Regional diving behavior of Pacific Arctic beluga whales *Delphinapterus leucas* and possible associations with prey

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ABSTRACT: Two populations of beluga whales *Delphinapterus leucas* in the Pacific Arctic make seasonal migrations to regions characterized by diverse bathymetry and hydrography, yet there is limited information contrasting behavior and foraging across regions. We used satellite-linked time−depth recorders attached to 30 belugas from 1997 to 2012 to infer the depths at which belugas forage seasonally and regionally. We also examined the correspondence between patterns of beluga diving and the vertical distribution of a primary prey species, Arctic cod *Boreogadus saida*, within the western Beaufort Sea. A suite of regional diving metrics revealed that beluga dive behavior varied among regions and sometimes between populations. Estimates of occupancy time at depth, in addition to maximum and modal dive depths for 6 h periods, suggested that Eastern Chukchi Sea and Beaufort Sea belugas were regularly diving to the seafloor in shallow shelf regions. Along slope margins and in the deep Canada Basin (>3000 m), specific portions of the water column were more frequently targeted. The greatest maximum daily dive depths were >900 m in the Canada Basin. Arctic cod were most abundant at 200−300 m in the western Beaufort Sea, and beluga dives within the survey area also most frequently targeted these depths. These results are consistent with a hypothesis that Arctic cod are a primary prey item for Pacific Arctic belugas and suggest that foraging belugas dive to depths that maximize prey encounters. In the context of a rapidly transforming Arctic ecosystem increasingly exposed to anthropogenic activities, our results quantify the ecological importance of key regions for these 2 populations.

KEY WORDS: Diving behavior · Arctic marine ecology · Foraging ecology · Optimal foraging · Chukchi Sea · Beaufort Sea · *Boreogadus saida*

INTRODUCTION

Optimal foraging theory predicts that predators aim to maximize energy intake through prey consumption while simultaneously minimizing the time and energy required to obtain that prey (Schoener 1971). For air-breathing marine predators, an additional constraint is imposed by oxygen demands limiting foraging time. Accordingly, predators minimize their transit time to optimize foraging before surfacing and recovering from oxygen depletion (Boyd 1997). In addition to physiological limits, other factors...
such as quality and distribution of prey affect dive duration (Thompson & Fedak 2001, Thums et al. 2013). Animals that frequently dive deep trade off foraging needs over the physiological demands of diving deeply (Davis 2014). As a result, depth layers of the water column where divers frequently visit or spend extended periods of time can indicate where foraging is focused (Laidre et al. 2003, Robinson et al. 2012).

Marine environments also present a complex suite of factors that may influence the dive behavior of marine predators. Dive depths can vary according to prey density at depth, as well as the habitat characteristics (e.g. bathymetry, hydrography, presence of sea ice) encountered. Indices of dive depth relative to ocean depth can indicate pelagic or benthic foraging (Jessopp et al. 2013, Watt et al. 2015), and maximum dive depth may identify specific foraging depths (Photopoulou et al. 2014). Understanding patterns of prey within the water column provides additional context for interpreting diving behavior where estimates of prey distribution, abundance, or density can inform a mechanistic understanding of marine mammal diving, foraging at depth, and habitat use (e.g. Palacios et al. 2013, Witteveen et al. 2015).

Beluga whales *Delphinapterus leucas* are generalist predators, occurring in a number of habitat types and feeding on diverse prey (Laidre et al. 2008). In the Pacific Arctic, some belugas migrate thousands of kilometers seasonally as the annual sea ice recedes and advances (Richard et al. 2001, Suydam et al. 2001, Suydam 2009). Two beluga populations, the Beaufort Sea (BS) and Eastern Chukchi Sea (ECS), winter in the northern Bering Sea before migrating during spring into the Beaufort and Chukchi Seas (Frost & Lowry 1990) as well as the deep (>3000 m) Canada Basin where they are generally spatially and temporally segregated during summer to fall (Hauser et al. 2014). Managed as separate stocks with distinct mitochondrial DNA signatures (O’Corry-Crowe et al. 1997, Allen & Angliss 2014), spatial overlap between populations is greatest in fall, when BS belugas initiate westward migration to the Chukchi Sea ahead of the ECS westward migration (Hauser et al. 2014). Both populations commence southward passage through the Bering Strait in November. These belugas use different habitats over the range of their seasonal migrations, and each region is characterized by complex hydrography, diverse topography, seasonal sea ice fluctuations, and freshwater input that presumably influence prey distributions and foraging arenas for diving belugas (Weingartner et al. 1998, Day et al. 2013).

The limited information on BS beluga diving behavior suggests that most dives are ‘square-shaped’ to the seafloor in Beaufort Sea waters 15 to 600 m deep, or ‘V-shaped’ to 700–900 m in deep (>600 m) areas over the Canada Basin (Richard et al. 1997). Recent analyses of ECS beluga diving suggest that regional differences exist in the proportion of time spent at depth, modal depths, and dive durations (Citta et al. 2013). Looking across inshore shelf (<75 m), slope (75–400 m), basin (>400 m), and Barrow Canyon regions, Citta et al. (2013) found that ECS shallow-type diving (<50 m) most often occurred in shelf waters compared to deep diving (with 1 mode at <50 m and another at >400 m) in the Canada Basin. However, intermediate diving (with 1 mode at <50 m and another near 250 m) was found in all but the shelf regions. ECS beluga diving in slope areas and the Barrow Canyon tended to coincide with depths where regional hydrographic conditions are commonly assumed to concentrate prey such as Arctic cod *Boreogadus saida*. Available stomach content and fatty acid diet information suggest that adult Arctic cod are a primary prey item of both BS and ECS belugas, although other fish (e.g. saffron cod *Eleginus gracilis*), cephalopods (primarily octopus), and benthic invertebrates (primarily crangonid shrimp or echiurid worms) have been recorded (Seaman et al. 1982, Loseto et al. 2009, Quakenbush et al. 2015). Arctic cod make up the bulk of surveyed fish biomass in the Pacific Arctic, yet have only been surveyed in the nearshore shelf, and more recently, slope waters of the Chukchi and Beaufort Seas (Parker-Stetter et al. 2011, Norcross et al. 2013, Logerwell et al. 2015).

Both BS and ECS populations are considered to be stable (Allen & Angliss 2014), although recent evidence suggests a decline in BS growth rates (Harwood et al. 2014). This decline is coincident with downward trends in body condition or reproduction of other Beaufort Sea marine predators that also primarily consume Arctic cod (Harwood et al. 2015). Harwood et al. (2014, 2015) suggested that these shifts in life history parameters are related to recent Pacific Arctic ecosystem changes (e.g. Grebmeier 2012) affecting prey, further indicating the importance of understanding foraging behavior and beluga relationships with prey.

Little work has been done to investigate inter-population differences in dive behavior for BS and ECS belugas, or to quantify how diving is related to foraging at depth. We used diving information from 30 beluga whales tagged with satellite transmitters linked to time–depth recorders to quantify differ-
ences in underwater behavior between populations and among regions. Our goal was to identify the depths and portions of the water column targeted by belugas within diverse oceanographic and bathymetric regions of the Pacific Arctic. We inferred foraging behavior for BS and ECS belugas based on the depths where belugas exhibited prolonged time at depth after accounting for the time spent traveling to and from each depth as well as modal and maximum dive depths. We then estimated whether diving was in close proximity to the seafloor or pelagic portions of the water column to indicate potential demersal or water-column foraging. Additionally, we examined patterns of ECS beluga diving relative to the vertical distribution of a primary prey item, Arctic cod, in the western Beaufort Sea. We compared a subset of ECS beluga diving data to the vertically integrated densities of Arctic cod from a fisheries acoustic survey (Parker-Stetter et al. 2011) and hypothesized that the number of beluga dives to target depths is positively related to the density of Arctic cod at the same depths. Although limited in spatial and temporal resolution, this analysis allowed us to explore the relationship between beluga diving behavior and prey abundance as well as to evaluate the use of our diving parameters as indicators of foraging behavior.

MATERIALS AND METHODS

Study area and region definition

The regions of the Pacific Arctic are characterized by complex hydrography and diverse bathymetry affecting habitat and ecosystem structure (Carmack & Wassman 2006; Fig. 1). The northern Bering Sea is comprised of the shallow Chirikov Basin (<75 m) and is connected to the Chukchi Sea via the narrow (~80 km wide) Bering Strait. The Chukchi Sea has complex, but shallow (<75 m everywhere but the northern perimeter), bathymetry characterized by several canyons and distinct water masses that can be strongly affected by wind-driven circulation (Danielson et al. 2014). The Alaska Coastal Current (ACC) flows from the northeast Chukchi Sea into the narrow, steeply sloped Alaska Beaufort Sea through Barrow Canyon (Pickart 2004), where wind forcing affects the presence of belugas and presumed foraging opportunities (Stafford et al. 2013). The warm, fresh ACC joins a shelfbreak jet along the Beaufort Sea slope, sometimes reaching the Canada Beaufort Sea (von Appen & Pickart 2012). Eddies spin off the shelfbreak jet into the deep, offshore Canada Basin (Spall et al. 2008), and upwelling along the slope is common (Pickart et al. 2013). Outflow from the Mackenzie River joins the shelfbreak jet and typically flows eastward over the shallow and wide Canada Beaufort sea shelf, forming a turbid plume over the Mackenzie Estuary (Carmack & Macdonald 2002). The plume continues into the steep-sloped Amundsen Gulf where wind-forced upwelling occurs near Cape Bathurst and a thermohalocline forms in mid-water depths.
We collected movement and diving behavior from BS and ECS belugas throughout the Pacific Arctic. To assess diving by belugas across the range of habitats in the Pacific Arctic, we designated 11 regions based on variable habitat types in the Pacific Arctic, previous descriptions of BS and ECS beluga distribution (Richard et al. 2001, Suydam et al. 2001, Suydam 2009), summer core areas (Hauser et al. 2014), and the 400 m isobath (Fig. 1). The 400 m isobath has been used in previous assessments of beluga diving (Citta et al. 2013) and was the maximum depth sampled by beluga tags in some years (Table S1 in the Supplement, available at www.int-res.com/articles/suppl/m541p245_supp.pdf). Bathymetry and hydrography affects habitat type (i.e. 'shallow shelf,' 'slope,' and 'deep pelagic') for each region, and belugas use each region at different times during seasonal migrations (Table 1).

### Beluga whale telemetry data


Table 1. Study regions (see Fig. 1) and number of tagged Eastern Chukchi Sea (ECS; total n = 26) and Beaufort Sea (BS; total n = 4) beluga whales Delphinapterus leucas in each region, including the number of 6 h maximum-depth periods. Habitat types vary according to bathymetry and hydrography. Median depth (range in parentheses) measured by the location of 6 h dive periods are also shown for each region. Minimum depths of 0 m correspond to locations very close to shore, based on a mean high water vertical datum (Jakobsson et al. 2012). The first to last day of the year and season (summer = July−August, fall = September−October, winter = November−April, spring = May−June) when a tagged whale was located in each region are indicated; --: no tagged whales occurred in the region.

<table>
<thead>
<tr>
<th>Region (habitat type)</th>
<th>ECS</th>
<th>No. tagged whales (no. 6 h dive periods)</th>
<th>BS</th>
<th>First–last day of year (season)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>No. tagged whales (no. 6 h dive periods)</td>
<td></td>
<td>First–last day of year (season)</td>
</tr>
<tr>
<td>Bering Sea (shallow shelf)</td>
<td>6</td>
<td>(1078) 34 (0−56) 311−142 (winter, spring)</td>
<td>2</td>
<td>(123) 75 (0−999) 335−107 (winter)</td>
</tr>
<tr>
<td>Chukchi Sea (shallow shelf)</td>
<td>26</td>
<td>(1612) 45 (0−1004) (year-round)</td>
<td>4</td>
<td>(541) 54 (0−995) (fall, winter)</td>
</tr>
<tr>
<td>ECS core: Kasegaluk Lagoon (shallow shelf)</td>
<td>23</td>
<td>(465) 3 (0−30) 178−317 (spring−fall)</td>
<td>3</td>
<td>(32) 145.5 (28−1702) (fall)</td>
</tr>
<tr>
<td>ECS core: Barrow Canyon (slope)</td>
<td>24</td>
<td>(1874) 150.5 (0−2429) 170−325 (spring−fall)</td>
<td>3</td>
<td>(140) 48.5 (0−943) 240−266 (fall)</td>
</tr>
<tr>
<td>Alaska Beaufort Sea (slope)</td>
<td>15</td>
<td>(221) 335 (0−999) 194−310 (summer, fall)</td>
<td>3</td>
<td>(62) 433 (0−976) 193−261 (summer, fall)</td>
</tr>
<tr>
<td>Canada Beaufort Sea (slope)</td>
<td>6</td>
<td>(86) 630.5 (189−987) 239−305 (fall)</td>
<td>4</td>
<td>(140) 333 (0−967) 189−239 (summer)</td>
</tr>
<tr>
<td>BS core: Mackenzie Estuary (shallow shelf)</td>
<td>--</td>
<td>-- -- --</td>
<td>4</td>
<td>(137) 17 (0−66) 189−239 (summer)</td>
</tr>
<tr>
<td>Amundsen Gulf (slope)</td>
<td>--</td>
<td>-- -- --</td>
<td>3</td>
<td>(238) 314 (1−644) 201−251 (summer)</td>
</tr>
<tr>
<td>East slope Canada Basin (slope)</td>
<td>6</td>
<td>(118) 636 (373−985) 214−271 (summer, fall)</td>
<td>--</td>
<td>-- -- --</td>
</tr>
<tr>
<td>BS core: Viscount Melville Sound (slope)</td>
<td>--</td>
<td>-- -- --</td>
<td>--</td>
<td>-- -- --</td>
</tr>
<tr>
<td>Canada Basin (pelagic deep)</td>
<td>22</td>
<td>(2417) 3248 (993−3907) 191−308 (summer, fall)</td>
<td>2</td>
<td>(274) 3108 (1025−3918) 210−260 (summer, fall)</td>
</tr>
<tr>
<td>Total</td>
<td>26</td>
<td>(7923) 147 (0−3907) 84.5</td>
<td>4</td>
<td>(1600) 84.5 (0−3918)</td>
</tr>
</tbody>
</table>

*aNo year-round tag deployments were available for BS beluga whales. Deployments ranged from Day 189 (early July) to 107 (mid-April) 

*bDate of last tag transmission, last date in region is unknown
manufactured by Wildlife Computers and consisted of ST-10, ST-16, SPLASH, and MK10transmitters. Most tags were deployed in early July, with tag duration varying by individual (Table 2). All tags transmitted continuously except 2 BS tags in 2005, which were programmed to transmit every 4 d to conserve battery life.

We obtained location data, associated error classes, and binned dive histograms from the Argos data system. The first 24 h of location and dive data post-tag deployment were excluded to eliminate any potential behavioral effects resulting from capture and tagging. Speed (maximum 6.4 km h⁻¹, Richard et al. 2001) and angle filters were used to eliminate unlikely locations (Freitas et al. 2008). The tags contained pressure transducers which sampled whale depths at a resolution of 0.5 (2005–2012 tags), 2 (1997 tags), or 4 m (1998–2002 tags). Satellite data transmission limitations required diving data to be compressed into four 6 h histograms per day. Dive data comprised 3 types of histograms: the number of dives to maximum depth layers (‘maximum depth’ histogram periods), the proportional time within each depth layer (‘time-at-depth’ histogram periods), and number of dives to dive duration categories (‘duration’ histogram periods). We pre-specified depth layers (or ‘bins’) for each histogram type before deployment, but different bin thresholds were used among histogram types, populations, and tagging years (see Table S1) as tag technology improved and for different research purposes (e.g. see Richard et al. 1997, Citta et al. 2013). For analysis, we consolidated bins to the finest resolution available among all years and populations: 0–10, 10–50, 50–100, 100–200, 200–300, 300–400, >400 m for maximum-depth 6 h periods; 0–10, 10–50, 50–100, 100–200, 200–400, >400 m for time-at-depth 6 h periods; and 0–1, 1–3, 3–6, 6–9, 9–20, >20 min for duration 6 h periods. We sampled time-at-depth layers differently than maximum-depth layers in 1 year (2005), so slightly coarser time-at-depth layers were necessary when consolidated for analysis (Table S1). A depth of at least 2 or 4 m (for 2007–2012 tags and 1997–2005 tags, respectively) was required before a dive was registered on the tag. We used a correlated random walk model to estimate geographic locations at the beginning of each 6 h period based on observed locations and associated Argos spatial error (Johnson et al. 2008). Six ECS tags from 2007, 2010, and 2012 also provided maximum daily dive depth (to a maximum of 1000 m), recorded as the maximum absolute depth measured by the tag within the previous 24 h. Thus, maximum daily dive depths were point estimates for a 24 h period measured with higher resolution than the dive data summarized in 6 h maximum-depth periods. We assigned the daily location from Hauser et al. (2014) to each maximum daily dive depth.

Table 2. Beluga whales Delphinapterus leucas tagged in the Beaufort Sea (BS, n = 4) and Eastern Chukchi Sea (ECS, n = 26) populations near the Mackenzie River Estuary, Canada, and Point Lay, AK, USA, respectively. Adult (ADU) or immature (IMM) reproductive status was visually assessed based on size, coloration, and presence of a calf if possible. Tag duration refers to the number of days dive data were collected and used in the present study

<table>
<thead>
<tr>
<th>Year</th>
<th>Tag ID</th>
<th>Sex</th>
<th>Length (cm)</th>
<th>Reproductive status</th>
<th>Capture date</th>
<th>Tag duration (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1997</td>
<td>97-2118</td>
<td>F</td>
<td>374</td>
<td>ADU</td>
<td>26 July</td>
<td>127</td>
</tr>
<tr>
<td></td>
<td>97-25846</td>
<td>M</td>
<td>374</td>
<td>IMM</td>
<td>29 July</td>
<td>83</td>
</tr>
<tr>
<td>2005</td>
<td>05-57591</td>
<td>F</td>
<td>275</td>
<td>IMM</td>
<td>5 July</td>
<td>283</td>
</tr>
<tr>
<td></td>
<td>05-57593</td>
<td>F</td>
<td>350</td>
<td>ADU</td>
<td>10 July</td>
<td>157</td>
</tr>
<tr>
<td>ECS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1998</td>
<td>98-11035</td>
<td>M</td>
<td>440</td>
<td>ADU</td>
<td>26 June</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>98-2284</td>
<td>M</td>
<td>432</td>
<td>ADU</td>
<td>28 June</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>98-11036</td>
<td>M</td>
<td>398</td>
<td>ADU</td>
<td>29 June</td>
<td>98</td>
</tr>
<tr>
<td></td>
<td>98-2285</td>
<td>M</td>
<td>415</td>
<td>ADU</td>
<td>29 June</td>
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<tr>
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<td>98-2282</td>
<td>M</td>
<td>414</td>
<td>ADU</td>
<td>1 July</td>
<td>58</td>
</tr>
<tr>
<td>1999</td>
<td>99-11035</td>
<td>M</td>
<td>418</td>
<td>ADU</td>
<td>30 June</td>
<td>85</td>
</tr>
<tr>
<td></td>
<td>99-11036</td>
<td>F</td>
<td>266</td>
<td>IMM</td>
<td>30 June</td>
<td>73</td>
</tr>
<tr>
<td></td>
<td>99-11037</td>
<td>M</td>
<td>424</td>
<td>ADU</td>
<td>30 June</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td>99-11041</td>
<td>M</td>
<td>424</td>
<td>ADU</td>
<td>30 June</td>
<td>84</td>
</tr>
<tr>
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<td>01-2093</td>
<td>M</td>
<td>381</td>
<td>ADU</td>
<td>3 July</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td>01-2094</td>
<td>F</td>
<td>359</td>
<td>ADU</td>
<td>3 July</td>
<td>17</td>
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<tr>
<td></td>
<td>01-11038</td>
<td>F</td>
<td>316</td>
<td>IMM</td>
<td>5-Jul</td>
<td>145</td>
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<tr>
<td></td>
<td>01-11041</td>
<td>M</td>
<td>324</td>
<td>IMM</td>
<td>5 July</td>
<td>145</td>
</tr>
<tr>
<td></td>
<td>01-2280</td>
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<td>335</td>
<td>IMM</td>
<td>5 July</td>
<td>107</td>
</tr>
<tr>
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<td>01-11037</td>
<td>M</td>
<td>340</td>
<td>ADU</td>
<td>7 July</td>
<td>131</td>
</tr>
<tr>
<td></td>
<td>01-228</td>
<td>M</td>
<td>320</td>
<td>IMM</td>
<td>7 July</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>01-2282</td>
<td>M</td>
<td>373</td>
<td>ADU</td>
<td>7 July</td>
<td>35</td>
</tr>
<tr>
<td>2002</td>
<td>02-11036</td>
<td>M</td>
<td>320</td>
<td>IMM</td>
<td>7 July</td>
<td>66</td>
</tr>
<tr>
<td></td>
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<td>IMM</td>
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<td>80</td>
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<tr>
<td></td>
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<td>IMM</td>
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<td>62</td>
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<tr>
<td>2007</td>
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<td>ADU</td>
<td>1 July</td>
<td>132</td>
</tr>
<tr>
<td></td>
<td>07-36516</td>
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<td>ADU</td>
<td>1 July</td>
<td>125</td>
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<tr>
<td></td>
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<td>ADU</td>
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<td>320</td>
</tr>
<tr>
<td>2010</td>
<td>10-22117</td>
<td>M</td>
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<td>IMM</td>
<td>30 June</td>
<td>157</td>
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<td></td>
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<td>345</td>
<td>ADU</td>
<td>30 June</td>
<td>99</td>
</tr>
<tr>
<td>2012</td>
<td>12-108772</td>
<td>F</td>
<td>328</td>
<td>ADU</td>
<td>9 July</td>
<td>303</td>
</tr>
</tbody>
</table>
Mapping beluga whale dive behavior

Beluga dive characteristics, including deepest (‘maximum’) and modal depth and longest dive duration, were identified for each 6 h period and mapped using ArcGIS 10.1 (ESRI). Prey searching and capture may occur throughout the water column, but we examined the deepest and modal depths in each 6 h period as indicators of where diving was focused as presumed foraging effort, both extracted from the maximum-depth histograms. Maximum dive depth has been used as a proxy for specific foraging depth in other marine mammals (e.g. Photopoulou et al. 2014), and we also examined modal depths to consider which depths were most frequently used. We assumed the longest dive durations, extracted from the duration histograms for each 6 h period, were associated with deeper dives indicative of foraging effort (e.g. Davis et al. 2013). However, dive depth and duration were not directly linked, and we cannot verify that longer dives were also deeper dives (Citta et al. 2013).

The maximum daily dive depths from the 2007–2012 ECS whales were also mapped to describe deep diving at a higher resolution than previously available or possible from the histogram data. We presumed that maximum daily depth also provided an indicator of the deepest depths at which foraging may occur. We compared maximum daily dive depth among regions using linear mixed effects (LME) models fit with maximum likelihood methods in the ‘nlme’ package (Pinheiro et al. 2013) in R (R Development Core Team 2012). Individual whales were treated as a random effect to account for repeated measures, and regions were considered a fixed effect. We excluded the Canada Basin eastern slope region from the LME due to small sample size.

The depth associated with each 6 h period’s location was extracted from the 500 m resolution International Bathymetric Chart for the Arctic Ocean (IBCAO; Jakobsson et al. 2012). Some dive locations (n = 74, 1 %) occurred south of the IBCAO extent in the northern Bering Sea, where we used the 1 arc-minute ETOPO1 global relief map (Amante & Eakins 2009). We also recorded whether the deepest and modal depths in a 6 h period overlapped with the depth of the seafloor (categorical yes/no), the proximity to the seafloor as a ratio of the modal depth in a 6 h period relative to seafloor depth (range 0–1), as well as the ratio of maximum daily dive depth to seafloor depth. On rare occasions, dives could be deeper than seafloor depth because of the matching of whale location, diving depth, and seafloor depth at different spatial and temporal scales. A ratio of 1 was assigned when the beluga dive depth equaled or exceeded the ocean depth for a given location. We assumed ratios >0.9 represented benthic diving that targeted the seafloor. Close proximity to the seafloor, measured by dive to seafloor depth ratios, has previously been used to indicate benthic foraging for seals and the closely-related narwhal Monodon monoceros (Jessopp et al. 2013, Watt et al. 2015). We assumed ratios <0.9 corresponded to pelagic dives targeting other depths in the water column.

Regional daily diving time budget model

A daily diving time budget model was used to assess how individual belugas allocated time among depth layers within each region. We followed the approach developed for narwhals by Laidre et al. (2003) to estimate time spent both transiting through (i.e. transit time) and within (i.e. occupancy time) each depth layer. We tested differences in regional occupancy time at depth, after eliminating estimated time transiting to and from each depth, to assess population, sex, and age class differences. We presumed that extended occupancy time at deeper depths is related to foraging (Laidre et al. 2003). The model was based on scaling 6 h dive periods up to a 24 h period and used all 3 histogram types (i.e. maximum-depth, time-at-depth, and duration histograms). Since transmission of dive data were sometimes fragmented by poor satellite reception, analyses were restricted to 6 h periods when we had complete records of all 3 histogram types. Similar to above, we used a consolidated depth layer classification scheme that matched both maximum-depth and time-at-depth histogram types across all years (see Table S1): 0–10, 10–50, 50–100, 100–200, 200–400, and >400 m.

The time budget model required estimation of several parameters within each region for each tagged whale. Each parameter will be described briefly, using the same nomenclature as Laidre et al. (2003). First, region-specific vertical transit time (TTj) was estimated as the average time spent transiting through each depth layer (j) per day. Vertical transit time included time spent ascending and descending through each depth layer and relied on first estimating a population- and depth-specific vertical transit speed (Sz) for each region (Heide-Jørgensen et al. 1998).

We estimated depth-specific vertical speeds for each population and region to account for potential behavioral differences among regions and between populations. Estimating a depth-specific vertical speed
required a depth layer \((j)\) to be isolated such that dives were counted in the previous \((j - 1)\) and next \((j + 2)\) depth layers to estimate the speed through layer \(j\). Sample size for vertical speed estimation was thus smaller than for the other daily time budget model parameters because there are fewer histograms that fit the constraint for isolated depth layers, particularly at each depth for all regions and both populations. This requirement limited the inclusion of regions for which a small number of samples were acquired, so we consolidated regions for vertical speed estimation for each population based on similar habitat types and used bootstrapping. The estimated population- and depth-specific vertical speeds for each habitat type were sampled with replacement 1000 times.

To estimate TT\(_j\) for each of the main regions, we applied the vertical speeds from the respective population and habitat type. Occupancy time (OT\(_j\)) within each depth layer was estimated by subtracting TT\(_j\) from the total time spent within each depth layer \(j\) each day. Resulting OT\(_j\) values were the mean number of minutes a whale spent within each depth layer per day after removing transit time to and from that depth. To estimate occupancy time on a per dive basis (average occupancy time per dive, AOT\(_{dive,j}\)), OT\(_j\) was divided by the average number of dives per day to depth layer \(j\).

For each region, we compared mean population responses of individual differences in AOT\(_{dive}\) using 2-way ANOVA. Factors included depth layers (i.e. 0–10, 10–50, 50–100, 100–200, 200–400, and >400 m), population (BS and ECS) and their interaction. Limited sample size of BS whales precluded analysis of differences between sex or age classes, but we also tested differences in AOT\(_{dive}\) between ECS sexes and between ages (i.e. adult and immature ECS whales) in additional ANOVAs. We applied a square-root transformation to each test of AOT\(_{dive}\) and evaluated model structure using Akaike’s information criterion (AIC). Applying the Bonferroni correction for multiple comparisons, statistical significance of ANOVAs was considered at \(\alpha = 0.002\).

### Estimation of the vertical distribution of Arctic cod density in the western Beaufort Sea

We estimated densities of age 1+ Arctic cod within each beluga whale depth layer from 38 kHz data collected during an acoustic-trawl survey in the western Beaufort Sea during 16 to 21 August 2008 (Parker-Stetter et al. 2011; see trackline in Fig. 1). Age 1+ Arctic cod (hereafter simply ‘cod’) constituted >99% of the Marinovich trawl (fishing dimensions 3–4 m vertical by 6 m horizontal) catches used to verify the species and size composition of fish within the midwater acoustic backscatter (\(n = 14\), Parker-Stetter et al. 2011), and cod was the dominant fish species across several recent surveys in the Beaufort Sea (Logerwell et al. 2015). The relatively small number of detections of age-0 Arctic cod (i.e. young of the year) or other species were not analyzed here and are not considered part of BS or ECS diets (Quakenbush et al. 2015). Integrated volume backscatter (area backscattering coefficient; \(s_a\) in m² m⁻²) of cod was exported in 10 m vertical depth bins, between 9.0 m below the surface and 0.5 m above the bottom, at a 500 m horizontal resolution. Density (no. fish m⁻²) was then calculated for each 10 m bin by dividing the integrated backscatter by a target strength of −52.7 dB re 1 m², estimated by Parker-Stetter et al. (2011). Cod densities in 10 m bins were then summed to vertical depth layers (i.e. 0–10, 10–50, 50–100, 100–200, 200–300, 300–400, >400 m) matching those used to identify the number of beluga dives made during 6 h periods (i.e. the 6 h maximum depth histograms). Cod densities along the survey trackline were next interpolated to create a smoothed surface of cod density for each beluga depth layer. We used ordinary kriging in the Geostatistical Analyst extension of ArcGIS 10.1, fitting a spherical model with no trend removal for each depth layer. Interpolation of cod density for each depth layer was limited to the bounds of the cod survey area other than the landward limit, which was set by the 100 m isobath used by Parker-Stetter et al. (2011) when estimating cod target strength. This landward limit was supported by beluga biology, since belugas detected by aerial surveys were most frequently located offshore of the 100 m isobath in the Alaska Beaufort Sea (e.g. Moore et al. 2000). We limited seaward interpolation of cod densities to the minimum depth threshold of each depth layer (e.g. the minimum isobath of the 200–300 m depth layer was 200 m) when interpolating densities of cod in the deeper depth layers (i.e. 200–300, 300–400, and >400 m), and we refer to these as the >200, >300, and >400 m cod ‘survey areas,’ respectively.

### Statistical analysis of ECS beluga dive behavior relative to Arctic cod density

We compared the pattern of beluga dive behavior to the depth-specific estimates of cod density within
the cod survey area. A subset of 192 ECS beluga dive records that occurred within the survey area were used in this analysis. Few (n = 4) BS 6 h dive periods occurred in the cod survey area, which precluded similar comparisons. We were particularly interested in diving activity targeting the 100–200, 200–300, and 300–400 m depth layers, because we inferred that foraging likely occurred at these depths. Specifically, ECS belugas had prolonged occupancy time as well as maximum and modal dives to these depths in Barrow Canyon and the Alaska Beaufort Sea regions coinciding with the survey area (see ‘Results’). These ‘target’ depths also match depths where fronts and upwelling occur in this area and promote the concentration of zooplankton, thereby presumably attracting beluga prey (Pickart et al. 2013, Stafford et al. 2013). Our goal was to examine whether the number of dives to these target depth layers corresponded with the depth layers at which cod were most abundant, indicating a relationship between inferred foraging dive depths and presumed prey patterns.

We used ECS beluga 6 h dive period locations that occurred within the cod survey area during July to October, when ECS belugas use Barrow Canyon and the Alaska Beaufort Sea (Hauser et al. 2014) and representing the period that is characteristic of when belugas dive to these target depth layers. We intersected the locations for each 6 h dive period within the >200 and >300 m cod survey areas with each interpolated cod density depth layer. Generalized linear mixed models (GLMMs) were used to model the number of dives to specific target depth layers (i.e. 100–200, 200–300, and 300–400 m) using the ‘lme4’ package in R (Bates et al. 2014). The number of dives to target depth layers was modeled using a Poisson error distribution and log link. Individual whales were considered a random effect to account for repeated measures and individual variation. We tested dive rates to 100–200 and 200–300 m depth layers in the >200 m survey area and also modeled dive rates to 200–300 and 300–400 m in the >300 m survey area. Explanatory variables for each GLMM included additive effects of cod densities in 10–50, 50–100, 100–200, and 200–300 m depth layers and log-transformed seafloor depth. We also included 2 additional predictor variables when modeling within the >300 m survey area that were not appropriate for models in the >200 m survey area: cod density in the 300–400 m depth layer, and a categorical variable identifying which depth layer contained the maximum cod density. Final model structure was determined via AIC model selection.

RESULTS

Four BS and 26 ECS beluga whales were tagged, resulting in 1600 and 7923 maximum depth 6 h histogram periods, respectively, within 10 Pacific Arctic regions (Tables 1 & 2). For the BS whales, 1561 time-at-depth and 1610 duration 6 h histogram periods were also collected. We collected 8075 time-at-depth 6 h histogram periods and 7823 duration 6 h histogram periods for ECS whales. Belugas from both populations ranged across all but 1 of the Pacific Arctic regions (Table 1, Fig. 2). No dive data were collected in the Viscount Melville Sound BS summer core area, although it is typically used by BS males in July and August (Hauser et al. 2014). The majority of BS 6 h dive periods were obtained from the Chukchi Sea shelf (34%), while the greatest percentage (31%) of ECS 6 h dive periods were collected from the deep Canada Basin, Barrow Canyon slope (24%), and Chukchi Sea shelf (21%; Table 1). Seasonal presence varied among regions, and we are the first to document winter (November to April) and spring (May to June) behavior and locations of tagged belugas in the Bering Sea (Table 1, Fig. 2).

Spatial patterns of dive behavior

Geographically referenced dive data revealed differences in dive behavior among regions and between populations (Fig. 2). Maximum dive depth in 6 h periods was similar between populations for each region, ranging >400 m in deep pelagic habitats (i.e. Canada Basin) where maximum dive depth never reached the seafloor. In slope areas, maximum dive depths were often >400 m and typically (63 and 65% of 6 h periods for BS and ECS, respectively) reached the seafloor (Fig. 3). However, ECS belugas typically dove to mid-water depth layers (100–200 and 200–300 m) in the Barrow Canyon and Alaska Beaufort slope regions (Fig. 2). BS belugas also often dove to mid-water depths (200–300 m) in certain slope regions like Amundsen Gulf. Maximum dive depth in shelf areas was most frequently to 10–50 m (62% of 6 h periods) and to the seafloor (91% of 6 h periods) for ECS belugas. BS beluga shelf maximum dive depth was more frequently to 50–100 m (60% of 6 h periods) to typically reach the seafloor (89% of 6 h periods).

Modal depths of 6 h periods were most frequently in the <10 and 10–50 m depth layers for both populations, regardless of habitat type (see Fig. S1). Dives <10 m represented surface-oriented dives (Citta et al.
so we excluded 6 h periods with <10 m modal depths in subsequent modal dive depth analyses (resulting in n = 962 and n = 2808 BS and ECS 6 h periods, respectively). We most often observed modal dives in the 10–50 m depth layers for both populations in all habitat types (Fig. 3), but deeper dive depths were also common in some regions (Fig. 2).

ECS modal depths in the 10–50 m depth layer (95% of 6 h periods) were typically benthic dives (90% of 6 h periods) within shallow shelf habitats (Fig. 3). BS modal depths in shelf habitats were common in the 10–50 and 50–100 m depth layers (50 and 43% of 6 h periods, respectively), and were less frequently benthic (70% of 6 h periods) than those of ECS belu-
gas in shelf habitats. BS modal dive to seafloor depth ratios were typically pelagic in the northern and northwestern Chukchi Sea (Fig. 2). In slope and deep pelagic habitats, modal depths for both populations were typically not benthic for BS (63% and 100% of 6 h periods, respectively) or ECS belugas (59% and 100% of 6 h periods, respectively), occurring most often in the 10−50 m depth layer but also in deeper depths (Fig. 3). Deeper modal depths were common in 100−200 and 200−300 m depth layers for ECS belugas in slope (22% and 9% of 6 h periods, respectively) and basin (10% and 4%, respectively) habitats, especially in the Barrow Canyon and Alaska Beaufort Sea (Fig. 2). Additional modal depths for BS belugas were in the 50−100 and 300−400 m depth layers (23% and 8% of 6 h periods, respectively) in slope habitat, and >400 m in deep pelagic habitat (20% of 6 h periods).

Maximum dive durations within a 6 h period were greater for ECS than BS whales (Fig. 2). Maximum dive duration was frequently >20 min for ECS whales, while BS maximum dive durations were often 6–9 min throughout the Beaufort and Chukchi Seas and the Amundsen Gulf, compared to 9–20 min in the Canada Basin and the Bering Sea. Dive-depth and duration histograms were not directly linked within a 6 h period, so it was not possible to examine whether longer dives were also deeper dives.

Maximum daily depths recorded for 6 ECS tags deployed in 2007 to 2012 also varied regionally (LME, $F_{6,16} = 375.4, p < 0.0001$) and never exceeded the 1000 m limit of their tags (Fig. 4). The deepest maximum daily depth recorded was 956 m for an adult male, although 2 adult females also attained maximum daily depths >900 m. The ratio of the maximum daily dive depth to ocean depth indicated that these whales were diving to the seafloor at least daily in all regions but the Canada Basin where seafloor depths ranged >3000 m. Within the Canada Basin, ECS belugas generally dove in excess of 600 m at least once daily (mean = 741.4 m).

**Occupancy time at depth**

Depth-specific vertical speeds ($S_j$) increased with depth for both populations in each habitat type (Fig. 5) and were similar in magnitude to those estimated for High Arctic belugas and narwhals (Heide-Jørgensen et al. 1998, Laidre et al. 2003). There were no significant differences in $AOT_{dive,j}$ between populations within any region, but there were significant differences in $AOT_{dive,j}$ among depth layers in all but the shallow shelf regions (i.e. the Bering and Chukchi Seas; Table 3, Fig. 6). Similarly, no significant differences between ECS male and female or immature and adult whales occurred in any region except the Canada Basin, where adult whales spent significantly more time at 200–400 and >400 m depth layers than immature whales, presumably because adult whales have larger oxygen stores and aerobic capacity than smaller-bodied immature whales (Schreer & Kovacs 1997, Noren & Williams 2000). Occupancy times were greatest in the 200−400 m depth layer in the deep pelagic basin and some slope regions (Fig. 6). However, whales also spent more time in the 100−200 m depth layer in the Barrow Canyon and Alaska Beaufort slope regions.

**Vertical distribution of Arctic cod and beluga diving in the Alaska Beaufort Sea**

Age 1+ Arctic cod occurred throughout the water column during the fish survey, but the 200−300 m depth layer contained the highest cod densities of all depth layers regardless of water depth (see Fig. S2). This supports suggestions by Parker-Stetter et al. (2011) that cod aggregations extend into the Canada Basin in a horizontal pelagic layer as the seafloor drops steeply along the continental slope. Coincident with depth layers where cod were most dense, the greatest number of ECS beluga dives occurring within the cod survey area also targeted the 200−300 m depth layer other than dives closer to the surface in 10−50 m (Fig. 7). Maximum and modal dive depths also most frequently occurred in the 200−300 m depth layer, in addition to a mode near the surface at 10−50 m.

The number of ECS beluga dives to 100−200, 200−300, and 300−400 m target depth layers was significantly related to the vertical distribution of cod density (Table 4). Cod density in 200−300 m was a consistently significant predictor for dive rates to each target depth, but cod density at other depths and seafloor depth were also related to dive rate for some models. Dive rate to a specific target depth was always negatively correlated with cod density at that same depth.

**DISCUSSION**

For marine mammals, targeted diving depths or extended occupancy at depth are often assumed to indicate foraging or the presence of predictable prey resources that may vary regionally or seasonally
(Laidre et al. 2003, Aguilar Soto et al. 2008, Robinson et al. 2012). In our study, spatial variation in diving behavior for BS and ECS beluga whales suggested different foraging strategies among regions and, in some cases, between populations. In the absence of high-resolution dive data, lower-resolution satellite-relayed data loggers, such as those used in our study, can still reliably estimate foraging behavior (e.g. Heerah et al. 2015). We determined foraging based on the maximum duration of dives, maximum (at 6 h and daily time scales) and modal dive depths, the proximity of dives to the seafloor, and the time spent within each depth layer. In studies where foraging can be confirmed, the deepest and longest maximum
dive depths and durations are often associated with dives that have prey captures (e.g. Davis et al. 2013). The higher vertical speeds estimated here for deeper depths also support the assumption that foraging occurs at deeper depths. For example, high speeds at depth for some odontocetes coincide with foraging buzzes for particular prey types and the deepest parts of a dive (Aguilar Soto et al. 2008). Similarly, maximum dive depth serves as an indicator of foraging depth for benthic foragers (Photopoulou et al. 2014), and close proximity to the seafloor (or not) can be confirmed by comparing dive depth to water depth (Jessopp et al. 2013, Watt et al. 2015). Extended time at target depths has also been used to identify depth layers where foraging is expected to occur (Laidre et al. 2003). Our depth and duration histogram data could not be directly compared to indicate whether deeper dives were also longer.

Table 3. Results of ANOVAs (p-values) comparing regional average occupancy time per dive (AOT) among depth layers and between Eastern Chukchi Sea (ECS) beluga whales Delphinapterus leucas in the Chukchi and Bering Seas (a,b) shallow shelf, (c,d) slope, (e,f) deep pelagic Canada Basin, and (g) Mackenzie Estuary and Amundsen shelf and slope habitat types. Lines indicate the standard deviation of vertical transit speed. No ECS belugas were found in the Mackenzie Estuary or Amundsen Gulf. No samples of \( S_j \) were obtained for BS belugas in the 200–400 m depth layer in the Chukchi and Bering Seas (a) shallow shelf or (c) slope habitats, so the corresponding estimates from ECS whales were applied to the estimation of average occupancy time per dive (AOT) in these cases.

![Fig. 5. Estimated mean vertical transit speed (\( S_j \); bars) among depth layers (shown as the maximum depth threshold) for (a,c,g) Beaufort Sea (BS) and (b,d,f) Eastern Chukchi Sea (ECS) beluga whales Delphinapterus leucas in the Chukchi and Bering Seas (a,b) shallow shelf, (c,d) slope, (e,f) deep pelagic Canada Basin, and (g) Mackenzie Estuary and Amundsen shelf and slope habitat types. Lines indicate the standard deviation of vertical transit speed. No ECS belugas were found in the Mackenzie Estuary or Amundsen Gulf. No samples of \( S_j \) were obtained for BS belugas in the 200–400 m depth layer in the Chukchi and Bering Seas (a) shallow shelf or (c) slope habitats, so the corresponding estimates from ECS whales were applied to the estimation of average occupancy time per dive (AOT) in these cases.](image)
dives, but our estimates of depths with prolonged occupancy time corresponded well with maximum dive depths. Lastly, we estimated modal dive depths to identify the most common depths used by belugas.

Non-foraging diving behavior

Analysis of modal dive depths was complicated by the high frequency of shallow dives in the 0–10 and 10–50 m depth layers (e.g. see Fig. S1). Citta et al. (2013) found that ECS belugas typically had a mode at the surface and another in a deeper layer. Bimodal distributions of maximum-depth histograms were still common after excluding dives <10 m, and the deeper of the 2 modes were used in analyses. We considered all depths in estimating modal dive depths, but we focused on modal dives >10 m since we assumed dives in the upper 10 m were associated with surfacing behavior as in Citta et al. (2013). The 10–50 m depth layer was still often a primary mode, even when there may have been a second deeper mode. We assumed dives in the 10–50 m depth layer likely included surface-oriented behavior such as
travel or recovery diving. Richard et al. (1997) provided evidence that shallower dives represent dive recovery or migration, reporting that BS belugas in areas 15–600 m deep and >600 m would dive deeply followed by long periods with dives <50 m. Histograms and dive records from other deep-diving predators also indicate concentrated and prolonged periods at shallow depths, consistent with travel and dive recovery following deep foraging dives (e.g., Laidre et al. 2003, Arranz et al. 2011). However, we cannot exclude the possibility that foraging occurred at <50 m, especially in shallow shelf habitats where water depths are often ≤50 m, yet a deeper secondary mode is presumably associated with foraging in slope and deep pelagic habitats.

The coastal and shallow summer core areas of Kasegaluk Lagoon (mean depth ~7 m) and the Mackenzie River Estuary (mean depth ~25 m) are used by ECS and BS belugas, respectively, in spring to early summer during annual migrations (Hauser et al. 2014). Stomachs of belugas harvested during subsistence hunts near these regions are typically empty (Harwood & Smith 2002, Quakenbush et al. 2015), although intestines were not examined. Histological studies of liver and kidney tissues from whales harvested in the areas also indicate fasting (Woshner et al. 2002). It seems unlikely that belugas forage in these coastal core areas and rather use them for an annual spring/summer molt where warmer and fresher water, such as near estuaries, accelerates epidermal regrowth (St. Aubin et al. 1990). Calving also occurs in spring, and shallow areas may provide thermal benefits to calves or provide additional protection from killer whale Orcinus orca predation (Finley 1982).

In contrast, the Canada Basin is a uniquely deep (>3000 m), pelagic, and historically ice-covered habitat where both populations had maximum dive depths >400 m (see Figs. 2 & 3) and extended occupancy time at 200–400 and >400 m (see Fig. 6). The average depth of maximum daily dives by ECS belugas (741 m) also suggested deep, pelagic diving (see Fig. 4). Richard et al. (1997) reported that male BS dives in the Canada Basin were typically ‘V-shaped’ to 700–900 m for 15 to 20 min and accompanied by many dives <50 m, suggesting that these deep dives involved a whale reaching its maximum depth before immediately heading to the surface without searching for prey at depth. They hypothesized that whales may have used these dives to locate small breathing holes in the dense pack ice during ascent from depth and noted that V-shaped dives were not detected elsewhere. In addition to maximum dives >400 m in Canada Basin, we found a secondary mode at >400 m for BS belugas (see Fig. 3) that would support the description by Richard et al. (1997). ECS belugas may also have V-shaped dives based on their maximum dive depths, but ECS whales had additional modes at 100–200
and 200−300 m in the Canada Basin (see Fig. 3) that may be associated with foraging. Relatively little is known about the distribution or abundance of potential prey in the Canada Basin, but boundary currents along the Chukchi and Beaufort shelves create mesoscale eddies that infuse nutrients and zooplankton into the region that could fuel higher trophic levels (Llinas et al. 2009). A stratified front is located ~200−250 m between cold, dense Pacific water and warmer Atlantic water (Pickart 2004, Nikolopoulos et al. 2009) that could aggregate zooplankton, thereby attracting forage fish like Arctic cod (Logerwell et al. 2011). A dense layer of Arctic cod likely extends off the Beaufort Sea slope into the Canada Basin around ~250 m depth (Fig. S2; Parker-Stetter et al. 2011). Another front is located between ~600 and 1000 m as the Atlantic water transitions to Deep Arctic water (McLaughlin et al. 1996). Ultimately, the functional importance of the Canada Basin remains somewhat unclear and may differ between BS and ECS belugas.

### Benthic diving behavior

Our results indicated that both beluga populations primarily targeted either benthic or pelagic portions of the water column depending on the geographic region. Both maximum and modal dive depths suggested that both populations usually make benthic dives in shallow shelf habitats like the Chukchi and northern Bering Seas, but pelagic diving was more common in the deep Canada Basin (e.g. see Fig. 3). Stable isotope values from BS and ECS belugas suggest reliance on both benthic and pelagic prey (Dehn et al. 2006, Horstmann-Dehn et al. 2012), which fits the mixed pattern of benthic and pelagic diving among regions in our results. Benthic invertebrates (e.g. shrimp and echiurid worms) also dominated stomach contents of BS and ECS belugas sampled in spring and early summer when whales would have most recently been in the northern Bering or eastern Chukchi Seas (Quakenbush et al. 2015), further supporting a conclusion that benthic foraging occurs in

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<td>Depth of maximum cod density 200−300 m</td>
<td>−1.028</td>
<td>0.497</td>
<td>0.039</td>
</tr>
<tr>
<td></td>
<td>300−400 m</td>
<td>0.574</td>
<td>0.508</td>
<td>0.259</td>
</tr>
</tbody>
</table>
these shallow shelf habitats. The northern Bering and Chukchi Seas are considered productive benthos-dominated systems, with abundant and diverse benthic macrofauna (Grebmeier et al. 2006, Day et al. 2013) and benthic age 1+ Arctic cod (Norcross et al. 2013) that are potential beluga prey.

A small proportion of 6 h periods were characterized by non-benthic diving in the Chukchi and Bering Seas for BS belugas (see Fig. 2), which coincided with the deeper northern portions of the Chukchi Plateau and Herald Canyon. Although Herald Canyon is less affected by wind-driven fronts than Barrow Canyon in the eastern Chukchi Sea, easterly winds contribute to continental shelf waves over the western Chukchi Sea that can affect the transport of zooplankton and may have important effects on benthos as well as forage fish (Danielson et al. 2014). Relatively little is known about the distribution and abundance of potential beluga prey in the western Chukchi Sea. Similarly, circulation in the northwest Bering Sea fluctuates depending on wind stress, which is also a dynamic marginal sea ice zone in winter when BS belugas were present (Danielson et al. 2014). Few tags continued transmitting during winter in the Bering Sea, and additional work is needed to identify habitat use and foraging within this period.

In slope habitats, diving generally appeared to target the seafloor but also portions of the water column in certain regions. Maximum dive depths typically reached the seafloor (see Fig. 3) but varied depending on finer regions for the 2 populations (see Fig. 2). BS beluga diving focused on the seafloor in the Alaska Beaufort Sea and Barrow Canyon, while ECS belugas had more pelagic 6 h maximum and modal depths that corresponded to a frontal transition zone between Pacific and Atlantic water (Pickart 2004, Nikolopoulos et al. 2009) along the steep continental slope where age 1+ Arctic cod are most abundant (Parker-Stetter et al. 2011, Crawford et al. 2012). Both beluga populations made deeper dives in the Canada Beaufort slope region than in the Alaska Beaufort Sea, but rarely to the seafloor. Prolonged occupancy time at depth (see Fig. 6) as well as pelagic maximum and modal dive depths corresponded to the same depths of dense Arctic cod schools surveyed during summer and fall of 2006 to 2012 in the Canada Beaufort Sea and Amundsen Gulf (Geoffroy et al. 2015, Majewski et al. in press). However, Richard et al. (1997) reported 10 to 20 min ‘square-shaped’ dives to the seafloor by BS belugas in waters 15–600 m deep and assumed these were demersal foraging dives. These dives would typically be followed by several minutes at shallow depths or were conducted in batches trailed by multiple hours at <50 m. Nearly 37% of the modal dives we observed for BS belugas in slope waters reached the seafloor and may have represented similar square-shaped foraging dives. Our slightly different categorization of slope habitat as well as our sample of tagged whales may account for the apparent differences in our results. Of note, the sample of whales included in the study by Richard et al. (1997) was largely male and focused on use of the 600 m deep BS core area in Viscount Melville Sound compared to our largely female (and 1 immature male) sample that remained in Amundsen Gulf and the Canada Beaufort Sea before migrating west.

**Maximum dive duration**

ECS belugas almost universally had greater maximum dive durations than BS whales (see Fig. 2). The general similarities in diving behavior between populations is suggestive of comparable foraging patterns at depth within a given region, although spatiotemporal differences in BS and ECS movements (Hauser et al. 2014) would limit overlap and potential competition for prey at depth. Whales from the BS population are comparably sized to ECS whales (Suydam 2009), but our sample of tagged BS whales was biased toward females, with the only male being immature. Thus the difference in population-specific dive duration was likely due to the small sample of tagged BS whales. Additional tagging is needed to further compare dive durations between populations.

**Beluga dive patterns relative to the vertical distribution of cod**

Even with temporal mismatch between predator and prey sampling, significant relationships between predator diving and prey abundance can be detected (Kuhn et al. 2015). Despite asynchronous sampling between ECS beluga diving and cod density data, we assumed that observed patterns were representative of typical conditions. Several aspects of beluga behavior, diet, and regional hydrography support this assumption. Belugas are long-lived, social cetaceans with matrilineally derived migration routes and philopatry (O’Corry-Crowe 2008, Turgeon et al. 2012), suggesting that the same regions are used each summer by related social groups. ECS belugas exhibit strong interannual philopatry for areas in the
Barrow Canyon and Alaska Beaufort Sea, areas which overlap with the fish survey area (Hauser et al. 2014). Beluga diving in the area targeted depths where fronts occur (e.g. occupancy time, maximum dive depth, dive rates to target depths, Citta et al. 2013). These persistent stratified fronts function to concentrate secondary production that likely attracts prey like Arctic cod and thus likely promote foraging opportunities for belugas in this area (Stafford et al. 2013). A simultaneously occurring benthic trawl survey supported the high biomass of cod found here and also indicated an association between Arctic cod and fronts (Logerwell et al. 2011). However, there have been few systematic surveys of cod in the Alaska Beaufort Sea for comparison to the distribution we observed (e.g. Lowry & Frost 1981), and the area surveyed in this study likely did not fully sample the offshore extent of cod schools (Parker-Stetter et al. 2011). Yet age 1+ Arctic cod have dominated catches elsewhere in the eastern Chukchi and Beaufort Seas, especially on continental slopes that were consistently sampled in numerous years (e.g. Norcross et al. 2010, Crawford et al. 2012). Collectively, both pelagic and benthic surveys support our assumptions of cod density, suggesting a persistent pattern of high Arctic cod biomass in our survey area and throughout the Pacific Arctic (see also Logerwell et al. 2015).

In this study, depths of maximum and modal beluga dives were similar to depths with maximum and high mean cod densities (see Fig. 7). While recognizing that beluga dive behavior and cod distribution were not sampled concurrently, these results add weight to the use of our dive parameters as indicators of foraging behavior. The depth layer with the highest cod density (200–300 m) was a consistent, significant predictor of the number of ECS beluga dives (see Table 4) and suggested an overlap between beluga dive depths and high densities of Arctic cod, at least for ECS belugas in the survey area. Even though Arctic cod density consistently predicted the number of dives to target depths, we cannot exclude the possibility that other prey are pursued in this area. For example, belugas also dove to the seafloor and depth layers deeper than 200–300 m, although with much lower frequency.

We hypothesized that the number of beluga dives would increase with the density of cod, but we found a negative correlation between beluga dive rate to a target depth and cod density in that target or shallower depths. In contrast, there was a positive correlation with cod density at depths deeper than the target depth. This result suggests that belugas dive less frequently when cod are abundant because foraging is efficient at high prey densities, consistent with marginal value theorem predictions of optimal foraging (Charnov 1976). Empirical tests have demonstrated that other marine predators reduce foraging time in high-quality prey patches (e.g. Thums et al. 2013), and higher prey abundance improves foraging efficiency for deep divers (Goundie et al. 2015). For Pacific Arctic belugas, additional data are needed that incorporate simultaneous sampling of beluga behavior, cod distribution, and other potential prey to quantify the costs and benefits of beluga foraging on cod prey.

**CONCLUSIONS**

We used satellite-linked time−depth recorders to infer depths at which BS and ECS beluga whales forage throughout diverse regions of the Pacific Arctic. Maximum and modal dive depths suggested demersal foraging on shallow shelves like the Chukchi and northern Bering Seas in contrast to pelagic diving in the deep Canada Basin for both populations. While maximum dive depths often reached the seafloor in slope habitats, portions of the water column were more frequently targeted in certain regions. We also found that patterns of ECS beluga diving coincided with the depths at which Arctic cod were most abundant during a fisheries acoustic survey in the western Beaufort Sea, thereby providing additional support for our classification of foraging dives.

Our regional and population-specific estimates of foraging behavior provide important baseline information in the context of a rapidly transforming Arctic ecosystem that is increasingly exposed to anthropogenic activities (Grebmeier 2012). Industrial and other human activities (e.g. shipping and other vessels, mining, fishing, and tourism) are expanding and can potentially affect belugas directly (e.g. hearing impairment or stress response) or indirectly alter behavior (Reeves et al. 2014). Although both populations are considered stable stocks (Allen & Angliss 2014), recent evidence for declining growth, body condition, and blubber thickness suggests that ecosystem changes may be affecting belugas through reduced availability or quality of prey (Harwood et al. 2014, 2015). Thus, understanding the behavioral use in each region offers a benchmark for which changes in beluga behavior can be assessed, as well as information to identify ecologically significant areas that may warrant protection.
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