Marine mammal response to interannual variability in Monterey Bay, California

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ABSTRACT: The coastal upwelling ecosystem near Monterey Bay, California, is a productive yet variable ecosystem and an important foraging area for many mobile apex predators, such as marine mammals. Long-term studies are necessary to better understand how wide-ranging predators respond to temporal environmental variability; however, few of these studies exist. We conducted monthly shipboard line-transect surveys in Monterey Bay from 1997 to 2007. We identified 22 species of marine mammals, and calculated monthly and annual densities for the 12 most commonly sighted (focal) species. Species richness remained relatively constant (mean richness ± SE: 13.7 ± 0.396 species yr⁻¹) from 1997 to 2006. Focal species were most evenly distributed (Shannon’s equitability, $E_H = 0.820$) but least dense (mean density ± SE: 0.0598 ± 0.0141) during the anomalous upwelling conditions of 2005, and least even (1997 $E_H = 0.413$; 1998 $E_H = 0.407$) but dense (mean density ± SE: 1997: 0.433 ± 0.177; 1998: 0.438 ± 0.169 ind. km⁻²) during the 1997/1998 El Niño event. There were no statistically significant differences in the densities of marine mammal species between warmer and cooler years. The community and species-specific responses of marine mammals to warm-water years differed depending on the mechanism of oceanographic variability. During the 1997/1998 El Niño (a basin-wide event), marine mammals aggregated in nearshore areas, such as Monterey Bay, with relatively greater productivity than offshore regions, whereas during anomalous upwelling conditions of 2005 (a more localized oceanographic event), marine mammals redistributed away from Monterey Bay to areas less affected by the anomaly.

KEY WORDS: Upwelling · El Niño · Density · Diversity · California Current · Distance sampling

INTRODUCTION

Eastern boundary currents, such as the California Current, are among the most productive ecosystems in the world (Hickey 1979, Glantz & Thompson 1981). Productivity is driven locally by coastal upwelling (Barber & Smith 1981, Carr & Kearns 2003) and on a larger scale by oceanic circulation patterns. Seasonal upwelling in the California Current is initiated when northwest winds along the west coast of North America combine with the earth’s rotation to create offshore Ekman transport of surface waters, resulting in the movement of cool, nutrient-rich waters to the surface (Barber & Smith 1981, Huyer 1983, Service et al. 1998). Nutrients brought to the euphotic zone induce phytoplankton blooms, which increase productivity at multiple trophic levels (Hutchings et al. 1995, Pennington & Chavez 2000).

In addition to seasonal coastal upwelling, interannual variability resulting from El Niño or La Niña

Physical and biological oceanographic variables associated with upwelling and El Niño or La Niña events influence the density and diversity of apex predators, such as marine mammals (Sydeman & Allen 1999, Benson et al. 2002, Burtenshaw et al. 2004, Keiper et al. 2005, Lowry & Forney 2005). Researchers have reported increased species diversity and richness in the North Pacific during El Niño years, and attributed the increases to the northward movement of species typically associated with warmer waters (Benson et al. 2002, Worm et al. 2005, Hooff & Peterson 2006). Indeed marine mammal species richness patterns are most closely correlated with SST (Whitehead et al. 2010) and the trend of increased biodiversity in lower (tropical) latitudes is a well-documented phenomenon (Rohde 1992, Hillebrand 2004). For example, copepod communities were most diverse and evenly distributed in warm tropical and subtropical waters (Woodd-Walker et al. 2002). Organisms that traditionally reside in warmer, lower-latitude habitats appear to be expanding their range poleward as the world’s oceans warm, resulting in an observed (and anticipated) increase in biodiversity at higher latitudes (Hughes 2000, Beaugrand et al. 2002, Hyrenbach & Veit 2003, Whitehead et al. 2008). The response of these species to short-term variability, such as delayed and weakened seasonal upwelling (Snyder et al. 2003) or El Niño events, may be the best predictor of how these species will respond to long-term oceanographic change, such as ocean warming (Trathan et al. 2007).

Understanding how environmental variability affects species density and diversity is an important step in anticipating changes that may occur in species composition and ecosystem functioning over longer time periods.

Many researchers have studied the movements or distribution of marine mammal species throughout a season or for several years (Sydeman & Allen 1999, Friedlaender et al. 2006, Weise et al. 2006), but few long-term (decadal) studies exist. Our objective was to examine species-specific and community responses of marine mammals to interannual environmental variability. To accomplish this objective, we (1) documented changes in marine mammal density and diversity in Monterey Bay, California, throughout an 11-yr period (1997−2007) and (2) tested for differences in marine mammal density and diversity (richness and evenness) between years with warm and cool oceanic conditions. We expected that marine mammal species typically associated with cool waters would be present in greater densities during years dominated by cool oceanic conditions (strong upwelling, low SSTs), species typically associated with warm waters would be present in greater densities during years dominated by warm oceanic conditions (weak upwelling, high SSTs), and that there would be no significant differences in the densities of resident species in Monterey Bay during warm and cool years. We also expected greater marine mammal species richness (the number of different species present) and evenness (the relative abundances of each species present) in Monterey Bay during warm-water years as species associated with warm waters moved north into the bay, but regularly occurring species remained.

MATERIALS AND METHODS

Study area

Monterey Bay, located off the central California coast, is the largest bay (~1200 km²) on the west coast of the United States completely open to the ocean (Benson et al. 2002, Croll et al. 2005; Fig. 1). The Monterey Submarine Canyon, one of the largest canyons in the world (Shepard 1973), divides the bay into 2 nearly equal shallower shelves (up to 140 m deep and 10 to 15 km wide), with deeper waters over the canyon in the center of the bay (Greene et al. 2002; Fig. 1). Monterey Bay is influenced seasonally by a coastal upwelling plume that originates ~30 km to the north, near Point Año Nuevo (Rosenfeld et al. 2002; Fig. 1). Monterey Bay is influenced seasonally by a coastal upwelling plume that originates ~30 km to the north, near Point Año Nuevo (Rosenfeld et al. 1994). Upwelling winds off the central California coast usually begin in March and continue through August, with periods of wind relaxation (Send et al. 1987) becoming more frequent during July and August (Pennington & Chavez 2000). A short transitional oceanic period occurs from late August through November, when winds continue to relax and SSTs increase until a warmer, less productive winter Davidson Current season begins in December and persists through February (Skogsberg 1936, Skogsberg & Phelps 1946, Pennington & Chavez 2000).
Line transect field methods

Monterey Bay was divided into 7 transect lines that were surveyed for marine mammals throughout the 11-yr study (Benson et al. 2002, Croll et al. 2005; Fig.1). Transect lines were 10 to 25 km in length, and totaled ~126 km. The location of the first line was randomly chosen from a 3-min latitudinal range, after which each line was spaced 5.5 km apart for uniform coverage of the bay (Benson et al. 2002). From September 2006 until November 2007, the first line was no longer randomly selected and the same grid of 7 lines was surveyed. The entire survey area, ~909 km², included all of Monterey Bay and the waters off the Monterey Peninsula (except nearshore regions) beginning at the 55 m (30 fathom) isobath and extending WNW to 122.083° W longitude (Fig. 1). Surveys were completed during 2 consecutive days each month from May through November 1997 to 2006 at a ship speed of 18.5 km h⁻¹ (10 knots). Additional surveys were completed during 2 consecutive days in January and March 2003 to 2006. Surveys were conducted 1 day a month (5 transect lines totaling 82 km) during January, March, May, July, August, and November 2007.

Two observers stationed on top of the bridge (4.3 m above sea level, except for March and July 2007 when observers were 5.66 m above sea level) searched for marine mammals on each side of the vessel from the trackline to 90° abeam of the ship using Fujinon 7 × 50 binoculars with a compass and reticle scale in the oculars. A third center observer searched mainly with the naked eye along the trackline and near the ship (binoculars were used to aid in species identification), while a fourth person entered sightings into a laptop computer using the program SeeBird_WinCruz (Holland 2008) with direct input from the ship’s GPS. When a sighting occurred, all observers assisted with species identification (to the lowest taxonomic level) and abundance estimation. Time, latitude, longitude, species, number of individuals, cue (body, blow), method of detection (eye, binoculars), compass bearing, and number of reticle marks down from the horizon were recorded. Environmental conditions (fog or rain, visibility, wind direction and speed, swell direction and speed, horizontal and vertical sun position, and Beaufort sea state) were continually updated throughout the survey.

Density calculations

Radial distances of marine mammal groups to the trackline were obtained using binocular reticle measurements and the platform (observer eye) height using the formula of Lerczak & Hobbs (1998). Radial distances for sightings made using the land/ocean interface as a reticle reference rather than the true horizon (sky/ocean) were adjusted using the military analyst toolset in ArcMap (ArcGIS desktop version 9.2, ESRI). Perpendicular distance (x) was computed from the radial distance (r) and the angle (θ) between the trackline and the marine mammal group using trigonometry (x = r sinθ).

Monthly marine mammal densities were calculated from line transect data using the Multiple Covariate Distance Sampling (MCDS) analysis engine in Distance software (Buckland et al. 2001, Buckland et al. 2004, Thomas et al. 2006). Density calculations were based on Eq. (1):

\[
\hat{D} = \frac{\sum_{i=1}^{n} s_i / \hat{p}_i}{2wL}
\]

(1)

\[s_i\] is the number of individuals counted in encounter i, \[\hat{p}_i\] is the corrected detection probability, \[w\] is the half-aperture correction factor in kilometers, and \[L\] is the length of the transect line in kilometers.
where \( \hat{D} \) is the density estimate (ind. km\(^{-2}\)), \( n \) is the number of marine mammal groups detected (where a group is comprised of one or more individuals), \( s_i \) is the size of the \( i \)th group, \( w \) is the truncation distance and half-width of the transect, \( L \) is the total line length, and \( \hat{p}_i \) is the estimated probability of detecting the \( i \)th group (Buckland et al. 2001, 2004). Detection probability \( (\hat{p}_i) \) was estimated from the detection function \( g(x) = \) the probability of detecting a group, given that it is at distance \( x \) from the line), which was fitted to the perpendicular distances using Distance software. Sightings from May through November 1997 to 2007, and January and March 2003 to 2007 were used to obtain the detection function. A sequence of models with different sets of covariates and series expansion terms (cosine, simple or hermite polynomial) were used to find the best fit model (i.e., detection function), which minimized Akaike’s information criterion (AIC; Akaike 1973). Only statistically non-collinear covariates with a significant effect on perpendicular distance were considered as covariates in MCDS (Beaufort sea state, swell height, group size, and visibility). Visibility was the approximated distance to the horizon (≤6.44 km) and was affected by fog and haze. Only survey effort that occurred in acceptable Beaufort sea states (0 to 4) and swell heights (<2.4 m) were included in analyses. Additionally, Beaufort sea state was treated as a factor with discrete levels 0 to 4, whereas all other covariates were continuous variables (non-factors). Observer was not included as a covariate because there were more than 200 volunteer observers during the 11-yr study. Because more experienced observers were likely better at detecting marine mammals than less experienced observers, not including observer as a covariate added to the random error in the density estimates. The truncation distance was set for each species such that ~5% of the most distant sightings were excluded (Buckland et al. 2001).

Model convergence was more difficult to achieve using all combinations of covariates and adjustments (i.e., too many parameters) for species with fewer sightings. In these cases, the best model was chosen from models including only one covariate and no adjustments, or no covariates (null model). Additionally, to achieve model convergence using the hazard rate key function in MCDS, starting values for the hazard rate parameter estimates were manually selected using those calculated from the half normal model for the same species, with the power parameter coefficient set to 2. Although distance sampling methods assume certain detection on the trackline (i.e., \( g(0) = 1 \)), it is likely that some individuals were not seen (perception bias) or submerged (availability bias) as the vessel passed, resulting in an underestimation of true density. However, because the aim of this study was to compare relative densities of animals in the same area through time, consistent underestimation should not affect the results.

### Annual density and diversity

Once monthly density estimates were obtained, mean annual focal species density and individual species densities were calculated for 1997 to 2007 using monthly density estimates for May through November. Focusing on these months enabled us to document the presence of seasonal summer migrants in addition to regularly sighted resident species, and capture the time period when the region is most productive. Mean focal species density was defined as the mean annual density of the 12 most abundant species with enough sightings to obtain density estimates.

Species richness and species evenness were calculated for 1997 to 2006 (2007 was excluded from diversity calculations due to reduced effort that year). Rather than using solely a diversity index, such as the Shannon-Weiner index, diversity measurements were divided into richness and evenness to discern which variable more strongly contributed to species diversity patterns. Species richness (\( S \)) was defined as the total number of marine mammal species identified each year, including rare species. Sightings that could not be identified to species were excluded from analyses. Species evenness was calculated for each year using the 12 most abundant species (focal species) with enough sightings to obtain density estimates. Species evenness was determined by first calculating the Shannon-Weiner index of diversity (\( H \)):

\[
H = -\sum_{i=1}^{S} (p_i)(\log p_i)
\]  

where \( p_i \) is the proportion of total sample belonging to the \( i \)th species. Shannon’s equitability (\( E_H \), evenness) was then calculated as:

\[
E_H = \frac{H}{\log S}
\]  

where \( S \) is the annual species richness of the 12 most abundant species. Shannon’s equitability index for species evenness quantifies the numerical equality of the annual densities of each of the 12 focal species. An \( E_H \) value of 1 indicates complete evenness (i.e., all 12 species were present in equal densities). Because of the difficulties associated with identifying common
dolphins in the field, long-beaked *Delphinus capensis* and short-beaked *Delphinus delphis* common dolphins were treated as 1 species for analyses.

**Environmental variables**

SST and upwelling indices (UIs) were used to categorize years as warm or cool. Mean monthly SSTs (°C) were calculated from a continuous record of temperature at 1 m depth using the Monterey Bay Aquarium Research Institute’s (MBARI) M1 mooring (www.mbari.org/) (Fig. 1). Mean monthly UIs (m³ s⁻¹ 100 m⁻¹; measures of wind-driven offshore Ekman transport derived from 6-hourly synoptic surface atmospheric pressure fields) from 36° N, 122° W were obtained from the Southwest Fisheries Science Center’s (SWFSC) Environmental Research Division’s (ERD) website (www.pfeg.noaa.gov/).

**Warm vs. cool years**

Species were classified as being associated with warm or cool waters in accordance with prior studies of marine mammal habitat associations off the US west coast (e.g. Forney & Barlow 1998, Keiper et al. 2005, Barlow & Forney 2007). Dall’s porpoise *Phocoenoides dalli*, northern right whale dolphin *Lissodelphis borealis*, and Pacific white-sided dolphin *Lagenorhynchus obliquidens* typically occur in cool temperate to sub-arctic waters and were thus classified as cool-water-associated species, whereas common dolphin *Delphinus* spp. and Risso’s dolphin *Grampus griseus* typically occur in tropical and warm temperate waters and were thus classified as warm-water-associated species. California sea lion *Zalophus californianus*, harbor porpoise *Phocoena phocoena*, harbor seal *Phoca vitulina*, and sea otter *Enhydra lutris* can be seen year-round off the central California coast and were classified as resident species. Blue whales *Balaenoptera musculus* and humpback whales *Megaptera novaeangliae* could not be clearly categorized as species associated with warm or cool water because they are wide-ranging predators that migrate seasonally between the tropics and higher latitudes. Blue and humpback whales are seasonal visitors to Monterey Bay, and are most abundant in the area during the summer and fall months.

To determine whether mean annual SST and UI affected annual marine mammal density and diversity, years were grouped into 2 categories based on similarities in physical oceanographic conditions: cool (low SSTs and high UIs) and warm (high SSTs and low UIs). Years with lesser UI and greater SST pairings (1998, 2004, 2005, and 2006) were categorized as warm years and years with greater UI and lesser SST pairings (1999, 2001, 2002, and 2007) were categorized as cool years (Fig. 2). Discriminate function analysis, performed using SYSTAT (version 12, SYSTAT Software), indicated that warmer and cooler year groupings were significantly different based on UIs and SSTs (n = 8, F = 29.847, p = 0.002, α = 0.05) and generated scores (−15.863 + 0.626 × SST − 0.864 × UI) to categorize years without a clear grouping as warm (1997) or cool (2000, 2003). Warm and cool year groupings agreed with annual oceanic conditions described in the California Cooperative Oceanic Fisheries (CalCOFI) reports (www.calcofi.org/).

Two distinct oceanographic events occurred during our study, resulting in anomalously warm conditions: a strong El Niño during 1997/1998, which affected the entire Pacific Ocean basin (Chavez et al. 2002), and delayed and weakened upwelling during the spring and summer of 2005, which affected only the northern California Current (Peterson et al. 2006). The effect of these 2 events on species composition, in addition to the effect of warm and cool years on
species composition, was examined using a binary similarity index, the Jaccard Index ($S_j$). The Jaccard index uses presence/absence data to determine the similarity between 2 samples, making it possible to include rare species (those without density estimates) in analyses. The range of possible Jaccard coefficients is from 0 (no similarity) to 1 (complete similarity); however, sample size and species richness can affect the maximum value of the coefficient (Wolda 1981).

Statistical analyses

Randomization tests (10,000 iterations) were performed, using Resampling Stats software (version 6, build 1, Resampling Stats, Inc.), to determine whether species richness, species evenness, mean (focal species) density, and individual species densities differed between cool and warm water years. Randomization tests were used rather than a parametric test (such as a t-test) to avoid violating the assumption of a normal error distribution and because t-tests are less robust with smaller sample sizes. Density and diversity estimates were considered to be significantly different if the observed mean difference was >95% of the randomized distribution. One analysis included all years surveyed ($n = 11$) and another included only the most disparate years ($n = 8$; 1997, 2000, and 2003 were excluded). Because 15 statistical comparisons were performed, we conducted a Benjamini-Hochberg false discover rate test (Benjamini & Hochberg 1995) using R software (R Development Core Team 2011) to control for the expected proportion of type I errors.

RESULTS

Density model selection

We identified 22 species of marine mammals during the 11-yr study (Table 1). MCDS results are presented for the 10 species with enough sightings ($n > 55$) to obtain density estimates (Table 2, Fig. 3). MCDS results also are presented for common dolphin and northern right whale dolphin from
**Table 2. Multiple Covariate Distance Sampling (MCDS) results for 12 focal marine mammal species, May to November 1997 to 2007, and January and March 2003 to 2007.** Species, number of observations (n), key function (Hr: hazard rate; Hn: half-normal), power parameter coefficient, intercept of the scale parameter, covariate coefficients for swell height (SH), visibility (V), group size (GS), and Beaufort levels 0–3 (B0–B3; for models with Beaufort, the intercept of the scale parameter is Beaufort 4), adjustment type (herm poly: hermite polynomial; cos: cosine) and number, adjustment coefficient, average probability of detection (P), 5% truncation distance (W), and goodness-of-fit Kolomogorov-Smirnov p-value (α = 0.05) from the model used to obtain species density estimates. Values in parentheses are SE unless otherwise indicated. Common dolphin *Delphinus* spp. (n = 39) and northern right whale dolphin *Lissodelphis borealis* (n = 28) observations were pooled to fit a common model with additive swell height, group size, and species effects. The intercept value for *L. borealis* is the species coefficient. The baseline scale for *Delphinus* spp. is 0.8383 and for *L. borealis* it is 0.8383 × exp (1.251). Densities were calculated from individual detection functions, but pooled values are reported under the common dolphin heading. For scientific names of species see Table 1. ‘–’ indicates that the parameter was not included in the model used to obtain species density estimates.

<table>
<thead>
<tr>
<th>Common name</th>
<th>n</th>
<th>Key function</th>
<th>Power coeff.</th>
<th>Intercept</th>
<th>SH</th>
<th>V</th>
<th>GS</th>
<th>B0</th>
<th>B1</th>
<th>B2</th>
<th>B3</th>
<th>Adj. type (no.) coeff. (SE)</th>
<th>P</th>
<th>W (km)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>California sea lion</td>
<td>177</td>
<td>Hr</td>
<td></td>
<td>2.103</td>
<td>(0.664)</td>
<td>0.104</td>
<td>-0.071</td>
<td>0.092</td>
<td>–</td>
<td>0.751</td>
<td>0.578</td>
<td>0.238</td>
<td>0.055</td>
<td>herm poly (1)</td>
<td>0.351</td>
</tr>
<tr>
<td>Sea otter</td>
<td>334</td>
<td>Hn</td>
<td></td>
<td>0.087</td>
<td>(0.018)</td>
<td>0.085</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.549</td>
<td>0.851</td>
<td>0.831</td>
<td>0.406</td>
<td>–</td>
<td>0.470</td>
</tr>
<tr>
<td>Humpback whale</td>
<td>319</td>
<td>Hr</td>
<td></td>
<td>2.072</td>
<td>(1.621)</td>
<td>0.215</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.942</td>
<td>0.786</td>
<td>0.933</td>
<td>0.691</td>
<td>–</td>
<td>0.370</td>
</tr>
<tr>
<td>Harbor porpoise</td>
<td>249</td>
<td>Hr</td>
<td></td>
<td>3.907</td>
<td>(3.500)</td>
<td>0.162</td>
<td>-0.081</td>
<td>0.079</td>
<td>–</td>
<td>0.572</td>
<td>0.807</td>
<td>0.745</td>
<td>0.192</td>
<td>–</td>
<td>0.540</td>
</tr>
<tr>
<td>Harbor seal</td>
<td>168</td>
<td>Hr</td>
<td></td>
<td>4.276</td>
<td>(4.610)</td>
<td>0.104</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.943</td>
<td>0.437</td>
<td>0.137</td>
<td>0.172</td>
<td>–</td>
<td>0.637</td>
</tr>
<tr>
<td>Risso’s dolphin</td>
<td>144</td>
<td>Hr</td>
<td></td>
<td>2.564</td>
<td>(2.816)</td>
<td>0.157</td>
<td>–</td>
<td>–</td>
<td>0.180</td>
<td>0.012</td>
<td>0.283</td>
<td>0.282</td>
<td>0.334</td>
<td>–</td>
<td>0.468</td>
</tr>
<tr>
<td>Pacific white-sided dolphin</td>
<td>141</td>
<td>Hn</td>
<td></td>
<td>0.232</td>
<td>(0.039)</td>
<td>0.157</td>
<td>0.007</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>cos (2)</td>
<td>0.377</td>
<td>0.954</td>
</tr>
<tr>
<td>Dall’s porpoise</td>
<td>79</td>
<td>Hn</td>
<td></td>
<td>0.110</td>
<td>(0.032)</td>
<td>0.032</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.063</td>
<td>1.293</td>
<td>0.805</td>
<td>0.846</td>
<td>–</td>
<td>0.422</td>
</tr>
<tr>
<td>Blue whale</td>
<td>74</td>
<td>Hn</td>
<td></td>
<td>0.681</td>
<td>(0.064)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.501</td>
</tr>
<tr>
<td>Elephant seal</td>
<td>55</td>
<td>Hr</td>
<td></td>
<td>2.039</td>
<td>(0.619)</td>
<td>0.165</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.388</td>
</tr>
<tr>
<td>Common dolphin</td>
<td>67</td>
<td>Hn</td>
<td></td>
<td>0.838</td>
<td>(3.742)</td>
<td>0.321</td>
<td>–</td>
<td>0.001</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.519</td>
</tr>
<tr>
<td>Northern right whale dolphin</td>
<td>–</td>
<td></td>
<td></td>
<td>1.251</td>
<td>(0.647)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.672</td>
</tr>
</tbody>
</table>
a shared model with additive swell height, group size, and species effects (Table 2, Fig. 3). The best models chosen for California sea lion, elephant seal *Mirounga angustirostris*, harbor porpoise, humpback whale, and sea otter were those that minimized AIC. The best models chosen for the remaining species excluded models with the lesser AIC if those models violated assumptions or produced unexpected results. For example, models with numeric covariates with a coefficient counter to expectation were disregarded and the next best model was chosen. Thus models with a positive swell height coefficient (for harbor seal) or negative visibility coefficients (for Dall’s porpoise and common dolphin–northern right whale dolphin) were not selected. The swell height coefficient should have been negative because as swell height increased, the distance at which we could detect individuals or groups should have decreased. Similarly, the visibility coefficient should have been positive because as visibility increased, the distance at which we could detect individuals or groups should also have increased. Coefficients with a sign opposite of expectation likely occurred by chance alone, resulting from unequal sample size distribution across all levels of a covariate. Adjustment terms were not included in the final model for

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**Fig. 3.** Observed perpendicular distance (km) with fitted detection functions for 12 focal marine mammal species. Note different scales on x-axes. *Common dolphin and northern right whale dolphin histograms are the same because a shared model was used.
Risso’s dolphin density because the probability of detection at zero distance with a simple polynomial adjustment was >1, violating the assumption of certain detection on the trackline. The best half-normal model was chosen over hazard-rate models for Pacific white-sided dolphin and Dall’s porpoise to avoid fitting the spike in sightings at zero distance, likely because of responsive movement of these species towards vessels (Williams & Thomas 2007). The model minimizing AIC for blue whale included Beaufort coefficients that did not increase from Beaufort 4 to Beaufort 0 as was expected (i.e., this model did not reflect the fact that we were able to detect individuals or groups at further distances in lesser Beaufort sea states), thus the next best model was chosen. Beaufort coefficients mostly were positive, changing the scale of the detection function and increasing the distance at which objects could be detected.

**Annual density and diversity**

Densities of the 12 focal species varied among years (Fig. 4), although mean species richness remained relatively constant (mean ± SE: 13.7 ± 0.396 species yr⁻¹; Fig. 5). California sea lions, Dall’s

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**Fig. 4.** Annual densities (mean ± SE) for 12 focal marine mammal species identified in Monterey Bay, California, May to November 1997 to 2007. Note different scales on y-axes.
porpoises, harbor porpoises, harbor seals, humpback whales, Risso’s dolphins, and sea otters were sighted every year, and elephant seals were sighted every year but 2007 (Fig. 4). Northern right whale dolphins were absent in 1997, 1998, and 2007, and present in greatest densities during 1999 (Fig. 4). Pacific whitesided dolphins were present in greatest densities from 1999 to 2002, and decreased densities during 1997, 1998, and 2003 to 2007 (Fig. 4). Common dolphins were present in greatest densities and were the most abundant species of marine mammal during 1997 and 1998, but were sighted infrequently or absent in subsequent years (Fig. 4). Blue whales were present in greatest densities during 2003 and in least densities or absent from 1997 to 1999 and 2005 to 2007 (Fig. 4). Risso’s dolphins were present in decreased densities during 1997 and 1998, but were almost 10 times as dense during 2002 (Fig. 4). Species richness varied by only 4 species during the 10-yr period, and was 15 (greatest richness observed) during 1997, 1998, and 2005 (Fig. 5). Marine mammals were most evenly distributed, but least dense during 2005, and least even, but dense during 1997 and 1998 (Fig. 5). The greatest density of focal species occurred during 1997, 1998, and 2007 (Fig. 5).

Warm vs. cool years

Species richness, species evenness, and mean focal species density did not differ significantly between warm and cool years (Table 3). Most species were sighted during at least 1 warm and 1 cool water year (Table 1). The community composition of marine mammals was more similar between warm and cool years ($S = 0.818$) than it was between El Niño and anomalous upwelling years ($S = 0.579$ for a community comparison between 1997/1998 and 2005). No species had significantly greater densities during cool or warm water years; however, Pacific whitesided dolphins and northern right whale dolphins had near significantly greater densities in cool years (Table 3). Performing the same analyses using only the 8 most disparate years (used to develop the discriminate function) did not yield significant results (results not shown).

DISCUSSION

Marine mammals are wide-ranging predators that respond to environmental conditions over a large spatial area. Monterey Bay is a small region within the larger California Current, which is a temporally and spatially dynamic system. Temporal environmental variability between 1997 and 2007 was most extreme during the 1997/1998 El Niño and 2005 anomalous upwelling events. Although 1997, 1998, and 2005 were all categorized as warm years (in our analyses for Monterey Bay and in CalCOFI reports), the community response of marine mammals differed between the 1997/1998 El Niño and 2005 anomalous upwelling events. These differences most likely resulted from the disparate spatial scales at which El Niño and upwelling occurred. The 1997/1998 El Niño affected the entire Pacific Ocean basin, whereas
anomalous upwelling was more localized off the west coast of North America. Thus it is important to consider the physical factors leading to anomalous oceanographic events to understand and predict how marine mammals will respond to environmental variability.

### Focal species density

The lack of statistically significant differences in mean focal species density between warm and cool years likely occurred because individual and community responses of marine mammals differed between the 1997/1998 El Niño and the 2005 anomalous upwelling events. During 2005, the delayed onset of upwelling resulted in warmer than average spring and summer SSTs, decreased primary production and zooplankton abundance (Mackas et al. 2006, Schwing et al. 2006, Barth et al. 2007), and reduced catches of mid-trophic-level fish species (Brodeur et al. 2006) off the west coast of North America. Although the annual UI value for 2005 was not extremely low, it was the delayed onset of upwelling that had substantial impacts on the ecosystem. The biological effects of the anomalous oceanic conditions of 2005 were limited to central California through southern British Columbia (northern California Current; Mackas et al. 2006). Thus, it is possible that decreased density of marine mammal species in Monterey Bay during 2005 resulted from redistribution of more mobile species to areas outside the region affected by the upwelling anomaly. Indeed, densities of wider-ranging, more seasonally abundant species (e.g. blue whales, Dall’s porpoises, and Pacific white-sided dolphins) decreased in Monterey Bay during 2005, whereas densities of resident species (e.g. California sea lions, harbor porpoises, harbor seals, and sea otters) remained similar to or increased from the previous year (point estimates only).

Increased densities of marine mammals in Monterey Bay during the 1997/1998 El Niño can be attributed to the marked increased density of common dolphins during those years, in addition to the aggregation of cetacean species in narrow nearshore areas of increased productivity, such as Monterey Bay (Benson et al. 2002). Although productivity was reduced in Monterey Bay during the 1997/1998 El Niño event compared with other years, there was still sufficient nutrient availability nearshore to support some primary production (Kudela & Chavez 2000, Chavez et al. 2002). Therefore, during basin-wide decreases in productivity (e.g. El Niño events), mobile top predators may be more likely to move from offshore to nearshore areas and less likely to redistribute north–south, to the extent that may have occurred during the anomalous upwelling conditions.

Given the response of marine mammals to the basin-wide 1997/1998 El Niño event, it is possible that marine mammals in the California Current will

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<table>
<thead>
<tr>
<th>Warm year mean</th>
<th>n</th>
<th>Cool year mean</th>
<th>n</th>
<th>Obs. mean diff.</th>
<th>Adj. p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species richness</td>
<td>13.800 (0.800)</td>
<td>5</td>
<td>13.600 (0.245)</td>
<td>5</td>
<td>0.200</td>
</tr>
<tr>
<td>Species evenness</td>
<td>0.575 (0.090)</td>
<td>5</td>
<td>0.650 (0.054)</td>
<td>5</td>
<td>0.075</td>
</tr>
<tr>
<td>Density (focal species)</td>
<td>0.244 (0.076)</td>
<td>5</td>
<td>0.305 (0.089)</td>
<td>6</td>
<td>0.061</td>
</tr>
<tr>
<td>Blue whale</td>
<td>0.003 (0.0002)</td>
<td>5</td>
<td>0.011 (0.005)</td>
<td>6</td>
<td>0.008</td>
</tr>
<tr>
<td>California sea lion</td>
<td>0.821 (0.309)</td>
<td>5</td>
<td>2.298 (1.056)</td>
<td>6</td>
<td>1.477</td>
</tr>
<tr>
<td>Common dolphin</td>
<td>1.489 (0.917)</td>
<td>5</td>
<td>0.129 (0.086)</td>
<td>6</td>
<td>1.360</td>
</tr>
<tr>
<td>Dall’s porpoise</td>
<td>0.066 (0.014)</td>
<td>5</td>
<td>0.061 (0.015)</td>
<td>6</td>
<td>0.005</td>
</tr>
<tr>
<td>Elephant seal</td>
<td>0.016 (0.006)</td>
<td>5</td>
<td>0.008 (0.002)</td>
<td>6</td>
<td>0.007</td>
</tr>
<tr>
<td>Harbor porpoise</td>
<td>0.065 (0.011)</td>
<td>5</td>
<td>0.097 (0.021)</td>
<td>6</td>
<td>0.032</td>
</tr>
<tr>
<td>Harbor seal</td>
<td>0.055 (0.015)</td>
<td>5</td>
<td>0.065 (0.012)</td>
<td>6</td>
<td>0.010</td>
</tr>
<tr>
<td>Humpback whale</td>
<td>0.037 (0.007)</td>
<td>5</td>
<td>0.052 (0.008)</td>
<td>6</td>
<td>0.016</td>
</tr>
<tr>
<td>Northern right whale dolphin</td>
<td>0.012 (0.010)</td>
<td>5</td>
<td>0.075 (0.024)</td>
<td>6</td>
<td>0.063</td>
</tr>
<tr>
<td>Pacific white-sided dolphin</td>
<td>0.126 (0.062)</td>
<td>5</td>
<td>0.553 (0.149)</td>
<td>6</td>
<td>0.427</td>
</tr>
<tr>
<td>Risso’s dolphin</td>
<td>0.177 (0.077)</td>
<td>5</td>
<td>0.255 (0.137)</td>
<td>6</td>
<td>0.079</td>
</tr>
<tr>
<td>Sea otter</td>
<td>0.065 (0.016)</td>
<td>5</td>
<td>0.049 (0.016)</td>
<td>6</td>
<td>0.016</td>
</tr>
</tbody>
</table>

Table 3. Randomization test results (10,000 iterations) for differences in species richness, species evenness, focal species density, and individual species densities between warm and cool years. Values presented are means (SE), sample size (n), observed mean difference, and probability of obtaining results that include the observed mean difference with p-values adjusted using the Benjamini-Hochberg false discover rate test (Adj. p). For scientific names of species see Table 1.
respond to global warming in a similar manner. Thus increased density and decreased species evenness may occur in Monterey Bay as Earth’s climate changes. Large groups of typically warm-water-associated species may move into Monterey Bay and aggregate nearshore where productivity is greater, increasing predation pressure on mid-trophic-level species that may already be stressed from other climate change-related factors.

Diversity

The lack of statistically significant differences in species evenness between warm and cool years is supported by Jaccard similarity results, which indicate that the community composition of marine mammals was more similar between warm and cool years than it was between the warm years of the 1997/1998 El Niño and the 2005 anomalous upwelling events.

The lack of a statistically significant difference in species richness between warm and cool years and the similarity of species richness among all years are indications that species richness may not be the appropriate measure of diversity for this study. Species richness did not vary much among years because the presence of regularly occurring species and different rare species totaled approximately the same richness every year. Therefore, despite changes in the species composition among years, the total number of species sighted was similar for all years. Additionally, both evenness and Jaccard similarity results indicated that there were differences in species composition between 1997/1998 and 2005, whereas species richness was identical in all 3 years. Therefore a measure of species composition, rather than species richness, may provide more insight into marine mammal diversity patterns for future studies.

Individual species densities

Although there were no statistically significant differences in the densities of individual species during cool or warm years, the greater (near significant) densities of Pacific-white sided dolphins and northern right whale dolphins in Monterey Bay during cool years may have biological significance. These 2 cold-temperate species likely moved south with cool waters during years dominated by cool conditions. Barlow & Forney (2007) found similar increased abundances of these species in waters off California during the cool-water year of 1996, but reported no consistent variation in the abundance of Risso’s or common dolphins with warm or cool water years. We also found no significant differences in the densities of Risso’s or common dolphins during warm years and attribute this result to the fact that these species did not respond consistently to similarly grouped years. The conspicuous increase in common dolphin densities reported in this study during the 1997/1998 El Niño event did not occur again in subsequent warm years (during 2004, 2005, and 2006 no common dolphins were sighted). It is possible that during the 1997/1998 El Niño event, large groups of common dolphins moved north or inshore into Monterey Bay in search of prey, whereas during 2004, 2005, and 2006 common dolphins remained in their normal habitat further south (southern California to Mexico) or offshore because those areas were largely unaffected by the upwelling anomalies along the coast (Peterson et al. 2006).

We expected our results would differ slightly from those of Barlow & Forney (2007) because they sampled a much larger region of the California Current at a coarser spatial (including waters off California, Oregon, and Washington) and temporal scale. However, when our results are similar, it may be a good indication of the extent and conditions under which Monterey Bay can be used as an indicator of the larger California Current System. For example, during cool-water years, increased abundances of Pacific white-sided dolphins and northern right whale dolphins in Monterey Bay may indicate increased abundances of these species throughout the California Current. However, Barlow & Forney (2007) found increased abundances of Dall’s porpoise in the California Current during cool-water years and we did not find a similar increase in Monterey Bay. Both our study and that of Barlow & Forney (2007) found decreased abundances of blue whales in 2005 and attribute this decrease to a redistribution of individuals outside of the study region. Thus, Monterey Bay may act as an indicator of the larger California Current for some cool-water associated species, but results are not consistent across species.

Marine mammals respond not only to environmental variability, but also to the distribution of their prey, which responds to oceanographic conditions. Marine mammals typically move to areas where prey is more abundant. For example, densities of Risso’s dolphins (which feed almost exclusively on squid) increased conspicuously in Monterey Bay during 2002, concurrent with a marked increase in abundance of Humboldt squid Dosidicus gigas (Zeid-
berg & Robison 2007). Additionally, the market squid (Loligo opalescens) fishery in Monterey Bay during 2002 had a near-unprecedented record-setting year (CDFG 2003). Thus the increase in Risso’s dolphin density in Monterey Bay during 2002 may have been a direct result of increased squid availability that year, and squid (as with other prey) are extremely responsive to changes in environmental conditions (Jackson & Domeier 2003, Zeidberg et al. 2006).

### Summary and conclusions

By tracking a community of top predators in Monterey Bay during an 11-yr period, we examined interannual changes in marine mammal density and diversity as related to temporal environmental variability. We found no statistically significant differences in density and diversity when comparing warm and cool water years, likely a result of the large amount of variability in the system. However, the long-term nature of the study enabled us to document the response of marine mammals to both local (2005 anomalous upwelling) and basin-wide (1997/1998 El Niño) events. We found that marine mammals responded differently to these warm-water years depending on the mechanism initiating the variability. During basin-wide decreases in productivity (e.g. El Niño events), mobile top predators may be more likely to move from offshore to nearshore areas and aggregate in regions with relatively greater productivity, whereas during more localized decreases in productivity (e.g. anomalous upwelling), marine mammals may redistribute to regions north and south of the area affected by the anomalous conditions. Climate change may affect marine mammals in a similar manner as the 1997/1998 El Niño event, and lead to increased densities of marine mammals in Monterey Bay as the oceans continue to warm. A better understanding of the specific processes that induce warm and cool water years will provide additional insight into the response of marine mammals to future environmental variability.

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