Role of the bowhead whale as a predator in West Greenland

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ABSTRACT: We examined the spatial and temporal linkage between primary production, zooplankton distribution and density, and bowhead whale Balaena mysticetus foraging behavior in Disko Bay, West Greenland using concurrent ship-based oceanographic and net sampling together with instrumentation of whales with satellite-linked transmitters and dive recorders. Estimates of bowhead whale abundance were used in a bioenergetic model to calculate the potential consumption of zooplankton during their 4 mo stay in Disko Bay. Between 2001 and 2006, 30 whales were fitted with satellite transmitters that provided information on daily movements, and 14 whales were tracked with archival time–depth or time–depth–fluorescence recorders that provided detailed dive data. Simultaneous data were collected on water column structure, phytoplankton and zooplankton density, taxa and biomass at 25 stations south of Disko Island in 2003, 2005 and 2006. After the retreat of annual winter sea ice, bowhead whales explored a limited area along the south coast of Disko Island and had high interannual site fidelity. Mean dive depths varied between 53 (SD = 33) and 109 (SD = 41) m but maximum dive depths were >400 m. Most dives targeted the bottom and dive durations >40 min were observed for several whales. Available prey for bowhead whales was dominated by calanoid copepods, with Calanus finmarchicus, C. glacialis and C. hyperboreus occurring at 90 to 100% of all stations between 0 and 50 m and contributing 78 ± 25% of the total biomass. Bottom sampling for epizooplankton in 2006 resulted in unprecedented densities of C. finmarchicus, several orders of magnitude higher than at any other depth. Bioenergetic modeling indicated the population consumes ~220 US t of zooplankton per day or >21 000 t during the 4 mo stay in Disko Bay. Although the total biomass of zooplankton in the upper 50 m of the water column theoretically could support this predation level, benthic zooplankton densities and behavioral data suggest whales target pre-ascension stage epibenthic copepods in high density patches.

KEY WORDS: Bowhead whale · Foraging · Zooplankton · Calanus · Greenland

INTRODUCTION

A fundamental concept in marine ecology is that the fitness of a predator depends on the spatial and temporal synchrony with the production of its prey (Cushing 1974). This is especially true in seasonal environments where the cycle of temperature and solar radiation only allow for short production periods. The spring bloom in high Arctic marine ecosystems is characterized by relatively short, efficient food chains where the large Calanus copepod population grazes directly on the primary producers. After the spring sea ice recession, many marine predators exploit Calanus spp. for nutritional gain and build up of fat deposits. The largest zooplankton predator in this ecosystem is the bowhead whale Balaena mysticetus. Recent estimates...
of bowhead whale abundance off the coast of West Greenland suggest over a thousand whales visit the region between February and May each year (M. P. Heide-Jørgensen unpubl. data). Several hundred whales concentrate in the Disko Bay region (Fig. 1) where they feed intensively before migrating across Baffin Bay in May and June. Satellite tracking studies of bowhead whales have documented specific fidelity to the region south of Disko Island (Heide-Jørgensen et al. 2003, 2006). The highly predictable arrival and departure dates of these whales each spring has been observed dating back to the 1700s (Eschricht & Reinhardt 1866).

The Disko Bay population of bowhead whales is part of a larger segregated population that ranges into the Canadian high Arctic, Foxe Basin, and Baffin Bay (Heide-Jørgensen et al. 2006). Over 85% of the whales that visit Disko Bay are females and most are >14 m in length (Greenland Institute of Natural Resources unpubl. data). Despite this, few calves and juvenile whales have been observed, which is consistent with observations made during commercial whaling (Eschricht & Reinhardt 1866).

Disko Bay is located between the sub-Arctic waters of southwest Greenland and the high Arctic waters of Baffin Bay. It is influenced by both the northbound warm West Greenland current of Atlantic Ocean origin and the southbound currents of polar origin in Baffin Bay. The annual sea ice forms early in winter and reaches peak coverage in March. It retreats in April and May and the bay is free of ice by June. During winter, the water column is well mixed and the lack of daylight and ice coverage prevents net growth of the phytoplankton. However, in spring the growth of phytoplankton is exponential and fuels a trophic web that culminates with bowhead whales. Disko Bay is relatively deep (>400 m) and sedimentation of phytoplankton begins shortly after the peak bloom phase (Pedersen et al. 2006); however, frequent extreme wind mixing events reintroduce nutrients from below the pycnocline and refuel the phytoplankton in the surface layer.

Few studies have been conducted on the trophic coupling between whales and their prey, given the dynamic nature of the marine environment and the difficulty in observing or quantifying concurrent whale behavior and prey availability (Croll et al. 2005, Tynan et al. 2005). In the high Arctic, long-term climate change and interannual climate variability are expected to have strong indirect effects on apex predators through trophic decoupling (Laidre et al. 2007). However, a quantitative baseline understanding of the relationship between predators and their environment is required before these effects can be predicted. The primary purpose of this study was to obtain information on bowhead whale movements and diving behavior during the spring foraging period. These data were combined with energetic demands and simultaneous oceanographic and zooplankton sampling to understand how distribution, biomass composition and spatial variability in prey affect the springtime foraging success of the Arctic’s largest zooplankton predator.

**MATERIALS AND METHODS**

**Tracking instrumentation.** Thirty Bowhead whales were fitted with satellite tags (Telonics ST-15 and ST-16, or Wildlife Computers, SPOT1, SPOT2 and SPOT3 tags) in Disko Bay, West Greenland (Fig. 1). Whales were approached in small boats and tags were pushed under the skin into the blubber layer with an 8 m long fiberglass pole (for details see Heide-Jørgensen et al. 2003, 2006) or tags were deployed with an air gun (modified pneumatic line thrower). Tags were deployed in April and May 2001 to 2006 and location data were obtained from polar orbiting...
satellites via the ARGOS data collection system. A biopsy was taken from each animal and length was estimated visually.

Another 14 bowhead whales were also fitted with archival time-depth-recorders (TDRs; MK9, Wildlife Computers) in April or May of 2002, 2003 and 2005 and with WetLabs FLNTUB fluorometers in April or May of 2005 and 2006. Fluorometers and TDRs were mounted on cylindrical shaped floats (~28 × 8 cm) with a VHF transmitter (Telenics) and a satellite tag (Wildlife Computers SPOT4) for instrument recovery. Floats were attached to whales using a flat 20 × 25 mm stainless steel anchoring system modified from traditional Inuit harpoon heads (4 mm thick). All instrument packages were deployed using the pole. While the whale was pursued, the float tags were held in a PVC housing mounted to the pole. Once the harpoon head was imbedded in the blubber of the whale, the float was released from the housing. The float was tethered to a 1.5 m long stainless steel wire with a corrosive magnesium bolt that released the float from the whale after a predetermined period. Floats were located and recovered by means of the real-time ARGOS satellite and a fine scale VHF search conducted with directional antennas mounted on a 13.6 m boat (RV ‘Porsild’, Arctic Station, University of Copenhagen) or from a small dinghy. The TDR sampled pressure (depth) and temperature every second and had a total memory of 16 MB and the FLNTUB sampled fluorescence, pressure (depth), turbidity and temperature every second and had a memory of ~65 000 readings.

**Location and dive data analysis.** Average daily positions were calculated from good quality ARGOS data (LC-0 or better) for each whale. Area use during the residency period in Disko Bay, including all days after instruments were fitted to the whales up to their departure from Disko Bay towards Baffin Island (defined as north of 70° N), was estimated using the 95, 75 and 50% fixed kernel home range with least squares cross validation using the Animal Movement extension in ArcView (Hoodge & Eichenlaub 1997). All analyses were conducted with an equal area projection and the area where land overlapped home ranges was subtracted from the home range estimates.

Data from archival dive tags were downloaded for processing and analysis. Drift in the depth values on the TDRs was corrected using the software Instrument Helper v. 0.3 (Wildlife Computers) including both ‘Zero-Offset Correction’ and ‘Dive Analysis’ to produce summary statistics for each dive. The minimum depth to be considered a dive was set at 12 m (approx. the length of a whale) with a surface error of 2 m. Bottom time was defined as 90% of dive depth. FLNTUB data were converted to measurement units using WetLabs ECOView software and processed using the same parameters as the TDR using Jensen software (Multi-Trace Dive Analysis).

Maximum dive depth, dive duration, ascent and descent rates, and proportional time at depth were calculated for each dive for each individual from the raw data collected by the TDR. ANOVA and t-tests with a significance level of 0.05 were used to contrast diving between individuals.

**Zooplankton and water column sampling.** During 5 to 7 May 2003, 25 systematically dispersed stations south of Disko Island in the bowhead whale concentration area (Fig. 1) were sampled for oceanographic properties and zooplankton distribution biomass using a WP-2 plankton net (200 µm mesh and 29 cm radius) from RV ‘Porsild’. Vertical profiles of salinity, temperature, and chl a fluorescence were taken at all stations using a Seabird 25 conductivity, temperature and depth (CTD) meter. At each station, the plankton net was lowered to 50 m depth and vertically hauled towards the surface at a constant rate of approximately 10 m s⁻¹ while the vessel was stopped. After each tow, the net was rinsed on deck and zooplankton were transferred to plastic jars and preserved in ~2% buffered formalin (final concentration).

Between 22 April and 1 May 2003, 9 stations west of Qeqertasuaq were selected, comprising a subset of the larger grid to focus on the areas with the highest densities of feeding bowhead whales. The zooplankton was sampled using a subsmersible pump (900 l min⁻¹, HOMA-H500, DIFRES-design) equipped with a flow-meter (Hydrobios), conical net (50 µm mesh size) and a nonfiltering cod end. Samples were collected in 4 depth dependent strata. Subsamples were taken using a sample splitter, and the number of animals and their stages were recorded. The samples were immediately preserved in buffered formalin (~2% final concentration). Additionally in 2005, a single standard station located 1.85 km (1 nautical mile) south of Qeqertasuaq (69°15'N, 53°33'W) was monitored weekly between late February and June. Both CTD casts and zooplankton were sampled once a wk through winter and spring, and time series of temperature, salinity, and phytoplankton and zooplankton biomass were obtained.

In 2006, bottom samples of zooplankton were sampled in 3 target coastal areas based on direct observations of bowhead whale feeding. These areas were less than 1 km from shore (Sorte Sand, Laksebugten and Fortune Bay), generally no more than 140 m in depth, and located in areas where bowheads predictably made feeding dives. The samples in 2006 were taken using a Hydrobios Multinet (300 µm mesh size) in 5 depth strata with the final strata lying as close to the bottom as possible. Strata were selected by dividing
the water column into equal segments based on the
maximum depth in the area. The net was lowered to
the bottom and left for 15 min, and then hauled
towards the surface at a constant rate of approximately
10 m s\(^{-1}\). In addition, a single sample was taken with a
WP-2 plankton net covering the whole water column.
The samples were fixed and quantified as described
previously.

In all years, mesozooplankton were sorted and iden-
tified to the lowest taxonomic level or developmental
stage possible. Samples were split to obtain sample
sizes of approximately 500 individuals. Each species or
category was enumerated for each tow and prosome
lengths were measured on up to 10 specimens for each
given species or category. Total abundance was esti-
mated by counting the number of individuals in each
taxa within a single aliquot and scaling up to the total
number of aliquots in each tow. Abundance and length
information was used to calculate biomass in each cat-
egory at each station (mg C m\(^{-3}\)) based on values ob-
tained from the literature (Thor et al. 2005). In 2005
these calculations were conducted at 50 m increments
in the water column down to maximum depths of
200 m. Biomass values were obtained from Hirche &
Mumm (1992) for Calanus species (stages CI–CV) and
Metridia longa, from Berggreen et al. (1988) for Acar-
tia spp., from Klein Bretler et al. (1982) for Pseudo-
calanus species, and from Sabatini & Kiørboe (1995) for
Microcalanus, Oithona and Oncaea species. Carbon
content of eggs was obtained from Kiørboe et al. (1985)
and nauplii stages from Hygum et al. (2000). For non-
copepod species, carbon biomass was estimated for
Appendicularia (Uye 1982), Cirripedia (Turner et al.
2001), Euphausiacea (Pedersen et al. 2005), Gast-
ropoda (Hansen & Ockelmann 1991), Polychaeta
(Hansen 1999), Amphipoda (Hirche & Mumm 1992),
Decapoda myis (Uye 1982), and Chaetognata (Uye
1982). Rare taxa were excluded from biomass calcu-
lations, but were enumerated in general categories for
2003 (Table 2).

Spatial gradients in mean copepod (Calanus,
Metridia, Pseudocalanus and Oithona spp.) biomass
were explored with an inverse distance weighted spa-
tial interpolation model with a search radius >10
neighbors (ESRI ArcINFO 9 Geostatistical Analyst) in
2003 and 2005. In 2003 krigging was conducted using
all grid stations (n = 25) 0 to 50 m deep. Total potential
zooplankton biomass was examined with respect to
bathymetry and focal area use of tagged whales. Krig-
ging in the 2005 focal region was based on the 9 grid
stations and 1 standard station 0 to 50 m deep (Fig. 1).
The number of stations >50 m deep in 2005 was too
low for spatial interpolation.

**Bioenergetic modeling.** A stage and mass struc-
tured population model was developed for bowhead
whales in Disko Bay using proportional stage classes
following directed and opportunistic observations of
whales over a 5 yr period. Age classes of bowhead
whales in Disko Bay in winter and spring were
modeled as 0% juveniles, 20% immatures and
80% adults based on 6 yr of field observations. Sex
ratios were set at 85:15 (F:M) based on genetic biopsy
studies (Greenland Institute of Natural Resources
unpubl. data). A median body mass of 60000 kg was
assumed for sexually mature males and females
(Reeves & Leatherwood 1985), and a median body
mass for immatures was assumed to be half of that, or
30000 kg.

A generalized bioenergetic model was developed to
quantify the gross energetic requirements of the vari-
ous population segments. The model was used to esti-
mate the population’s daily energy requirements
(kcal d\(^{-1}\)) by extrapolating recent population abundance
estimates of 250 whales (CV = 0.35) for a region
corresponding to the home range in Disko Bay (Heide-
Jørgensen & Acquarone 2002). Basal metabolic rate
(BMR in kcal d\(^{-1}\)) was calculated as:

\[
BMR = 70 \times W^{0.75} \tag{1}
\]

where \(W\) = the body mass in kg (Kleiber 1975). Addi-
tional energy required for activity, growth and repro-
duction was added into the model as proportions of
basal metabolic rate using the simple bioenergetic model framework:

\[
ER = \frac{BMR(A + G + R)}{DE \times AE} \tag{2}
\]

where \(ER\) = energy requirement in kcal d\(^{-1}\) for an indi-
vidual, \(A\) is an activity metabolic multiplier, \(G\) is a
growth multiplier, \(R\) is the reproduction cost multiplier for adult females (costs of fetal metabolism, fetal
growth and lactation), \(DE\) is the digestive efficiency (digestible energy) and \(AE\) is the assimilation effi-
ciency (digestibility of dry matter).

Activity metabolism (A) assumes that active metabo-
lim is a constant multiple of basal metabolic rate (Ken-
consists of foraging behavior, movement or migration,
or age or sex-specific behavior, and has been esti-
mated to be 2 to 5 times the BMR in cetaceans (Lock-
yer 1981, Kenney et al. 1997). A mean of 2.5 was used
following Kenney et al. (1997).

Energy requirements for growth of body tissues and
blubber, which decrease with increasing age until
physical maturity, can be 110 to 300% of the
maintenance energy required for adults (Innes et al.
1987). Therefore, growth in body mass or production
was modeled as additional needs (varying with age)
and calculated as a proportion of BMR. We let \(G\) range from 1.0 to 2.0 for the immature stage classes
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and set $G$ to 0 for adult stage classes, assuming physical maturity was reached and there were no further growth requirements.

Energy for reproduction ($R$) for adult females was estimated for the costs of pregnancy (energy for fetal metabolism and fetal growth) and lactation. Bowhead whales were assumed to have a calf every 3 yr; consequently, within a given year an ‘average’ reproducing adult female in the model was pregnant, lactating or resting. A composite energy requirement for pregnancy was based on a fetus assumed to have a $BMR$ proportional to that of an adult per kg (Yasui & Gaskin 1986), or a mean mass of 6000 kg, estimated as half the birth weight based on 3.5 m long fetus (Reeves & Leatherwood 1985) proportional to adult mass–length ratio. The cost of fetal growth (14 mo or 420 d gestation period) was calculated using Brody’s (1945) equation for the heat increment of gestation, equal to $4400 M^{1.2}$, where $M$ is the mean birth mass (13 000 kg). The cost of lactation was assumed to be 3 times the combined cost of fetal growth and basal metabolism (Yasui & Gaskin 1986). This resulted in an estimated cost of lactation of approx. twice the basal metabolic rate for a non-lactating adult (Winship et al. 2002). The costs of pregnancy and lactation were combined to create a value for $R$ of 0.6 for the adult female portion of the bowhead population.

Digestive efficiency ($DE$) was assumed to be 0.9 following that for other marine mammals and assimilation efficiency ($AE$) for cetaceans was assumed to be 0.8 (Lockyer 1981). The energetic value of the Calanus dominated zooplankton was assumed to be 5 kJ g$^{-1}$ (1.18 kcal g$^{-1}$) with a mean size of $1.0 \times 10^{-4}$ g per individual copepod.

Parameter uncertainty was addressed with Monte Carlo simulations where parameter values were randomly selected from sampling distributions that best described their uncertainty. Activity and growth multipliers were sampled from a uniform distribution where $A$ varied between 2 and 3 for all stage classes and $G$ varied between 1 and 2 for immatures. Population size was drawn from a lognormal distribution following Heide-Jørgensen & Aquarone (2002). Approximately 10 000 simulations were used to calculate a mean caloric requirement and confidence intervals for the spring feeding period. These prey demands were used to estimate the total amount of zooplankton consumed by the population.

**RESULTS**

**Area use of satellite tagged whales**

The 30 bowhead whales fitted with satellite tags were all 12 m or greater in length. Average daily positions were obtained for individual whales tagged in April and May from 2001 to 2006 (Fig. 2). In all years, area use was focused on the southern coast of Disko Bay between 2001 and 2006. Data were calculated from (a) 5 whales tagged in 2001, (b) 6 in 2002, (c) 3 in 2003, (d) 11 in 2005 and (e) 5 in 2006 using average daily positions while whales remained in Disko Bay before departing on the migration to Canada past 70°W.
Island, extending offshore about 50 km. The bathymetry within home ranges varied widely and covered areas ranging from a steep coastal slope to trenches of 400 m depth.

In 2001, the movements of 5 tagged whales were focused along the south coast of Disko Island close to shore and the area use ranged up to 3652 km² (95% kernel) during 4 to 21 May (Table 1, Fig. 2a). In 2002, the area use extended farther west to include Disko Fjord and a larger part of the coast of south Disko Island and covered 4502 km² (95% kernel) during 7 to 14 May (Table 1, Fig. 2b). In 2003, whales used a region similar to that used in 2001; however, it was slightly larger, up to 4089 km² (95% kernel), during 10 to 21 May (Table 1, Fig. 2c). In 2005, whales used a region similar to that used in 2001; however, it was slightly larger, up to 6897 km² (95% kernel) during 16 April to 17 May (Fig. 2d). In 2006, the home range included some area use north of Disko Island; however, whales still concentrated along the south coast of Disko Island, using up to 35 028 km² (95% kernel) during 6 April to 31 May (Table 1, Fig. 2e).

Diving behavior

Fourteen archival tags (TDRs and FLNTUBs) were deployed and retrieved on bowhead whales between 2002 and 2006 (Table 2). Tag attachment lasted between 1 h 45 min and 54 h 41 min, with an average attachment time of 11 h 40 min for all 14 samples. Most instruments were located and recovered 1 to 2 d after release from the whale with the ARGOS location data and VHF tracking. Some instruments were retrieved with mud on the instrument package, suggesting recent bottom contact.

Intra- and interanimal variability in diving behavior was high in all years. Mean interwhale dive depths ranged from 53 (SD = 35) to 109 m (SD = 41) (Table 2). Dive depths ranged from 12 m to 487 m for individual whales, and the maximum depth reached by each individual generally exceeded 150 m (80% of tagged whales made at least 1 dive >150 m). Significant differences in individual mean dive depth were detected (F<sub>13,607</sub> = 4.25, p < 0.001). Mean dive durations ranged from 3 (SD = 2) to 18 (SD = 9) min, yet the maximum dive duration recorded was 48 min. Significant interwhale differences were detected in dive duration (F<sub>13,607</sub> = 18.0, p < 0.001). Descent rates and ascent rates between individuals ranged widely (Table 2) and were significantly different (Descent F<sub>13,594</sub> = 22.1, p < 0.0001, Ascent F<sub>13,554</sub> = 23.4, p < 0.001). Maximum observed ascent rates were higher than maximum

<table>
<thead>
<tr>
<th>Year</th>
<th>No. whales</th>
<th>Area use (km²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>50%</td>
</tr>
<tr>
<td>2001</td>
<td>5</td>
<td>829</td>
</tr>
<tr>
<td>2002</td>
<td>6</td>
<td>751</td>
</tr>
<tr>
<td>2003</td>
<td>3</td>
<td>335</td>
</tr>
<tr>
<td>2005</td>
<td>11</td>
<td>684</td>
</tr>
<tr>
<td>2006</td>
<td>5</td>
<td>3494</td>
</tr>
</tbody>
</table>

Table 2. *Balaena mysticetus*. Estimated kernel area use of satellite tagged bowhead whales in Disko Bay between 2001 and 2006. See Fig. 2 for spatial distribution.

Table 1. *Balaena mysticetus*. Summary dive statistics collected from TDRs and FLNTUBs deployed on 14 bowhead whales from May 2002 to 2006 in Disko Bay, West Greenland. TDR: Wildlife Computers Time Depth Recorder, FLNTUB: WetLabs Fluorometer. Dive depth in m, dive duration in min, ascent and descent rates in m s⁻¹ (SD). Unknown sex (U) due to absence of a biopsy sample from the animal.
observed descent rates. Dive duration ranged widely for a given maximum dive depth (Fig. 3a) and the two were poorly correlated. Furthermore, post-dive surface time was not significantly correlated with dive depth or dive duration for any individual (Figs. 3b–c).

Surfacing intervals tended to be short when active diving was observed and over 50% of post-dive surface intervals, regardless of destination depth, were <4 min. Occasionally, diving activity ceased and whales spent extended periods at the surface. These periods of apparent inactivity were characterized by few dives below 12 m and ranged from 30 min to >6 h in duration.

Dive shapes could be classified into 2 general types. Typical feeding dives were characterized by a rapid descent to a particular depth, fidelity to that depth (±5 m) during the bottom time portion of the dive, and then a rapid ascent back to the surface. These foraging dives were generally U-shaped (Fig. 4) with a large proportion of time spent near the bottom and often at the maximum depth in the area. The other type of dive identified resembled a deep V-shaped search dive that traversed a large portion of the water column (generally >100 m) with minimal time spent on the bottom. Often a series of search dives preceded a series of foraging dives with remarkable fidelity to a certain depth.

**Zooplankton and water column sampling in 2003**

Stations were evenly distributed in a grid network between 8 and 50 km off the southwestern corner of Disko Island (Fig. 1). Zooplankton samples were collected from a standardized volume of water (13 m³) between 0 and 50 m, and whenever possible CTD casts were taken to 100 m. CTD data demonstrated a characteristic water column stratification for polar seas, with low saline (33.1 to 33.5 psu) and high temperature (−0.7 to 1.0°C) water masses at the surface extending down to the pycnocline (20 m). There was no sign of deeper mixing in the water column; however, the deep advection of warm saline water was evident below 70 m. Surface temperature was approximately 1°C and declined to about 0.25°C at depths of 20 m before warming again to 1.5°C at 100 m.

Zooplankton taxa were pooled into categories representing both the largest proportions of biomass in the study area and the primary prey species for bowhead whales (Table 3). Across the 2003 grid network, cope-
Pods dominated the biomass with *Calanus finmarchicus*, *C. glacialis* and *C. hyperboreus* occurring at 90 to 100% of all stations, contributing on average 78% (SD = 25) of the total biomass (interstation range 30 to 97%). Among the copepods, *C. finmarchicus* contributed a disproportionate amount of the biomass, outnumbering *C. glacialis* and *C. hyperboreus* at least by a factor of 4. *Calanus* eggs and nauplii were found at all stations. Copepod species from other genera, including *Metridia*, *Oithona*, *Pseudocalanus*, *Microcalanus* and *Acartia*, also contributed a large proportion of the total biomass (Table 3). Non-copepod species found frequently included those of *Cirripedia*, *Euphausiacea*, *Amphipoda* and *Gastropoda*, although these taxa were only identified in nauplii, egg or larval stages and contributed minimally to the total biomass. Mean station biomass measures were normally distributed and ranged between 4 and 233 mg C m\(^{-3}\). Average interstation biomass for all zooplankton between 0 and 50 m was 79 mg C m\(^{-3}\) (SD = 52).

Interpolated spatial gradients in biomass in 2003 were higher in the eastern and offshore part of the bay, with lower biomass occurring in the western part and at shallow depths (Fig. 5). The low biomass in the western portion of the bay (ranging from 4 to 50 mg C m\(^{-3}\)) spatially coincided with the 50% probability home range in 2003 (Fig. 2c). Biomass levels >200 mg C m\(^{-3}\) were found in areas where few satellite positions were received from whales in 2003.

### Zooplankton and water column sampling in 2005

The standard station in 2005 demonstrated the temporal trend in the development of primary production in Disko Bay. Low chl \(_a\) values were recorded in the upper 50 m of the water column until late March, after which phytoplankton biomass increased exponentially and peaked in mid-April at 14 mg chl \(_a\) m\(^{-3}\) (Fig. 6a). Copepod biomass peaked between the end of April and early May at 2500 mg C m\(^{-2}\) (Fig. 6b). The temporal trend of the ascension of copepods to the upper 50 m was essentially linear and densities reached an asymptote at levels of ~2000 mg C m\(^{-2}\) by June.

The additional focal area sampled in 2005 corresponded to the region used heavily by feeding whales. Biomass values in 2003 and 2005 were compared for 9 stations (Figs. 5 & 7). Biomass to 50 m was lower in 2005 than in 2003 (between 1 and 16 mg C m\(^{-3}\)); however, increasing trends with increasing distance from the coast were found in both years. Copepod biomass was 5 to 15 times higher at depths >150 m where whales concentrated diving activity (Fig. 8). This was the same region where biomass above 50 m was negatively correlated with whale occurrence in 2003 (Fig. 5).

### Zooplankton and water column sampling in 2006

In 2006, 3 sites (Sorte Sand, Fortune Bay and Laksebugten) were sampled for zooplankton concentrations from the surface to the bottom in 15 m increments and in 5 replicates (Fig. 1). At all sites the biomass of copepods and *Pandalus* sp. increased by several orders of magnitude from the surface of the water column (<1 mg C m\(^{-3}\)) to an average biomass of 353 mg C m\(^{-3}\) at 40 m on the bottom (Fig. 9). *Pandalus* sp. was the dominant species found in samples obtained between 30 and 75 m, but between 75 and 115 m *C. finmarchicus* was found in quantities several

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**Table 3. Species identified at 25 stations during tows for zooplankton in the upper 50 m in Disko Bay, West Greenland, May 2003.**

<table>
<thead>
<tr>
<th>Plankton taxa</th>
<th>Included in biomass calculations</th>
<th>Occurrence (% of Stns) in 2003</th>
<th>Mean biomass (SD) (mg C m(^{-3}))</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Copepoda</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Calanus finmarchicus</em></td>
<td>X</td>
<td>100</td>
<td>49 (39)</td>
</tr>
<tr>
<td><em>Calanus glacialis</em></td>
<td>X</td>
<td>90</td>
<td>2.8 (2.3)</td>
</tr>
<tr>
<td><em>Calanus hyperboreus</em></td>
<td>X</td>
<td>100</td>
<td>12.3 (14.9)</td>
</tr>
<tr>
<td><em>Pseudocalanus</em> spp.</td>
<td>X</td>
<td>100</td>
<td>0.5 (0.3)</td>
</tr>
<tr>
<td><em>Oithona</em> spp.</td>
<td>X</td>
<td>100</td>
<td>0.1 (0.03)</td>
</tr>
<tr>
<td><em>Metridia</em> longa</td>
<td>X</td>
<td>50</td>
<td>0.3 (1.1)</td>
</tr>
<tr>
<td><em>Oncaea</em> spp.</td>
<td>X</td>
<td>23</td>
<td>&lt;0.01 (nd)</td>
</tr>
<tr>
<td>Other Copepoda*</td>
<td>X</td>
<td>83</td>
<td>&lt;0.01 (nd)</td>
</tr>
<tr>
<td>Copepod eggs</td>
<td>X</td>
<td>100</td>
<td>&lt;0.01 (nd)</td>
</tr>
<tr>
<td>Copepod nauplii</td>
<td>X</td>
<td>100</td>
<td>0.1 (0.1)</td>
</tr>
<tr>
<td><strong>Other invertebrates</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Cirripedia</em></td>
<td>X</td>
<td>100</td>
<td>0.4 (0.5)</td>
</tr>
<tr>
<td><em>Euphausiacea</em></td>
<td>X</td>
<td>100</td>
<td>1.2 (1.9)</td>
</tr>
<tr>
<td><em>Decapoda</em></td>
<td>X</td>
<td>87</td>
<td>0.05 (0.1)</td>
</tr>
<tr>
<td><em>Amphipoda</em> (Hyperiidea)</td>
<td>X</td>
<td>100</td>
<td>1.5 (1.6)</td>
</tr>
<tr>
<td><em>Gastropoda</em></td>
<td>X</td>
<td>100</td>
<td>&lt;0.01 (nd)</td>
</tr>
<tr>
<td><em>Chaetognata</em></td>
<td>X</td>
<td>93</td>
<td>10.4 (12.6)</td>
</tr>
<tr>
<td><em>Polychaeta</em></td>
<td>X</td>
<td>93</td>
<td>0.04 (0.1)</td>
</tr>
<tr>
<td><em>Appendicularia</em></td>
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<td></td>
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<tr>
<td><em>Hydrozoa</em></td>
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<tr>
<td><em>Echinodermata</em></td>
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<tr>
<td><em>Ostracoda</em></td>
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<tr>
<td><em>Ctenophora</em></td>
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<tr>
<td><em>Protozoa</em></td>
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<tr>
<td><em>Isopoda</em></td>
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<tr>
<td><em>Arachnidae</em></td>
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<tr>
<td><em>Nemertini</em></td>
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<tr>
<td><em>Bivalvia</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Fish larvae and eggs</em></td>
<td>X</td>
<td>33</td>
<td>0.02 (0.04)</td>
</tr>
</tbody>
</table>

*a* Includes *Microcalanus* spp. and *Acartia* spp.
orders of magnitude higher than in all other prey categories or depths (Fig. 9).

**Bioenergetic model**

The bioenergetic model produced a daily energy requirement of $2.67 \times 10^8$ kcal d$^{-1}$ (SE = $9.7 \times 10^5$) for the 250 (CV = 0.35) bowhead whales in Disko Bay. The adult female portion of the population had the highest energy requirements, $1.2 \times 10^8$ kcal d$^{-1}$ (SE = $4.4 \times 10^5$) with adult male stage class following them at $9.8 \times 10^7$ kcal d$^{-1}$ (SE = $3.6 \times 10^7$ kcal d$^{-1}$). Daily energetic needs of the population were met with approximately 223 t (SE = 0.8) of zooplankton. Thus, the estimated biomass of zooplankton required by the total population of bowheads over the 4 mo period was 26 798 t (SE = 97), assuming energetic requirements for an individual are met every day in Disko Bay.

The mean biomass of zooplankton in the upper 50 m in May 2003 was estimated as 79 mg C m$^{-3}$. Using a carbon content of 40% of dry weight (Karnovsky et al. 2003), the biomass estimate of zooplankton corresponds to $2.6 \times 10^7$ mg dry weight. An energy density of 26 kJ g dry weight (Karnovsky et al. 2003) results in 0.001 kg m$^{-3}$ of copepods in the upper 50 m of the water column, or approximately 122 770 t of zooplankton in the 2600 km$^2$ sampled area. Thus, if bowheads were to feed only in the upper 50 m of the water column, they would (at the current population size) remove approximately 22% of the pelagic biomass in spring.

**DISCUSSION**

**Bowhead foraging behavior**

The U-shaped feeding dives observed in this study are characteristic for baleen whales exploiting patches of zooplankton (Baumgartner & Mate 2003, Baumgartner et al. 2003). The remarkable fidelity to a specific depth suggests that once high or sufficient densities of zooplankton are located, the dive duration at that depth is maximized and the same depth is targeted repeatedly. In this study, target depths were highly variable given that bowhead whale foraging...
occurred on the seabed in an area with a large gradient in bottom depths. Whales actively foraged at 80 to 200 m, corresponding well to the bottom contours along the slope. The functional relationship between U-shaped feeding and V-shaped searching dives probably facilitates an active search for discrete layers of zooplankton before initiation of foraging dives.

In this study, bowheads dove to a maximum of 487 m, and in many cases whales targeted depths of >200 m repeatedly. In the Beaufort Sea, bowhead whales have been reported to make shallower dives (50 to 100 m) for calanoid copepods on the bottom (Krutzikowsky & Mate 2000). The dive of 487 m recorded during this study is the deepest dive measured for a bowhead whale.

Dive durations measured for foraging bowhead whales were surprisingly longer than those reported for other baleen whales. Dives lasted up to 48 min and in several cases whales repeatedly dove for 30 to 40 min at a time with post-dive times spent at the surface lasting <2 min. Mean dive durations in this study (5 to 18 min) are similar to those for the right whale *Eubalaena glacialis* (12 min); however, maximum values are much higher than those for right whales...
(16 min) (Baumgartner & Mate 2003). The fin whale *Balaenoptera physalus* and the blue whale *B. musculus* have shorter foraging dive durations (mean, 6 to 7 min) and it has been suggested that continuous filter feeding in bowhead and right whales lowers metabolic rates and enables these species to maintain higher dive durations (Croll et al. 2001).

Krutzikowsky & Mate (2000) reported bowhead whale dive durations exceeding 60 min in the Beaufort Sea. Furthermore, bowhead whales have been reported to stay submerged for >80 minutes when harpooned (Scoresby 1820, Scammon 1874), and this species may have a diving capacity far beyond that of other baleen whales (Figs. 3a & 4).

Bowhead whales appear to have large flexibility in their use of the water column. There was no correlation between dive depth and duration and no correlation with post-dive surface interval (i.e. recovery time). This is in contrast to strong positive correlations between dive duration and dive depth for right whales that focus on dense patches of zooplankton (Baumgartner & Mate 2003).

Bowhead whales in Alaska feed between spring and fall in the water column and on the bottom or near-bottom on larger amphipods and euphausiids (Lowry 1993, Lowry et al. 2004). Lowry (1993) and Lowry et al. (2004) found hundreds of prey species in the stomachs of harvested bowhead whales between Point Barrow and Kaktovik and differences in the seasonal proportion of species in stomachs was consistent with seasonal patterns of zooplankton abundance.

**Bowhead prey items**

Bowhead whales have been postulated to filter feed on zooplankton at depths just below the euphotic zone after the ascension of the secondary production. Therefore, the lack of relationship between high densities of prey in the upper 50 m of the water column and bowhead dive behavior and area use was surprising. The major portion of the spring zooplankton biomass in the upper 50 m of Disko Bay consists of Calanoid copepods, primarily *Calanus finmarchicus*, a North Atlantic species that successfully reproduces and recruits in polar ecosystems (Pedersen et al. 2005, Thor et al. 2005). Studies of interannual variation in the upper 50 m of the water column in Disko Bay indicate that zooplankton biomass can vary by >30% (Thor et al. 2005). The predictability of prey for bowhead whales in surface waters is therefore low. Bowhead whales occupy a seasonally dynamic environment with brief optimal foraging conditions. Thus, maintaining a foraging strategy that targets the densest and most predictable prey concentrations (i.e. near seabed zoo-plankton) offers insurance against interannual fluctuations in prey abundance at the surface. Bowheads ingest about 10 times as much phytoplankton as copepods while foraging (750 mg C m⁻³ of phytoplankton vs. 79 mg C m⁻³ of *Calanus*); however, the extent of the importance of diatoms in the diet of bowhead whale is unknown.

**Bowhead energetics**

Uncertainties in metabolic rates, feeding time for baleen whales and energetic densities of zooplankton (Costa et al. 2006) inhibit the ability to accurately model caloric needs. Several unknown and estimated parameters make bioenergetic considerations for large baleen whales simply an approximation.

Bowheads migrate seasonally between local Arctic hot spots with high densities of zooplankton. Studies of the movements of bowhead whales from satellite telemetry show intense and localized use of a single site for 3 to 6 wk interspersed with long distance movements (>1000 km) occurring over 1 to 2 wk (Heide-Jørgensen et al. 2006). Travel speeds are too high (up to 200 km d⁻¹) for foraging to be feasible during these long distance movements, and whales traverse offshore deep areas, regions that contain little to no zooplankton. Thus, it is possible that bowhead whale caloric needs are not met on a daily basis; instead, they are obtained during intense foraging periods with little to no feeding between sites. Their large body size likely acts as a buffer from the variability in pelagic productivity and allows the species to move long distances over long periods of time between regions of patchy and ephemeral productivity.

Technically, the densities of zooplankton above the 50 m depth in Disko Bay are sufficient to support the energetic demands of the population. However, the question remains as to the amount of water that needs to be filtered to meet these caloric needs and if zooplankton densities are even sufficiently high enough to warrant filter feeding by bowheads. Given the bioenergetic estimates, bowheads would need to filter more than 200 × 10⁶ m³ of water daily, or 22% of the total volume of water that exists between 0 and 50 m in their home range, during a 4 mo period. Densities of copepods rapidly increase with depth, especially below 150 m, and copepod densities are orders of magnitude higher near or on the bottom. The dive data indicate that bowhead whales make deep dives, following the highly complex bottom contours of Disko Bay in areas where epibenthic densities of copepods are dramatically greater (Figs. 8a, b & 9). Given the requirement to strain enormous quantities of water, large whales have likely evolved to exploit their zooplankton prey in
regions with high density aggregations (see Baumgartner & Mate 2003, Croll et al. 2005). Little is known about the fine scale spatial variability of these patches or how much time bowhead whales must spend searching for them.

Affinity of whales to the southern coast of Disko Island has been known for hundreds of years (Eschricht & Reinhardt 1866). It is likely that this particular area affords unparalleled feeding opportunities due to physical patterns in water temperatures, upwelling, currents or high densities of lipid rich Arctic zooplankton species (Lee et al. 2006). Eschricht & Reinhardt (1866) reported on departure dates of bowhead whales in Disko Bay between 1780 and 1837 with a mean date for the last sighting of whales of 1 June (range, 26 April to 25 June). Bowhead whales in this study departed between 10 and 21 May, precisely following observations from the past 2 centuries. Bowhead whales depart from Disko Bay before the ascension and peak densities of copepods in the water column (Fig. 6b); this has also been noted for whales in the Bering Sea (Lowry 1993).

Bowhead whales in West Greenland focus foraging effort on benthic zooplankton in coastal zones, the spatial distribution of which probably results from a combination of coastal upwelling and complex bottom topography funneling or channeling prey. The steep shelf off the coast of Disko Island (0 to 400 m across 4 km) makes a large range of complex and deep depths available to whales over a small horizontal area, and Disko Bay is known as one of the most productive areas in West Greenland (Heide-Jørgensen and Laidre 2004).

In this study, the combination of observations of bowhead whale foraging behavior collected by satellite and archival telemetry combined with intensive and local-scale whale foraging behavior collected by satellite and archival telemetry combined with intensive and local-scale observations of phytoplankton and zooplankton (Heide-Jørgensen et al. 2007), it is likely the springtime foraging conditions encountered by bowhead whales in Disko Bay will be altered. Biotic factors like competition and predation become more important at higher levels of the food chain; thus, long-term studies of the ecosystem, including physical and multiple trophic levels, are required to develop a holistic understanding of how climate change will affect bowhead whales.

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**LITERATURE CITED**


Eschricht DF, Reinhardt J (1866) Om Nordhvalen (Balaena mysticetus L.) navnlig med Hensyn til dens Udbredning i Fortiden og Nutiden og til dens ydre og indre Saerkjender. K. Danske Videnskabernes Selskabs Skrifter, Series 5.
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