otters, aquatic sloths, and bears given their relatively low generic diversity and categorical data with little variation. Relevant feeding and habitat data collected for these taxa (Supplementary Information, https://doi.org/10.6086/d14671) allow a few general comments. Since their first appearance in the Eocene, sireniens have been identified as herbivores feeding on aquatic plants. Prorastomids possessed low crowned, rounded tooth cusps and were probably browsers that fed on floating aquatic plants and to
a lesser extent sea grasses, the primary food of modern dugongs (Donning, 2001). Later dugong-gids evolved tusks and bilophodont postcanine teeth for feeding on sea grass rhizomes. In response to cooler climates, members of the hydrodamaline lineage are hypothesized to have fed on kelp which replaced (to a large extent) sea grasses (Pyenson and Vermeij, 2016). Since the oldest known sirenian locality in Africa consists of lake deposits, sirenians like cetaceans may have first evolved in freshwater (Clementz et al., 2006; Benoît et al., 2013). Based on isotope data, Clementz et al. (2003) concluded that desmostylians were aquatic herbivores that, like sirenians, foraged on sea grasses in estuarine and coastal freshwater systems. Coincident with the evolution of large sirenians in the North Pacific there is evidence to suggest that this region served as the origin of kelps (Laminariales) (Pyenson and Vermeij, 2016). Another herbivore, the aquatic sloth (*Thalassocnus* spp.), evolved in the South Atlantic (Peru and Chile) and fed on sea grasses in marine coastal habitats (de Muizon et al., 2004). The aquatic ursid/pinnipedimorph *Kolponomos* occupied a specialized crushing/suction feeding guild consuming various hard benthic invertebrates (e.g., sea urchins, clams, and abalones) during the Miocene in the North Pacific (Tseng et al., 2016). More recent occupants of this niche are the otters *Enhydra* and *Enhydritherium* and the sea mink *Neovison*. Paleoenvironmental evidence indicates that they occupied a broad range of habitats including freshwater as well as estuarine and coastal marine environments.
Evolution of Feeding Guilds

Results of this study allow preliminary inferences about the evolution of feeding guilds in marine mammals. Most fossil and extant marine mammals fall into one of four feeding guilds: biting/crushing, suction, filter, and grazing. By far the most common method of feeding for both cetaceans and pinnipedimorphs was a biting/crushing strategy using pointed teeth to consume fish. For pinnipedimorphs, this strategy evolved during the late Oligocene. Most stem taxa lived in coastal marine habitats, which is also the dominant habitat of extant pinnipeds. The distinctive suction feeding habit of the modern walrus feeding on mollusks in a benthic habitat evolved in later diverging walruses beginning in the late Pliocene (3 Ma). Earlier stem walruses were more generalist feeders consuming fish and employing a biting strategy. Extinct desmatophocid pinnipeds exhibited varied diets, with Allodesmus possessing adaptations associated with specialized suction feeding and Desmatophoca spp. identified as more generalized biting/crushing/suction feeders. The filter feeding niche is primarily occupied at present by a single pinniped species, the crabeater seal Lobodon carcinophagus, which employs multi-cusped teeth as a sieve to filter zooplankton from the water.

The biting/crushing feeding guild strategy is exhibited by stem whales (archaeocetes) approximately 50 Ma. Early whales inhabited riverine and estuarine environments and later dispersed to occupy marine coastal and pelagic habitats. One of the more interesting results of the ASR of prey capture in whales is that filter/suction feeding in whales appeared early in the evolution of mysticetes (e.g., Coronodon), at approximately 30 Ma. The shift toward bulk-based filter feeding as the sole feeding strategy began with eomysticetids that employed baleen to filter fish and zooplankton. Odontocetes are characterized by several feeding strategies including biting/crushing/suction, biting/crushing, and specialized suction feeding. Specialized suction feeding evolved multiple times in extinct stem odontocetes (e.g., Inermorosstrum), physteroids, ziphids, and some delphinoid odontocetes (e.g., Monodon, Delphinapterus).

The specialized crushing/suction feeding niche was occupied by the extinct aquatic carnivore Kolponomos in the Miocene and the extant sea otter Enhydra and its fossil relatives in the Pliocene. Marine herbivores (desmostylians, sirenians, and aquatic sloths) filled the grazing feeding guild with stem sirenians appearing first in the late Eocene (50 Ma). There are strong geographical differences in the occupants of this niche. Aquatic sloths were restricted to the South Pacific while desmostylians occupied the North Pacific. Sirenians were more broadly distributed in the past, especially in the subtropical tropical western Atlantic and Caribbean regions, with the large bodied hydromodaline lineage occupying temperate to sub-polar waters.

Several previous studies of feeding guild evolution in marine mammals have shown the importance of body size as an important variable (e.g., Churchill et al., 2015; Pyenson and Vermeij, 2016; Slater et al., 2017). The next step involves the incorporation of quantitative data (e.g., cranial morphology) with dietary and feeding mode data as has been done with other marine vertebrates (Friedman, 2012; Motani et al., 2015; Kelley and Motani, 2015; Pyenson and Vermeij, 2016). Only then will it be possible to identify patterns in skull and tooth morphology and better discriminate trophic groups among clades. It is also important to consider these patterns in the context of the geographic and temporal distributions of marine mammal faunas. Mapping these patterns onto phylogenies will enable testing of previous hypotheses of trophic convergence, niche diversity, and replacement.

The Future: Marine Mammals and Ecosystems

By analyzing the past and present feeding diversity of marine mammals, we can make predictions about the future. In addition to changes in global climate and ocean productivity, human interactions since the Late Pleistocene have strongly influenced marine mammals and their ecosystems. Several marine mammals, notably Steller’s sea cow (Hydromaline gigas) and the Caribbean monk seal (Neomonachus tropicalis) have been hunted to extinction (Turvey and Risley, 2006; Baisre, 2013). The extinction of other species such as the Yangtze river dolphin, Lipotes vexillifer, has been attributed to habitat degradation (Pyenson, 2009). Historical records document the extensive hunting of manatees, Trichechus spp., by early colonists in the Americas and dugongids by Aboriginal Australians (Estes et al., 2016). Anthropogenic climate change and resultant expansions and contractions of habitat are responsible for distributional shifts as well as changes in feeding ecology. For example, Pyenson and Lindberg (2011) showed that changes in Pleistocene sea level cycles resulted in fluctuations in available benthic feeding areas. These scientists suggested that gray whales survived Pleistocene glacial maxima by feeding...
outside of the Sea of Okhotsk and Bering region in the North Pole, which were ice covered, employing a more generalized feeding strategy. Given this change in available habitat, gray whales likely fed on fish during the Pleistocene, rather than benthic amphipods consumed by the living gray whale. Changing food web dynamics, such as the historic interactions between killer whales and sea otters or baleen whales (Estes et al., 1998) and Steller’s sea cow-sea otters and kelp (Estes et al., 2016), would benefit from further exploration in the fossil record.

Another area of investigation with important implications for modern marine ecosystems is the evolution of fossil marine mammal communities, little studied thus far. A notable exception is identification of the co-occurrence of species, or sympatry, among dugongids evolving in separate time periods and geographic regions, the late Oligocene in Florida, the early Miocene in India, and the early Pliocene in Mexico. Resource partitioning among fossil sirenians was examined in these communities defined on the basis of dugongid body size and ecomorphological differences such as rostral deflection and tusk morphology (Vélez-Juarbe et al., 2012).

Also worthy of consideration is the decline of marine mammals that have been attributed to predators, such as the decline of Pliocene mysticetes that has been linked to the appearance and diversification of their likely predators, carcharodontid sharks (Lindberg and Pyenson, 2006).

These examples point up the importance of collecting fossil data as fossils provide critical, unique reference points for understanding changes in marine mammal communities and diversity through time.

CONCLUSIONS

Marine mammals have evolved four feeding strategies to capture prey underwater: biting, filter, suction, and grazing. Distinct skull and soft tissue morphologies have been identified and associated with each feeding strategy, and future studies would benefit from more comparative and quantitative approaches to the study of feeding structures, styles, and performance. By analyzing the evolution of feeding in marine mammals, we confirm that the earliest feeding guild in the late Eocene was occupied by both herbivores (grazing sirenians) and carnivores (biting/crushing archaeocetes). Morphological and paleoecological evidence indicate that a biting feeding mode was predominant among Miocene cetaceans and pinnipeds whereas other feeding styles, notably the specialized crushing of hard-shelled invertebrates, had various occupants in the past, including a phocid seal and an aquatic ursid, and this niche has been represented since the Pleistocene by sea otters. The filter feeding niche appears to have originated perhaps as early as the late Eocene although details of the tooth to baleen transition remain contentious as does the broader ecological context. Large marine herbivores (desmostylians and sirenians) were continual occupants of the grazing niche with the addition of aquatic sloths in the South Pacific.

Understanding how drivers of past diversity, both biotic and environmental, continue to operate today helps shape our efforts to conserve and manage these remarkable mammals of the sea. Major changes in productivity of the world’s oceans was a primary driver of feeding diversification in both pinnipeds and cetaceans. The peak in diversity of major lineages of marine mammals occurred during the mid-Miocene and has been associated with the Mid-Miocene Climate Optimum. Later changes in diversity can be related to the Messinian salinity crisis, especially for those taxa occupying the Mediterranean, and to Pleistocene sea level changes that affected habitat and prey availability for coastal dwelling pinnipeds and cetaceans. Ecological replacements are more difficult to recognize in the absence of a good, well-sampled fossil record. Perhaps the best example is replacement of desmostylians by sirenians in the North Pacific that occurred in the late Miocene. Evidence for trophic convergence during this same time comes from sea grass feeding aquatic sloths in the South Pacific that appear to be the ecological homologue of desmostylians in the Pacific. Disentangling the effects of ecosystem changes requires the development of rigorous hypotheses that incorporate biotic as well as environmental drivers of ecosystem change. Additionally, further study of the prey of marine mammals would enable understanding of the co-evolution of predator-prey communities through time.

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APPENDICES

All appendices are grouped in a zipped file available at https://palaeo-electronica.org/content/2020/3136-feeding-in-marine-mammals.

APPENDIX 1.

Details and sources trees for Pinnipedimorpha (1a) and Cetacea (1b) used for ASR.

APPENDIX 2.

Complete results of statistical analyses.

APPENDIX 3.

Ancestral State Reconstruction of prey capture strategies in Pinnipedimorpha. Tree topology as in Appendix 1a.

APPENDIX 4.

Ancestral State Reconstruction of tooth pattern and cusp shape in Pinnipedimorpha. Tree topology as in Appendix 1a.

APPENDIX 5.

Ancestral State Reconstruction of habitat preference in Pinnipedimorpha. Tree topology as in Appendix 1a.

APPENDIX 6.

Ancestral State Reconstruction of prey capture strategies for Cetacea. Tree topology as in Appendix 1b.

APPENDIX 7.

Ancestral State Reconstruction of tooth pattern and cusp shape in Cetacea. Tree topology as in Appendix 1b.

APPENDIX 8.

Ancestral State Reconstruction of prey type preference in Cetacea. Tree topology as in Appendix 1b.

APPENDIX 9.

Ancestral State Reconstruction of habitat preference in Cetacea. Tree topology as in Appendix 1b.