Feeding Kinematics, Suction, and Hydraulic Jetting Performance of Harbor Seals (*Phoca vitulina*)

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Abstract

The feeding kinematics, suction and hydraulic jetting capabilities of captive harbor seals (Phoca vitulina) were characterized during controlled feeding trials. Feeding trials were conducted using a feeding apparatus that allowed a choice between biting and suction, but also presented food that could be ingested only by suction. Subambient pressure exerted during suction feeding behaviors was directly measured using pressure transducers. The mean feeding cycle duration for suctionfeeding events was significantly shorter (0.15 ± 0.09 s; P<0.01) than biting feeding events (0.18 ± 0.08 s). Subjects feeding inwater used both a suction and a biting feeding mode. Suction was the favored feeding mode (84% of all feeding events) compared to biting, but biting comprised 16% of feeding events. In addition, seals occasionally alternated suction with hydraulic jetting, or used hydraulic jetting independently, to remove fish from the apparatus. Suction and biting feeding modes were kinematically distinct regardless of feeding location (in-water vs. on-land). Suction was characterized by a significantly smaller gape (1.3 \pm 0.23 cm; P<0.001) and gape angle (12.9 \pm 2.02°), pursing of the rostral lips to form a circular aperture, and pursing of the lateral lips to occlude lateral gape. Biting was characterized by a large gape (3.63±0.21 cm) and gape angle (28.8±1.80°; P<0.001) and lip curling to expose teeth. The maximum subambient pressure recorded was 48.8 kPa. In addition, harbor seals were able to jet water at food items using suprambient pressure, also known as hydraulic jetting. The maximum hydraulic jetting force recorded was 53.9 kPa. Suction and hydraulic jetting where employed 90.5% and 9.5%, respectively, during underwater feeding events. Harbor seals displayed a wide repertoire of behaviorally flexible feeding strategies to ingest fish from the feeding apparatus. Such flexibility of feeding strategies and biomechanics likely forms the basis of their opportunistic, generalized feeding ecology and concomitant breadth of diet.

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Introduction

Secondarily adapted aquatic tetrapods (e.g., marine turtles, penguins, and marine mammals) represent an evolutionarily interesting experiment in organismal adaptation. The aquatic environment has imposed strong selection pressures on all aquatic vertebrates, particularly for feeding and locomotion. This has also been true for terrestrial vertebrates re-invading aquatic habitats. Raptorial feeding is the ancestral condition among all gnathostomes, but the evolution of suction feeding is an especially important feeding mode among actinopterygians, as well as chondrichthyans [1-5]. The terrestrial ancestors of marine mammals are thought to have exhibited a raptorial-type feeding mode [6]. Therefore, part of their transition back to the sea likely involved an independent and secondary emergence of new mechanisms for underwater feeding. Our knowledge regarding these adaptations to underwater feeding and the feeding repertoire of aquatic mammals is limited to only a few species and much work needs to be conducted to understand feeding adaptations of marine mammals from a comparative perspective. For example, it is not well known how widespread suction feeding is among aquatic mammals, and in the case of pinnipeds, a generalist (pierce) biting feeding mode is often evoked to describe their ancestral feeding mode [6]. The lack of feeding performance data for secondarily aquatic tetrapods is due to the fact that they spend considerable time at depth foraging and direct observation of prey capture is rare.

The field of feeding functional morphology among secondarily aquatic tetrapods is still in its infancy but has benefitted greatly from investigations of fish and amphibian feeding biomechanics [4,7–11]. Among secondarily aquatic tetrapods suction feeding specialists, such as walruses (*Odobenus rosmarus*), bearded seals (*Erignathus barbatus*), pygmy sperm whales (*Kogia* sp) and belugas (*Delphinapterus leucas*), the primary mechanism of generating subambient pressure is the rapid depression and retraction of the tongue and hyoid apparatus [12–18]. This rapid depression of the tongue and the hyoid apparatus can be observed externally (gular

depression) in feeding pinnipeds and cetaceans [15,17-19]. The rapid change in intraoral volume generates significant subambient pressure. In these specialists, the tongue is wide, thick and pistonlike [13,14,16]. It has been suggested that suction specialists also possess an enlarged hyoid apparatus [20-22], which presumes a hypertrophied hyolingual musculature and more forceful contraction. While this may be true for some taxa, it does not hold for all suction feeders. For example, Bloodworth and Marshall [16] compared the hyolingual musculature of pygmy and dwarf sperm whales (suction specialists) and bottlenose dolphins (a ram feeder), as well as the feeding kinematics in these species [19], and found no evidence of increase muscle tension capability in suction specialists. Instead modified orofacial morphology is thought to be of greater importance [16], as well as shape of the mandible and head [23-25]. Shape of the mouth and the occlusion of lateral gape are important for generating and orienting suction forces for prey capture [5]. Walruses and bearded seals have broad skulls [26,27] and muscular snouts [28,29] which they use to purse and create a circular aperture rostrally, and occlude lateral gape by pursing the lateral margins of the lips and mouth. Salamanders occlude lateral gape using labial lobes [10,30,31], actinopterygian fish use membranous labial lips that span the upper and lower jaws [32] and suction feeding elasmobranchs occlude lateral gape using labial cartilages [4,33,34]. In addition, the vaulted upper palate of suction feeding specialists is hypothesized to increase the rapid volume change, when the tongue is rapidly depressed, to further maximize subambient pressure generation [13,26]. Rapid lower jaw opening, while maintaining small gape, can also contribute to subambient pressure generation in some species (bearded seals) [17] due to buccal expansion as long as a circular aperture is maintained and the lips occlude the lateral sides of the mouth.

Understanding prey-capture tactics and feeding performance are important considerations in trophic ecological questions since such behavior can determine prey choice due to energetic constraints [35-37]. The primary methods of studying pinniped foraging ecology have been through indirect methods (e.g., stomach content and fecal analyses), and more recently through the use of animal-born cameras [37-44] and other instrumentation. While such underwater footage can reveal much regarding hunting tactics, the actual moment of prey capture is often obscured and performance measurement of prey capture in the wild is limited. Although the marine mammal literature is rich in correlations among anatomy and feeding modes, studies that have collected direct empirical data of marine mammal feeding performance are limited. Among odontocetes, only belugas (Delphinapterus leucas), pygmy and dwarf sperm whales (Kogia breviceps, K. sima) and longfinned pilot whales (Globicephala melas) have been demonstrated to use suction as a primary feeding mode, whereas only bottlenose dolphins (Tursiops truncatus), and Pacific white-sided dolphins (Lagenorhynchus obliquidens) have been shown to primarily use a ram or raptorial feeding mode [18,19], although this is likely a common feeding mode among many odontocetes. Sirenian feeding biomechanics and food handling have received some attention; their feeding involves the use of modified vibrissae, or bristles, for gathering vegetation and transporting it to the cheek-teeth for mastication, but no evidence for suction has been reported [45-47]. Data for pinnipeds are also few, but a mix of kinematic and performance data exists for bearded seals (Erignathus barbatus) [17], leopard seals (Hydrurga leptonyx) [48], and walruses (Odobenus rosmarus) [13,49]. In marine mammals subambient pressure, or suction, is primarily generated by the rapid retraction and depression of the tongue via the hyolingual apparatus [20-23].

Pinnipeds are apex predators that comprise an important component of marine food webs. They exhibit an interesting range of feeding modes that include suction, biting, grip-and-tear, and filter feeding [50-53]. However, among pinnipeds only walruses, bearded seals, and leopard seals [13,17,48,49] have been demonstrated to use suction, although indirect evidence suggests many more species use suction [51,53]. Given the functional requirements of capturing prey underwater, and the dense and viscous physical properties of water, one would predict that the adaptations for prev acquisition would converge widely with other obligate underwater feeding vertebrates. That is, suction feeding should be a widespread feeding mode among pinnipeds and other marine mammals. Although phocids are thought to have a greater suction capability than otariids and otariids are presumed to predominantly use biting for capturing prey [51], there are few data to support or refute these claims. However, it is becoming clear that marine mammals use multiple feeding modes depending upon the need [17,48]. Although recent data have demonstrated high performance values for suction feeding in bearded seals and walruses, these species are specialists among pinnipeds and may not represent the more generalized trophic ecology of many pinnipeds.

Among phocids, harbor seals (Phoca vitulina, Linnaeus, 1758) are the most widely distributed species globally and exhibit an opportunistic and generalized feeding ecology. Up to five subspecies are recognized: P. v. vitulina (Linneaus, 1858), P. v. concolor (DeKay, 1842), P. v. mellonae (Doutt, 1942), P. v. richardii (Gray, 1864), and P. v. stejnegeri (Allen, 1902) [54,55], but there is much disparity in the morphological and genetic data and many of these subspecies are unsupported, or require further assessment. Harbor seals are generalists that feed upon a wide diversity of small- to medium-size fishes (e.g., herring, anchovy, cod, hake, trout, smelt, shad, scorpionfish, rockfish, prickleback, greenling, sculpin, capelin, sandlance, salmon, and flatfish), and a variety cephalopods, and invertebrates (mostly crab and shrimp species but also mollusks) [56-62]. Foraging can occur adjacent to haulouts, along rivers, or ~ 50 km offshore from haulouts [63]. They can forage for food at considerable depths (\sim 500 m) or in shallow water [55,64]. Diets vary regionally and seasonally, and prey availability is likely the driving force behind diet composition, however, the diet of different age classes also varies [65]. Despite the reported diversity of diet, often a few items will make up the bulk of seasonal diets, which vary with locality [65]. Some evidence suggests that harbor seals use several feeding modes including suction and biting [37]. Therefore, harbor seals are ideal candidates to test hypotheses regarding feeding performance of secondarily aquatic mammals that have implications for both proximal and ultimate questions. The objectives of this study were to investigate the feeding performance of harbor seals to begin to determine the range of their behavioral repertoire for capturing prey, and to test the hypothesis that suction feeding is their primary feeding mode, as opposed to biting. In addition, the hypothesis that rapid jaw opening, in addition to rapid tongue and hyoid depression, is correlated with suction feeding was tested. Since suction appears to be an important feeding mode, subambient (suction) and suprambient pressure (hydraulic jetting) generated by harbor seals were directly measured to test the hypothesis that harbor seal pressure generating capability is similar to values reported for bearded seals [17].

Table 1. Experimental Subjects.

Subject	Birth year	Age (years)	Body Length (cm)	Mass (kg)	Birth Location
Filou	2006	4	130	68.4	captivity
Luca	2002	8	143	84.5	captivity
Malte	1999	11	141*	108.5	captivity
Nick	1999	11	161	102	captivity
Bill	1998	12	150	94	captivity
Henry	1997	13	151	75	captivity
Sam	1994	16	153	89	captivity
Marco	1982	28	143	85.5	unknown

Body length follows American Society of Mammalogists standards [77].

*This subject lacks the small tail in between the hind flippers (average tail length: 8 cm).

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Materials and Methods

Subjects

This study was conducted at the Cologne Zoo (Cologne, Germany) and at the Marine Science Center at the University of Rostock (Rostock, Germany). Eight adult male harbor seals participated in this study (see Table 1 for details regarding subjects); the seals were well trained and were eager to participate in these novel tasks. All work was approved by Texas A&M University's Institute of Animal Care and Use Committee Animal Use Protocol # 2010–67, and was conducted in accordance with the European Communities Council Directive of 24 November 1986 (86/609/EEC). All work at the Marine Science Center at the Cologne Zoo and at the University of Rostock was approved by both institutes.

Feeding Platform

Feeding apparatuses were constructed to present subjects with food items (cut herring and sprat) in a controlled and repeatable research design. A plexiglass feeding panel was inserted vertically into the feeding apparatus with the feeding surface parallel to an underwater camera's perspective (Fig. 1). Nine holes, 3.3 cm in diameter, were drilled through the plexiglass in three rows and three columns, 2 cm apart. Pieces of fish were presented to the subjects in two ways, simultaneously. Cut fish were pushed through the holes such that portions of fish projected out of the feeding surface, and were accessible to the subjects. In addition, fish were cut to fit within recessed plexiglass cylinders (inner diameter = 3.8 cm; length = 5.7 cm) that were positioned behind five of the holes through the plexiglass. Holes (1 cm diameter) were drilled through the back of each plexiglass cylinder to allow water flow. The feeding apparatus was placed in the water, suspended

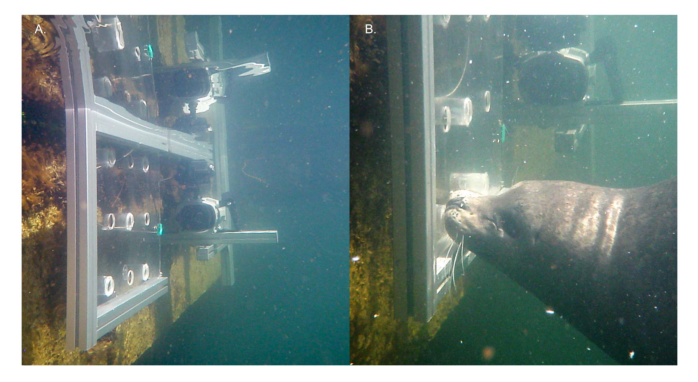


Figure 1. Experimental Feeding Platform. A. Feeding platform in place in the enclosure. B. Harbor seal feeding from feeding apparatus. doi:10.1371/journal.pone.0086710.g001

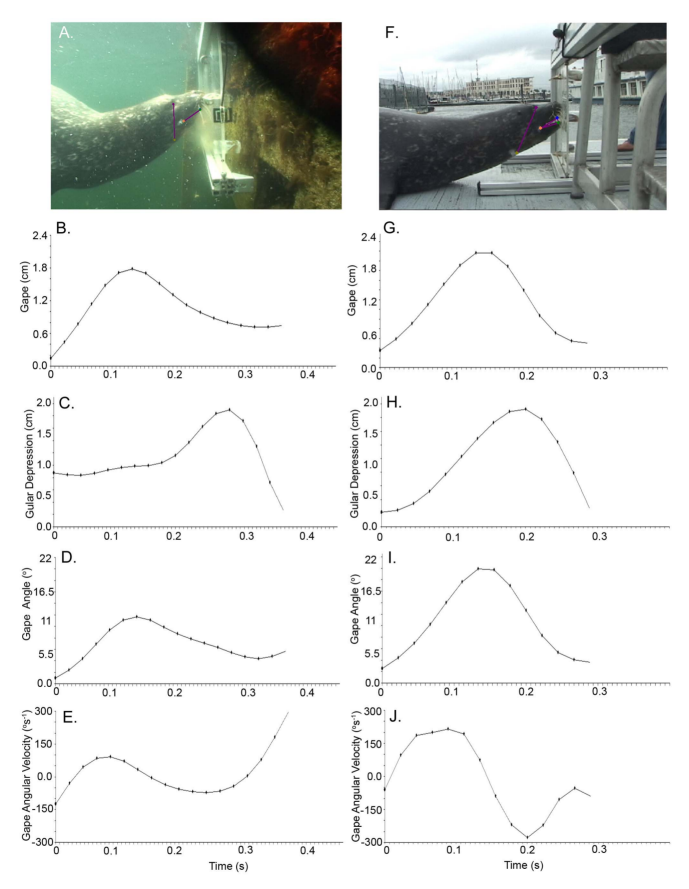


Figure 2. Representative Kinematic Profiles of Suction vs. Biting. A. Frame from video during in-water suction feeding trial with overlaid spatial model stick figure. B. Plot of Gape (cm) for a single suction feeding trial. C. Plot of Gular Depression (cm) for a single suction feeding trial.

D. Plot of Maximum Gape Angle (degrees) for a single suction feeding trial. E. Plot of Gape Angle Velocity (degrees/s; opening and closing) for a single suction feeding trial. F. Frame from video during on-land biting feeding trial with overlaid spatial model stick figure. G. Plot of Gape (cm) for a single biting feeding trial. H. Plot of Maximum Gape Angle (degrees) for a single biting feeding trial. I. Plot of Gular Depression (cm) for a single biting feeding trial. J. Plot of Gape Angle Velocity (degrees/s; opening and closing) for a single biting feeding trial. J. Plot of Gape Angle Velocity (degrees/s; opening and closing) for a single biting feeding trial. doi:10.1371/journal.pone.0086710.g002

just below the surface, in the vertical plane (see Fig. 1). In addition, feeding trials were also conducted with the feeding apparatus out of the water. Subjects hauled out pool-side and were allowed to feed from the apparatus in the same manner as during the in-water trials.

By presenting food items projecting from holes in the feeding surface and within recessed cylinders during feeding trials, subjects were forced to make several choices when presented with food. Upon encountering food items, their first choice was whether to consume fish projecting from the holes or fish residing within the recessed cylinder. If seals chose to consume fish projecting from holes in the plexiglass feeding surface, then they had to choose whether to consume the piece of fish by biting and removing the fish with their teeth, or to use suction. If a subject chose to consume a piece of fish in the recessed cylinder, then its only option was to use suction to obtain that food item. In this way it was possible to determine whether subjects used biting or suction as their primary feeding mode. To challenge subjects, numerous pieces of fish were packed into recessed cylinders to elicit maximum suction generation. We used both kinematic and pressure data to categorize each feeding event as suction or biting. A hermetically sealed pressure catheter was placed through one of the cylinders to directly measure suction forces simultaneously with kinematic events (see below).

Feeding Events, Kinematic Variables and Analyses

One hundred forty-four feeding events were recorded from eight subjects. From this dataset, ninety-one feeding events (from fifty-one feeding trials) from five subjects met the criteria for kinematic analyses. To meet our kinematic criteria, the cranial landmarks of each subject and food items had to be visible within each video frame during an entire feeding event, and rotation of the body around the longitudinal axis had to be minimal ($<10^\circ$). Harbor seals had a propensity to roll as they fed but could successfully feed in any orientation. This necessitated conducting many more feeding trials than were ultimately used in the

Kinematic Variable	Suction	Biting	Р
		29	·
Max. Gape (cm)	1.3±0.23	3.6±0.21	0.001
Time to Max. Gape (s)	0.07±0.01	$0.08\!\pm\!0.04$	0.01
Maximum Gape Angle (°)	12.9±2.02	28.2±1.8	0.001
Time to Max. Gape Angle (s)	0.07±0.01	0.08 ± 0.04	0.02
Max. GAOV (deg. s ⁻¹)	197.2±42.04	561±37.61	0.001
Time to Max. GAOV (s)	0.05±0.01	0.04±0.01	0.172
Max. GACV (deg. s ⁻¹)	159.1±42.05	544.2±37.61	0.001
Time to GACV (s)	0.11 ± 0.01	0.13 ± 0.01	0.001
Max. Gular Depression (cm)	1.4±0.08	0.9±0.07	0.001
Time to Max. Gular Depression (s)	0.14±0.01	0.07±0.01	0.001
Feeding Cycle Duration (s)	0.15±0.09	0.18±0.08	0.01

Values are means \pm S.D., N = 5, 91 feeding events, GAOV = Gape Open Angle Velocity, GACV = Gape Angle Close Velocity.

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kinematic analyses. The number of feeding events used in the kinematic analyses was nearly equal between in-water and out-ofwater trials (44 in-water vs. 47 out-of-water events). Subjects were videotaped at 60 Hz using a Sony camcorder within an underwater video housing (Fig. 1). Prior to feeding trials, zinc oxide landmarks were placed on the subject's lips, jaws, and head to provide high contrast landmarks for digitizing. Homologous high-contrast landmarks were digitized frame-by-frame for motion analysis using Motus 9.0 motion analysis software system (Vicon, Denver, CO, USA). Digitized points were placed within spatial models and used to calculate kinematic variables.

The kinematic variables listed below were selected to determine the behavioral repertoire of prey capture, characterize the feeding mode of harbor seals, test the hypothesis that rapid jaw opening contributes to subambient pressure generation, and build upon our comparative feeding performance dataset for marine mammals [17-19]. Kinematic variables measured were: (1) maximum gape, the maximum distance from maxillary tip to mandibular tip; (2) time to maximum gape, the duration from when the lower jaw began to open until maximum gape; (3) maximum gape angle, the maximum angle from the maxillary tip to corner of the mouth to mandibular tip; (4) time to maximum gape angle; the duration from when the lower jaw began to open until maximum gape angle; (5) maximum opening gape angle velocity, the greatest angular rate of lower jaw opening; (6) time to maximal opening gape angle velocity, the duration from when the lower jaw began to open until maximum gape angle velocity was achieved; (7) maximum closing gape angle velocity, the greatest angular velocity during lower jaw closure; (8) time to maximum closing gape angle velocity, the duration from when the lower jaw began to close until maximum gape angle velocity was achieved; (9) maximum gular depression, the greatest increase in distance from the eye to external rostral border of the hyoid; and (10) time to maximum gular depression, the duration from start of gular depression to maximum gular depression. Total feeding cycle duration was also calculated.

I	Table	3.	Loadings	for	Principal	Components	Axes	1_3
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Kinematic Variable	PC 1	PC 2	PC 3
Max. Gape (cm)	0.87150	-0.26014	0.21490
Time to Max. Gape (s)	0.74441	0.45444	-0.17919
Maximum Gape Angle (°)	0.89261	-0.26282	0.25726
Time to Max. Gape Angle (s)	0.73908	0.44899	-0.17911
Max. GAOV (deg. s ⁻¹)	0.82027	-0.31931	-0.27337
Time to Max. GAOV (s)	0.38675	0.64345	-0.17341
Max. GACV (deg. s ⁻¹)	0.82532	-0.34010	0.21580
Time to GACV (s)	0.75757	0.27698	-0.20563
Max. Gular Depression (cm)	-0.24685	0.53656	0.70944
Time to Max. Gular Depression (s)	-0.36336	0.66035	0.37378
Feeding Cycle Duration (s)	0.75588	0.32745	-0.11137

Log₁₀ transformed data, N=5, 91 feeding events, GAOV=Gape Open Angle Velocity, GACV=Gape Angle Close Velocity. doi:10.1371/journal.pone.0086710.t003

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	Max. Gape	lime to Max. Gape	Max. Gape Angle	lime to Max. Gape Angle Max. GAOV	Max. GAOV	I Ime Max. GAOV	Max. GACV	IIMe Max. GACV	Max. סטומר Depress	i ime Max. Gular Depress	Feeding Cycle Duration
Max. Gape	1.0000	0.4684	0.8942	0.5192	0.7947	0.1201	0.8099	0.5620	-0.2153	-0.3877	0.5336
Time to Max. Gape	0.4684**	1.0000	0.4760	0.9460	0.4053	0.4791	0.4635	0.5501	-0.0281	-0.0694	0.6181
Max. Gape Angle	0.8942**	0.4760**	1.0000	0.4727	0.8782	0.1857	0.8785	0.5656	-0.2290	-0.3481	0.5340
Time to Max. Gape Angle	0.5192**	0.9460**	0.4727**	1.0000	0.4117	0.4986	0.4068	0.5264	-0.0191	-0.0937	0.5850
Max. GAOV	0.7947**	0.4053**	0.8782**	0.4117**	1.0000	0.1409	0.7774	0.4708	-0.1659	-0.4500	0.4634
Time Max. GAOV	0.1201	0.4791**	0.1857	0.4986**	0.1409	1.0000	0.0853	0.4346	0.1341	0.1276	0.4082
Max. GACV	0.8099**	0.4635**	0.8785**	0.4068**	0.7774**	0.0853	1.0000	0.4734	-0.2627	-0.3992	0.4468
Time Max. GACV	0.5620**	0.5501**	0.5656**	0.5264**	0.4708**	0.4346**	0.4734**	1.0000	-0.1696	-0.0494	0.8069
Max. Gular Depression	-0.2153*	-0.0281	-0.2290*	-0.0191	-0.1659	0.1341	-0.2627*	-0.1696	1.0000	0.5079	-0.0917
Time Max. Gular Depress.	-0.3877**	-0.0694	-0.3481**	-0.0937	-0.4500**	0.1276	-0.399**	-0.0494	0.5079**	1.0000	-0.0702
Feeding Cycle Duration	0.5336**	0.6181**	0.5340**	0.5850**	0.4634**	0.4082**	0.4468**	0.8069**	-0.0917	-0.0702	1.0000
Log ₁₀ transformed data, GAOV = Gape Open Angle Velocity, GACV = Gape Angle Close Velocity, $* = P < 0.05$, $** = P < 0.01$, Bold = positive correlation, Italics = negative correlation, N = 5, 91 feeding events. I doi:10.1371/journal.pone.0086710.t004	ape Open Angle Ve 1004	elocity, GACV = G	ape Angle Close V	elocity, *=P<0.0)5, **= P<0.01, F	30ld = positive c	orrelation, Italics	s = negative corr	elation, N=5, 91	feeding events.	

compared to the mean maximum biological gape and gape angle for all animals in the study. Each subject was digitally photographed while opening their mouth to their widest extent at the command of a trainer. Maximal biological gape and gape angle was measured from scaled digital photographs using ImageJ (National Institutes of Health, Bethesda, MD, USA). **Pressure Measurements** A total of 667 pressure measurements from 144 feeding events and eight subjects were recorded simultaneously with kinematic events. Whereas numerous kinematic feeding events from several individual subjects did not meet the criteria for inclusion of kinematic analyses, many more of the pressure measurements did

Maximum gape and gape angle during feeding events was

events. Whereas numerous kinematic feeding events from several individual subjects did not meet the criteria for inclusion of kinematic analyses, many more of the pressure measurements did meet the criteria for inclusion for pressure analyses. Therefore many more pressure measurements were collected than kinematic feeding events. Pressure measurements were collected using a Millar MPC-500 catheter pressure transducer connected to a transducer control box (Millar TCB-600; Houston, TX, USA) and a Biopac MP150 portable electrophysiological recording system (Biopac, Oleta, CA, USA). Pressure data were saved to a laptop using Acknowledge software (Biopac, Oleta, CA, USA). To synchronize kinematics with pressure data, we used an electronic device that generated a square wave pattern and corresponding flashing pattern of dual LED lights. The LED display was affixed to the feeding platform and recorded by the camcorder. The square wave pattern was recorded as a second channel simultaneously with pressure data collection in Acknowledge. This allowed synchronization of the suction feeding behavior with pressure measurements in Acknowledge. The pressure transducer was calibrated using the control box, but also in the laboratory under a range of known pressure regimes. Prior to each feeding trial a pressure transducer was placed through the back of a recessed cylinder (that contained food) so that the tip of the transducer, where the recording element was located, projected \sim 1 cm beyond the plexiglass feeding surface. This allowed the pressure sensor to be just at the seal's lips, or slightly within the oral cavity, during feeding. This distance was verified visually during the feeding trials and from video footage. Only suction force data in which the tip of the transducer was at this location were analyzed. In addition, only pressure measurements that exceeded $\pm 0.1 \text{ V} (\sim 5 \text{ kPa})$ in magnitude were included in the data analysis. The maximum amplitude and duration of every subambient and suprambient pressure event was measured. Prior work on bearded seal pressure generation [23] demonstrated that suction generated when feeding from the apparatus did not differ from suction values collected during hand feeding.

Statistics

Normality of data was tested using a Kolmogorov-Smirnov test. If normality was not met, then the data were \log_{10} transformed. Levene's test was used to test the assumption of homogeneity of variances. When both variance and normality requirements were met, one-way analyses of variance (ANOVAs) were performed to determine significant differences ($\alpha \leq 0.05$) of kinematic variables and pressure data during feeding trials for the categorical treatment of biting vs. suction behaviors averaged from all five subjects that met the criteria for inclusion in the statistical analyses. A Principal Component (PC) analysis on correlations was performed on the kinematic data as a tool to explore the correlation of kinematic variables. A PC analysis on correlations was used instead of PC on covariances since our variables have different units of scale, and a PC analysis on the correlation matrix is a way of standardizing such variables. Pearson's correlation

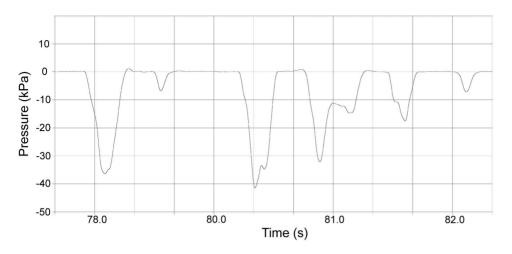


Figure 3. Pressure Traces of Subambient Pressure Data. A series of six suction events of varying magnitude. doi:10.1371/journal.pone.0086710.g003

analysis assessed the positive or negative correlation of the timing and displacement variables of feeding events. All statistical tests were conducted using JMP 9.0 (SAS Institute, Cary, North Carolina, USA).

Results

Biting vs. Suction Behavior

While all subjects used suction and biting when feeding underwater, suction was favored extensively (84% of all feeding events) compared to biting (16%). However, biting was still an important feeding mode. Food items that projected from a hole (i.e., not within a recessed cylinder) were usually consumed first using suction, but food items were also removed by grasping the food with the mouth and teeth (biting), and removing the fish from the apparatus. Food items placed within a recessed cylinder were removed using suction. These food items were more difficult to acquire and required more effort to consume. On a few occasions subjects were able to consume a recessed piece of fish by grasping and pulling on a small piece of fin with their incisors, just enough to place pursed lips on it, and remove the food using suction. Fish placed within the recessed cylinders also initiated hydraulic jetting behavior, which was often alternated with suction generation, but could also be used independently. Since this behavior is related to suction it is not consider to be one of the four feeding modes often described [30]. Hydraulic jetting was evident by water flow and turbulence appearing at the back of the recessed wells. When only a single piece of fish was left in a cylinder, alternation of suction with hydraulic jetting caused the food item to oscillate back-andforth within the recessed cylinder until it came free and was consumed. After consuming fish, subjects routinely expelled water from the corner of their mouths after a suction event, which was evident from water turbulence emanating from the caudal lips and mouth corner, as well as the bulging out of the lips and soft tissue in this region. In general, subjects became proficient at consuming all fish pieces, regardless if they were recessed or not.

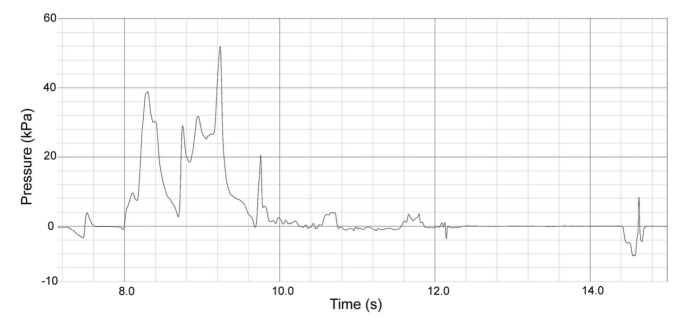


Figure 4. Pressure Traces of Suprambient Pressure Data. A series of seven hydraulic jetting events. doi:10.1371/journal.pone.0086710.g004

Suction feeding could be clearly distinguished from biting by the reduced gape from casual observation and by kinematic analyses (see below). Suction feeding was characterized by pursing of the lips to form a circular aperture, sealing of the lips to occlude lateral gape, and a cranial-to-caudal wave of motion in the gular region indicating hyolingual depression. A biting feeding mode was characterized by increased gape, lip curling with exposed teeth, and the lack of marked gular depression. The recessed cylinders of the feeding apparatus were designed to make food extraction difficult so that maximum suction and hydraulic jetting performance could be measured. Holes in the plexiglass were slightly narrower than the inner diameter of the plexiglass cylinder, which created a ridge (0.7 to 1 cm) between the cylinder and the location where seals placed their muzzle. To extract fish, enough pressure had to be employed to pull the food item up and over this ridge. In addition, numerous pieces of fish were packed into each cylinder in an effort to make extraction difficult and elicit powerful suction forces. The alternating use of suction with hydraulic jetting was successful because movement of the food items increased the chance that the food item could be lifted over the ridge. The appearance of bubbles from turbulence flowing from the back of the recessed cylinders toward the subject's lips during suction events, alternating with cloudy plumes of minute fish particles and scales exiting the back of the recessed wells during hydraulic jetting events, in addition to direct pressure measurements (see below), was further evidence that suction and hydraulic jetting were being used. Feeding trials on-land elicited only biting behavior (100%). Biting events on land involved subjects grasping projecting fish items with their teeth, removing them from the apparatus and ingesting the food. Occasionally fish were bit in half and consumed, leaving pieces of fish in the apparatus.

Vibrissal Use

Mystacial vibrissae were used during all feeding trials regardless of location. During on-land feeding trials, it appeared that subjects used vision in addition to other senses since the eyes often remained open. During in-water feeding trials, subjects always fed with their eyes closed and appeared to rely more on active touch sensation using mystacial vibrissae. Seals were observed to use the largest of the mystacial vibrissae to explore the edges of the feeding apparatus and to locate pieces of fish protruding from holes in the plexiglass surface. However, for food held within recessed cylinders, subjects would systematically locate the center of each cylinder by sweeping the plexiglass surface with their most medially located vibrissae and allow these whiskers to protrude into each cylinder. If a food item was still located within a cylinder, these whiskers could often touch a food item. This would elicit suction behavior and if the food item was difficult to remove it would then elicit hydraulic jetting behaviors, often alternating with suction behavior, until the food items(s) were removed and consumed.

Feeding Kinematics

As characterized in bearded seals [23], four feeding phases were differentiated: (I) preparatory, (II) jaw opening, (III) gular depression, and (IV) jaw closing were observed regardless of the feeding mode. Phase I began at the onset of jaw opening and ended when gape increased by greater than 0.2 cm field⁻¹ (1 field = 60 Hz) and the jaws rapidly opened. Phase II began when gape increased by \geq 0.2 cm field⁻¹ and persisted until maximum gape. Phase III began when gular depression increased by \geq 0.2 cm field⁻¹. This phase overlapped with Phases II and IV, persisted the longest in duration, and concluded when gular depression returned to its original position, which was often at the

end of the feeding event. Phase IV began at maximum gape and concluded when the jaws closed. The timing of maximum gular depression during suction feeding events always followed maximum gape or coincided with maximum gape. Little to no gular depression was observed during biting feeding events, whether seals were feeding in-water or on-land. The mean durations for Phases I–IV were, 0.02 ± 0.001 s, 0.08 ± 0.04 s, 0.16 ± 0.10 s, and 0.11 ± 0.07 s, respectively.

The mean feeding cycle duration for suction events was significantly shorter $(0.15 \pm 0.09 \text{ s S.E.}; P < 0.01)$ than the mean feeding cycle duration during biting feeding events $(0.18\pm0.08 \text{ s};$ Fig. 2, Table 2). This reflected divergent kinematic profiles between suction and biting feeding events. Suction was characterized by a significantly smaller gape $(1.3\pm0.23 \text{ cm}; P<0.001)$ and gape angle $(12.9\pm2.02^\circ; P<0.001)$ compared to biting events, which was characterized by a large gape (3.630±0.21 cm) and gape angle $(28.8 \pm 1.80^{\circ}; P < 0.001)$. Gape and gape angle during suction or biting events was significantly less than maximum biological gape and gape angle (P<0.001). Mean maximum biological gape and gape angle from all seals that participated in the study was 10.1 ± 0.80 cm and $69.7\pm2.66^{\circ}$, respectively. Time to maximum gape and gape angle were significantly shorter during suction events $(0.07\pm0.01 \text{ s each respectively}; P<0.01 \text{ and}$ P < 0.02, respectively) compared to biting events (0.08 ± 0.01 s each respectively). Maximal gape angle opening velocity (GAOV) was significantly slower during suction events (197.2±42.04°s⁻ P < 0.001) than during biting events (561.6±37.61°s⁻¹). However, time to GAOV for suction (0.05 ± 0.01) vs. biting (0.04 ± 0.01) events was not significantly different. Maximal gape closing angle velocity (GACV) was significantly slower during suction events P<0.001) $(159.1 \pm 42.05^{\circ}s^{-1})$ than biting events $(544.2\pm37.61^{\circ}s^{-1})$; time to GACV was significantly longer for suction $(0.11\pm0.01 \text{ s}; P<0.001)$ compared to biting events (0.13 ± 0.01) . Gular depression was significantly greater, and time to maximum gular depression was significantly longer, during suction feeding events $(1.4\pm0.08 \text{ cm}, 0.14\pm0.01 \text{ s}, \text{ respectively};)$ P < 0.001) compared to biting events (0.9±0.07 cm, 0.07±0.01 s, respectively). Figure 2 depicts representative kinematic profiles for suction and biting; kinematic variables of biting and suction feeding are summarized in Table 2.

Principal component (PC) analysis on correlations of logtransformed data demonstrated that the first 3 PC axes characterized 78% of the variation of harbor seal feeding kinematics (PC1 = 49.986%, PC2 = 18.949%, PC3 = 9.072%; Table 3). High loadings on PC axis 1 identified most kinematic variables, with the exception of maximum gular depression and time to maximum gular depression. PC axis 2 identified all timing kinematic variables with the exception of time to maximum gape angle closing velocity. In addition PC axis 2 had a high loadings\ for maximum gular depression. PC axis 3 had high loadings for only maximum gular depression and time to maximum gular depression, indicating distinct differences in suction vs. biting feeding events. Pearson's correlation analysis further supported the difference between suction and biting feeding kinematics and is summarized in Table 4. As shown by the PC analysis, the Pearson's correlation analysis demonstrated that most, but not all, kinematic variables were positively correlated. However, gular depression and time to gular depression were distinct in that they were negatively correlated with most kinematic variables.

Suction and Hydraulic Jetting Pressures

Direct measurement of sub- and suprambient pressure generation during feeding events supported the observational data that harbor seals primarily used suction (subambient pressure generation) when feeding in-water. However, hydraulic jetting (suprambient pressure generation) was also employed. Frequency data from pressure traces show that subjects in this study employed suction 90.5% of the time and hydraulic jetting 9.5% of the time. Pressure data demonstrate that suction events were composed of an expansive phase, during which the maximum subambient pressure was reached for that event, followed by a compressive phase, during which pressure returned to baseline levels (Fig. 3). A preparatory phase was not observed. Similarly, pressure data from hydraulic jetting (Fig. 4) demonstrated only an expansive phase during which maximum suprambient pressure was recorded, followed by a compressive phase, during which pressure values returned to baseline. As with subambient pressure traces, a preparatory phase was never observed. The maximum subambient pressure recorded was 48.6 kPa and the maximum suprambient pressure recorded was 53.9 kPa. The mean duration of subambient pressure events $(0.5\pm0.3 \text{ ms})$ was significantly longer (P<0.001) than the mean duration of suprambient pressure events $(0.35 \pm 0.25 \text{ ms}).$

Discussion

This study provides the first detailed kinematic and physiological data of feeding performance in harbor seals. The data support our hypothesis that suction is the primary underwater feeding mode in harbor seals. However, biting is also an important feeding mode, and likely more important than for suction feeding specialists such as bearded seals and walruses. The primary finding of this study was that suction was used predominantly underwater, but subjects were also capable of using hydraulic jetting either alone or in conjunction with suction feeding. The combined use of suction and hydraulic jetting for feeding has been documented previously for bearded seals [17], walruses [13,49] and leopard seals [48]. Walruses excavate bivalves using hydraulic jetting to clear the sediment and then employ suction to remove all or part of the bivalve [13,66]. The use of suction and hydraulic jetting have also been observed in cetacean suction feeding specialists such as pygmy and dwarf sperm whales [19], belugas [18] and pilot whales [15,18]. All of these species, including harbor seals (this study), used hydraulic jetting either alone or alternating with suction feeding. The behavior of hydraulic jetting is likely similar to that of suction generation, but where the tongue and hyoid are elevated, instead of retracted, to produce suprambient pressures rather than subambient pressures. Although similar behaviorally, the underlying mechanisms (i.e., muscles of tongue retraction vs. protraction) may differ. However whether this is manifested in sub- vs suprambient pressure performance differences currently remains unclear. In this study, and in others [17,18], maximum subambient and maximum suprambient pressures were similar in magnitude (48.6 kPa and 53.9 kPa, respectively). Harbor seals demonstrated several feeding mechanisms and modes that were used to capture prey.

The hypothesis that rapid jaw opening is correlated with suction feeding in harbor seals was not strongly supported in this study. Although harbor seals employed several of the same mechanisms to generate subambient pressure that are employed by other suction feeding specialists (closure of lateral gape, pursing of lips to form a circular aperture, and rapid hyolingual depression [13,17,48,49], they did not employ a preparatory phase for suction that likely increases the change in intraoral volume that is related to greater subambient pressure development as reported for bearded seals [17]. This is likely related to the lack of a vaulted palate, observed in other pinnipeds specialized for suction [13,17,51,53,55] in harbor seals. Nor was rapid jaw depression

(as measured by GAOV), which can also contribute to subambient pressure generation via further buccal expansion in conjunction with hyolingual depression, high compared to other marine mammal suction feeders, such as pygmy sperm whales (GAOV = 293 deg s⁻¹) [19] and bearded seals (GAOV = 205 deg s⁻¹) [17]. However, it is becoming apparent that the underlying mechanism of suction feeding is not homogeneous among all species for which data are available, and not all suction feeding specialists employ rapid jaw depression (i.e., belugas, GAO-V = 119°s⁻¹ [18]). As a result, the data in this study show that although harbor seals can create significant subambient pressures, their suction and hydraulic jetting performance are not at the same level for pinniped suction feeding specialists such as bearded seals (91.2 kPA [13]) and walruses (91.2 kPa [25] and 118 kPa [49]).

Harbor seals in this study used the large lateral mystacial vibrissae for exploration of the large scale features of the feeding apparatus, but then shifted to using the small medially located mystacial vibrissae for more refined and discrete tactile exploration. In particular, these small medially located mystacial vibrissae were used to locate the center of each cylinder and to protrude into each cylinder to touch recessed food items, if possible. Our observation that harbor seals use different regions of the mystacial vibrissae during feeding supports the results of more focused active touch performance studies [67,68] in which harbor seals used the smaller medial mystacial vibrissae for detailed size discrimination, but within a different context. This active touch exploratory pattern is likely a typical pattern of how harbor seals explore new objects in their environment. Such exploratory behavior and use of different regions and size of mystacial vibrissae has also been observed in California sea lions [69], manatees and dugongs [45,47], walruses [70], and rodents [71–73] and likely represents a generalized mammalian pattern of tactile exploration.

Data from this study show that harbor seals have a wide repertoire of feeding strategies that include biting, suction, and hydraulic jetting. Behavioral observations also showed that harbor seals were flexible and creative in extracting food items from the feeding apparatus. Biting in-water appeared to involve smaller gapes than biting on-land, but this currently remains unclear and needs to be investigated further. Such flexibility of feeding strategies and biomechanics likely form the basis of their opportunistic, generalized feeding ecology and concomitant breadth of diet. Although there were many similarities in the kinematic profile of harbor seals compared to bearded seals, harbor seals lacked certain behaviors (i.e., lack of preparatory phase prior to suction feeding and slower jaw depression) that could have increased their subambient pressure generation. However, the subambient pressures produced were still significant and are likely strong enough to be an important feeding mode for harbor seals. In addition, like bearded seals, it is probable that harbor seals can use the substrate and the geometry of the habitat to passively increase their suction, and possibly their hydraulic jetting, capability. The positive effect of the substrate to passively increase the suction capability of foragers has been documented in both chondrichthyan and teleost fishes [74–76] and is the result of the conservation of momentum of water flow [76]. This strategy likely has important trophic implications, such as increasing preycapture efficiency, for any suction feeding marine mammal, as it does for benthic foraging fishes.

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References

- Lauder GV (1985) Functional morphology of the feeding mechanism in lower vertebrates. In: Duncker HR, Fleischer G, editors. Functional Morphology of Vertebrates. New York: Springer-Verlag. pp. 179–188.
- Lauder GV, Shaffer HB (1993) Design of feeding systems in aquatic vertebrates: major patterns and their evolutionary interpretations. In: Hanken J, Hall Bk, editors. The Skull, Vol. 3: Chicago Chicago Press. pp. 113–149.
- Higham TE, Day SW, Wainwright PC (2005) Multidimensional analysis of suction feeding performance in fishes: fluid speed, acceleration, strike accuracy and the ingested volume of water. J Exp Biol 209: 2713–2725.
- Wilga CD, Motta PJ, Sanford CP (2007) Evolution and ecology of feeding in elasmobranchs. Integr Comp Biol 47: 55–69.
- Wainwright PC, Day SW (2007) The forces exerted by aquatic suction feeders on their prey. J R Soc Interface 4: 553–560.
- Berta A, Sumich JL, Kovacs KM (2006) Marine Mammals: Evolutionary Biology. 2nd Edition. San Diego: Academic Press. 560 pp.
- Lauder GV (1980) The suction feeding mechanism in sunfishes (Lepomis): an experimental analysis. J Exp Biol 88: 49–72.
- Muller M, Osse JWM, Verhage JHG (1982) A quantitative hydrodynamical model of suction feeding in fish. J Theor Biol 95: 49–79.
- Lauder GV (1985) Aquatic feeding in lower vertebrates. In: Hildebrand M, Bramble DM, Liem KF, Wake DB, editors. Functional vertebrate morphology. Cambridge: Harvard University Press, USA. pp. 210–229.
- Deban SM, Wake DB (2000) Aquatic feeding in salamanders. In: Schwenk, K, editor. Feeding: form, function, and evolution in tetrapods. San Diego: Academic Press. pp. 65–94.
- Wainwright P, Carroll AM, Collar DC, Day SW, Higham TE, et al (2007) Suction feeding mechanics, performance, and diversity in fishes. Integr Comp Biol 47: 96–106.
- Gordon KR (1984) Models of tongue movement in the walrus (Odobenus rosmarus). J Morph 182: 179–196.
- Fay FH (1982) Ecology and biology of the Pacific walrus, Odobenus rosmarus divergens Illiger. Washington D.C.: Fish & Wildl. Serv. Publ. No. 74, 279 pp.
- Kastelein RA, Staal C, Terlouw A, Muller M (1997) Pressure change in the mouth of a feeding harbor porpoise (*Phocoena phocoena*). In: Read AJ, Wiepkema PR, Nachtigall PE, editors. The biology of the harbor porpoise. Woerden, The Netherlands:DeSpil Publishers. pp. 279–291.
- Werth AJ (2000) A kinematic study of suction feeding and associated behavior in the long-finned pilot whale, *Globicephala melas* (Traill) Mar Mamm Sci 16: 299– 314.
- Bloodworth B, Marshall CD (2007) A functional comparison of the hyolingual complex in pygmy and dwarf sperm whales (*Kogia breviceps & K. sima*), and bottlenose dolphins (*Tursiops truncatus*). J Anat 211: 78–91.
- Marshall CD, Kovacs K, Lydersen C (2008) Feeding Kinematics, Suction, and Hydraulic Jetting Capabilities in Bearded Seals (*Erignathus barbatus*). J Exp Biol 211: 699–708.
- Kane EA, Marshall CD 2009. Comparative feeding kinematics and performance of odontocetes: belugas, Pacific white-sided dolphins, and long-finned pilot whales. J Exp Biol 212: 3939–3950.
- Bloodworth B, Marshall CD (2005) Feeding kinematics of Kogia and Tursiops (Odonticeti: Cetacea): Characterization of suction and ram feeding. J Exp Biol 208: 3721–3730.
- Werth AJ (2007) Adaptations of the cetacean hyolingual apparatus for aquatic feeding and thermoregulation. Anat. Rec 290: 546–568.
- Heyning JE, Mead JG (1996) Suction feeding in beaked whales: morphological and observational evidence. Nat Hist Mus LA County Contrib Sci 464: 1–12.
- Reidenberg JS, Laitman JT (1994) Anatomy of the hyoid apparatus in Odontoceti (toothed whales): specializations of their skeleton and musculature compared with those of terrestrial mammals. Anat Rec 240: 598–624.
- Werth AJ (2000b) Marine mammals. In: Schwenk K, editor. Feeding: form, function, and evolution in tetrapods. San Diego: Academic Press. pp. 475–514.
- 24. Werth AJ (2006a) Mandibular and dental variation and the evolution of suction feeding in Odontoceti, J Mammal 87: 579–588.
- Werth AJ (2006b) Odontocete suction feeding: experimental analysis of water flow and head shape. J Morphol 267: 1415–1428.
- King JE (1972) Observations on phocid skulls. In: Functional Anatomy of Marine Mammals, Vol 1. R.J Harrison, ed. London: Academic Press. pp. 81– 115.
- Kastelein RA, Gerrits NM (1990) The anatomy of the Walrus head (Odobenus rosmarus). Part 1: The skull. Aq Mamm 16: 101–119.
- Kastelein RA, Gerrits NM, Dubbeldam JL (1991) The anatomy of the Walrus head (*Odobenus rosmarus*) Part 2: Description of the muscles and of their role in feeding and haul-out behavior. Aq Mamm 17: 156–180.

Author Contributions

Conceived and designed the experiments: CDM. Performed the experiments: CDM SW WH FH GD. Analyzed the data: CDM AM BK. Contributed reagents/materials/analysis tools: CDM GD. Wrote the paper: CDM.

- Marshall CD, Kovacs K, Lydersen C (2001) Morphology of the Bearded Seal (*Erignathus barbatus*) Muscular-Vibrissal Complex. 14th Biennial Conference on the Biology of Marine Mammals.
- Lauder GV, Shaffer HB (1986) Functional design of the feeding mechanism in lower vertebrates: unidirection and bidirectional flow systems in the tiger salamander. J Linn Soc (Zool.) 88: 277–290.
- Lauder GV, Reilly SM (1988) Functional design of the feeding mechanism in salamander: causal bases of ontogenetic changes in function. J Exp Biol 134: 219–233.
- Lauder GV (1979) Feeding mechanisms in primitive teleosts and in the halecomorph fish *Amia calva*. J Zool (Lond) 187: 543–578.
- Motta PJ, Hueter RE, Tricas TC, Summers AP (2002) Kinematic analysis of suction feeding in the nurse shark, *Ginglymostoma cirratum* (Orectolobiformes, Ginglymostomatidae). Copeia 1001: 24–38.
- Matott MP, Motta PJ, Hueter RE (2005) Modulation in feeding kinematics and motor pattern of the nurse shark *Ginglymostoma cirratum*. Environ Biol Fishes 74: 163–174.
- Emlin JM (1966) The role of time and energy in food preference. Am Nat 100: 611–617.
- Schoener TW (1971) Theory of feeding strategies. Ann Rev Ecol Syst 2: 369– 404.
- Bowen WD, Tully D, Bones DJ, Bulheier BM, Marshall GJ (2002) Prey dependent foraging tactics and prey profitability in a marine mammal. Mar Ecol Progr Ser 244: 235–245.
- Marshall GJ (1998) Crittercam: an animal-borne imaging and data logging system. Mar Tech Soc J 32: 11–17.
- Davis RW, Fuiman LA, Williams TM, Collier SO, Hagey WP, et al (1999) Hunting behavior of a marine mammal beneath the Antarctic fast ice. Science 283: 993–996.
- Parrish FA, Craig MP, Ragen TJ, Marshall GJ, Buhleier BM (2000) Identifying diurnal foraging habitat of endangered Hawaiian monk seals using a sealmounted video camera. Mar Mamm Sci 16: 392–412.
- Parrish FA, Abernathy K, Marshall GJ, Buhleier BM (2002) Hawaiian monk seals (*Monachus schauinslandi*) foraging in deep-water coral beds. Mar Mamm Sci 18: 244–258.
- Ponganis PJ, Van Dam RP, Marshall G, Knower T, Levenson DH (2000) Subice foraging behavior of emperor penguins. J Exp Biol 203: 3275–3278.
- Davis RW, Fuiman LA, Williams TM, Collier SO, Horning M, et al (2003) Classification of Weddell seal dives based on 3-dimensional movements and video-recorded observations. Mar Ecol Progr Ser 264: 109–122.
- 44. Mitani Y, Sato K, Ito S, Ceron MF, Siniff DB, et al (2003) A methods fore reconstructing three-dimensional dive profiles of marine mammals using geomagnetic intensity data: results from two lactating Weddell seals. Polar Biol 26: 311–317.
- Marshall CD, Huth GD, Edmonds VM, Halin DL, Reep RL (1998) Prehensile use of perioral bristles during feeding and associated behaviors of the Florida manatee (*Trichechus manatus latirostris*). Mar Mamm Sci 14: 274–289.
- Marshall CD, Huth GD, Edmonds VM, Halin DL, Reep RL (2000) Foodhandling ability and feeding-cycle length of manatees feeding on several species of aquatic plants. J Mamm 81: 649–658.
- Marshall CD, Maeda H, Iwata M, Furuta M, Asano A, Rosas F, et al (2003) Orofacial morphology and feeding behaviour of the dugong, Amazonian, West African and Antillean manatees (Mammalia: Sirenia): functional morphology of the muscular – vibrissal complex. J Zool 259: 1–16.
- Hocking DP, Evan AR, Fitzgerald EMG (2013) Leopard seals (*Hydrurga leptonyx*) use suction and filter feeding when hunting small prey underwater. Polar Biol 36: 211–222.
- Kastelein RA, Muller M, Terlouw A (1994) Oral suction of a Pacific walrus (Odobenus rosmarus divergens) in air and under water. Zeitsch Säugetierk 59: 105– 115.
- Klages NTW, Cockcroft VG (1990) Feeding behaviour of a captive crabeater seal. Polar Biol 10: 403–404.
- King JE (1983) Seals of the world, 3rd ed. Ithaca :Cornell University Press. 240 pp.
- Marshall CD (2009). Feeding Functional Morphology. In: Perrin WF, Würsig B, Thewissen JGM, editors. Encyclopedia of Marine Mammals San Diego: Academic Press. pp. 406–413.
- Adam PJ, Berta A (2002) Evolution of prey capture strategies and diet in the Pinnipedimorpha (Mammalia, Carnivora). Oryctos 4: 83–107.
- Rice DW (1998) Marine Mammals of the World: Systematics and distribution. Special Publication No. 4. The Society for Marine Mammalogy. 231 pp.

- Burns JJ (2009) Harbor and spotted seal. In: Perrin WF, Würsig B, Thewissen JGM, editors. Encyclopedia of Marine Mammals. San Diego: Academic Press. pp. 533–542.
- Bigg MA (1981) Harbour seal, *Phoca vitulina* Linnaeus, 1758 and *Phoca largha* Pallas, 1811. In: Ridgway SH, Harrison RJ, editors. Handbook of Marine Mammals, Vol. 2, Seals. London: Academic Press. pp. 1–27.
- Hoover AA (1988) Harbor seal. In: Lentfer JW, editor. Selected Marine Mammal of Alaska: Species Accounts with Research and Management Recommendations. Washington DC: Marine Mammal Commission. pp. 125– 157.
- Sharples RJ, Arrizabalaga B, Hammond PS (2009) Seals, sandeels and salmon: diet of harbour seals in St. Andrews Bay and the Bay Estuary, southeast Scotland. Mar Ecol Prog Ser 390: 265–276.
- Kavanagh AS, Cronin MA, Walton M, Rogan E (2010) Diet of the harbour seal (*Phoca vituina*) in the west and south-west of Ireland. J Mar Biol Assoc UK 90: 15– 17–1527.
- Thomas AC, Lance MM, Jeffries SJ, Miner BG, Acevedo-Gutiérrez A (2011) Harbor seal foraging response to a seasonal resource pulse, spawning Pacific herring. Mar Ecol Prog Ser 441: 225–239.
- Brown SL, Bearhop S, Harrod C, McDonald RA (2012) A review of spatial and temporal variation in grey and common seal diet in the United Kingdom and Ireland. J Mar Biol Assoc UK 92: 1711–1722.
- 62. Bromaghin JF, Lance MM, Elliott EW, Jeffries SJ, Acevedo-Gutiérrez A, et al (2013) New insights into the diets of harbor seals (*Phoca vitulina*) in the Salish Sea revealed by analysis of fatty acid signatures. Fish Bull 111: 13–26.
- Peterson SH, Lance MM, Jeffries SJ, Acevedo-Gutiérrez A (2012) Long Distance Movements and Disjunction Spatial Use of Harbor Seals (*Phoca vitulina*) in the Inland Waters of the Pacific Northwest. PLoS ONE: e39046. doi:10.1371/ journal.pone.0039046.
- Eguchi T, Harvey JT (2005) Diving behavior of the Pacific harbor seal (*Phoca vitulina richardii*) in Monterey Bay, California. Mar Mamm Sci 21: 283–295.

- Germain LR, McCarthy MD, Koch PL, Harvey JT (2012) Stable carbon and nitrogen isotopes in multiple tissues of wild and captive harbor seals (*Phoca* vitulina) off the California coast. Mar Mamm Sci 28: 542–560.
- Kastelein RA, Mosterd P (1989) The excavation technique for molluscs of Pacific Walrusses (*Odobenus rosmarus divergens*) under controlled conditions. Aquat Mamm 15: 3–5.
- Dehnhardt G, Kaminski A (1995) Sensitivity of the mystacial vibrissae of harbor seals (*Phoca vitulina*) for size difference of actively touched objects. J Exp Biol 198: 2317–2323.
- Grant R, Wieskotten S, Wengst N, Prescott T, Dehnhardt G (2013) Vibrissal touch sensing in the harbor seal (*Phoca vitulina*): how do seals judge size? J Comp Physiol A 199: 521–533.
- 69. Dehnhardt G, Dücker G (1996) Tactual discrimination of size and shape by a California sea lion (Zalophus californianus) An Learn Behav 24: 366–374.
- Kastelein RA, van Gaalen MA (1988) The sensitivity of the vibrissae of a pacific walrus (Odobenus rosmarus). Aq Mamm 14: 123–133.
- Brecht M, Preilowski B, Merzenich MM (1997) Functional architecture of the mystacial vibrissae. Behav Brain Res 84: 81–97.
- Hartmann M (2001) Active sensing capabilities of the rat whisker system. Auton Robots 11: 249–254.
- Grant RA, Sperber A, Prescott TJ (2012) The role of orienting in vibrissal touch sensing. Frontiers Behav Neurosci 6: 39.
- Carroll AM, Wainwright PC, Huskey SH, Collar DC, Turingan RG (2004) Morphology predicts suction feeding performance in centrarchid fishes. J Exp Biol 207: 3873–3881.
- Gibb AC, Ferry-Graham L (2005) Cranial movements during suction feeding in teleost fishes: Are they modified to enhance suction production? Zool 108: 141– 153.
- Nauwelaerts S, Wilga CD, Sanford CP, Lauder GV (2007) Substrate passively improves suction feeding in benthic sharks. Published online Nov. 28, 2006. J Roy Soc Interface 2: 341–345.
- 77. American Society of Mammalogists Committee on Marine Mammals (1967) Standard Measurements of Seals. J Mamm 48: 459–462.