

RESEARCH ARTICLE

How harbour seals (*Phoca vitulina*) encode goals relative to landmarks

Eric Maaß and Frederike D. Hanke*

ABSTRACT

Visual landmarks are defined as objects with prominent shape or size that distinguish themselves from the background. With the help of landmarks, animals can orient themselves in their natural environment. Yet, the way in which landmarks are perceived and encoded has previously only been described in insects, fish, birds, reptiles and terrestrial mammals. The present study aimed to provide insight into how a marine mammal, the harbour seal, encodes goals relative to landmarks. In our expansion test, three harbour seals were trained to find a goal inside an array of landmarks. After diagonal, horizontal or vertical expansion of the landmark array, the search behaviour displayed by the animals was documented and analyzed regarding the underlying encoding strategy. The harbour seals mainly encoded directional vector information from landmarks and did neither search arbitrarily around a landmark nor used a rule-based approach. Depending on the number of landmarks available within the array, the search behaviour of some harbor seals changed, indicating flexibility in landmark-based search. Our results present the first insight into how a semi-aquatic predator could encode landmark information when swimming along the coastline in search of a goal location.

KEY WORDS: Spatial strategies, Navigation, Orientation, Expansion test, Marine mammal

INTRODUCTION

In all moving organisms, the need to remember the locations of foraging sites, sleeping grounds, nests and even items not immediately visible in the environment is vital. Under many circumstances, organisms are guided to goal locations by landmarks. Landmarks are defined as objects with specific characteristics such as a prominent shape or size that clearly contrast from the background (Yesiltepe et al., 2021). The use of visual landmarks for goal localization has been documented in multiple species including bees (Cartwright and Collett, 1983), birds (e.g. Cheng, 1989; Cheng and Sherry, 1992; Spetch, 1995), fish (Burt and Macias Garcia, 2003), dogs (Fiset, 2007), rodents (e.g. Cook and Tauro, 1999), turtles (Lopez et al., 2001) and several non-human primates (e.g. MacDonald et al., 2004; Marsh et al., 2011). In contrast, the role of landmarks and landmark orientation has not yet been experimentally studied in marine mammals, although the behaviour of wild animals has already been assumed to be based on landmarks. Matsumura et al. (2011) speculated that

wild elephant seals close to the coast were guided by landmarks in the final phase of migrating back to their natal beach. Grey seals crossing the channel switched their navigational strategy when they were reaching familiar areas close to the coast in which local cues such as landmarks might have guided their journeys (Chevaillier et al., 2014).

In this study, we aimed at describing whether and how the position of a goal is memorized in respect to landmarks by a marine mammal, the harbour seal. Harbour seals that commute between the coast and the open ocean appear to be very suitable subjects for assessing the role of landmarks, as previous studies revealed an extraordinary ability to return to previous haul-out places along the coast after foraging in deeper waters (Brown and Mate, 1983; Stewart, 1984; Suryan and Harvey, 1998; Steingass et al., 2019; Vance et al., 2021). While in the open ocean, landmarks may not be continuously available; however, the coastline offers many landmarks, such as rock formations, sandbanks and anthropogenic structures, that can be used for orientation or specifically for homing.

We used a classic experimental approach to study the use of landmarks and the underlying strategies in harbour seals, the expansion test. This experimental paradigm was previously established including numerous animals (Wehner and Räber, 1979; Cartwright and Collett, 1983; Spetch et al., 1996; Spetch et al., 1997; MacDonald et al., 2004; Poti et al., 2005, 2010; Marsh et al., 2011). First, the subject is trained to locate a hidden item or goal within an array of landmarks. Afterwards, the array is then expanded, meaning that the distances between the landmarks are modified. The geometrical relationship may remain constant in some but may change in other expansion schemes. The peak search areas of the animals are subsequently analysed to unravel the underlying strategy (Marsh et al., 2011).


There are at least three different strategies describing how a landmark is used for orientation and navigation (Marsh et al., 2011). In the first strategy, landmarks can serve as beacons (Fig. 1); thus, the organisms search for a goal near an individual landmark in an undirected way. This beacon strategy was described for rats (Cook and Tauro, 1999), turtles (Lopez et al., 2001), monkeys (Poti et al., 2005) and human children (MacDonald et al., 2004).

Another group of animals seems to encode distance and direction between a goal and one or multiple landmarks (Fig. 1). This second strategy has been called the directional vector strategy, and it is defined as averaging of familiar directional vectors between a goal and a landmark (Cheng, 1989; Cheng et al., 2006). It can be differentiated from the undirected search of a beacon strategy in that the animals search in relation to a single landmark, but they combine multiple landmarks or the entire landmark array to determine the direction and length of the vector (Marsh et al., 2011). This type of landmark use has been documented in gerbils (Collett et al., 1986), pigeons (Spetch et al., 1996, 1997) and primates (Poti et al., 2005, 2010).

Those organisms that apply a third strategy, the rule-based strategy, operate with the configuration of an entire array of landmarks and

University of Rostock, Institute for Biosciences, Neuroethology, Albert-Einstein-Str. 3, 18059 Rostock, Germany.

*Author for correspondence (frederike.hanke@uni-rostock.de)

 F.D.H., 0000-0002-1737-3861

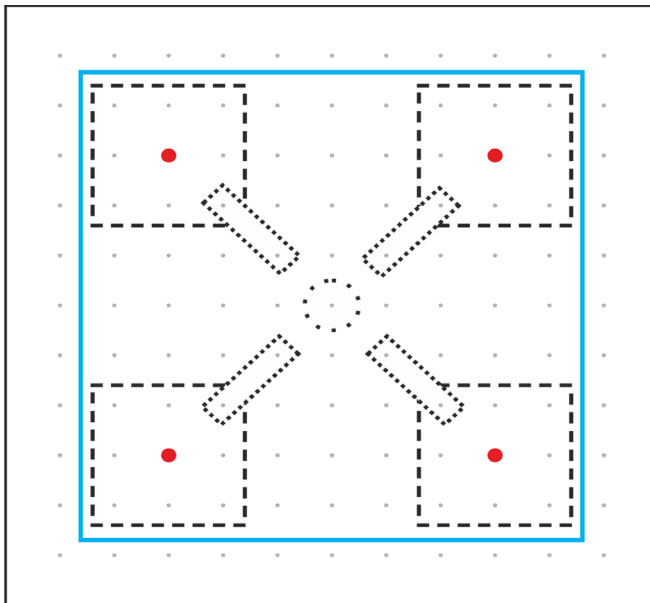


Fig. 1. Search areas predicted by a beacon strategy (squared dashed lines), a rule-based strategy (dotted circle) and a directional vector strategy (dotted rectangles) in an expansion test. Red dots represent the landmarks within a four-landmark array, and the blue line indicates the landmark boundary area (modified after Marsh et al., 2011; Poti et al., 2010).

encode the position of the goal in relation to all available landmarks (Fig. 1). So far, only adult humans have been documented to use this strategy ad hoc (MacDonald et al., 2004). Studies on landmark use in bees indicated that the responses of bees also followed a rule-based approach. However, their search behaviour might also be explained by comparing a 2D snapshot of the landmarks with images stored in memory (Cartwright and Collett, 1983). Interestingly, some birds and primates, among others, seem to be able to learn this strategy when trained in paradigms that forced the animals to rely on the configuration of the array (Jones et al., 2002; Poti et al., 2010).

In our study, we designed an expansion experiment to unravel the strategy of landmark use by harbour seals by first using an array of four landmarks (experiment 1). Subsequently, in experiment 2, we reduced the number of landmarks within the array to two landmarks and ultimately to a single landmark to determine whether the seals' strategy would change with less goal-defining information available.

MATERIALS AND METHODS

Experimental animals

The experiment was conducted with three adult male harbour seals (*Phoca vitulina* Linnaeus 1758) named 'Nick' (21 years old; length: 173 cm; mean mass: 121 kg), 'Filou' (14 years old; length: 165 cm; mean mass: 109 kg) and 'Moe' (14 years old; length: 151 cm; mean mass: 91 kg) at the Marine Science Center of the University of Rostock, Germany. All seals had previously participated in numerous different scientific experiments (e.g. Kowalewsky et al., 2006, Schulte-Pelkum et al., 2007, Byl et al., 2016, Niesterok et al., 2017, Krüger et al., 2018, Maaß and Hanke, 2021). They were housed with nine other harbor seals, two California sea lions (*Zalophus californianus*) and a South African fur seal (*Arctocephalus pusillus pusillus*) in a seawater enclosure. The seals were mainly fed freshly thawed cut herring (*Clupea harengus*) and sprats (*Sprattus sprattus*). During the experiment and the general training, the animals received 1–5 kg of fish a day depending on season and motivation, meaning eagerness to participate during training and during experiments. We performed experiments 3 to 4 days a week. The experiment took place in an enclosure (7×12 m) separated from the main enclosure.

The experiments carried out in this study were in accordance with the European Communities Council Directive of 22 September 2010 (2010/63/EU) and the German Animal Welfare Act of 2006. The individuals used in the study were not subject to pain, suffering or injury; therefore, no approval or notification was required.

Experimental setup

The experimental setup consisted of a ring station (Fig. 2A) that served as the starting point for the animal in each trial. This station

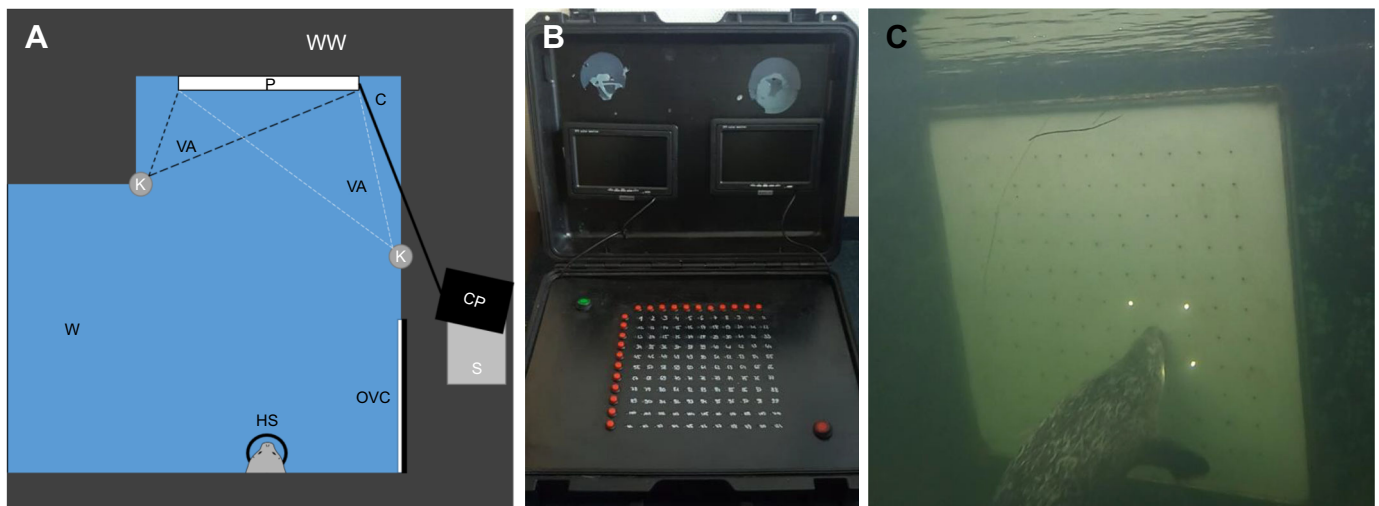


Fig. 2. Experimental setup. (A) Schematic top view of the experimental basin, in which the experiment took place, with the walkway (WW) and the water area (W). The experimenter sat on the walkway (position S) behind an opaque visual cover (OVC) to avoid secondary cueing and set the landmark array with the help of the control panel (CP) connected to the submerged LED panel (P) with a cable (C). At the beginning of a trial, the seal was stationing in a hoop station (HS). Upon a signal, it swam towards the submerged LED panel (P), indicating with its snout where it assumed the goal was. The LED panel was within the viewing angle (VA) of three cameras mounted on two mountings (K; two cameras on the right-hand side) which allowed to oversee the response behaviour of the seal at the panel as well as to control stimulus presentation. (B) Control panel with which the specific LED landmark array could be set on the submerged LED panel from a distance. (C) Submerged LED panel in a training situation with seal Nick giving a response at the goal location.

was positioned opposite to a 2×2 m integral foam wall (Figs 2A,C and 3) with integrated LED lights, which served to present the stimuli. The wall was fully submerged with the upper rim 20 cm below the water surface. In total, 121 LED lamps (Luckylight, Shenzhen, China; Ø 10 mm, 8000 mcd, cold white, radiation angle 20 deg) were inserted in the wall in 11 rows and 11 columns (Figs 2C and 3). The LEDs were 15 cm apart from each other; the outermost LEDs were 25 cm apart from the aluminium frame surrounding the wall. Every LED was connected to a control panel (Fig. 2B) installed at a distance of 5 m to the wall. The control panel served as a miniature version of the LED wall equipped with 22 light switches, which allowed control of the LEDs from afar. Two cameras (Eyoyo 1000 TVL Waterproof Camera, Eyoyo Shenzhen, Guangdong, China) on aluminium mountings were placed to the left (2 m away) and right (3 m away) of the LED wall and served to observe the animals' performances, displayed on two LCD monitors during the experiment. A third camera (GoPro Hero 7 Black Edition, GoPro, San Mateo, CA, USA) on the right aluminium mounting recorded the experiment for later analysis. During the experimental sessions, the experimenter hid behind an opaque visual cover to avoid secondary cueing. The influence of secondary cues from the experimenter was additionally prohibited as the seal swam away from the experimenter when indicating its response at the LED wall.

General experimental procedure

After entering the enclosure, the animal was asked to swim to and rest in its ring station. At the same time, the experimenter hid behind the opaque visual cover next to the control panel. After the experimenter had switched on the specific landmark array of the respective trial, the seal was indicated to leave its station by a short

whistle and had to approach the wall to indicate its response by touching the position where it assumed the goal was with its snout (Fig. 2C). After every correct response, the animal received up to three (pieces of) fish from the experimenter. An incorrect response was answered by the German word for no ('nein'), and no reward was given. After the feedback, the animal had to swim back to its station for a new trial to begin. The duration of the inter-trial interval was approximately 60–90 s.

Experiment 1

Stimulus

The stimulus presented was an LED array consisting of four lit LEDs (Figs 1–3). The task for the animal was to find the goal in the middle of the array. For each trial, the LED array configuration was varied in its absolute position on the LED wall following a pre-set schedule. A total of 81 target locations could be chosen for each trial. In order to systematically vary the position of the LED array, we divided the wall into four quadrants and an overlapping area (Fig. 3). During a session, the LED array was placed four (during testing) or six (in training) times in each quadrant and the overlapping area, resulting in a session of 20 or 30 trials, respectively.

Pre-training

Pre-training started with the animal swimming from its ring station towards the panel touching a target held at the goal location by an assistant from above the array. Over the course of pre-training, the response target was successively reduced in size. During these familiarization trials, a correct answer was defined as the animal swimming to the target ball and touching it with its snout for 3 s. After successfully completing 10 correct trials per target in succession, trials without an assistant and target ball were

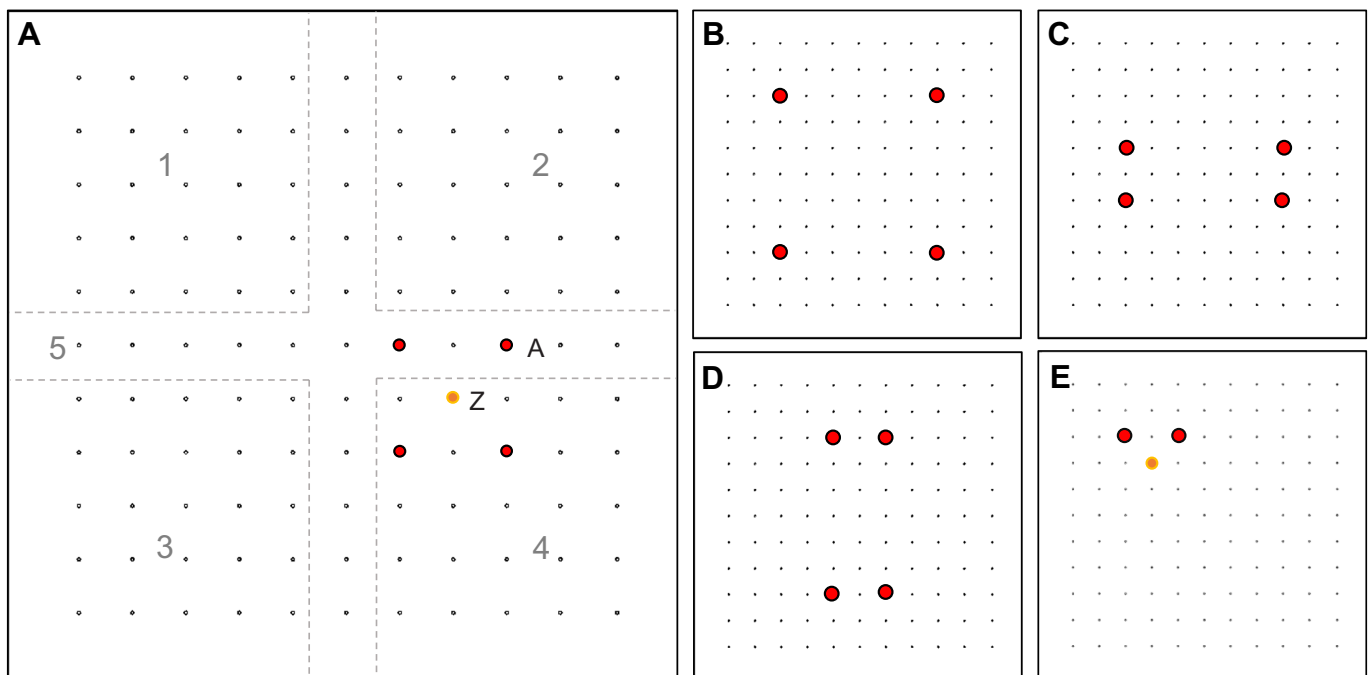


Fig. 3. Landmark array on the LED panel during baseline and test trials in experiment 1 and 2. (A) One-hundred and twenty-one LED lights were attached to a plastic foam wall. In every trial of experiment 1, four LEDs (red dots) were lit and served as landmark array (A). The task of the animal was to touch an unlit LED, the goal (Z; for representation, this unlit LED is here marked by a yellow dot; however, during the experiment, Z remained unmarked) in the middle of the landmark array. The position of the array was shifted to all quadrants (1–4) and the overlap area (5); see Materials and Methods for details. (B–D) The different types of expansion the seals experienced during the test trials of experiment 1: (B) diagonal expansion, (C) horizontal expansion and (D) vertical expansion. (E) Landmark array of experiment 2 as presented during baseline trials. The array consisted of two lit LEDs. The goal the animals needed to respond to is marked with a yellow dot; however, during the experiment the goal remained unmarked.

interspersed. The number of interspersed trials varied between 5 and 25 trials, depending on the animals' performance and motivation, meaning if the seal continued to respond even without assistance and was eager to participate in the training, more trials without assistance were conducted, in comparison with sessions in which the animal was responding more hesitantly without guidance and was generally cooperating less well.

Training phase

In the training phase, the LED array was presented, and the animal was required to touch the goal location with its snout. An incorrect answer was defined as the animals touching elsewhere on the LED wall. Training was continued until the animal reached a learning criterion of 80% correct choices in two consecutive sessions.

Testing phase

During the testing phase, test trials were interspersed into the session. In test trials, the landmark array was expanded either diagonally, horizontally or vertically. Diagonal expansion resulted in the LEDs of the array to be 90 cm apart from each other, instead of 30 cm apart as during baseline trials (Fig. 3B). During horizontal expansion, the two landmarks on the right and left kept their position relative to each other; however, these two pairs were moved 90 cm apart horizontally (Fig. 3C). In vertical expansion, the two upper and the two lower LEDs kept their position, but those two pairs were moved 90 cm apart vertically (Fig. 3D).

During the testing phase, the sessions consisted of 19 baseline trials and one test trial. The test trial consisted of one of the expansions and was interspersed at random; however, it was never included as first and last trial of the 20-trial session. Baseline trials were ended by feedback from the experimenter, either reinforcement or a verbal no. No feedback was given in the test trials. We performed 10 test trials for each expansion, resulting in 30 test sessions overall. We kept the number of expansion trials per session small, as we were interested in the spontaneous instead of a learned reaction of the seals to the expansion.

Experiment 2A

Stimulus

In this experiment, two landmarks were lit in every trial (Fig. 3E). The landmarks were aligned in the horizontal dimension of the search space. The goal was located between the landmarks but at a perpendicular distance away from and below the line connecting the two landmarks.

Training and testing phase

The training phase was conducted as described for experiment 1. In the testing phase, test trials were interspersed in which the two-landmark array was expanded in the left–right dimension of the search space, meaning the distance between the landmarks was increased. After expansion, the two landmarks were 90 cm apart from each other, instead of 30 cm as during training. In each testing session, the LED array was placed in each quadrant four times and five times in the overlapping area, resulting in 19 baseline trials and one additional test trial. The position of the LED array in the test trial was chosen at random, but over the course of the sessions, the position occurred equally often in the quadrants and the overlapping area, which resulted in six test trials per area. Again, no feedback or reward was given for the seals' answers in test trials. Altogether, 30 sessions were run, resulting in 30 responses to the expanded array per animal.

Experiment 2B

After completing the testing phase of experiment 2A, we conducted a brief follow-up test. In these sessions, test trials with a single landmark were interspersed into the baseline trials with a two-landmark array to determine how the seals would respond to a further reduction of the number of landmarks. We conducted two sessions with 25 baseline trials and 5 test trials.

Data analysis

We performed all statistical tests with an alpha level of 0.05 in Microsoft Excel (Version: Office 2019; Redmond, WA, USA) and IBM SPSS (v.26; International Business Machines Corporation Armonk, NY, USA). During analysis, we focused on the first choices the animals made when performing the control and test trials in all phases of the experiment; it needs to be noted that the seals hardly (only two to six times in each experiment) gave second responses. In order to unravel the underlying strategy of landmark perception, we performed an analysis similar to that of Marsh et al. (2011). According to their analysis, the three landmark-based strategies predict specific hypothetical goal-locations, with corresponding peak search areas. Because Marsh and colleagues could not differentiate between the beacon strategy and the directional vector strategy owing to an overlap of the hypothetical goal locations, we redefined the goal locations to clearly separate them for the beacon and the directional vector strategies (Fig. 1). We then determined the frequency of searches that fell into each of the hypothetical goal locations and performed binomial tests to determine whether the answers of the animal that were directed towards each area differed from what would be expected by chance. Similar to Marsh et al. (2011), we compared the frequency of searches per area with the expected frequency of searches in the areas according to the number of possible goal locations (see Fig. 1) in the area (1 goal location for the rule-based strategy=1% chance, 8 or 4 goal locations for the vector strategy=10% or 5% chance, and 28 or 14 goal locations for the landmark strategy=34% or 17% chance in experiment 1 or experiment 2). Our analysis assumes that a random or indirect search would target any LED in the area of the respective strategy.

RESULTS

Experiment 1

The seals needed 746 trials in 37 sessions (Nick), 995 trials in 34 sessions (Filou) and 1725 trials in 59 sessions (Moe) to meet the learning criterion in the training phase. In the testing phase, the seals chose the goal location of the landmark array with 87.7% (Nick), 91.9% (Moe) and 97.6% (Filou) of the choices in the baseline trials. During the expansion trials, the seals directed all their searches to locations inside the landmark boundary area (Fig. 1). Inside the landmark boundary area, irrespective of the type of expansion, all three seals prioritised their searches in the regions predicted by the directional vector strategy more than expected by chance (binomial test: $P < 0.05$; Fig. 4, Table 1). No animal directed its search according to a rule-based strategy, which, in our configuration, would have resulted in choosing the centre of the expanded array. While focusing their searches in the surrounding of landmarks, Moe and Filou mostly responded to an LED that adopted the same angle and distance to a landmark as the goal during the baseline trials and the trials in the training sessions (Fig. 4). However, the seals favoured LEDs at the training angle and distance to different landmarks. Filou preferred the LED defined by the training vector from the top-right landmark, whereas Moe also preferred to answer at the position defined by the training vector but with respect to the

Table 1. Number of searches in the expansion test trials of experiments 1 and 2 in the hypothetical goal locations predicted by the three different strategies (beacon, vector, rule-based strategy) for the three different types of expansion (diagonal, horizontal, vertical expansion)

Experiment	Subject	Expansion	Hypothetical goal locations			
			Beacon	Vector	Rule-based	Other
1	Filou	Diagonal	3	6	0	1
		Horizontal	2	8	0	0
		Vertical	3	6	0	1
		Overall	8	20	0	2
	Moe	Diagonal	2	5	0	3
		Horizontal	1	9	0	0
		Vertical	2	8	0	0
		Overall	5	22	0	3
	Nick	Diagonal	3	5	0	2
		Horizontal	2	8	0	0
		Vertical	6	4	0	0
		Overall	11	17	0	2
2	Filou	Horizontal	12	18	0	0
	Moe	Horizontal	12	18	0	0
	Nick	Horizontal	19	10	0	1

Numbers written in *italics* indicate percentages higher than expected by chance ($\alpha=0.05$). Note that the chance level was different for each of the landmark strategies in the two experiments (see Materials and Methods, Data analysis). 'Other' defines any position chosen by the seals not in line with the goal locations predicted by the beacon, vector or rule-based strategy.

top-left landmark. Both animals thus responded as in the training phase and with the same distance to the landmarks but orientated to different landmarks. Nick, in contrast, favoured two different positions, one defined by the training vector, and one defined by a length of 15 cm with an angle of 45 deg counter-clockwise from the training vector.

The animals' responses were predominantly related to the two uppermost landmarks. Filou and Nick selected a location in the upper half of the wall in all of their searches. With 85% of his responses to the upper half of the wall, even Moe mainly directed his search to the upper two locations and only went to locations in the lower half of the configuration wall three times; then the seal gave responses with respect to the lower landmarks consistent with its responses to the upper landmarks.

Experiment 2

All animals needed only two training sessions including 60 trials to complete the learning criterion for experiment 2. In the testing phase, 92.3% (Nick), 96.3% (Moe) and 99.1% (Filou) of the baseline trials were directed to the goal location of the unexpanded landmark array. In the testing phase of experiment 2A, all seals prioritised their searches in the regions predicted by the vector and beacon strategies more than expected by chance (binomial test: $P<0.05$; Fig. 5, Table 1). No search was ever in line with rule-based searching, i.e. to the middle of the array or in triangular form.

In this experiment, Filou again preferably chose to respond at a single vector from a landmark (Fig. 5A), thus searching for the goal at the same vector as in the baseline/training condition, but he did not discriminate between the left and right landmarks; instead, he always searched at the same vector irrespective in relation to which landmark. On the contrary, Moe's searches were directed to locations defined by three different vectors: the training vector, a vector 45 deg counter-clockwise to the training vector with a length of 21 cm and a vector 45 deg clockwise to the training vector with a length of 15 cm with almost the same frequency (Fig. 5A). Nick again, as in experiment 1, favoured the training vector and the vector 45 deg clockwise to the training vector with a length of 15 cm.

In experiment 2B, all seals maintained a high performance as 95.5% of the baseline trials were in the correct location of the

unexpanded landmark array for Moe and Nick. Filou did not make any mistakes at all in the baseline trials. In the control trials, Filou responded at a location defined by the training vector relative to the landmark in 90% of the trials (Fig. 5B). In contrast, Moe's and Nick's responses were distributed over locations defined by the training vector and by a vector pointing to the LED directly underneath the landmark. Both animals responded with the training vector in 50% of the trials (Fig. 5B).

DISCUSSION

In this study, it was determined how harbour seals encode positional information in respect to landmarks. The seals learnt the experimental paradigm within 746–1725 trials. For comparison, orangutans needed several thousand trials to acquire the basic task in a comparable study (Marsh et al., 2011). The relatively fast acquisition process in harbour seals supports findings from previous studies that had revealed excellent access to as well as high performance in visuo-spatial tasks (Renouf and Gaborko, 1989; Mauck and Dehnhardt, 2007).

In the testing phase of the first experiment, in which the four-landmark array was expanded, the seals mostly showed responses to locations in the dimension parallel to the shift and no shift in searching in the perpendicular dimension. The search behaviour of the seals was consistent with a directional vector strategy as previously described for non-human primates and gerbils, among others (Collett et al., 1986; MacDonald et al., 2004; Poti et al., 2005, 2010). The seals mostly kept the same distance and angle towards a landmark that they had experienced during training; they chose the goal in line with the training vector. Filou mainly applied one vector, the appropriate vector to locate the goal with respect to the top-right landmark, irrespective to which landmark, which was most apparent in experiment 2B. Moe even chose three different training vectors depending on the specific landmark he was targeting. Moreover, all seals responded inside the landmark array. Overall, these observations stress the high directionality of their response behaviours; their responses were clearly more directed than predicted by the alternative strategy, the beacon strategy.

The ability to memorize and apply a vector would allow seals to relocate a specific goal with respect to (a) landmark(s) precisely. The application of a directional vector strategy would furthermore

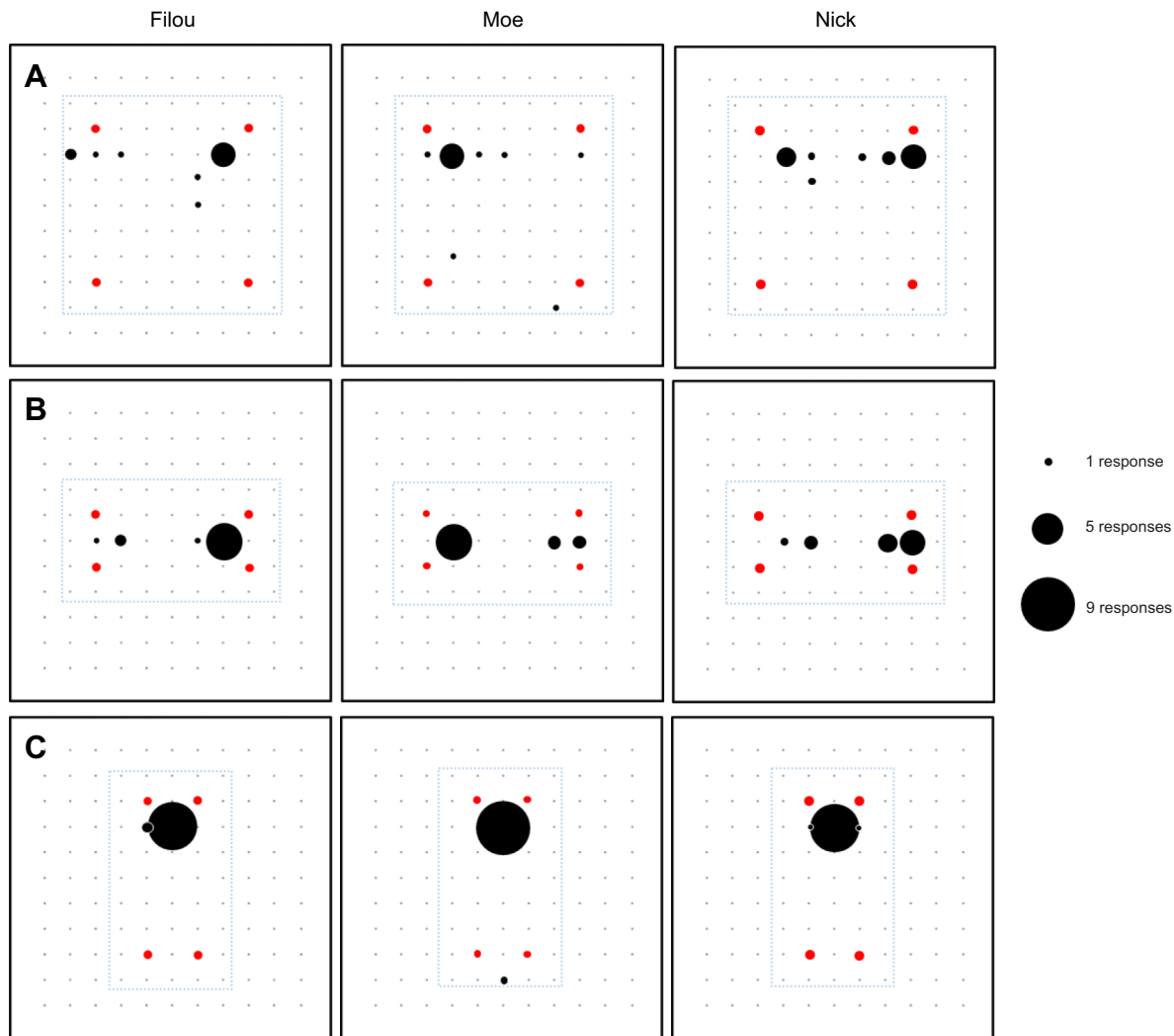


Fig. 4. Response behaviour of seals Filou (left), Moe (center) and Nick (right) in respect to different types of expansion of the four-landmark array (experiment 1). (A) Diagonal expansion, (B) horizontal expansion and (C) vertical expansion. Black circles represent the responses of the seals with the number of responses per position coded by the size of the circles: the largest circle represents the highest number of responses at a position, as indicated in the key. Conventions as in Figs 1 and 3; the array is always shown in the middle of the LED wall, although its position was varied across the LED wall over trials.

enable seals to use landmarks for piloting. When encountering (a) landmark(s), seals would be required to determine the correct, previously memorized/learned vector, including directional as well as distance information, with respect to the landmark(s), leading the seals to the next station on its journey and/or finally towards its end-goal. This piloting strategy would benefit from the previously reported abilities of seals to estimate distances (Maaß and Hanke, 2021) and to keep a straight path (Vance et al., 2021). Our results thus allow the formulation of new hypotheses on landmark orientation or orientation/navigation in general to be tested in the future in an attempt to explain the well-documented navigational abilities of seals that are commuting between the open ocean and the coast.

The response behaviour in the baseline trials of experiment 1 shows that the animals must have identified individual as well as groups of landmarks inside the array; the correct identification of the middle of the array requires the determination of upper versus lower landmarks and left versus right landmarks. For this identification process, the seals could have used cues, such as the setup's position in the water column, the relative position of the seal to the setup during stationing/approaching, and their own position in the water

column. These cues were available in our experiment. However, it needs to be stressed that, in our experiment, the aforementioned cues did not interfere with the experimental paradigm, as only the landmarks defined the goal precisely, thus the seals were forced to use the LED landmark array to solve the task.

In the test trials, the seals were mainly answering in the upper half of the panel. The focus of the seals to the upper landmarks might result from the asymmetry of the seals' visual field in the vertical meridian (Hanke et al., 2006). Owing to their dorsal eye position, harbour seals have a large dorsal, but only a small ventral, visual field (see Fig. S1). Thus, when approaching the panel, the two upper landmarks remained within in the visual field longer than the two lower landmarks. Thus they localized the goal with respect to the upper two landmarks that defined the goal most precisely. When transiting to experiment 2A, the seals' responses clearly indicate that the top two landmarks provide enough orientation cues to be used for goal localization.

The analysis of the results obtained in the two-landmark array experiment revealed that the response behaviour was in line with the directional vector and the beacon strategy but did not correspond with a rule-based approach. The number of responses in line with a beacon

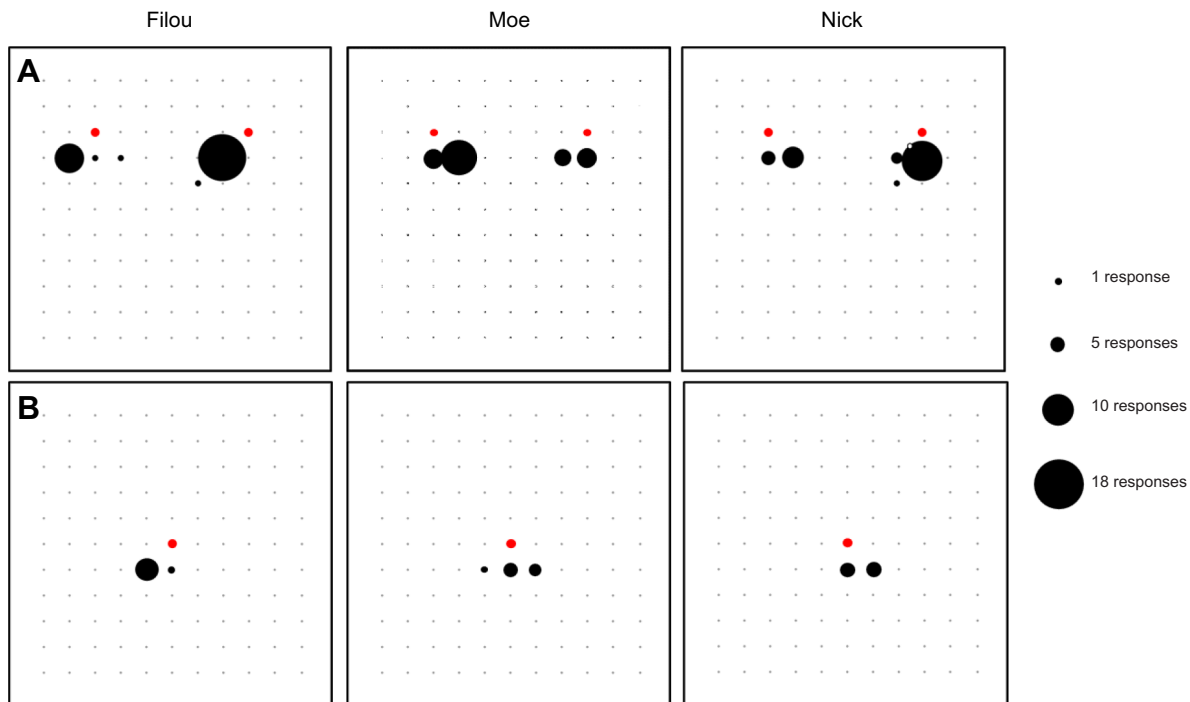


Fig. 5. Response behaviour of seals Filou (left), Moe (center) and Nick (right). Data are shown for the expansion trials of (A) experiment 2A and (B) experiment 2B. Results are visualized as in Fig. 4 and the conventions of all other figures. Please note that in experiment 2A, each seal performed 30 test trials and one test trial was interspersed in a session of 20 trials, while in experiment 2B each seal performed five test trials in each of the two sessions of 30 trials.

strategy increased in experiment 2 in comparison to experiment 1; Nick even predominantly answered in line with a beacon strategy. Thus, with reduced landmark information, it seemed more difficult for the seals to obtain/memorize the angular information of the goal versus the landmark. In conclusion, the amount of information available in the environment determines the strategy chosen by the seals and the accuracy of the search behaviour. Flexibility in landmark-based search is vital, allowing the seals to optimize their search in respect to the information available.

From experiment 1 to experiment 2, the seals slightly or clearly shifted their search strategy. Differential use of search strategies in different experimental conditions has already been documented for human children and capuchin monkeys, for example (MacDonald et al., 2004; Poti et al., 2005). Whereas the human children seem to choose a strategy depending on their age (towards using a rule-based strategy when adult), the capuchin monkeys switched their strategy according to the complexity of the task. In contrast to our seals, the primates used a beacon strategy when confronted with a four-landmark configuration but shifted to a directional vector strategy when confronted with a two-landmark configuration. This discrepancy needs to be worked on in future experiments.

In all our experiments, the harbour seals did not implement a rule-based approach in the sense of ‘find the middle’ or ‘complete the triangular form’ to find the goal in the landmark array, which would have resulted in a higher frequency of searches in the respective positions of the expanded array. Adult humans responded according to a rule-based strategy during expansion by answering directly in the middle of the array or by maintaining a triangular shape in tests with two landmarks, which they also expressed verbally when asked about the strategy they had followed during testing (Spetch et al., 1996, 1997; MacDonald et al., 2004). Even though the seals did not spontaneously use a rule-based approach in the current study, seals might be capable of using such an approach when forced to rely on a

rule with a different experimental paradigm, in line with previous studies including birds and primates (Spetch et al., 1997; Poti et al., 2005). When these organisms were asked to respond to the middle of two landmarks that varied in inter-landmark distance, they adopted a rule-based strategy (Kamil and Jones, 1997; Jones et al., 2002; Spetch et al., 2003; Poti et al., 2010). A comparable experiment conducted with harbour seals could reveal whether seals also switch to a rule-based strategy depending on context/task. This context-dependent shift of strategies seems possible, as it would be in line with experimental evidence just mentioned and as the seals showed a change of their response behaviour with the modifications of the landmark array from experiment 1 to experiment 2.

In conclusion, we showed that harbour seals can learn to locate a goal with the help of landmarks and that they preferably choose the vector(s), including direction and distance information, relative to (a) landmark(s) memorized during training. However, the encoding of goals with respect to landmarks is adjusted with respect to the specific environment as indicated by the context-dependent shifts in search strategy, a flexibility that seems to be adaptive in a complex environment.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: E.M., F.D.H.; Methodology: E.M.; Validation: E.M.; Formal analysis: E.M.; Investigation: E.M.; Resources: E.M., F.D.H.; Data curation: E.M.; Writing - original draft: E.M.; Writing - review & editing: E.M., F.D.H.; Visualization: E.M.; Supervision: F.D.H.; Project administration: F.D.H.; Funding acquisition: E.M., F.D.H.

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References

- Brown, R. F. and Mate, B. R.** (1983). Abundance, movements, and feeding habits of harbor seals, *Phoca vitulina*, at Netarts and Tillamook Bays, Oregon. *Fish. Bull.* **81**, 291-301.
- Burt, T. and Macias Garcia, C.** (2003). Amarillo fish (*Girardinichthys multiradiatus*) use visual landmarks to orient in space. *Ethology* **109**, 341-350. doi:10.1046/j.1439-0310.2003.00876.x
- Byl, J. A., Miersch, L., Wieskotten, S. and Dehnhardt, G.** (2016). Underwater sound localization of pure tones in the median plane by harbor seals (*Phoca vitulina*). *J. Acoust. Soc. Am.* **140**, 4490-4495. doi:10.1121/1.4972531
- Cartwright, B. and Collett, T. S.** (1983). Landmark learning in bees. *J. Comp. Physiol.* **151**, 521-543. doi:10.1007/BF00605469
- Cheng, K.** (1989). The vector sum model of pigeon landmark use. *J. Exp. Psychol. Anim. Behav. Process.* **15**, 366. doi:10.1037/0097-7403.15.4.366
- Cheng, K. and Sherry, D. F.** (1992). Landmark-based spatial memory in birds (*Parus atricapillus* and *Columba livia*): the use of edges and distances to represent spatial positions. *J. Comp. Psychol.* **106**, 331. doi:10.1037/0735-7036.106.4.331
- Cheng, K., Spetch, M. L., Kelly, D. M. and Bingman, V. P.** (2006). Small-scale spatial cognition in pigeons. *Behav. Process.* **72**, 115-127. doi:10.1016/j.beproc.2005.11.018
- Chevallier, D., Karpitchev, M., McConnell, B. J., Moss, S. and Vincent, C.** (2014). Can gray seals maintain heading within areas of high tidal current? Preliminary results from numerical modeling and GPS observations. *Mar. Mamm. Sci.* **30**, 374-380. doi:10.1111/mms.12024
- Collett, T., Cartwright, B. A. and Smith, B. A.** (1986). Landmark learning and visuo-spatial memories in gerbils. *J. Comp. Physiol. A* **158**, 835-851. doi:10.1007/BF01324825
- Cook, R. G. and Tauro, T. L.** (1999). Object-goal positioning influences spatial representation in rats. *Anim. Cogn.* **2**, 55-62. doi:10.1007/s100710050024
- Fiset, S.** (2007). Landmark-based search memory in the domestic dog (*Canis familiaris*). *J. Comp. Psychol.* **121**, 345-353. doi:10.1037/0735-7036.121.4.345
- Hanke, W., Römer, R. and Dehnhardt, G.** (2006). Visual fields and eye movements in a harbor seal (*Phoca vitulina*). *Vision Res.* **46**, 2804-2814. doi:10.1016/j.visres.2006.02.004
- Jones, J. E., Antoniadis, E., Shettleworth, S. J. and Kamil, A. C.** (2002). A comparative study of geometric rule learning by nutcrackers (*Nucifraga columbiana*), pigeons (*Columba livia*) and jackdaws (*Corvus monedula*). *J. Comp. Psychol.* **116**, 350. doi:10.1037/0735-7036.116.4.350
- Kamil, A. C. and Jones, J. E.** (1997). The seed-storing corvid Clark's nutcracker learns geometric relationships among landmarks. *Nature* **390**, 276-279. doi:10.1038/36840
- Kowalewsky, S., Dambach, M., Mauck, B. and Dehnhardt, G.** (2006). High olfactory sensitivity for dimethyl sulphide in harbour seals. *Biol. Lett.* **2**, 106-109. doi:10.1098/rsbl.2005.0380
- Krüger, Y., Hanke, W., Miersch, L. and Dehnhardt, G.** (2018). Detection and direction discrimination of single vortex rings by harbour seals (*Phoca vitulina*). *J. Exp. Biol.* **221**, jeb170753. doi:10.1242/jeb.170753
- Lopez, J., Gómez, Y., Rodríguez, F., Broglio, C., Vargas, J. and Salas, C.** (2001). Spatial learning in turtles. *Anim. Cogn.* **4**, 49-59. doi:10.1007/s100710100091
- Maaß, E. and Hanke, F. D.** (2021). Distance estimation in reproduction tasks in a harbor seal (*Phoca vitulina*). *Water* **13**, 938. doi:10.3390/w13070938
- Macdonald, S. E., Spetch, M. L., Kelly, D. M. and Cheng, K.** (2004). Strategies in landmark use by children, adults, and marmoset monkeys. *Learn. Motiv.* **35**, 322-347. doi:10.1016/j.lmot.2004.03.002
- Marsh, H. L., Spetch, M. L. and Macdonald, S. E.** (2011). Strategies in landmark use by orangutans and human children. *Anim. Cogn.* **14**, 487-502. doi:10.1007/s10071-011-0382-9
- Matsumura, M., Watanabe, Y. Y., Robinson, P. W., Miller, P. J. O., Costa, D. P. and Miyazaki, N.** (2011). Underwater and surface behavior of homing juvenile northern elephant seals. *J. Exp. Biol.* **214**, 629-636. doi:10.1242/jeb.048827
- Mauck, B. and Dehnhardt, G.** (2007). Spatial multiple-choice matching in a harbour seal (*Phoca vitulina*): differential encoding of landscape versus local feature information? *Anim. Cogn.* **10**, 397-405.
- Niesterok, B., Krüger, Y., Wieskotten, S., Dehnhardt, G. and Hanke, W.** (2017). Hydrodynamic detection and localization of artificial flatfish breathing currents by harbour seals (*Phoca vitulina*). *J. Exp. Biol.* **220**, 174-185. doi:10.1242/jeb.148676
- Poti, P., Bartolomei, P. and Saporiti, M.** (2005). Landmark use by *Cebus apella*. *Int. J. Primatol.* **26**, 921-948. doi:10.1007/s10764-005-5330-6
- Poti, P., Kanngiesser, P., Saporiti, M., Amiconi, A., Bläsing, B. and Call, J.** (2010). Searching in the middle: capuchins' (*Cebus apella*) and bonobos' (*Pan paniscus*) behavior during a spatial search task. *J. Exp. Psychol. Anim. Behav. Process.* **36**, 92. doi:10.1037/a0015970
- Renouf, D. and Gaborko, L.** (1989). Spatial and visual rule use by harbour seals (*Phoca vitulina*). *Biol. Behav.* **14**, 169-181.
- Schulte-Pelkum, N., Wieskotten, S., Hanke, W., Dehnhardt, G. and Mauck, B.** (2007). Tracking of biogenic hydrodynamic trails in harbour seals (*Phoca vitulina*). *J. Exp. Biol.* **210**, 781-787. doi:10.1242/jeb.02708
- Spetch, M. L.** (1995). Overshadowing in landmark learning: touch-screen studies with pigeons and humans. *J. Exp. Psychol. Anim. Behav. Process.* **21**, 166. doi:10.1037/0097-7403.21.2.166
- Spetch, M. L., Cheng, K. and Macdonald, S. E.** (1996). Learning the configuration of a landmark array: I. Touch-screen studies with pigeons and humans. *J. Comp. Psychol.* **110**, 55. doi:10.1037/0735-7036.110.1.55
- Spetch, M. L., Cheng, K., Macdonald, S. E., Linkenhoker, B. A., Kelly, D. M. and Doerkson, S. R.** (1997). Use of landmark configuration in pigeons and humans: II. Generality across search tasks. *J. Comp. Psychol.* **111**, 14. doi:10.1037/0735-7036.111.1.14
- Spetch, M. L., Rust, T. B., Kamil, A. C. and Jones, J. E.** (2003). Searching by rules: pigeons' (*Columba livia*) landmark-based search according to constant bearing or constant distance. *J. Comp. Psychol.* **117**, 123. doi:10.1037/0735-7036.117.2.123
- Steingass, S., Horning, M. and Bishop, A. M.** (2019). Space use of Pacific harbor seals (*Phoca vitulina richardii*) from two haulout locations along the Oregon coast. *PLoS ONE* **14**, e0219484. doi:10.1371/journal.pone.0219484
- Stewart, B. S.** (1984). Diurnal hauling patterns of harbor seals at San Miguel Island, California. *J. Wildl. Manag.* **48**, 1459-1461. doi:10.2307/3801821
- Suryan, R. M. and Harvey, J. T.** (1998). Tracking harbor seals (*Phoca vitulina richardii*) to determine dive behavior, foraging activity, and haul-out site use. *Mar. Mamm. Sci.* **14**, 361-372. doi:10.1111/j.1748-7692.1998.tb00728.x
- Vance, H., Hooker, S. K., Mikkelsen, L., Van Neer, A., Teilmann, J., Siebert, U. and Johnson, M.** (2021). Drivers and constraints on offshore foraging in harbour seals. *Sci. Rep.* **11**, 1-14. doi:10.1038/s41598-021-85376-2
- Wehner, R. and Räber, F.** (1979). Visual spatial memory in desert ants, *Cataglyphis bicolor* (Hymenoptera: Formicidae). *Experientia* **35**, 1569-1571. doi:10.1007/BF01953197
- Yesiltepe, D., Conroy Dalton, R. and Ozbil Torun, A.** (2021). Landmarks in wayfinding: a review of the existing literature. *Cogn. Process* **22**, 369-410. doi:10.1007/s10339-021-01012-x

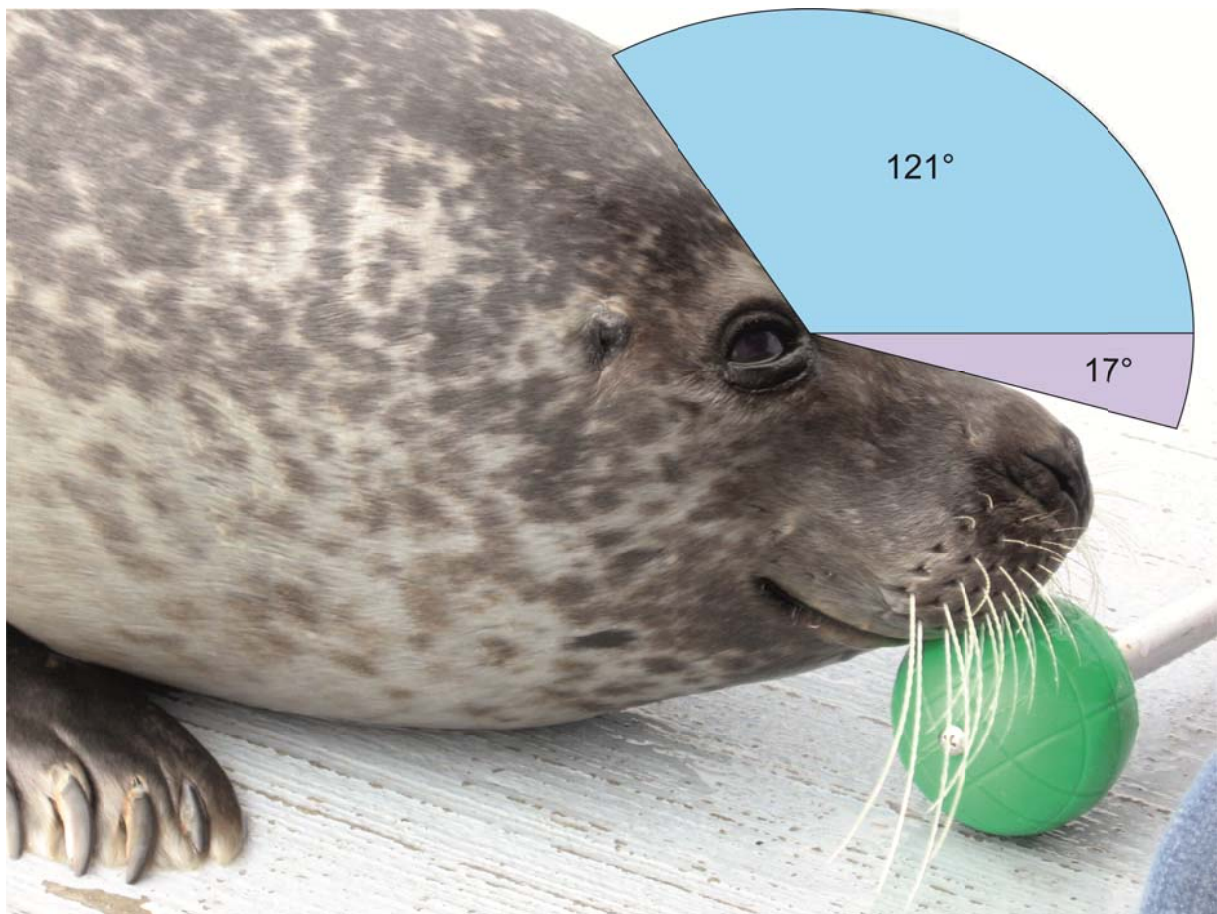


Fig. S1. Dynamic visual field of a harbor seal. Harbor seals have a large dorsal but only small ventral dynamic visual field. When eye movements are prohibited, the dorsal visual field is still extending over 69 deg. The ventral visual field is reduced to 12 deg without eye movements (data taken from Hanke et al. 2006)