

Title	Abundance, horizontal and vertical distribution of epipelagic ctenophores and scyphomedusae in the northern Bering Sea in summer 2017 and 2018 : Quantification by underwater video imaging analysis
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Citation	Deep Sea Research Part II Topical Studies in Oceanography, 181-182, 104818 https://doi.org/10.1016/j.dsr2.2020.104818
Issue Date	2020-12
Doc URL	http://hdl.handle.net/2115/87790
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Туре	article (author version)
File Information	Maekakuchi_MS.pdf



2	Abundance, horizontal and vertical distribution of epipelagic ctenophores and scyphomedusae in the
3	northern Bering Sea in summer 2017 and 2018: quantification by underwater video imaging analysis
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17 Abstract

18	We examined the abundance and horizontal and vertical distributions of epipelagic ctenophores and
19	scyphomedusae in the northern Bering Sea using an underwater video camera during July of 2017
20	and 2018. The effects of environmental and biological parameters on the distribution of these
21	species were evaluated by generalized additive modelling (GAM). In 2017, the dominant
22	ctenophore, Bolinopsis infundibulum, was mainly distributed in the north and west of St. Lawrence
23	Island (SLI), and their vertical distribution varied with the region but not by the time of day. We
24	found that <i>B. infundibulum</i> was distributed in the upper pycnocline north of SLI, but below the
25	pycnocline west of SLI. Biological interactions with other gelatinous zooplankton may explain
26	these regional differences in vertical distribution; GAM analysis revealed a negative interaction
27	between B. infundibulum and the large scyphomedusa, Chrysaora melanaster, which occurred in the
28	upper layer in the west of SLI. <i>B. infundibulum</i> may avoid that layer to reduce feeding
29	competition. For the ctenophore, <i>Beroe</i> sp., vertical and horizontal distributions were similar to
30	those of <i>B. infundibulum</i> , and GAM analysis also revealed a positive interaction for both species.
31	As <i>B. infundibulum</i> is an important prey of <i>Beroe</i> sp., a prey-predator interaction may result from
32	their similar horizontal and vertical distributions. Standing stocks of epipelagic ctenophores and
33	scyphomedusae in 2018 were low compared to those in 2017, by a factor of 1/20 (C. melanaster)
34	and1/90 (Beroe sp.). This might be due to annual differences in water mass in this region, in that

35	the thermal conditions characterized by a high abundance of the dominant <i>B. infundibulum</i> in 2017
36	(<2 and >8°C) were absent in 2018. As this drastic decrease in standing stock in 2018 was
37	apparent for both ctenophores and scyphomedusae, food availability was hypothesized to be poor
38	that year.
39	
40	Keywords:
40 41	Keywords: Ctenophores, Scyphomedusae, Northern Bering Sea, Vertical distribution, Horizontal distribution,
	·

1. Introduction

46	Recently, increases in the abundance of large ctenophores and scyphomedusae have been
47	reported in various oceans worldwide, likely due to human alternation of marine environments and
48	climate change (Purcell et al., 2007; Condon et al., 2013; Duarte et al., 2013). The main food
49	sources of ctenophores and scyphomedusae in higher latitudes are mesozooplankton, especially
50	copepods (Brodeur et al., 2002; Purcell et al., 2010). Ctenophores and scyphomedusae are thus
51	competitors of the planktivorous fishes, and also act as predators upon larval fishes; thus, the
52	abundance of large gelatinous zooplankton can have a great effect on fish stocks (Brodeur et al.,
53	2002; Purcell et al., 2007; Robinson et al., 2014). As large gelatinous zooplankton are composed
54	mainly of water and require less organic material for their body composition, they can respond to
55	environmental changes more rapidly than crustacean zooplankton, and their biomass can vary
56	dramatically between years (Falkenhaug, 1996; Brodeur et al., 2008).
57	The northern Bering Sea, the target area of the present study, is a transit region for the
58	warmer waters of southern origin (Alaskan Coastal Water, Bering Shelf Water, Anadyr Water),
59	which intrude into the western Arctic Ocean (Shimada et al., 2006; Sasaki et al., 2016; Danielson et
60	al., 2017). A polynya forms south of St. Lawrence Island (SLI), the largest island in this region,
61	and is characterized as being ice-free, even in winter (Grebmeier and Cooper, 1995). Recently, a
62	drastic decrease in the ice-covered area and an early ice retreat in the northern Bering Sea has been

64	Additionally, changes in zooplankton biomass, a northern shift of the fish community, and a mass
65	mortality of seabirds occurred during the winter of 2017/2018 and spring/summer of 2018
66	(Cornwall, 2019; Duffy-Anderson et al., 2019; Huntington et al., 2020). Under conditions of
67	greater variability in the environment and marine ecosystem, the amount and distribution of
68	ctenophores and scyphomedusae were also expected to change. However, ecological information on
69	these species is presently scarce for this region.
70	The methods used to quantify ctenophores and scyphomedusae have several limitations
71	(Graham et al., 2003, Uye et al., 2017). Traditional sampling using a plankton net tow is hampered
72	by patchy spatio-temporal distributions, relatively large body size, low abundance, and net
73	avoidance, leading to inevitable underestimation of their biomass and species diversity (Youngbluth
74	and Båmstedt, 2001; Graham et al., 2003; Raskoff et al., 2005; Uye et al., 2017). The fragile
75	bodies of ctenophores and scyphomedusae are also heavily damaged by net towing (Graham et al.,
76	2003; Raskoff et al., 2005: Uye et al., 2017). Large-volume trawl nets have been used; however,
77	using these nets requires large effort and cost compared to plankton sampling. Moreover, changes
78	in the mouth area and collection efficiency varies with mesh size and towing speed. These nets
79	cause serious damage to the fragile bodies of gelatinous zooplankton, making quantitative collection
80	difficult for ctenophores and scyphomedusae (Graham et al., 2003, Uye et al., 2017). To overcome

63 reported (Comiso et al., 2008; Parkinson and Comiso, 2013; Stabeno and Bell, 2019).

81	these problems in the quantification of ctenophores and scyphomedusae, alternative non-capture
82	methods such as sonar cameras (Han and Uye, 2009), video cameras using a Remotely Operated
83	Vehicle (Båmstedt and Martinussen, 2015), and visual monitoring using ships and airplanes (Purcell
84	et al., 2000) have been used.
85	In the Bering Sea, Brodeur et al. (2017) reported horizontal distributions, and seasonal and
86	annual changes in ctenophores and scyphomedusae based on data collected by trawl nets, although
87	there was the paucity of data for the northern Bering Sea. The vertical distributions of ctenophores
88	and scyphomedusae in the southeastern Bering Sea have been observed using a video camera
89	mounted on an ROV (Brodeur, 1998; Brodeur et al., 2002). Annual variations in the biomass of the
90	large scyphomedusa, Chrysaora melanaster, over the southeastern Bering Sea shelf has previously
91	been analysed using Generalized Additive Models (GAM) to explore which environmental variables
92	might explain the variability (Brodeur et al., 2008). Although the northern Bering Sea is
93	characterized by large interannual changes in the ice-covered area and timing of the ice retreat, few
94	studies have investigated interannual changes in ctenophores and scyphomedusae in this region.
95	In the present study, we quantified the horizontal and vertical distributions, and annual
96	changes in the abundances of ctenophores and scyphomedusae using an underwater video camera in
97	the northern Bering Sea during the summers of 2017 and 2018. Interactions among vertical,
98	horizontal, and annual changes in the abundance of ctenophores and scyphomedusae were assessed

- 99 with environmental parameters (depth, temperature, salinity, diel period, year, and location relative
- 100 to the pycnocline) and biological parameters (mesozooplankton biomass, other species of
- 101 ctenophores and scyphomedusae) by GAM analysis.

- **103 2. Material and methods**
- 104 2.1. Field observation
- 105 Imaging data for ctenophores and scyphomedusae were collected by vertical casts of a frame camera

106 at 21 (2017) and 14 (2018) stations located between 63°00'-66°44'N and 166°30'-174°50'W in the

- 107 northern Bering Sea. These sampling stations were occupied by the T/S Oshoro-Maru during 9–22
- 108 July 2017 and 2–12 July 2018 (Fig. 1). Imaging data down to 50 m were collected by dead-slow
- 109 (0.1 m s⁻¹) vertical deployment of an underwater video camera (Marine Arkas, Kowa Co. Ltd.)
- mounted within a stainless frame of 1.0×1.0 m bottom and 1.5 m depth (Fig. 2B). A charge-
- 111 coupled device (CCD) camera (NTSC PA-290) was equipped with a f2.9 lens that had 0.035 lux
- 112 light sensitivity. The final resolution of each digital frame was 768 (horizontal) by 494 (vertical)
- 113 pixels. The *in situ* images were monitored from the ship. To evaluate diel changes in vertical
- distribution, observations were made at 1 h intervals at one station (St. 10) during 19:40–7:10, 13–14
- 115 July 2017 (total number of samples = 11) (Appendix A). Observations were made both day and
- night at three additional stations (St. 14, 20, and 23) in 2017. At each station, temperature and

117	salinity were measured using a Conductivity Temperature Depth (CTD) sensor (SBE911, Sea-Bird
118	Electronics, Inc.). To evaluate mesozooplankton biomass, a vertically-stratified tow of a 60 cm
119	opening-closing net (mesh size: 100 μ m) (Kawamura, 1989) was conducted from the sea-surface to
120	the thermocline, and from the thermocline to near-bottom. Mesozooplankton samples were
121	preserved in 5% (v/v) borax-buffered formalin seawater.
122	The underwater camera (Marine Arkas, Kowa Co. Ltd.) was equipped with two halogen
123	lights (JCD100V-150W) with 150W and 3300 lumen luminous flux and a pressure-depth sensor
124	(model P193-010-45, SENSIT., Co. Ltd., Hampshire, UK) with a precision of $\pm 0.25\%$ FS. A
125	picture of the frame camera is presented in Fig. 2B. Using the bottom observation frame as a
126	guide, the camera was able to image an observational area of 1.2×0.8 m (Fig. 2B, C). On the side
127	of the frame, a current fin $(0.8 \times 0.3 \text{ m})$ was attached so that the horizontal current was flowing in
128	one direction along the diagonal of the bottom observation frame. Sinkers (20 kg each) were set at
129	the four corners of the frame. We measured the wire angle, and, because of the heavy weight of the
130	camera and frame, the angles were less than 5° at each cast. Underwater videos were transferred
131	on-board through a tether cable and recorded using an HDD and DVD recorder (Toshiba RD-X4).
132	The recording method was MPEG2. The depth data were displayed on a captured image and were
133	recorded using video imaging data. For examples of the video images, see video supplemental
134	materials captured for Cast 14 at St. 10 on 14 July 2014 (Video 1) and for Cast 27 at St. 19 on 19

135 July 2017.

136

137 *2.2. Quantification of ctenophores and scyphomedusae*

138	In the laboratory on land, all ctenophores and scyphomedusae within the observation frame (0.8 m x $$
139	1.2 m) were identified and counted in 2 m vertical intervals from the recorded video. As the frame
140	camera was towed at a speed of 0.1 m s ⁻¹ , video images obtained at 2 m depth intervals corresponded
141	to 20 s (= $2.0/0.1$). Depth data were expressed with 0.1 m accuracy, and the error for estimated
142	depths obtained by the sonar- equipped ship was less than 0.5 m. To evaluate the flow rates of
143	horizontal currents, the diagonal passage time (Pt , s) of ctenophores, scyphomedusae and marine
144	snow within the observation field was measured at 10 m intervals. The <i>Pt</i> was not varied with the
145	targets. For real images of the video, see video supplemental materials. Note that we measured
146	current speed every 10 m constantly and applied these current data for jellyfishes quantified with 2
147	m interval.
148	To calculate individual density, observation volume over 2 m intervals (V, m^3) was
149	calculated from the following equation (Fig. 2C):
150	$V = 2 \times 1.2 \times 0.8 \times 20 \times 1/Pt,$

151 where, $2 \times 1.2 \times 0.8$ is the volume of the observation field (height × width × depth, m³). Thus, 1/Pt

152 represents the changes in viewing within 1 s. As a 2 m vertical movement of the video camera

153 required 20 s., " $20 \times 1/Pt$ " represents the change in view caused by horizontal advection within a 2 154 m observation distance. Swimming speeds of ctenophores and scyphomedusae may also affect 155 their quantification. The swimming speed (V: cm s⁻¹) of ctenophores is known to be a function of 156 diameter (*Dia*: cm): V = 0.12 + 0.04 *Dia* (Cowan and Houde, 1992). If we assume 15-20 cm 157 diameter, the swimming speed of the ctenophore would have been 0.72-0.92 cm s⁻¹. From Pt, the 158 horizontal current speed was calculated to be ca. 0.5-2.0 m s⁻¹. Thus, since horizontal current speed 159 was generally faster than the swimming speed of ctenophores and scyphomedusae, the effect of their 160 swimming speeds on the quantification of their numbers was ignored. 161 Ctenophores and scyphomedusae were quantified for both the descent (D) and ascent(A)162 of the tows, and the data from both directions corresponded well ($D = 0.857 \times A$, $r^2 = 0.742$, ρ 163 <0.0001, n = 561) (Appendix B). Thus, we calculated the mean descent and ascent abundance for 164 each depth. The settling volume for mesozooplankton samples collected by the closing net was 165 measured with 0.1 mL accuracy and expressed as their biovolume biomass (mL m⁻³). 166 167 2.3. Statistical analysis 168 To examine diel changes in vertical distribution, the abundances at depths of 10, 25, 50, 75, and 90% 169 (D10%, D25%, D50%, D75%, and D90% respectively, Pennak, 1943) were calculated for all stations at

170 which day-night observations were made (St. 10, 14, 20, 23, in 2017). To evaluate diel vertical

171	migration, $D_{50\%}$ was compared between the day ($n=7$) and night ($n=4$) using a Mann Whitney U-test
172	at St. 10. For the remaining stations (St. 14, 20, 23) where day and night sampling was conducted,
173	the Kolmogorov-Smirnov test was used to evaluate diel changes in vertical distribution (Sokal and
174	Rohlf, 1995).
175	The effects of environmental and biological parameters on the distribution of ctenophores
176	and scyphomedusae were analysed by GAM. The densities of ctenophores and scyphomedusae
177	were applied as response variables, and environmental and biological parameters, such as
178	hydrography (temperature, salinity), depth, day-night, year, upper/lower pycnocline,
179	mesozooplankton biomass, and the densities of other ctenophores and scyphomedusae were applied
180	as explanatory variables. The pycnocline was defined as the depth at which the seawater density
181	was higher than that at the 5 m depth by 0.1 kg m ⁻³ (Danielson et al., 2011). For the GAM analysis,
182	R software with "mgcv" package was used (Wood, 2017).
183	
184	3. Results
185	3.1. Hydrography
186	Cross-sectional distributions of temperature and salinity along each line transect in the northern
187	Bering Sea during 2017 and 2018 are shown in Figs. 3 and 4, respectively. In 2017, water

188 temperatures ranged from -1.26 to 11.6°C and salinities from 30.2 to 32.9 psu, respectively (Fig. 3).

189	In most locations, the upper layer was characterized by warm temperatures and low salinity. No
190	pycnocline developed north of 66°N, near the Bering Strait, whereas a pycnocline was observed
191	around 6-32 m in the western and southern regions of SLI, with substantial differences in
192	temperature between the upper and lower layers. Conversely, in 2018, temperature and salinity
193	ranged from -0.21 to 12.8°C and from 28.3 to 32.9 psu, respectively (Fig. 4). Pycnocline
194	development was much weaker in 2018 than in 2017. The differences in temperature between the
195	upper and lower layers of the pycnocline were smaller in 2018 compared with 2017.
196	
197	3.2. Horizontal distribution of ctenophores and scyphomedusae
198	Horizontal distributions of ctenophore and scyphomedusa standing stocks (ind. m ⁻²) at each
199	sampling station in 2017 and 2018 are shown in Fig. 5. Three taxa of large gelatinous zooplankton
200	were commonly observed: the ctenophores, Bolinopsis infundibulum and Beroe sp., and the
201	scyphomedusae, Chrysaora melanaster (Fig. 2A). Among these, B. infundibulum was the most
202	numerous species in both 2017 and 2018. In 2017, the standing stock of <i>B. infundibulum</i> was 0–
203	35.6 ind. m ⁻² and was the greatest north and west of SLI. The maximum abundance of C .
204	melanaster and Beroe sp. occurred at 0.689 and 0.567 ind. m ⁻² , respectively, and they were abundant
205	west and south of SLI, and north of SLI to the Bering Strait, respectively. In 2018, standing stocks
206	of all species were much lower than those in 2017, and their mean abundance in 2018 was $1/90$ (C.

melanaster)-1/20 (Beroe sp.) of those in 2017 (Fig. 5). 207

209	3.3. Diel changes in the vertical distribution of <i>Bolinopsis infundibulum</i>
210	Diel changes in the vertical distribution of <i>B. infundibulum</i> were examined at four stations in 2017
211	(Fig. 6). At St. 10, where multiple observations were made, day and night $D_{50\%}$ (mean ± 1 sd) was
212	6.97 ± 4.62 and 1.92 ± 0.32 m, respectively. Although the night $D_{50\%}$ was 5 m shallower than the
213	day $D_{50\%}$, this difference was not significant ($p > 0.05$, U-test), suggesting that there was no diel
214	vertical migration (DVM). For the three stations with only one day-night observation (St. 14, 20,
215	23), diel differences were detected at St. 14 and 20 (St. 14: $p < 0.005$, St. 20: $p < 0.001$,
216	Kolmogorov-Smirnov test); however, the diel pattern varied with the station. A nocturnal ascent
217	occurred at St. 14, while a nocturnal descent occurred at St. 20. The diel changes in $D_{50\%}$ were 5.3
218	m (St. 14) and 5.5 m (St. 20). No significant change was detected between the day and night
219	vertical distributions at St. 23 ($p > 0.05$), likely due to the extremely low abundance at that station
220	(Appendix D). For details of each observation, see Appendix C (St. 10) and Appendix D (Sts. 14,
221	20, 23).
222	

- 223 3.4. Vertical distributions of ctenophores and scyphomedusae
- 224 In 2017, the vertical distribution of *B. infundibulum* showed a clear regional pattern (Fig. 7). This

225	species was distributed throughout the water column north of the Bering Strait (St. 1, 5), but was
226	distributed around or below the pycnocline south of the Bering Strait (St. 7, 9). B. infundibulum
227	was primarily distributed at shallower depths than the pycnocline north of SLI (St. 12, 13), while a
228	bimodal distribution with peaks in the upper and lower layers of the water column was observed
229	northwest of SLI (St. 16, 18). West of SLI (St. 19, 20, 24), this species was distributed below the
230	pycnocline.
231	In 2017, C. melanaster was distributed in a layer shallower than the pycnocline (Fig. 8).
232	In the region west and south of SLI, C. melanaster was distributed at shallower depths than the
233	pycnocline with a maximum density at 0.161 ind. m ⁻³ (St. 22, 6–8 m) for these regions.
234	Beroe sp. were distributed at depths shallower than the pycnocline south of the Bering
235	Strait (St. 6, 9), and both above and below the pycnocline north of SLI (St. 13, 14) (Fig. 9).
236	Conversely, they were distributed entirely below the pycnocline west and south of SLI (St. 18, 20,
237	22, 24).
238	
239	3.5. Interannual changes in ctenophores and scyphomedusae densities
240	Based on data from all sampling stations, the densities of the most dominant ctenophore (B .
241	<i>infundibulum</i>) in 2017 and 2018 are shown with the hydrography in Fig. 10. Annual changes were
242	observed between 2017 and 2018, where 2017 showed a bimodal distribution, with high abundances

at >8 and <2°C, and low abundance at intermediate temperatures (2–8°C). Conversely, the thermal range available in 2018 was mostly limited to 2–8°C and *B. infundibulum* showed very low densities within these temperature ranges.

246

247 3.6. GAM Environmental Relationships

248	In 2018, depth, temperature, salinity, and zooplankton all had significant effects on <i>B. infundibulum</i> ,
249	which were abundant at night and below the pycnocline; these factors were negatively related to C .
250	melanaster abundance, and positively related to Beroe sp. (Table 1). There was a significant
251	positive relationship between C. melanaster abundance and temperature, salinity, and zooplankton
252	biomass, and a negative relationship with <i>B. infundibulum</i> abundance. There was a significant
253	relationship between Beroe sp. abundance and depth, temperature, salinity, and zooplankton
254	biomass, and a positive interaction with <i>B. infundibulum</i> abundance. Smoothing spline regressions
255	between ctenophore and scyphomedusae abundance and environmental parameters with a significant
256	relationship are shown in Fig. 11. The abundance of <i>B</i> . <i>infundibulum</i> was high at depths <24 m,
257	temperatures of -1-2, 5.5-7.5, and 9-10°C, salinities of >30.7, 31.8-32.1, and >32.3 psu, and
258	zooplankton biomasses of 0.1–0.6, 1.4–3.0, and 3.4–4.1 mL m ⁻³ . The abundance of <i>C. melanaster</i>
259	was high at temperatures >5.5°C, salinities between 31.0 and 32.4 psu, and zooplankton biomasses
260	of <1.7, and 3.0–4.4 mL m ⁻³ . The abundance of <i>Beroe</i> sp. was high at depths <14 m, temperatures

261 <6.9°C, salinities of <30.4, 30.8–31.3, and 31.7–32.8 psu, and zooplankton biomasses between 0.2–

262 2.1 mL m⁻³.

204 4. Discussion	264	4. Disc	ussion
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- 265 *4.1. Quantification of ctenophores and scyphomedusae*
- 266 In the present study, we used an underwater video camera to collect quantitative data on ctenophores
- and scyphomedusae during day and night, even under dark conditions in the deepest layers using
- artificial light (3300-lumen luminous flux). Use of an underwater video camera to quantify
- 269 ctenophores and scyphomedusae has been somewhat limited in past studies by reduced visibility and
- 270 difficulty quantifying their abundance in high turbidity waters (Honda and Watanabe, 2007; Honda
- et al., 2016), but we did not encounter high turbidity water in our study region in either year as
- judged by by using the square frame at the bottom of the frame as a guide.
- 273 Several metrics have been presented to quantify data on ctenophores and scyphomedusae
- 274 obtained with an underwater video camera. These include individual number over each observation
- time within a certain depth range (ind. min⁻¹ or ind. hour⁻¹) (Toyokawa et al., 2003; Raskoff et al.,
- 276 2005; Honda and Watanabe, 2007; Raskoff et al., 2010), the individual number observed at each
- depth (number observed) (Purcell et al., 2010; Båmstedt and Martinussen, 2015), and species
- 278 composition within the total observed number throughout the water column (%) (Brodeur, 1998).

279	In the present study, we calculated the observed volume by multiplying the observation area (1.2 \times
280	0.8 m) by depth (2 m) and considered the change in view caused by horizontal advection within a 2
281	m observation (Fig. 2C). A similar calculation method has been used in previous studies.
282	Youngbluth and Båmstedt (2001) calculated volume by multiplying the observation area by vertical
283	depth. To account for horizontal advection, the current flow of the water mass was applied (Nogata
284	et al., 2009), and by measuring the horizontal current using a shipboard acoustic Doppler current
285	profiler (ADCP), Honda et al. (2016) were able to calculate volumes considering horizontal
286	advection. On our cruises, horizontal current was also measured by the shipboard ADCP.
287	However, vertical changes in the horizontal current speed were also observed in the present study.
288	Thus, we measured horizontal current speed at depth using the time taken for a particle (e.g. marine
289	snow) to pass through the observation frame at 10 m depth intervals. As we set the current fin for
290	the frame, the horizontal current was flowing diagonally; thus, it was possible to quantify the
291	horizontal current measurements in this study.
292	Båmstedt and Martinussen (2015) reported a maximum density of <i>B. infundibulum</i> of 2–5
293	ind. m ⁻³ at 0–50 m depths in a western Norwegian fjord. This value corresponds well with the
294	maximum density (3.58 ind. m ⁻³) observed in our study. For <i>C. melanaster</i> , a maximum density of
295	0.07 ind. m ⁻³ has been reported for the southeastern Bering Sea shelf (Brodeur, 1998), whereas in the
296	present study, we observed 0.16 ind. m ⁻³ . Thus, our use of underwater video to quantify individual

- density, considering horizontal advection, seems appropriate and comparable to the results ofprevious studies.
- 299
- 300 *4.2. Diel vertical migration of ctenophores and scyphomedusae*
- 301 For B. infundibulum, no diel changes in vertical distribution were observed at two stations (St. 10, 302 23), nocturnal ascent was observed at one station (St. 14), and nocturnal descent was found at one 303 station (St. 20). However, where changes were observed, the diel differences were small (5.3-5.5 304 m). Little information is available regarding the DVM of ctenophores (Vereshchaka, 2002; 305 Haraldsson et al., 2014; Júnior et al., 2015). The absence of DVM in ctenophores has been 306 attributed to their lacking organs that can detect light (Graham et al., 2001). However, it was 307 recently reported that ctenophores may possess an organ capable of detecting light (Haraldsson et al., 308 2014). 309 The DVM pattern observed for *B. infundibulum* varied with the station, and the magnitude 310 (5.3–5.5 m) was close to the sampling interval of this study (2 m), and relatively small compared to 311 the entire observational depth (50 m). Therefore, we conclude that B. infundibulum did not perform 312 extensive DVM in our study region. The DVM of Beroe spp. has been reported for the northeast 313 Atlantic and the south Brazilian Bight (Roe et al., 1984; Júnior et al., 2015), but in the present study, 314 because of the low abundance, DVM could not be evaluated for *Beroe* sp. Several studies have

315	reported the DV	VM of scyphomedusa	e (Youngbluth and	d Båmstedt, 2001;	Graham et al., 2001;
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- 316 Gorbatenko et al., 2009; Brodeur et al., 2017). For C. melanaster, no DVM had been reported in
- 317 the southwestern Bering Sea shelf region (Brodeur et al., 2017), and in the present study, C.
- 318 *melanaster* was distributed above the pycnocline during both day and night.
- 319
- 320 *4.3.* Horizontal and vertical distribution of ctenophores and scyphomedusae
- 321 4.3.1. Bolinopsis infundibulum
- 322 The vertical distribution of *B. infundibulum* varied with the region and was distributed above the
- 323 pycnocline north of SLI, and below the pycnocline west of SLI. The region west of SLI was
- 324 characterized by an extremely cold water mass (<0°C) below the pycnocline (Fig. 3), and the
- 325 occurrence of *C. melanaster* above the pycnocline (Fig. 8). Therefore, the deep distribution of *B*.
- 326 infundibulum in the west of SLI could have been caused by both physical oceanographic factors and
- 327 biological interaction factors. We were unable assess the relative contributions of these variables.
- 328 A polynya was present in 2017 (Grebmeier, J.M. pers. comm.), and the loss of saline and
- dense brine water during the formation of ice may have forced *B. infundibulum* to be distributed
- 330 below the pycnocline. Several ctenophores, Mnemiopsis leidyi, Pleurobrachia spp., and Beroe
- 331 spp., do not appear to cross a strong pycnocline, and instead distribute either above or below the
- 332 pycnocline (Roe et al., 1984; Vereshchaka, 2002; Haraldsson et al., 2014). When a strong

333	pycnocline developed west of SLI, <i>B. infundibulum</i> was restricted to below the pycnocline and did
334	not appear to migrate upward across the pycnocline. <i>B. infundibulum</i> can live under cold ($<0^{\circ}$ C)
335	conditions (Raskoff et al., 2005), and may be able to survive under the cold conditions that occur
336	below the pycnocline in this region.
337	Due to predator-prey interactions, the occurrence of large scyphomedusae C. melanaster
338	above the pycnocline west of SLI may have caused <i>B. infundibulum</i> to avoid that layer. GAM
339	analysis revealed a negative interaction between these two species. C. melanaster feeds on
340	gelatinous zooplankton (Purcell, 1991), and a prey-predator interaction between C. melanaster and
341	Bolinopsis spp. has been noted off Japan (Kinoshita et al., 2006); thus, the distribution of B.
342	<i>infundibulum</i> that we observed may have been a behavioural avoidance of predation by C.
343	melanaster. Indeed, in the Nordic fjord, B. infundibulum has been shown to remain below the
344	pycnocline to avoid predation from a large predatory scyphomedusae, Cyanea capillata (Båmstedt
345	and Martinussen, 2015). However, in the Canada Basin, in the western Arctic Ocean, the
346	coexistence of C. melanaster and B. infundibulum within the same depth layer has been reported
347	(Raskoff et al., 2005; Purcell et al., 2010).
348	Although we identified two possible factors that may explain horizontal changes in the
349	vertical distribution of <i>B. infundibulum</i> , physical oceanographic factors and biological interaction
350	factors, we cannot conclude which factor is most important in determining the regional changes in

352	Oyashio Current in the western subarctic Pacific Ocean (Toyokawa et al., 2003) and Canada Basin,
353	in the western Arctic Ocean (Raskoff et al., 2005; Purcell et al., 2010). These findings suggest that
354	B. infundibulum exhibits much flexibility in terms of their vertical distribution and ability to adapt to
355	various environments and regions.
356	
357	4.3.2. Chrysaora melanaster and Beroe sp.
358	We found that C. melanaster was distributed above the pycnocline at most stations, and the GAM
359	analysis revealed a significant positive interaction between C. melanaster abundance and
360	temperature (p <0.01, Table 1, Fig. 11). Previous studies have reported that <i>C. melanaster</i> is
361	distributed above the pycnocline (Brodeur, 1998; Brodeur et al., 2002; Raskoff et al., 2005;
362	Gorbatenko et al., 2009; Radchenko, 2013; Brodeur et al., 2017). Raskoff et al. (2005) reported
363	that the above pycnocline distribution of C. melanaster may be explained by their feeding on
364	copepods and gelatinous zooplankton, which are more abundant above the pycnocline. Radchenko
365	(2013) noted that the standing stock of zooplankton is more important than the thermal condition
366	above the pycnocline in explaining the vertical distribution of this species. Previous studies have
367	shown that C. melanaster feed on crustaceans, gelatinous zooplankton, larvae of walleye pollock,

vertical distribution. B. infundibulum has been reported to occur at depths up to 1250 m in the

351

368 ostracods, and decapod larvae in the Bering Sea (Brodeur et al., 2002; Zavolokin et al., 2008;

369	Ruzicka et al., 2020). Video images from the present study showed that hydromedusae Aglantha
370	<i>digitale</i> was abundant above the pycnocline (Maekakuchi unpublished data). Since A. digitale is
371	also an important food item for C. melanaster (Radchenko, 2013), we suggest that the observed
372	distribution of C. melanaster may be related to the abundance of their food items, such as copepods
373	and gelatinous zooplankton, above the pycnocline.
374	Standing stocks of C. melanaster were highest in the southern region in this study (south
375	of SLI). <i>C. melanaster</i> is very abundant in the southeastern Bering Sea shelf where it can exert a
376	substantial effect on other pelagic animals and marine food web structure in that region (Brodeur et
377	al., 2002, 2008), and C. melanaster has been reported in lower abundances farther north, in the
378	Canada Basin of the western Arctic Ocean (Raskoff et al., 2005; Purcell et al., 2010). These
379	observations suggest that the high standing stocks of C. melanaster in the southern region of this
380	study may reflect the regional differences in standing stocks.
381	The GAM analysis revealed a positive relationship between Beroe sp. and the abundance
382	of <i>B. infundibulum.</i> Beroe sp. were mainly distributed above the pycnocline, but they were
383	distributed below the pycnocline in the west of SLI, which parallels the regional pattern of vertical
384	distribution for <i>B. infundibulum</i> (Fig. 9). We suggest that predator-prey relationships may be the
385	cause of this parallel regional vertical distribution pattern of <i>Beroe</i> sp. with those of <i>B. infundibulum</i> .
386	Beroe spp. has been shown to feed on B. infundibulum (Greve, 1970; Purcell, 1991), and the vertical

387	distribution	of Beroe	cucumis ha	is been	documented	to l	be similar	to 1	that	of their	prey	(Falkenhau	g,
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- 388 1996; Bandara et al., 2016). However, as noted for *B. infundibulum, Beroe* sp. may be restricted in
- their distribution to below the pycnocline due to highly saline water, so there could also be a
- 390 physical oceanographic mechanism affecting their distribution.
- 391
- 392 *4.4. Interannual changes in ctenophores and scyphomedusae*
- 393 This study was conducted in the same location and season over two consecutive years and standing
- 394 stocks of ctenophores and scyphomedusae were much lower in 2018 than in 2017, by a factor of
- 395 1/20 to 1/90. The biomass of large scyphomedusae in the southeastern Bering Sea has been shown
- to fluctuate annually due to climate variability (Brodeur et al., 2017). There, the biomass of large
- 397 scyphomedusae, which is dominated by C. melanaster, increased 20-fold from 1975 to 2000, then
- decreased to one-third of the maximum after 2001, possibly due to climate-induced changes in the
- lower trophic levels (Brodeur et al., 2008). In warm years, ice melts quickly, leading to late
- 400 pycnocline development and phytoplankton bloom initiation; thus, smaller copepods such as
- 401 *Pseudocalanus* and *Acartia* dominate the zooplankton biomass (Hunt et al., 2011), which may result
- 402 in poor food conditions for C. melanaster (Brodeur et al., 2008). Conversely, in cold years, the
- 403 phytoplankton bloom initiates earlier, and large copepods, such as *Calanus*, dominate the
- 404 zooplankton biomass and these provide sufficient food conditions for the survival and growth of

405	ephyrae of C. melanaster, thus leading to an increase in scyphomedusae biomass (Brodeur et al.,
406	2008). Large changes in biomass over a short period have been observed in other regions. For
407	example, the biomass of ctenophores and scyphomedusae along the Kurile Islands in the Western
408	Pacific Ocean increased 10-fold from 2011 to 2012; intrusions from the Bering Sea and Okhotsk Sea
409	was invoked to explain this high biomass (Radchenko, 2013). Thus, there are large annual
410	fluctuations in the biomass of ctenophores and scyphomedusae, which are related to climate change
411	and oceanographic conditions.
412	Observations of water masses with high (>8°C) and low (<2°C) temperature conditions
413	that were characterized by a high abundance of <i>B. infundibulum</i> in 2017 were much more limited in
414	2018, and a decrease in optimal thermal conditions for <i>B. infundibulum</i> (>8 and <2°C) may explain
415	their very low abundance that year. Brodeur et al. (2017) reported that the biomass of <i>C</i> .
416	melanaster was high around SLI during both warm and cold periods. This suggests that around
417	SLI, conditions are suitable for the growth of <i>C. melanaster</i> polyps. Regarding annual differences
418	in the oceanographic conditions of this region between 2017 and 2018, the sea ice retreated in April
419	during 2018, which was approximately 1 month earlier than in 2017 (see Appendix F which is
420	derived from Arctic Data archive System (ADS) (https://ads.nipr.ac.jp/). As previously noted,
421	early ice retreat may induce late phytoplankton blooms, the dominance of small-sized copepods, and
422	low productivity, severely affecting food availability for <i>C. melanaster</i> (Brodeur et al., 2008). As

423	the sea ice began to retreat faster in 2018 compared with 2017, food conditions were likely to have
424	been poor for the survival and growth of the ephyrae of <i>C. melanaster</i> . The ephyrae of <i>C</i> .
425	melanaster grow in the spring to become medusae in summer and reach their peak level of biomass
426	in autumn (Zavolokin et al., 2008). Thus, the lower numbers of <i>C. melanaster</i> in 2018 may be
427	related to poor food conditions for their ephyra larvae, which in turn, may be related to the early ice
428	retreat that year.
429	The ctenophores, B. infundibulum and Beroe sp., which dominated in this region, spend
430	their entire life cycle in the plankton. Thus, annual differences in food conditions are experienced
431	during their early juvenile life-history phases. The main food for small ctenophores and ephyrae of
432	scyphomedusae is copepods (Purcell, 1991). Thus, the timing of the ice retreat underlies annual
433	differences in food conditions: early ice retreat leads to low productivity with a dominance of small
434	copepods, whilst late ice retreat leads to high productivity due to the dominance of large copepods
435	(Brodeur et al., 2008). For ctenophores and scyphomedusae in this region, low standing stocks in
436	2018 may be related to low productivity of large copepods, which was caused by the early ice retreat
437	in that year.
438	
439	Acknowledgments

440 We thank the crew members of the T/S Oshoro-Maru and our collaborators, especially Toru

441	Hirawake, for assistance in collecting the video and CTD data. Constructive comments from
442	Richard D. Brodeur and one anonymous reviewer significantly improved the manuscript and are
443	highly appreciated. This work was conducted by the Arctic Challenge for Sustainability (ArCS)
444	Project (Program Grant Number JPMXD130000000) and ArCS II. Part of this study was
445	supported by a Grant-in-Aid for Scientific Research 20H03054 (B), 19H03037(B), 18K14506 (Early
446	Career Scientists), and 17H01483 (A) from the Japan Society for the Promotion of Science (JSPS).

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590 Table and figure captions

591	Table 1	. Results of generalized additive models (GAM) based on the abundance of ctenophores
592		and scyphomedusae (Bolinopsis infundibulum, Chrysaora melanaster, Beroe sp.), and
593		environmental parameters: depth, temperature, salinity, the effect of pycnocline (U: upper),
594		day/night, and species interactions of other jellyfishes. +: positive, -: negative, *: $p < 0.05$,
595		**: $p < 0.01$, ***: $p < 0.001$. Detailed patterns between each parameter are presented in
596		Fig. 11.
597	Fig. 1.	Location of the stations used to observe ctenophores and scyphomedusae by frame camera
598		in the northern Bering Sea in 9–22 July 2017 (left) and 2–12 July 2018 (right). Numbers in
599		italics denote the depth strata in meters.
600	Fig. 2.	Captured images (A): <i>Bolinopsis infundibulum</i> (a), <i>Chrysaora melanaster</i> (b), <i>Beroe</i> sp. (c).
601		Frame camera (B): video camera (a), current fin (b), weight (20 kg x 4) (c), observation
602		frame (d), halogen light (x 2) (e), electronic data cable (f). Schema shows the calculation
603		of the observed volume (C). For details, see the text.
604	Fig. 3.	Temperature (upper) and salinity (lower) cross-sections at each line set in the northern Bering
605		Sea during 9–22 July 2017. The location of each line is shown in the upper-right map.

606	Fig. 4.	Temperature (upper) and salinity (lower) cross-sections at each line set in the northern
607		Bering Sea during 2–12 July 2018. The location of each line is shown in the upper-right
608		map.
609	Fig. 5.	Horizontal distribution of standing stock (ind. m ⁻²) of <i>Bolinopsis infundibulum</i> (left),
610		Chrysaora melanaster (middle), and Beroe sp. (right) in the northern Bering Sea during 9-
611		22 July 2017 (upper) and 2–12 July 2018 (lower).
612	Fig. 6.	Diel changes in the vertical distribution of <i>Bolinopsis infundibulum</i> at four stations (St. 10,
613		14, 20, 23) in the northern Bering Sea during 9–22 July 2017. Thick bars represent the
614		distribution core $(D_{25\%}-D_{75\%})$ split with the distribution centre $(D_{50\%})$. Thin bars show the
615		ranges of $D_{10\%}$ and $D_{90\%}$. Horizontal black bars at the top indicate night-time samplings.
616		Shaded zones represent the sea bottom. Triangles represent pycnocline depths. Details of
617		each observation are presented in Appendix C (St. 10) and Appendix D (Sts. 14, 20, 23).
618	Fig. 7.	Vertical distribution of temperature, salinity, and Bolinopsis infundibulum in the northern
619		Bering Sea during 9–22 July 2017. Triangles represent pycnocline depths.
620	Fig. 8.	Vertical distribution of temperature, salinity and Chrysaora melanaster in the northern
621		Bering Sea during 9–22 July 2017. Triangles represent pycnocline depths.
622	Fig. 9.	Vertical distribution of temperature, salinity and Beroe sp. in the northern Bering Sea
623		during 9–22 July 2017. Triangles represent pycnocline depths.

37

624	Fig. 10. T-S diagrams of all stations in 2017 (left) and 2018 (right) (A). Abundance (ind. m ⁻³) of
625	Bolinopsis infundibulum at 2 m intervals is shown by bubble plots on T-S diagrams (B).
626	Fig. 11. Result of the generalized additive model (GAM) based on abundance anomalies of three
627	large gelatinous zooplankton with environmental parameters.
628	Appendix A (Table). Data for ctenophores and scyphomedusae observations in the northern Bering
629	Sea during 9–22 July 2017 and 2–12 July 2018. One cast required approximately 20 min.
630	To evaluate day-night differences, 11 observations were made at 1 h intervals at St. 10 from
631	19:40 13 July to 7:10 14 July 2017. During that period, seven day (19:40, 21:15, 23:12,
632	0:15, 6:10, 7:10) and four night (1:12, 2:13, 3:12, 4:13) casts were made. Observed
633	volume (m ³) and total counts of each species are also shown for each cast.
634	Appendix B (Figure). Scatter plot on the density of <i>Bolinopsis infundibulum</i> quantified during
635	descent and ascent.
636	Appendix C (Figure). Diel changes in the vertical distribution of <i>Bolinopsis infundibulum</i> at St. 10
637	during 13–14 July 2017. Open and solid markers represent day and night, respectively.
638	Appendix D (Figure). Day (open) and night (solid) vertical distribution of <i>Bolinopsis infundibulum</i>
639	at St. 14, 20, 23 during 17–21 July 2017.
640	Appendix E (Figure). Contour plots for abundance anomalies of three large gelatinous zooplankton
641	on a T-S diagram.

642	Appendix F (Figure).	Sea ice concentration on April 1st in 2017 (left) and 2018 (right).	Images
643	were downloa	ded from the Arctic Data archive System (ADS) (https://ads.nipr.ac	.jp/).
644	Video supplemental m	aterial 1: Example of video image captured at St. 10 (cast 14) on 14	July
645	2017.		
646	Video supplemental m	aterial 2: Example of video image captured at St. 19 (cast 27) on 19	July

647 2017.

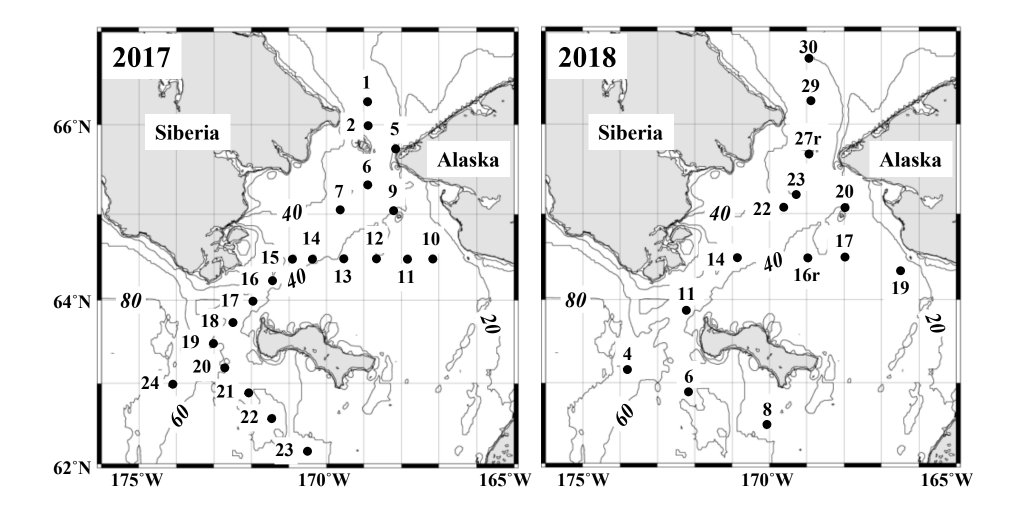
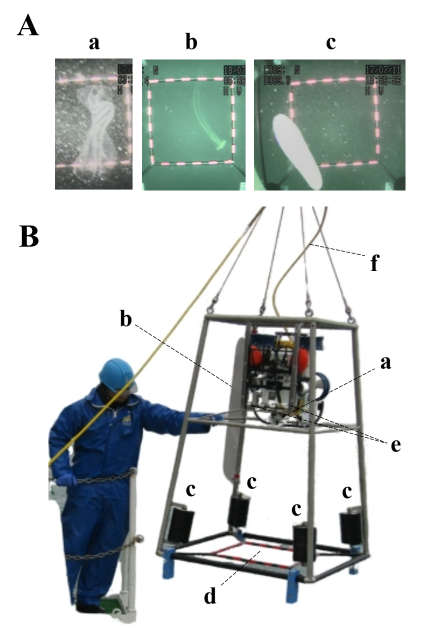
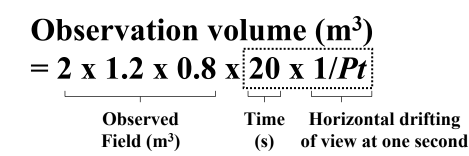


Fig. 1. Location of the stations used to observe ctenophores and scyphomedusae by frame camera in the northern Bering Sea in 9–22 July 2017 (left) and 2–12 July 2018 (right). Numbers in italics denote the depth strata in meters.





Pt: passage time within the observation field (s)

Effect of horizontal advection by current

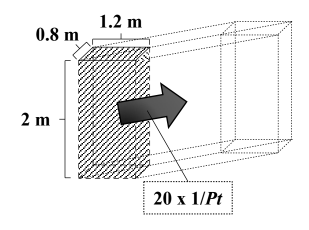
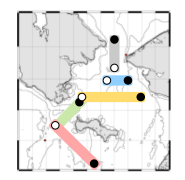


Fig. 2. Captured images (A): *Bolinopsis infundibulum* (a), *Chrysaora melanaster* (b), *Beroe* sp. (c). Frame camera (B): video camera (a), current fin (b), weight (20 kg x 4) (c), observation frame (d), halogen light (x 2) (e), electronic data cable (f). Schema shows the calculation of the observed volume (C). For details, see the text.

С



2017

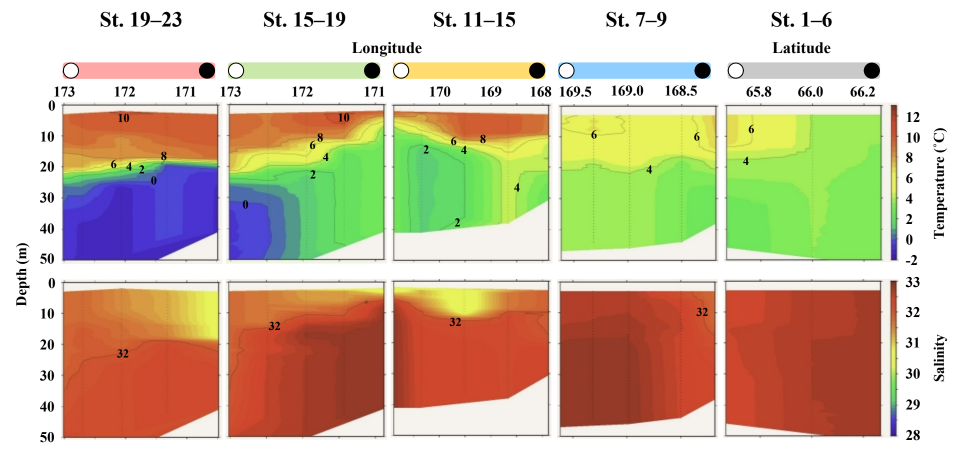
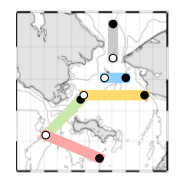


Fig. 3. Temperature (upper) and salinity (lower) cross-sections at each line set in the northern Bering Sea during 9–22 July 2017. The location of each line is shown in the upper-right map.





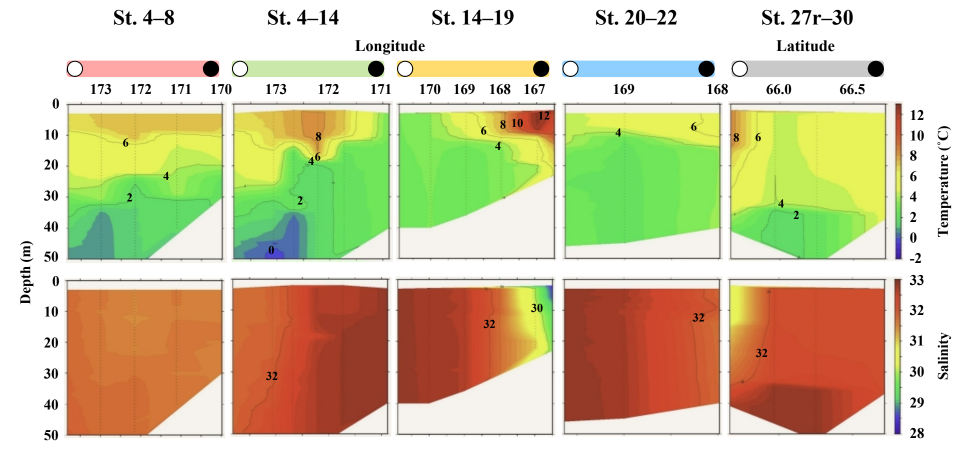


Fig. 4. Temperature (upper) and salinity (lower) cross-sections at each line set in the northern Bering Sea during 2–12 July 2018. The location of each line is shown in the upper-right map.

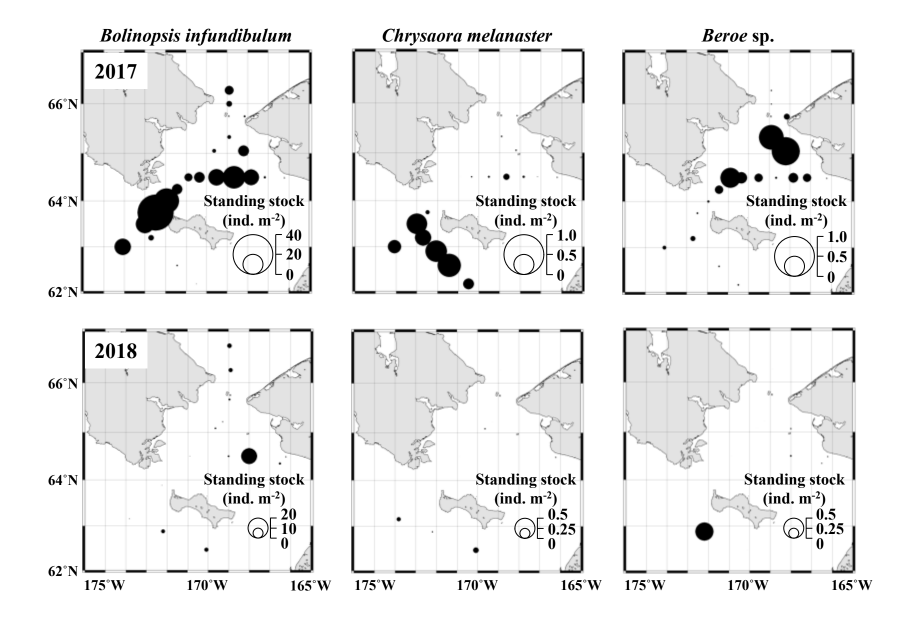


Fig. 5. Horizontal distribution of standing stock (ind. m⁻²) of *Bolinopsis infundibulum* (left), *Chrysaora melanaster* (middle), and *Beroe* sp. (right) in the northern Bering Sea during 9–22 July 2017 (upper) and 2–12 July 2018 (lower).

Local time

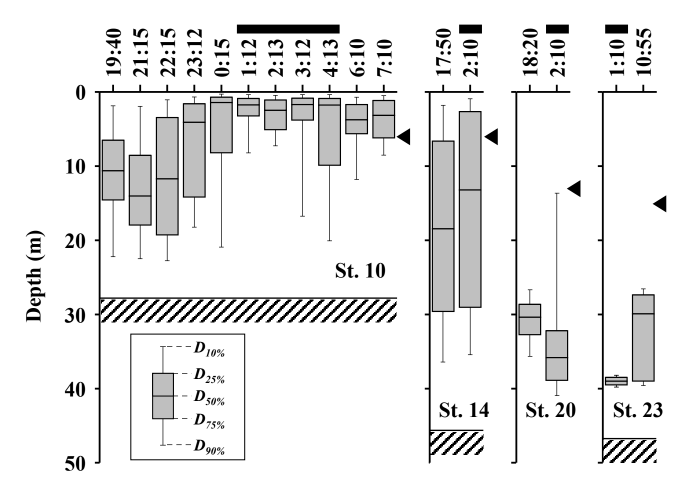


Fig. 6. Diel changes in the vertical distribution of *Bolinopsis infundibulum* at four stations (St. 10, 14, 20, 23) in the northern Bering Sea during 9–22 July 2017. Thick bars represent the distribution core $(D_{25\%} - D_{75\%})$ split with the distribution centre $(D_{50\%})$. Thin bars show the ranges of $D_{10\%}$ and $D_{90\%}$. Horizontal black bars at the top indicate night-time samplings. Triangles represent pycnocline depths. Details of each observation are presented in Electronic Supplement 1 (St. 10) and 2 (Sts. 14, 20, 23).

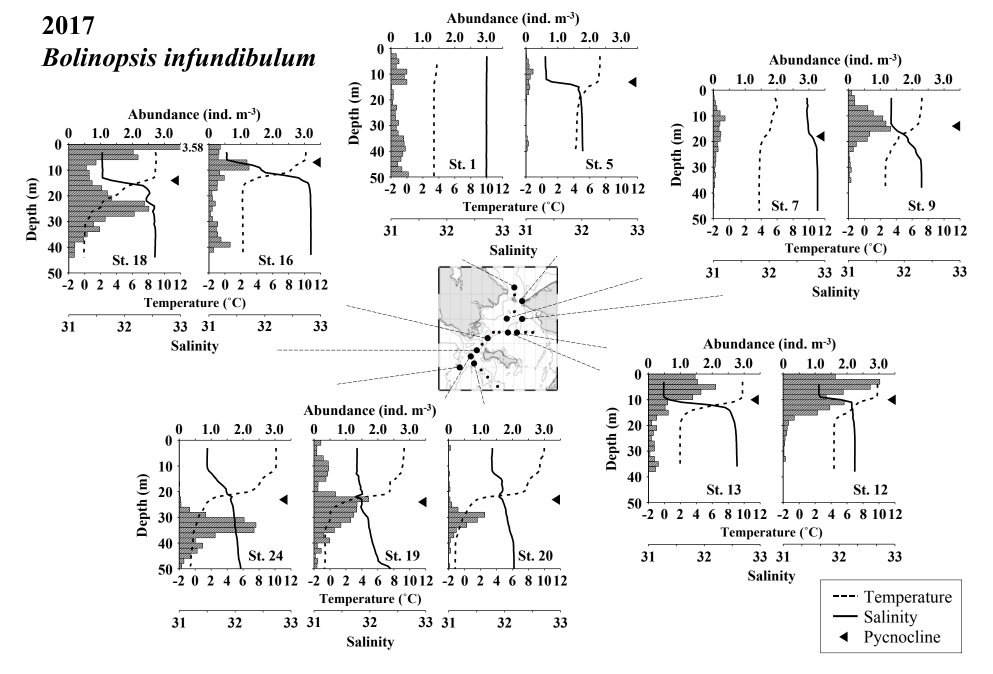


Fig. 7. Vertical distribution of temperature, salinity, and *Bolinopsis infundibulum* in the northern Bering Sea during 9–22 July 2017. Triangles represent pycnocline depths.

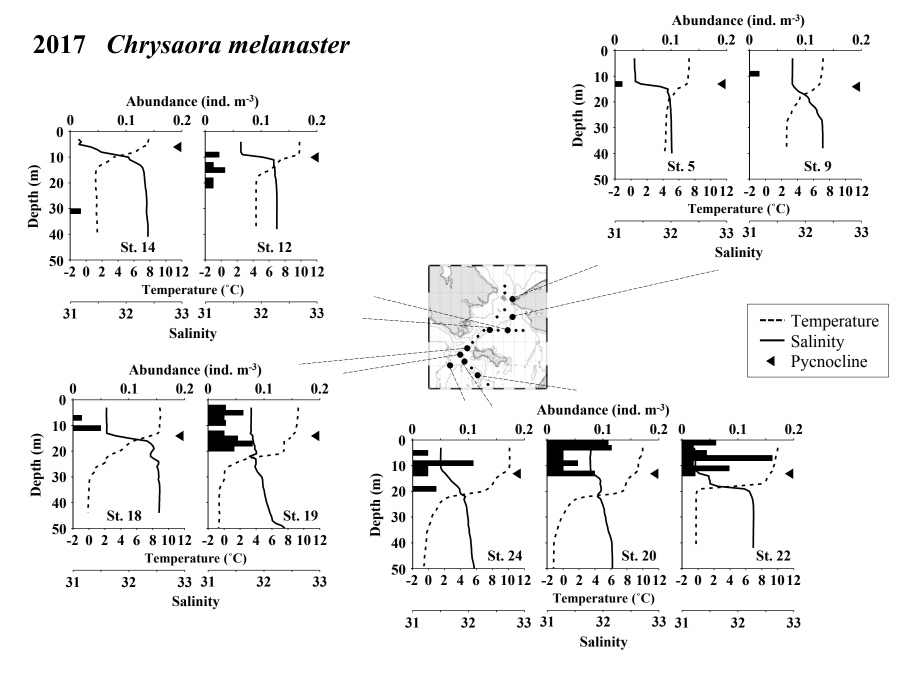


Fig. 8. Vertical distribution of temperature, salinity and *Chrysaora melanaster* in the northern Bering Sea during 9–22 July 2017. Triangles represent pycnocline depths.



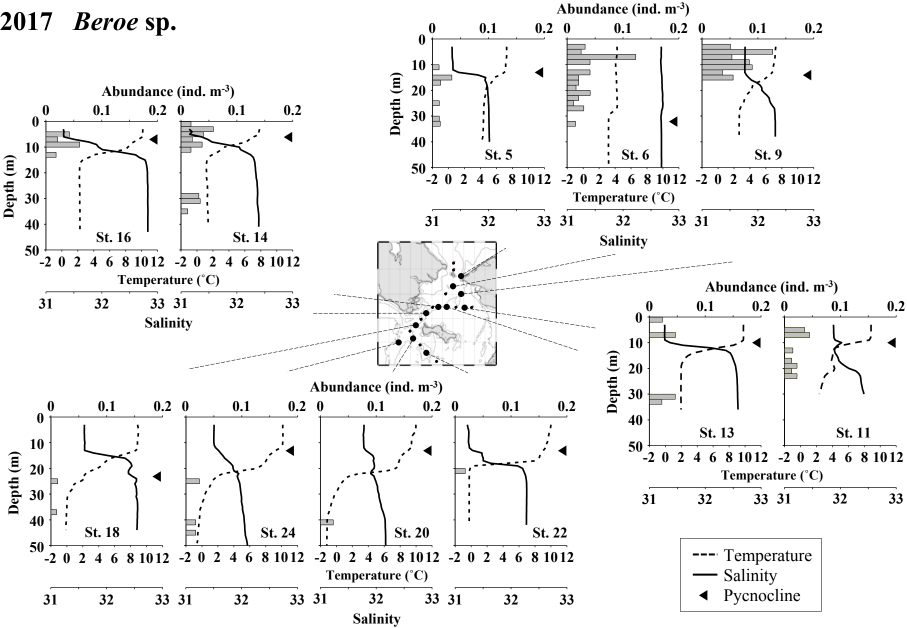


Fig. 9. Vertical distribution of temperature, salinity and *Beroe* sp. in the northern Bering Sea during 9–22 July 2017. Triangles represent pycnocline depths.

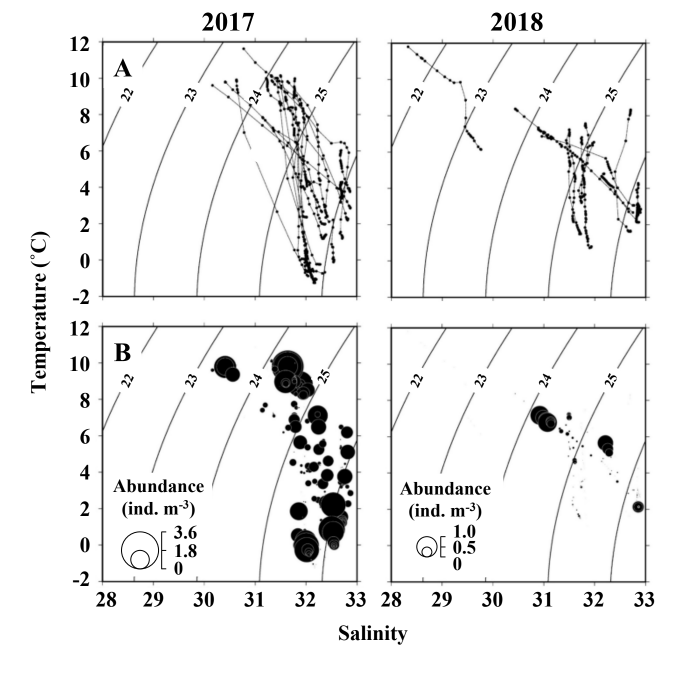


Fig. 10. T-S diagrams of all stations in 2017 (left) and 2018 (right) (A). Abundance (ind. m⁻³) of *Bolinopsis infundibulum* at 2 m intervals is shown by bubble plots on T-S diagrams (B).

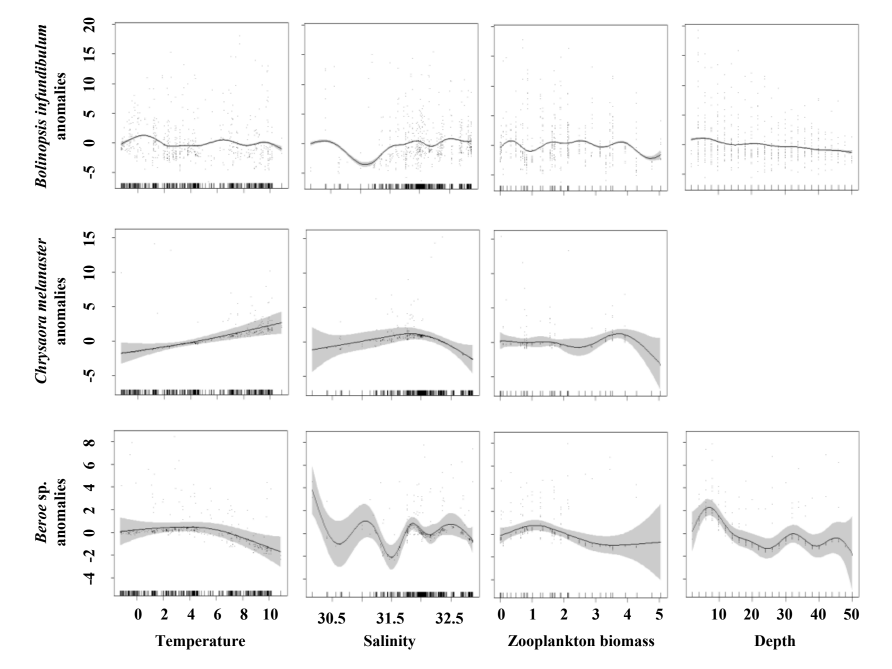
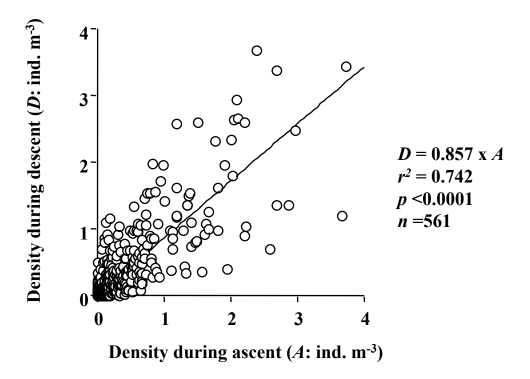


Fig. 11. Result of generalized additive model (GAM) based on abundance anomalies of three large gelatinous zooplankton with environmental parameters.

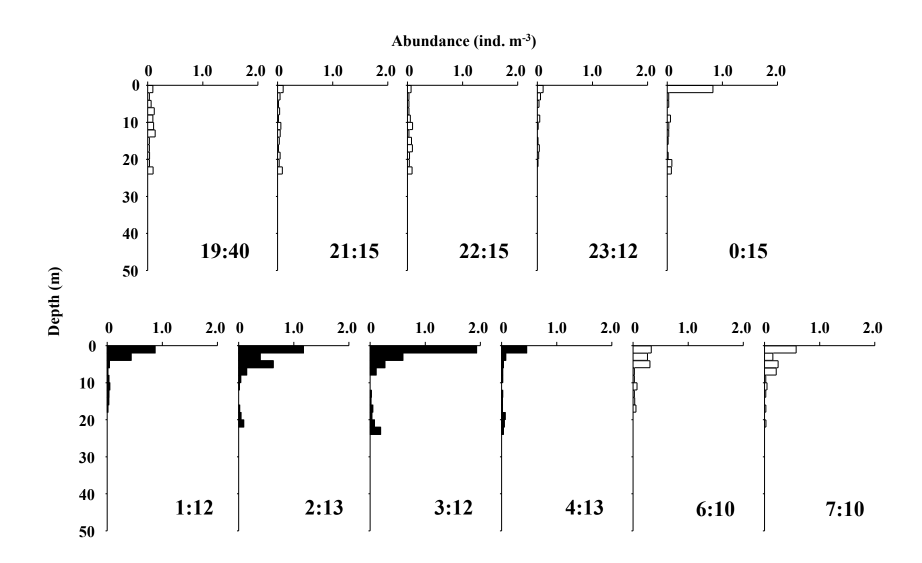
Appendix A. Data for ctenophores and scyphomedusae observations in the northern Bering Sea during 9-22 July 2017 and 2-12 July 2018. One cast required approximately 20 min. To evaluate day-night differences, 11 observations were made at 1 h intervals at St. 10 from 19-40 13 July to 7:10 14 July 2017. During that period, seven day (19-40, 21:15, 23:12, 0:15, 6:10, 7:10) and four night (1:12, 2:13, 3:12, 4:13) casts were made. Observed volume (m²) and total counts of each species during descent frank asserved for a for a star at also shown.

								Descent				Ascent			
Year	Station		Position Lat. (N)	Lon. (W)	Local time Date	Hour	— Day/night	Observed	Number of counted		Observed	Number of counted			
	(depth m)	Cast						volume (m3)	B. infundibulum	C. melanaster	Beroe sp.	volume (m ³)	B. infundibulum	ı C. melanaster	Beroe sp.
017															
	1 (57)	1	66 16'	168°54'	9 July	13:00	Day	677.8	221	0	1	688.9	111	0	0
	2 (53)	2	66°00'	168°54'	10 July	3:40	Night	1105.4	215	0	0	1069.1	177	0	1
	5 (44)	3	65°45'	168°09'	11 July	9:00	Day	701.9	59	1	2	818.1	17	0	5
	6 (55)	4	65°20'	168°54'	11 July	16:30	Day	780.5	77	0	10	946.0	113	1	13
	7 (51)	5	65°03'	169°38'	12 July	8:45	Day	995.0	89	0	0	934.6	100	0	0
	9 (42)	6	65°03'	168°12'	13 July	10:00	Day	697.8	262	1	20	620.8	163	0	8
	10 (28)	7	64°30'	167°10'	13 July	19:40	Day	496.0	48	0	2	492.6	15	0	3
	10 (28)	8	64°30'	167°10'	13 July	21:15	Day	547.2	35	0	9	475.4	15	0	7
	10 (28)	9	64°30'	167°10'	13 July	22:15	Day	428.4	16	0	1	340.7	14	0	5
	10 (28)	10	64°30'	167°10'	13 July	23:12	Day	478.2	20	0	9	274.3	3	0	3
	10 (28)	11	64°30'	167°10'	14 July	0:15	Day	508.4	25	0	5	438.0	58	0	2
	10 (28)	12	64°30'	167°10'	14 July	1:12	Night	459.5	30	0	0	297.4	35	0	5
	10 (28)	13	64°30'	167°10'	14 July	2:13	Night	452.6	62	0	5	272.0	73	0	2
	10 (28)	14	64°30'	167°10'	14 July	3:12	Night	342.6	93	0	7	297.1	44	1	7
	10 (28)	15	64°30'	167°10'	14 July	4:13	Night	430.4	18	0	3	357.0	27	0	6
	10 (28)	16	64°30'	167°10'	14 July	6:10	Day	282.9	32	0	5	250.6	15	0	4
	10 (28)	17	64°30'	167°10'	14 July	7:10	Day	285.3	43	0	3	305.9	18	0	2
	11 (34)	18	64°30'	167°50'	16 July	20:12	Day	615.9	400	1	6	504.9	300	1	4
	12 (42)	19	64°30'	168°40'	17 July	3:50	Night	710.2	549	3	2	475.3	320	2	0
	13 (40)	20	64°30'	169°31'	17 July	11:40	Day	425.4	161	0	5	361.3	271	1	1
	14 (46)	21	64°30'	170°21'	17 July	17:50	Day	609.2	227	1	5	651.0	279	0	10
	14 (46)	22	64°30'	170°21'	18 July	2:10	Night	724.2	146	0	4	815.5	304	1	5
	15 (45)	23	64°30'	170°53'	18 July	10:50	Day	677.3	187	0	17	521.8	116	0	4
	16 (47)	24	64°15'	171°26'	18 July	16:40	Day	680.5	206	0	2	569.9	197	0	5
	17 (53)	25	64°00'	171°57'	18 July	21:50	Day	935.0	783	0	0	889.6	487	0	2
	18 (48)	26	63°45'	172°29'	19 July	3:50	Night	870.1	1026	1	2	880.8	861	3	0
	19 (65)	27	63°30'	173°00'	19 July	10:55	Day	661.7	324	6	0	522.8	283	7	0
	20 (66)	28	63°12'	172°42'	19 July	18:20	Day	658.7	79	7	0	561.1	109	9	1
	20 (66)	29	63°12'	172°42'	20 July	2:10	Night	731.5	69	3	4	678.7	130	3	4
	21 (54)	30	62°54'	172°04'	20 July	10:35	Day	548.4	0	7	0	480.3	0	7	0
	22 (47)	31	62°35'	171°26'	20 July	18:45	Day	520.7	12	17	1	427.5	18	3	0
	23 (46)	32	62°10'	170°30'	21 July	1:10	Night	518.2	1	1	0	404.7	0	5	0
	23 (46)	33	62°10'	170°30'	21 July	10:55	Day	330.2	2	3	1	510.5	1	6	0
	24 (76)	34	63°00'	174°05'	22 July	1:30	Night	645.3	295	6	1	527.1	251	2	2
018					-		-								
	4 (76)	35	63°09'	173°50'	2 July	14:08	Day	1144.5	3	2	0	928.4	3	3	0
	6 (56)	36	62°53'	172°12'	3 July	1:34	Night	875.9	44	0	5	928.1	94	0	13
	8 (36)	37	62°28'	170°05'	3 July	12:50	Day	515.9	56	3	0	334.1	31	0	0
	11 (55)	38	63°53'	172°15'	4 July	14:58	Day	794.8	1	0	0	750.9	2	0	0
	14 (46)	39	64°30'	170°54'	5 July	1:20	Night	704.3	1	0	0	646.8	0	0	0
	17 (35)	40	64°30'	168°00'	5 July	14:54	Day	279.9	10	0	0	487.6	33	0	0
	19 (28)	41	64°21'	166°30'	5 July	21:56	Day	294.7	0	0	0	247.6	0	0	0
	20 (45)	42	65°04'	168°00'	6 July	6:48	Day	512.0	11	0	0	656.3	15	0	0
	22 (52)	43	65°05'	169°39'	6 July	16:40	Day	588.7	6	0	0	392.4	2	0	0
	23 (49)	44	65°13'	169°18'	7 July	21:58	Day	711.0	39	0	0	706.5	91	0	0
	30 (42)	45	66°44'	168°58'	11 July	0:50	Day	652.9	14	0	0	475.6	17	0	0
	29 (56)	46	66°15'	168°54'	11 July	10:18	Day	575.4	10	0	0	571.8	22	0	0
	27 (56) 27r (46)	40	65°40'	168°58'	11 July	21:37	Day	397.1	59	0	0	619.5	51	0	0
	271 (40) 16r (41)	48	64°30'	169°00'	12 July	9:30	Day	305.4	164	0	0	244.7	36	0	0

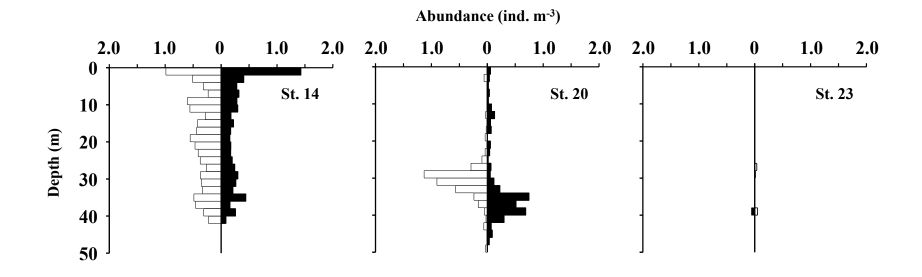




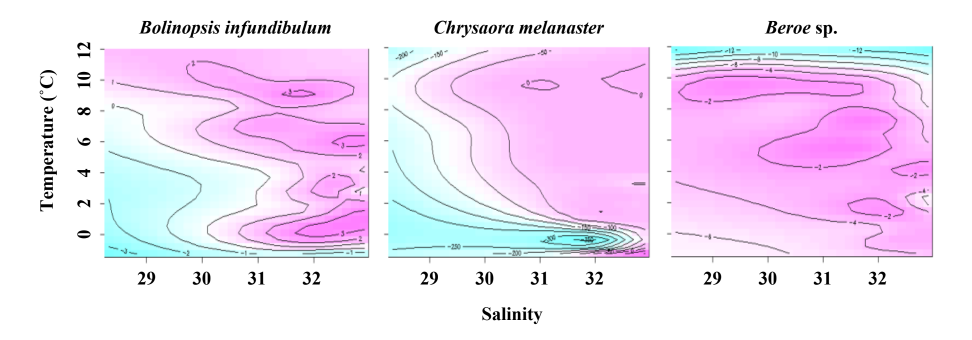
Appendix B. Scatter plot on the density of *Bolinopsis infundibulum* quantified during descent and ascent.



Appendix C. Diel changes in the vertical distribution of *Bolinopsis infundibulum* at St. 10 during 13–14 July 2017. Open and solid markers represent day and night, respectively.



Appendix D. Day (open) and night (solid) vertical distribution of *Bolinopsis infundibulum* at St. 14, 20, 23 during 17–21 July 2017.

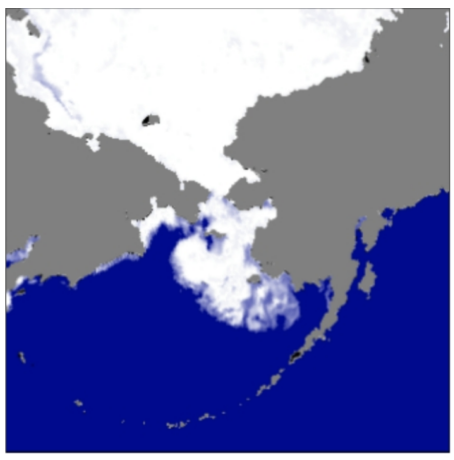


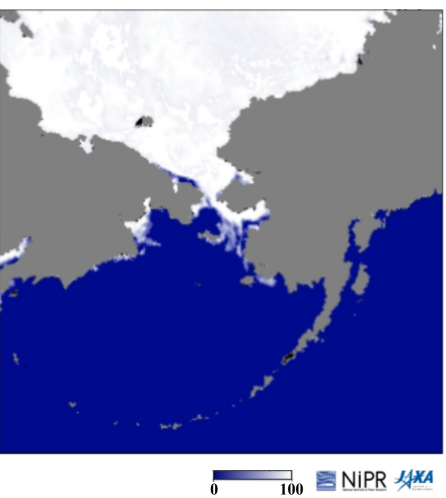
Appendix E. Contour plots for abundance anomalies of three large gelatinous zooplankton on a T-S diagram.

Sea ice concentration

2017/04/01

2018/04/01





Appendix F. Sea ice concentration on April 1st in 2017 (left) and 2018 (right). Images were downloaded from Arctic Data archive System (ADS) (https://ads.nipr.ac.jp/).