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- 2 Year-long benthic measurements of environmental conditions indicate high sponge
- 3 biomass is related to strong bottom currents over the Northern Labrador shelf
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30 Key words: deep-sea sponges, sponge grounds, benthic-pelagic coupling, organic matter

31 transport, tidal dynamics, nutrients

#### 33 Abstract

34 Deep-sea sponge grounds are distributed globally and are considered hotspots of biological 35 diversity and biogeochemical cycling. To date, little is known about the environmental constraints that control where deep-sea sponge grounds occur and what conditions allow high 36 37 sponge biomass to develop in the deep sea. Here, we characterize oceanographic conditions at 38 two contrasting high- and low-sponge-biomass sites off the northern Labrador Shelf in 39 Canadian waters. Unique data for the region were collected by year-long benthic lander deployments equipped with current meters, turbidity and chlorophyll-a sensors, and sediment 40 41 traps. Additionally, the regional oceanography was described by analysing vertical conductivity/salinity-temperature-depth (CTD) and Argo float profiles for the Northern 42 43 Labrador Shelf from 2005 to 2022, including those from the CTD casts taken at the benthic lander stations. Benthic fauna stable isotopes were analysed to identify potential food sources. 44 Our results revealed strong ( $0.26 \pm 0.14$  m s<sup>-1</sup>; mean  $\pm$  SD) semidiurnal tidal currents at the 45 high-sponge-biomass site, but twofold weaker currents ( $0.14 \pm 0.08$  m s<sup>-1</sup>; mean  $\pm$  SD) at the 46 low-sponge-biomass site. These tidal currents cause periodic temperature fluctuations, 47 48 sediment resuspension, intense vertical flows across the slope, which during spring, contribute 49 to transport of organic material to the seafloor during a diurnal tidal cycle. Periodic fluctuations 50 in bottom water temperature confirm the amplified transport across the shelf break at the highsponge-biomass site. The high-sponge-biomass area is situated where the Hudson Strait 51 52 Ouflow, the Irminger Current, and the West Greenland Current converge, which could lead to 53 downwelling. Bottom silicate concentrations were increased at the high-biomass sponge 54 ground due to advection of silicate-rich bottom water from Baffin Bay. Finally, the arrival of 55 chlorophyll-a rich material in spring at both the low- and high-sponge-biomass sites 56 demonstrated tight benthic-pelagic coupling prior to the onset of stratification. Mass fluxes of trapped material were higher and consisted of less degraded material at the high-sponge-57 58 biomass site. Stable isotope signatures indicated that soft corals (Primnoa resedueformis) fed on suspended particulate organic matter, while massive sponges (Geodia spp.) likely utilized 59 60 additional food sources. Our results imply that benthic fauna at the high-sponge-biomass site 61 benefit from strong tidal currents, which increases food supply, and favourable regional ocean currents that increase nutrient concentration in bottom waters. 62

#### 64 1 Introduction

Sponges are an ancient group of sessile filter feeders capable of pumping large quantities of 65 66 water through their bodies (Vogel, 1977; Bergquist, 1978; Leys et al., 2011), thereby 67 exchanging significant amounts of particulate- and dissolved organic matter and nutrients with 68 the water column (e.g., van Duyl et al., 2008; Maldonado et al., 2012; Kahn et al., 2015; Rix et al., 2016). In the deep sea, sponges can form dense aggregations, known as sponge grounds, 69 70 which are considered hotspots of macrofaunal diversity and abundance (Klitgaard, 1995; Buhl-71 Mortensen et al., 2010; Beazley et al., 2013; McIntyre et al., 2016), carbon- and nutrient cycling 72 (Kutti et al., 2013; Cathalot et al., 2015; Maldonado et al., 2020a), and benthic-pelagic coupling 73 (Pile and Young, 2006). Sponge grounds are often classified as Vulnerable Marine Ecosystems 74 (VMEs) as defined by the Food and Agriculture Organization of the United Nations (FAO, 75 2009). They form complex habitats that provide breeding grounds and shelter for commercially important fish species, increasing demersal fish biomass and diversity (Kenchington et al., 76 77 2013; Kutti et al., 2015; Meyer et al., 2019).

78 Deep-sea sponge ecosystems are currently under threat from anthropogenic disturbances such 79 as deep-water bottom trawling and climate change. Pham et al. (2019) found that large 80 quantities of sponges (~4% of total stock) have been removed by bottom trawling from sponge 81 grounds on the Flemish Cap. Deep-sea sponges are especially vulnerable to bottom fishing due 82 their longevity and slow growth (Leys and Lauzon, 1998; Hogg et al., 2010). Benthic trawling 83 reduces the density and diversity of deep-sea sponge grounds (Colaço et al., 2022), and 84 recovery of disturbed sponge habitats can take decades to centuries (Vieira et al., 2020). Recent 85 studies suggest that climate change also impacts deep-sea benthic fauna (Brito-Morales et al., 86 2020; Jorda et al., 2020). For example, modelling predicted that the suitable area for Vazella 87 pourtalesii on the Scotian Shelf would increase four-fold in the coming years due to warming 88 of colder waters around its current habitat (Beazley et al., 2021). Nevertheless, research on the 89 effect of climate change on deep-sea sponges is still in its infancy and to predict its effects on 90 sponge grounds, a better understanding of the environmental conditions that favour their 91 occurrence is needed.

In the past decades, research on deep-sea sponges has focussed on their physiology and feeding behaviour (e.g., Leys and Lauzon, 1998; Yahel et al., 2007; Kahn et al., 2015; Kazanidis et al., 2018; Maier et al., 2020b; Bart et al., 2021; de Kluijver et al., 2021), and assessing their spatial distributions using habitat suitability models (Knudby et al., 2013; Howell et al., 2016; Beazley et al., 2018; Murillo et al., 2018). More recently, data on the environmental conditions where sponge grounds are found have been gathered using long-term measurements from lander-

98 mounted equipment. These data indicate that sponge grounds are commonly found in areas with internal waves (Davison et al., 2019) and comparatively strong tidal currents which flush 99 100 the seafloor with oxygen and nutrient-rich water, and with a high suspended particle matter 101 load near the seabed (Roberts et al., 2018; Hanz et al., 2021a, 2021b). The spatial distribution of sponge grounds is also linked to gamete/larval dispersal and food availability (Abelson and 102 Denny, 1997; Robertson et al., 2017). In addition, sponges can alter the hydrodynamic 103 104 conditions of the benthic boundary layer by increasing the bottom roughness, creating conditions favourable for larval recruitment and suspended particle deposition (Culwick et al., 105 106 2020). These studies show that sponge grounds can be found in areas having a variety of environmental conditions, but little is known of the mechanisms controlling their spatial 107 108 distribution or what controls their biomass.

109 The Canadian Atlantic continental shelf breaks and upper slopes, including the northern Labrador Shelf, host extensive sponge grounds (Kenchington et al., 2010; Knudby et al., 2013). 110 Sponge assemblages occur over a large depth range (200 - 2875 m) and are often aligned along 111 depth contours with presumably similar environmental conditions (Murillo et al., 2012; 112 113 Knudby et al., 2013; Steve W. Ross, pers. obs.). On the northern Labrador Shelf and upper slope, sponge assemblages consist mostly of *Geodia* spp. and glass (hexactinellid) sponges 114 (Kenchington et al., 2010) but with strongly varying sponge biomass among areas. Therefore, 115 this region provides an interesting setting to study which environmental conditions favour high 116 117 sponge biomass and to provide insight into the factors that drive the spatial distribution of sponge assemblages on the eastern Canadian Shelf. Furthermore, research on present 118 environmental conditions on the seafloor is timely as the Labrador Shelf region is one of the 119 fastest warming large marine ecosystems globally (~1 °C decade<sup>-1</sup>; Belkin, 2009), and 120 121 according to ensemble-based climate change prediction, critical water mass properties there, including temperature, particulate organic carbon, pH, and aragonite saturation, are likely to 122 123 change substantially by 2100 (Puerta et al., 2020). Therefore, any analysis of the contemporary 124 conditions provides a baseline or a benchmark for referencing future ocean and ecosystem 125 conditions. This study presents a valuable reference dataset for the upper slope of the Northern 126 Labrador Shelf against which future changes can be evaluated.

To obtain a better understanding of the environmental conditions and ecosystem functioning of sponge grounds on the upper slope of the northern Labrador Shelf, this study specifically aimed to examine at the high- and low-sponge-biomass sites: (i) differences in ocean dynamics, seawater properties and mixing regimes, (ii) the annual dynamics of near-bed environmental and hydrodynamic conditions, and (iii) differences in organic matter flux and food sources for 132 sponges and associated macrofauna. This study is the first to collect year-long hydrodynamic

- and environmental data simultaneously at a high- and a low-biomass sponge ground.
- 134 2 Material and methods

#### 135 2.1 Oceanographic setting and the study area

136 The study area comprises the northern Labrador Shelf and upper slope and extends from the 137 south-eastern Hudson Strait outflow region to the base of the Labrador slope (Figure 1A). This region is known for intense mixing and water mass transformation (Dunbar, 1951; Kollmever 138 et al., 1967; Griffiths et al., 1981; Drinkwater and Jones, 1987; Yashayaev, 2007) and four 139 distinct flow components can be identified (Figure 1A; Smith et al., 1937; Yashayaev, 2007; 140 141 Straneo and Saucier, 2008; Curry et al., 2011, 2014): first, the cold and relatively fresh Arctic 142 outflow, passing through the Davis Strait via the Baffin Island Current (BIC), enters the region 143 from the north as Arctic Water (AW) and Baffin Bay Water (BBW; Sherwood et al., 2021); 144 second, the West Greenland Current (WGC) approaches our study site from the northeast; third, Irminger Water (IW), a warmer and saltier water mass, can often be seen underneath the WGC, 145 146 usually below 150 m depth; and fourth, Hudson Strait outflow water which enters the region from the west. The resulting aggregated boundary current joins the Labrador Current (LC) 147 148 flowing southward along the Labrador Shelf/slope, effectively forming and maintaining a baroclinic transition between the less-saline shelf water and the more-saline deep-basin water 149 150 (Yashayaev, 2007).

151 The northern Labrador Shelf hosts multiple sponge grounds with contrasting sponge community composition, density, and biomass (Kenchington et al., 2010; Dinn et al., 2020). 152 153 We selected a high-sponge-biomass site (HSB; 410 m depth) in the north and a low-spongebiomass site (LSB; 558 m depth) in the south of the study area (Table S1:; Figure 1B), 154 155 approximately 130 km apart. The seafloor at HSB was characterized by large-sized massive 156 demosponges (e.g. *Geodia* spp.) and glass sponges (e.g. *Asconema* spp.), large gorgonian corals 157 (Primnoa resedueformis), and rock boulders (Figure 2 A & B; Kenchington et al., 2010; Dinn et al., 2020). At LSB the seafloor mainly consisted of sediment, boulders, and small sponge 158 159 structures (e.g., Mycale spp.; Figure 2 C & D). The HSB lander was located on the shelf on a 2° slope and slope aspect was directed northwest at 60°. The LSB lander was located on the 160 upper slope, east of the shelf break, on a 7° slope and aspect was directed southeast at 105° 161 (Figure S1). The west-to-east slope angle was directed downhill, and north-to-south slope angle 162 163 was directed uphill at both lander sites (Figure S2).





165 Figure 1: Map of the study area with (A) the general circulation pattern (Curry et al., 2014). Cold Artic Water (AW) flows 166 southward through the Davis Strait and continues as the surface-intensified Baffin Island Current. The warmer, more saline 167 West Greenland Slope Current (WGC) of North Atlantic origin largely follows the continental slope in the depth range 150 -168 800 m and is deflected westward at approximately 64° N. Cold and fresh water leaves Hudson Strait and joins the BIC and 169 WGC to form the offshore branch of the Labrador Current (Straneo and Saucier, 2008). (B) Location of lander deployments 170 and CTD-casts, with sponge biomass (in kg  $m^{-1}$ ) based on Kenchington et al. (2010). Dotted line boxes indicate the shallow 171 shelf and deeper slope stations at both sites. HSB = high-sponge-biomass transect (red symbols), LSB = low-sponge-biomass 172 transect (blue symbols).

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Figure 2: Images of benthic lander deployment sites, at the high-sponge-biomass (HSB) site (A,B) and low-sponge-biomass
(LSB) site (C, D). Photographs were taken by drop camera at LSB and by ROV at HSB. ROV image credits:
ArcticNet/CSSF/DFO, CSSF = Canadian Scientific Submersible Facility, DFO = Department of Fisheries and Oceans.

- 180 2.2 Sampling methodology
- 181 2.2.1 Near-bed lander deployment

182 Landers were deployed during research cruise Amundsen 2018 leg 2c (27 July 2018) and 183 retrieved during research cruise Amundsen 2019 leg 1b (4 July 2019). The landers were each 184 equipped with a 2 mHz ADCP (upward-looking, Nortek Aquadopp), a sediment trap, and chlorophyll-a (chl-a) and optical backscatter sensors (Wetlabs – FLNTU; Table S1:). The 185 186 ADCPs collected an ensemble average of the 3D velocity field and echo intensity (acoustic 187 backscatter signal) every 600 seconds over one year along with pressure, temperature and data 188 from altitude sensors including heading, pitch, and roll. The ADCP was mounted 2 m above 189 the bottom, the blanking distance was 1.14 m, and the ADCP was programmed to measure 190 velocities at the first bin closest to the transducer head. Velocity data were recorded in beam 191 coordinates and transformed to ENU coordinates (East, North, Up) after recovery using the transformation matrix provided by the manufacturer. The chl-a and optical backscatter sensors 192 193 were programmed to measure every 600 seconds over the one-year period. Sediment traps (PPS 4/3, Technicap Inc.) with a surface area of 0.05 m<sup>2</sup> were equipped with twelve bottles for 194 195 particle collection and mounted 2 m above the bottom. Collection started at 15/08/2018 and 196 lasted until the end of the deployment. Different time intervals of bottle rotation were set to 197 increase sampling resolution during spring and summer months. The bottles rotated every 15 198 days from mid-August to mid-September 2018, every 30 days from mid-September to midNovember 2018, every 60 days from mid-November to mid-March 2019, then every 30 days
from mid-March to mid-May 2019, and every 15 days again from mid-May to mid-July 2019.
Prior to deployment, a 4% solution of formalin in brined seawater (40 psu) was added to each
bottle.

203 2.2.2 Water column and benthic sampling

204 Conductivity-Temperature-Depth (CTD) casts were performed over two cross-shelf transects 205 at LSB and HSB (Coté et al., 2018; Figure 1; Table S1:). Two CTD casts were carried out on the shelf and three over the slope, where the third or middle cast was performed above each 206 207 benthic lander deployment. The CTD-Rosette water column profiling and sampling package was equipped with a Seabird SBE 911plus system, which contained sensors to measure 208 temperature (Seabird SBE 3plus), conductivity (Seabird SBE 4), pressure (Paroscientific 209 210 Digiquartz®), dissolved oxygen (Seabird SBE 43), fluorescence (Seapoint), and a rosette water sampler with 12 Niskin bottles (12L each). CTD data were processed and "cleaned" with the 211 Sea-Bird SBE Data Processing software (Guillot, 2018). Water samples were taken from 212 213 Niskin bottles at five depths (5 m, 50 m, mid-water, 50 m above bottom, 10 m above bottom) for the determination of nutrients ( $NH_4^+$ ,  $NO_2^- + NO_3^-$ ,  $PO_4^{3-}$ ,  $SiO_2$ ), and suspended particulate 214 215 organic matter (sPOM).

216 Benthic macrofauna samples for stable isotope analysis were collected at the two lander 217 locations using a rock dredge on retrieval of the benthic landers (Coté et al., 2019; Table S2). 218 A description of the species found at the two locations can be found in Coté et al. (2019). The 219 rock dredge (7 mm mesh size) was deployed in "drift" mode at HSB, with a maximum speed 220 of two knots (~4 km h<sup>-1</sup>) for 10-20 minutes, and "tow" mode at LSB, with the ship moving at 221 one knot for 10 minutes. On deck, the dredge was rinsed, and the catch was subsampled and deposited in fish totes (64 L). The remaining material was sieved through a 2 mm mesh for 222 analysis of invertebrates and fishes. The total catch was photographed and preserved for species 223 224 identification and quantification. Samples for stable isotopes were frozen (-20 °C) for further analysis at the Netherlands Institute for Sea Research (NIOZ). 225

#### 226 2.2.3 Regional temperature and salinity profiles

To explore the regional oceanography on the northern Labrador Shelf and upper slope, vertical CTD profiles collected within the water depth range 330 - 2575 m (Figure S3) were extracted from the NOAA NODC World Ocean Dataset and profiling Argo float Global Argo Data Repository archives (Kieke and Yashayaev, 2015; Yashayaev and Loder, 2017). A similar approach was used in Kenchington et al. (2017). We used data collected between 2005 and 2022. Data from the Argo float profiles (N = 1472) were used to determine the seasonal 233 variability in temperature and salinity along the northwest Labrador shelf break. Specifically,

seawater properties were assessed of the corresponding water layers to the depth of the benthic

landers (LSB = 350 - 450 m, HSB = 550 - 650 m depth). Argo float profiles below ~59° N latitude were considered LSB and above as HSB. Temperature and salinity values were detrended for interannual variability using an 8<sup>th</sup> degree least-square polynomial fit.

#### 238 2.3 Laboratory analysis

Water column nutrient concentrations were analysed with a SEAL QuAATro analyser (Bran + Luebbe, Norderstedt, Germany) following standard colorimetric procedures. POM samples were freeze-dried, weighed, and analysed for organic carbon content, total nitrogen content, and  $\delta^{13}$ C using an elemental analyser (Flash 1112, THERMO Electron Corporation) coupled to an isotope ratio mass spectrometer (EA-IRMS, DELTA-V, THERMO Electron Corporation).

Sediment trap samples were filtered through a 1 mm sieve to remove large particles and 245 swimmers, then split into five sub-samples using a McLane WSD-10 rotary splitter, rinsed with 246 demineralized water to remove salts and formalin and subsequently freeze-dried and weighed 247 (Newton et al., 1994; Mienis et al., 2012). Lipids were extracted and analysed following the 248 method of Kiriakoulakis et al. (2004). Briefly, samples were spiked with internal standard 249 250  $(5\alpha(H)$ -cholestane), extracted by sonication in dichloromethane:methanol (9:1; x3). The 251 solvent was removed and samples were first trans-methylated (Christie, 1982) and then treated 252 with bis-trimethylsilyltrifluoroacetimide: trimethylsilane (99:1; 30-50 µL; 60 °C; 1 h) prior to 253 analysis by gas chromatography-mass spectrometry (GCMS). GCMS analyses were conducted using a GC Trace 1300 fitted with a split-splitless injector and column DB-5MS (60m x 254 255 0.25mm (i.d.), with film thickness 0.1 µm, non-polar stationary phase of 5% phenyl and 95% methyl silicone), using helium as a carrier gas (2 mL min<sup>-1</sup>). The GC oven was programmed 256 after 1 minute to rise from 60°C to 170°C at 6°C min<sup>-1</sup>, then from 170°C to 315°C at 2.5 °C 257 min<sup>-1</sup> and was then held at 315 °C for 15 min. The eluent from the GC was transferred directly 258 259 via a transfer line (320 °C) to the electron impact source of a Thermoquest ISQMS single 260 quadrupole mass spectrometer. Typical operating conditions were: ionisation potential 70 eV: source temperature 215°C; trap current 300 µA. Mass data were collected at a resolution of 261 262 600, cycling every second from 50–600 Daltons and were processed using Xcalibur software.

Compounds were identified either by comparison of their mass spectra and relative retention
indices with those available in the literature and/or by comparison with authentic standards.
Quantitative data were calculated by comparison of peak areas of the internal standard with

those of the compounds of interest, using the total ion current (TIC) chromatogram. The relative response factors of the analytes were determined individually for 36 representative fatty acids, sterols and an alkenone using authentic standards. Response factors for analytes where standards were unavailable were assumed to be identical to those of available compounds of the same class.

271 Sponges and other benthic fauna collected using a rock dredge were freeze-dried and homogenized with a pestle mortar/ball mill. Subsamples (ca. 10 mg) were transferred into 272 silver cups and acidified by addition of dilute HCL (2%, 5%, and 30%) to remove carbonates. 273 Organic carbon and  $\delta^{13}C$  were analysed on acidified subsamples, and total nitrogen and  $\delta^{15}N$ 274 275 were analysed on non-acidified subsamples using an Electron Analyser coupled to an Isotope Ratio Mass Spectrometer (Thermo flash EA 1112).  $\delta^{13}$ C and  $\delta^{15}$ N isotope values are expressed 276 in parts per thousand (‰) relative to the standards for carbon  $({}^{13}C/{}^{12}C = 0.0111802)$  and 277 nitrogen  $({}^{15}N/{}^{14}N = 0.0036782)$ , respectively. 278

#### 279 2.4 Data processing

280 The transformation of beam coordinates to ENU coordinates for the ADCP data was carried out in Matlab (MATLAB, 2010), and all other data processing steps used R using various R 281 packages (Wickham, 2007, 2016; Grolemund and Wickham, 2011; Neuwirth, 2014; signal 282 283 developers, 2014; Michna and Woods, 2019; Pedersen, 2019; R Core Team, 2019; Wickham and Bryan, 2019; Wilke, 2019; Kelley and Richards, 2020; Stoffer, 2020; Vaughan and 284 Dancho, 2020; Xie, 2020; Lovelace et al., 2022). Occasionally, pitch and roll data from the 285 ADCP sensor at HSB were shifted for a small period of the deployment, implying the lander 286 was occasionally moving a bit. However, removing these datapoints did not change the 287 outcome of any of the analyses, statistical tests, or descriptive statistics and these datapoints 288 289 were therefore retained in the HSB time series. Chl-a (in  $\mu g L^{-1}$ ) and turbidity (in NTU) concentrations were calculated from ping counts as described in the manual of the 290 291 manufacturer. To investigate connectivity in environmental variables between the two benthic 292 landers, and correlations between hydrodynamic and environmental conditions, a cross-293 correlation analysis with time lag was performed. Spectral analysis on lander data was 294 performed to examine recurring patterns or periodicity in the time-series data (e.g. Shumway 295 et al., 2000), and coherence analysis was carried out to assess correlation in periodicity between 296 landers and variables (Bloomfield, 2004). Spectral and coherence analyses were based on a 297 Fourier transformation on unfiltered data (Bloomfield, 2004). Prior to these analyses, time 298 series data were smoothed using modified lowpass Daniell filters (Bloomfield, 2004), to 299 remove periodicities shorter than 3 hours. The magnitude and direction of ADCP-recorded tidal

300 currents were analysed with least-squares harmonic analysis, using the t\_tide MATLAB 301 toolbox (Pawlowicz et al., 2002). Bottom currents and direction were compared to model 302 derived barotropic tidal currents, retrieved from the Oregon State University (OSU) Tidal 303 Inversion Software (OTIS; Egbert and Erofeeva, 2002). Sea-ice cover above the two benthic 304 landers was extracted from weekly ice charts (Canadian Government, 2022). Statistics are 305 presented as means  $\pm$  standard deviations. Slope aspect was estimated for each lander by taking 306 the wider topography into account (Gille et al., 2004).

307 3 Results

#### 308 3.1 Seawater properties over the northern Labrador Shelf and upper slope and

#### 309 regional oceanography

310 The CTD casts, performed July 2018, revealed a difference in seawater properties between the two transects (Figure 3; Figure S3). The surface water at the time of survey was relatively warm 311  $(2 - 6 \,^{\circ}\text{C})$  and fresh (31.2 to 33.8 psu) yet showing a significant offshore increase in 312 temperature and salinity. From the surface to the depth of 20-70 m, depending on the transect 313 314 and location, temperature decreased to sub-zero or near-zero at the shelf locations, to 3 °C at 315 the slope locations, and then increased again to 2.8 °C at 250 m depth on the shelf and to 4.3 °C 316 at 150 m on the slope. The temperature changes from cooling to warming with depth signify the Cold Intermediate Layer (CIL). Salinity in the CIL increased nearly monotonically with 317 318 depth across all stations. The stations at LSB were more saline overall than those at the 319 matching water depths on the HSB transect.

The oxygen concentration was highest in the surface waters (0 - 50 m) on the shelf and 320 321 decreased with depth at all CTD stations (Figure 4A). The bottom oxygen concentrations at the 322 lander stations were, for both transects, relatively depleted compared to the deep water CTD transects at similar depths. Concentrations of nitrate, phosphate, and silicate were lowest above 323 324 the thermocline  $(0 - 3 \mu M)$  and increased with depth, while ammonium and nitrite were higher near the surface than at depth (Figure 4B & C, Figure S4). The HSB station exhibited relatively 325 326 high nitrate, phosphate, and silicate concentrations at 10 and 50 metres above bottom compared 327 to similar depths at shelf and deep stations (Figure 4B & C, Figure S4). This increased nutrient concentration in the bottom waters was also apparent at the LSB station, but to a lesser degree. 328 329 Chl-a profiles showed a deep chlorophyll maximum along both transects at 50 m, and nearzero concentrations in the bottom waters (Figure S3D). Particulate organic carbon (POC) 330 concentrations were highest in the surface waters  $(8 - 38 \mu mol POC L^{-1})$  and on the shelf 331

- 332 (Figure S5). POC concentrations decreased with depth, and concentrations 10 m above bottom
- 333 were 1.48  $\mu$ mol POC L<sup>-1</sup> at HSB, and 5.95  $\mu$ mol POC L<sup>-1</sup> at LSB.
- The HSB lander was located in an area where three (surface) currents converge (Figure 5A).
- 335 Strong surface currents (>0.24 m s<sup>-1</sup> on average) carry water from the Hudson strait towards
- the Labrador shelf break, where this current meets two others that, respectively, flowed toward
- 337 the HSB site from the north and northeast. On convergence, the currents followed the
- bathymetry of the Labrador shelf break or upper slope southwardly.
- 339 The seawater in the region of HSB was warmer and less saline than around LSB for both depth
- 340 ranges within which the landers were deployed (Figure 5B & C; Figure S6). Bottom water
- 341 temperature shows a steeper decrease in February at LSB compared to HSB (Figure 5C).
- 342 Temperature and salinity show higher scatter at HSB than LSB throughout the season, but
- 343 variability in temperature is highest at HSB in February/March (Figure 5B & C).



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Figure 3: Hydrographic conditions in the study area: (A) temperature, (B) salinity and (C) temperature – salinity (TS) plots for the two transects. LSB = low-sponge-biomass, HSB = high-sponge-biomass. Depths of landers are indicated by the horizontal grey lines in A and B. Temperature and salinity profiles in A and B only show top 600 m, while TS plots include the entire water column.



350Figure 4: Oxygen (A), nitrate (B), and silicate (C) concentration profiles for the two transects. HSB = high-sponge-biomass351site, LSB = low-sponge-biomass.



Figure 5: A) general circulation pattern in the Labrador Sea based on drifter data from 1995 - 2020. The lander locations
are indicated by the coloured dots. B) seasonal temperature signal, from Argo float data, of the water layer in which HSB
lander is located. C) seasonal temperature signal of the water layer in which LSB is located.

#### 358 3.2.1 Near-bottom current velocities

359 In general, bottom current speeds were higher at the HSB compared to the LSB station (Table 360 1; Figure 6). The eastward velocity (u) was directed more eastward at HSB than at the LSB site 361 and northward velocity (v) was comparable between sites and directed southward. The residual 362 current was south-easterly at HSB and south-south-westerly at LSB (Figure 7). Vertical 363 velocity (w) was on average upward and comparable between HSB and LSB, but the range in vertical velocity was higher at HSB (-0.35 to 0.32 m s<sup>-1</sup>) compared to LSB (-0.11 to 0.21 m s<sup>-1</sup>) 364 <sup>1</sup>). Bottom currents were twice as high at HSB than at the LSB (Table 1), and peak bottom 365 current speeds were 0.75 m s<sup>-1</sup> (HSB) and 0.65 m s<sup>-1</sup> (LSB), with the third quantile at 0.33 m 366 s<sup>-1</sup> (HSB) and 0.18 m s<sup>-1</sup> (LSB). The pressure signal, a proxy for sea surface height, showed 367 peaks in variance preserving spectrum periodicity at the semidiurnal (M2, S2, N2), and diurnal 368 369 tidal harmonics (K1, O1; Figure 8 A). Bottom current speeds showed semi-diurnal and spring-370 neap tidal patterns, with bottom currents peaking every fortnight for both sites (Figure 6 C; 371 Figure 8 B; Figure 11). The major axes of the semidiurnal tidal ellipses were directed in a 372 northwest-southeast direction at HSB and a north-south direction at LSB, and were aligned 373 with the continental shelf and slope, respectively (Figure 8D). The M2 and S2 major axes at the HSB station (0.28 m s<sup>-1</sup> and 0.05 m s<sup>-1</sup>) were a factor of five larger than the corresponding 374 magnitudes at the LSB station, whereas diurnal major axes were small ( $< 1 \text{ cm s}^{-1}$ ) and of 375 376 similar magnitude at both locations. Frequency distributions of spectral variance showed 377 highest variability in semidiurnal periodicity for bottom current components at both sites, but 378 the peak in the variance-preserved spectrum was higher at HSB than at LSB. Furthermore, 379 spectral density for the HSB bottom current components also peaked at shorter frequencies (3-380 6 h) and at the fourteen-day spring-neap tide (Figure 8B). In addition, a superimposed seasonal 381 pattern can be seen at both sites, where the bottom current speed gradually increased from July 382 2018 to March 2019 and decreased again from March 2019 to July 2019.

383 The residual current followed roughly the topography at both sites with, on average, a slight downward cross-slope current at HSB and an upslope current at LSB (Table 1). However, 384 385 frequency distribution of current direction shows at HSB bottom current is mostly directed at 150°, which is 30° downslope of the along-slope direction, and at LSB mostly directed at 190°, 386 which is 5° downslope of the along-slope direction (Figure 7 C & D). High downward velocities 387 were recorded during periods having south-easterly and north-westerly current direction at 388 389 HSB. High upward velocities at LSB were recorded when current direction was south or south-390 westerly (Figure 9).

392 Bottom temperature was slightly warmer at HSB compared to LSB and increased at both sites 393 (0.2 - 0.3 °C) during December and January (Figure 10). The benthic lander temperature signal 394 aligned well with the seasonal temperature pattern retrieved by Argo float profiles (Figure 5 B & C). Acoustic backscatter signal (ABS) was similar for the two stations (Table 1; Figure 10 395 396 B) and showed higher values in winter months. Chl-a remained low from October to early 397 March when a spring peak was observed for both landers (Figure 10 C). Maximum chl-a concentration was lower at HSB (2.24  $\mu$ g L<sup>-1</sup>) than at the LSB (5.41  $\mu$ g L<sup>-1</sup>). The HSB station 398 399 showed spring bloom conditions from mid-March to the end of May, while at the LSB station 400 the spring bloom lasted from mid-March to early May. Turbidity was comparable at the two sites, and was elevated at HSB from February to April, and at LSB from December to January. 401 402 Turbidity increased at high south-easterly current velocities at HSB and high southerly current velocities at LSB (Figure 9 C & D). The higher variability in chl-a and turbidity at the LSB 403 404 site over the year (Table 1) was caused by several peaks in chl-a and turbidity that were an 405 order of magnitude higher than average values (Figure S7).

406 Daily temperature fluctuations were higher at HSB than at LSB. Cross and along slope water 407 transport influenced bottom temperature. For example, in the first week of September, temperature decreased when the current was directed northwest and increased when the current 408 409 was directed southeast (Figure 9 A-D; Figure 11 A-E). Temperature showed a reoccurring tidal signal, with higher peaks in spectral density for the semidiurnal periodicity at HSB than at LSB 410 411 (Figure 8C). Cross-correlation showed that near-bottom temperatures (daily averaged) were correlated between the two landers with a lag of five days ( $R^2 = 0.52$ ). ABS (Acoustic 412 Backscatter Signal) increased often at the turning of the tide and at high south-easterly current 413 414 velocities at HSB (Figure 11F; Figure 9C & G). Strong along slope bottom currents, which are 415 slightly directed downslope, increased ABS and turbidity at LSB (Figure 11 F; Figure 9C & G). Cross-correlation showed ABS was weakly correlated with bottom current speed at HSB 416  $(R^2 = 0.34)$  and LSB  $(R^2 = 0.44)$ . During the spring bloom, bottom chl-a concentration 417 418 increased at strong south-easterly current velocities at HSB (Figure S8) and showed a periodic 419 reoccurring signal (Figure S9A).

Temperature, chl-*a*, ABS, turbidity, all showed a reoccurring tidal signal, with higher peaks in spectral density for the semidiurnal periodicity at HSB than at LSB (Figure 8C). Ice cover seemed to affect the peak of chl-*a* concentration at the seafloor (Figure S9). However, both sites were located at the sea-ice border in the study area and had highly variable sea-ice coverage. Only during January coverage was above 70% at both sites. The Hudson Strait froze

- 425 up in early December and opened again in early June. During the spring bloom, between the
- 426 end of March and early May, sea-ice coverage tended to be higher at HSB than at LSB (Figure
- 427 S9D).

428<br/>429Table 1: Summary statistics for the long-term near-bottom measurements. Values are given as mean  $\pm$  standard deviation.429HSB = high-sponge-biomass lander, LSB = low-sponge-biomass lander. ABS = acoustic backscatter signal.

Variable	HSB	LSB
u (eastward velocity; m s <sup>-1</sup> )	$0.05\pm0.22$	$-0.01 \pm 0.09$
v (northward velocity; m s <sup>-1</sup> )	$-0.07 \pm 0.16$	$-0.09 \pm 0.11$
w (vertical velocity; m s <sup>-1</sup> )	$0.03\pm0.05$	$0.02\pm0.03$
Bottom current speed (m $s^{-1}$ )	$0.26\pm0.14$	$0.14\pm0.08$
Temperature (°C)	$3.70\pm0.17$	$3.58\pm0.17$
Daily temperature variability ( $\Delta^{\circ}C d^{-1}$ )	$0.25\pm0.16$	$0.17\pm0.1$
ABS (counts)	$98.1\pm9.8$	$96.6 \pm 11.0$
Chl- <i>a</i> concentration ( $\mu$ g L <sup>-1</sup> )	$0.11\pm0.03$	$0.08\pm0.10$
Turbidity (NTU)	$0.20\pm0.10$	$0.21\pm0.27$
Across slope velocity (m s <sup>-1</sup> )	$0.01 \pm 0.13$	$-0.01 \pm 0.01$
Along slope velocity (m $s^{-1}$ )	$-0.08 \pm 0.23$	$-0.09 \pm 0.11$



432 433 434 Figure 6: Time series of the flow velocities with eastward u velocity (A), northward v velocity (B), vertical w velocity (C), and bottom current speed (D). Plots show the hourly averaged data as transparent lines and the seven-day rolling means as

solid lines.







441 Figure 8: Variance preserving spectra for (A) pressure, (B) bottom current speed, (C) temperature, acoustic backscatter

442 signal (ABS), and chl-a, and (D) resulting tidal current ellipses for the two dominant diurnal and semidiurnal tidal

<sup>443</sup> *harmonics derived from the unfiltered ADCP velocities.* 



444

Figure 9: Progressive vector plot for 1 September to 7 September for the high-sponge-biomass site, HSB (A, B, C, D) and
low-sponge-biomass site, LSB (E, F, G, H), with temperature (A, E), vertical velocity (B, F), acoustic backscatter (C, G),
and turbidity (D, H) as colour variable. The dotted line indicates the along-slope direction at both lander sites.





- 449 450 451 concentration in  $\mu g L^{-1}(C)$ , and turbidity in NTU (D). Plots C and D are limited on the y-axis to 1.25  $\mu g L^{-1}$  and 2.5 NTU,
- respectively, for clarity. Chl-a and turbidity data without the Y-axis cut-offs are plotted in Figure S9.



454 Figure 11: Expanded detail for the first week of September for the eastward velocity u (A), northward velocity v (B), vertical 455 velocity (C), bottom current speed (D), temperature (E), acoustic backscatter signal (ABS; F), turbidity (G), and chl-a 456 concentration (H).

The average mass fluxes were higher at HSB  $(2.46 \pm 1.76 \text{ g m}^{-2} \text{ day}^{-1})$  than at LSB  $(1.43 \pm 0.93 \text{ m}^{-2} \text{ day}^{-1})$ 458 g m<sup>-2</sup> day<sup>-1</sup>), with highest fluxes in winter (October to April) at both sites and lowest in spring. 459 Average POC fluxes were also higher at HSB  $(3.07 \pm 1.91 \text{ mmol C m}^{-2} \text{ d}^{-1})$  than at LSB (1.91 460  $\pm 0.71$  mmol C m<sup>-2</sup> d<sup>-1</sup>). Organic carbon content was at HSB highest in autumn/summer months 461 (~2%) and at LSB highest in autumn (2-4%; data not shown). Average C:N ratios were lower 462 463 at HSB ( $8.6 \pm 3.2$ ) than at LSB ( $10.8 \pm 2.7$ ) and were higher in winter and also in May 2018 (Figure 12C). The  $\delta^{13}$ C ratios of trapped material were in winter higher at HSB than at LSB, 464 and in summer higher at LSB than at HSB (Figure 12D). The  $\delta^{15}N$  of trapped material was 465 comparable between sites, although slightly higher at LSB. Winter  $\delta^{15}$ N values were higher 466 than spring values, and at LSB the September and summer samples showed increased  $\delta^{15}N$ 467 (Figure 12E). The lipid flux was slightly higher at LSB, with low values in winter and peak 468 values during the spring bloom (Figure 12F). Unsaturated alcohols comprised the largest 469 fraction of lipids at LSB, especially in autumn and winter (Figure S10B). Peak lipid flux in 470 471 April consisted of 25% polyunsaturated fatty acids (PUFAs) at HSB (Figure S10C). Sterols made up the largest fraction of total lipids at HSB and LSB in May (Figure S10D). The sterol 472 fraction was lower in spring at both sites. Swimmers were found in the sediment trap bottles, 473 474 especially in the autumn months at LSB. These consisted mostly of copepods (e.g., Calanus 475 sp.), mysids (e.g., Boreomysis sp.), amphipods (e.g., Eusiridae) and chaetognaths (i.e., arrow 476 worms). Numbers of trapped swimmers were lowest during winter at both sites. In addition, several large sponge spicules were found in the bottles at HSB, but not at LSB. 477



479Figure 12: Sediment trap content from the two benthic landers. HSB = high-sponge-biomass lander, LSB = low-sponge-480biomass lander. A) mass flux in g m<sup>-2</sup>d<sup>-1</sup>, B) organic carbon flux in mmol C m<sup>-2</sup> d<sup>-1</sup>, C) molar C:N ratio of trapped material,<br/>A)  $\delta^{13}C$  of trapped material, E)  $\delta^{15}N$  of trapped material, F) total lipid flux in  $\mu g m^{-2} d^{-1}$ .

#### 3.4 $\delta^{13}$ C and $\delta^{15}$ N isotopic ratios of benthic fauna and trapped material 482

The massive sponge Geodia spp. Sampled at HSB showed a distinct isotopic signal compared 483 to the other benthic organisms, with a relatively enriched  $\delta^{13}$ C (-18.55 ± 0.17 ‰) and a low 484  $\delta^{15}$ N (8.24 ± 0.16 ‰; Figure 13). The gorgonian coral Primnoa resedue formis had a  $\delta^{13}$ C (-485  $21.19 \pm 0.59$  ‰) and a  $\delta^{15}$ N (10.54  $\pm 0.33$  ‰), values that indicated a lower trophic level than 486 the Decapoda sp. ( $\delta^{13}$ C: -20.48 ± 0.31 ‰, and  $\delta^{15}$ N: 11.97 ± 0.43 ‰) and the glass sponge 487 Asconema sp. ( $\delta^{13}$ C: -20.27 ± 0.36 ‰, and  $\delta^{15}$ N: 12.57 ± 0.31 ‰). The sponge Mycale sp., 488 sampled at LSB, had a high  $\delta^{15}$ N isotopic ratio (13.05 ± 0.41 ‰), and a  $\delta^{13}$ C ratio of -19.47 ± 489 0.06 ‰. Sediment trap samples had the lowest  $\delta^{15}$ N and  $\delta^{13}$ C isotopic ratios, with only small 490 differences between HSB and LSB (Figure 12 D & E; Figure 13). 491

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494 495 Figure 13: Carbon and nitrogen isotope biplots of megafauna and sediment trap samples. HSB = high-sponge-biomass, LSB = low-sponge-biomass.

#### Discussion 496 4

497 In this study, hydrodynamic- and environmental conditions and food availability were compared at two contrasting high- and low-sponge-biomass sites along the northern Labrador 498 499 shelf break. More specifically, the aim of this study was to compare differences between the 500 two sites in terms of (i) seawater properties and regional hydrography (section 4.1), (ii) bottom 501 currents and environmental conditions, including seasonal variations over the course of a year

502 (section 4.2), and (iii) organic matter supply and food sources for benthic macrofauna (sections

503 4.3, 4.4, and 4.5).

#### 504 4.1 Hydrography and bottom nutrients on the northern Labrador Shelf and Slope

The northern Labrador Shelf and Labrador Slope is known to be subject to strong tidal forcing 505 506 which causes vertical mixing, high bottom current speeds (Griffiths et al., 1981; Drinkwater 507 and Jones, 1987), and reduced stratification compared to the more northerly Baffin Island Shelf (Lazier 1982; Sutcliffe et al. 1983; Drinkwater and Harding 2001). The results of our drifter 508 509 analysis confirm that around the HSB area three currents converge: the Hudson Strait Outflow, the Baffin Intermediate Current, and the West Greenland Current (Figure 5A; Smith et al., 510 511 1937; Yashayaev, 2007; Straneo and Saucier, 2008; Curry et al., 2011, 2014). These three currents transport, respectively, Hudson Strait Outflow Water, Arctic Water and/or Baffin Bay 512 513 (intermediate) Water, and Irminger Water towards the northern Labrador Shelf and upper 514 slope. Our CTD transects show characteristics of these water masses, and are similar to earlier 515 cross-shelf transects (Petrie et al., 1988; Fissel and Lemon, 1991; Drinkwater and Harding, 2001). The warmer and saltier water observed at HSB compared to LSB is likely caused by 516 517 Irminger Water (Figure 5 B & C). This water mass is relatively warm and salty ( $\Theta \sim 4.5$  °C and  $S \sim 34.9$  psu) and follows the Labrador slope in cyclonic direction beneath the cold water of 518 519 the West Greenland Current and above the upper slope (Lazier et al., 2002). Our findings confirm previous work which showed that Irminger Water is gradually cooled while moving 520 521 southward by mixing with the Baffin Island Current (Cuny et al., 2002). However, our Argo float profiles, in combination with the drifter profiles, indicate that the area around HSB might 522 523 play an important role transforming Irminger Water. For example, the 350-450 m depth layer 524 in the HSB area regularly showed presence of Irminger Water (>4.5 °C), while Irminger Water 525 was only sporadically measured at LSB (Figure 5B). Irminger Water might therefore be cooled and freshened in the area around HSB due to convergence and consequently mixing with the 526 527 Hudson Outflow and Baffin Island Current. Benthic lander bottom temperatures were well 528 correlated between the two sites, with a time lag of five days at the LSB site. This time lag corresponds to an along slope velocity of  $\pm 0.3$  m s<sup>-1</sup>, which is close to the mean bottom current 529 speeds measured at HSB (0.25 m s<sup>-1</sup>) and on the Labrador Slope (0.11 - 0.23 m s<sup>-1</sup>; Lazier and 530 531 Wright, 1993). This supports earlier findings on the Labrador Shelf that found a connection between the Hudson Strait outflow strength and the southern Labrador Shelf water salinity 532 533 (Sutcliffe et al., 1983; Myers et al., 1990).

534 Both the LSB and HSB lander sites show higher nutrient concentrations in the bottom water 535 compared with the other shelf/deep CTD stations, and this difference was more pronounced at 536 the HSB site. These observations are thought to be related to the sources of the bottom water 537 and circulation. Thus, intermediate water flows from Baffin Bay via the Davis Strait southward 538 along the continental slope (Curry et al., 2014). This water mass, referred to as Baffin Bay Water (BBW), contains higher nutrient concentrations (e.g.,  $41.6 \pm 25.5 \mu$ M Si(OH)<sub>4</sub>,  $18.5 \pm$ 539 2.6 µM NO<sub>3</sub>; Sherwood et al., 2021) due to *in situ* remineralization of deep water circulating 540 541 in the Baffin Bay basin (Jones et al., 1984; Tremblay et al., 2002; Lehmann et al., 2019). BBW 542 mixes with water masses on the Labrador Shelf and Slope and Hudson Strait outflow water 543 while flowing southward along the Labrador Slope, resulting in lower nutrient concentrations 544 at the LSB compared to the HSB (Figure S4). The absence of high nutrient concentrations at the shelf/deep CTD station at both sites supports this interpretation. The elevated nutrient 545 546 concentrations could be beneficial for benthic organisms, for example, deep-sea sponges, 547 which require silicic acid for spicule formation and skeletal growth (Whitney et al., 2005; 548 Maldonado et al., 2011, 2020b; López-Acosta et al., 2016). Published kinetic uptake curves, 549 describing silicic acid uptake rate versus concentration, suggests the concentration at the HSB 550 lander (13.6 µM) compared to LSB shelf (9.3 µM) would imply a higher silicic acid uptake rates of glass sponges at the HSB site of 39% for Axinella spp. and 40% for V. pourtalesii 551 552 (Maldonado et al., 2011, 2020b). Furthermore, elevated silicic acid concentrations on the 553 spatial scale of kilometres are thought to allow the persistence of sponge grounds and build-up 554 of (glass) sponge biomass over long timescales (Whitney et al., 2005; Maldonado et al., 2020a). 555 While it has been suggested that biogenic silica efflux from the sediments could cause higher bottom water concentrations of silicic acid (Maldonado et al., 2020a), this is unlikely for our 556 557 study sites. Namely, such an efflux from the sediments would be quickly advected away by the high bottom tidal currents, while nutrient concentrations were elevated up to 100 meters above 558 559 the bottom (Figure 4 B & C). Overall, our study shows that bottom water between the LSB and 560 HSB sites are connected, with higher nutrient availability at the HSB station, linked to large-561 scale circulation patterns.

#### 562 4.2 Bottom hydrodynamics and environmental conditions over a year

This study provides the first concurrent long-term measurements of hydrodynamic- and 563 564 environmental conditions at a high- and low-sponge-biomass site. Our measurements show 565 high bottom currents at both sites with distinct differences in tidal dynamics. Bottom current direction and tidal ellipses (northeast-southwest for HSB, north-south for LSB) align well with 566 bathymetry and the OTIS modelled barotropic tide (Figure S11). The tidal amplitude is around 567 five times higher at HSB than at LSB. This outcome is contrary to White (2003) who measured 568 569 high current speeds in areas where no sponges were recorded, and vice versa, at the Porcupine 570 Sea Bight. Although bottom currents are higher at HSB than at LSB (Table 1), the bottom

571 currents at LSB are still comparable with current speeds found at other sponge grounds on the 572 Scotian Shelf (mean:  $0.12 \text{ m s}^{-1}$ ; Hanz et al., 2021a) and on the Arctic mid-Atlantic ridge (mean: 573 0.14 m s<sup>-1</sup>; Hanz et al., 2021b). Hence, the high bottom tidal currents seem to be a more 574 prominent difference in the environmental conditions at HSB compared to LSB.

Bottom water temperatures at both sites (3.5 - 4 °C) are within range of values reported for 575 boreal deep-sea sponge grounds previously (<0-8 °C; Kutti et al., 2013; Howell et al., 2016; 576 577 Strand et al., 2017; Hanz et al., 2021b, 2021a). Bottom temperatures measured by the benthic landers followed a similar seasonal pattern as the Argo float temperature profiles. Temperature 578 increased gradually from summer until December, which is measured previously on the 579 580 Labrador upper slope and attributed to Irminger Water (Cuny et al., 2002). From a biological point of view, fluctuations in temperature over a year were in general low (<1 °C) and unlikely 581 582 to affect the sponge distribution in the study area. The temperature fluctuations in bottom water do however reveal clear differences between the two sites in terms of hydrography. Tidal 583 584 currents have a distinct effect on bottom temperature at both sites, and this effect depends on the season. For example, in the first week of September at HSB, the bottom temperature 585 586 decreased after water moves in a northwest direction and increased after the current changed to a south-easterly direction. As the lander was placed ~500 m from the shelf break (Figure S2 587 588 C&D), and bottom water could be transported ~5 km in the north-easterly direction in one 589 semidiurnal tidal cycle (Figure 9A), this means that colder bottom water is transported on to 590 the Labrador Shelf from beyond the shelf break to the HSB lander site. Furthermore, the tidal currents in the south-easterly direction bring warmer bottom water from the Labrador Shelf to 591 592 the HSB lander (Figure 9A). Colder bottom water temperatures were also observed when water 593 moved upslope at LSB (Figure 9C). Therefore, although higher variability in bottom water 594 temperature has been attributed to the presence of internal waves at other sponge grounds (Roberts et al., 2018; Davison et al., 2019), we attribute the variability in our study area to 595 596 tidal-induced cross-slope transport of bottom water. Nevertheless, high downward velocities 597 (>0.2 m s<sup>-1</sup>), which occurred while water was moving in a south-easterly direction sometimes 598 caused a drop in bottom temperature at HSB (Figure 9A), which suggests that colder water 599 from shallower depths mixed with bottom water.

500 Strong tidally-induced bottom currents can benefit the benthic community at the HSB site in 501 various ways. First, passive suspension feeders as the gorgonian *P. resedaeformis* benefit from 502 high horizontal currents through an increased particulate organic matter flux (Shimeta and 503 Jumars, 1991) and sponges (specifically glass sponges) could benefit from an increased water 504 flow rate through their body plan (Vogel, 1977), thereby increasing food availability. Second, 505 resuspension caused by high bottom current speeds could enhance organic matter availability in the benthic boundary layer and prevent smothering of sponges by sedimentation (Roberts et
al., 2018). Here, high along-slope bottom currents at both sites were associated with increased
ABS and turbidity, indicative of resuspension (Figure 9). However, the beneficial effect of
resuspension for sponge biomass is not yet fully understood, as reoccurring strong turbidity
flows (at LSB) could also prevent high sponge biomass from developing by smothering young
sponges when particles settle out (Klitgaard and Tendal, 2004).

The substrate at HSB consisted mostly of pebbles, cobbles, and boulders (Dinn et al., 2020) 612 and a qualitative assessment of the sediment type at LSB suggested the dominance of muddy 613 soft sediment (Coté et al., 2019; J. Vad, pers. com.). As higher bottom currents would increase 614 bed shear stress and thereby enhance resuspension (Lesht, 1979; Jones et al., 1998), we argue 615 616 that fine material is resuspended at HSB before its accumulation on the seafloor. This increases 617 availability of organic matter to benthic suspension feeders in the benthic boundary layer and prevent smothering. Resuspension has also been linked to high sponge biomass, although the 618 619 mechanisms behind this link are still unclear (Davison et al., 2019). Third, the interaction of 620 high bottom currents with rough topography causes turbulence and mixing of bottom waters 621 (Witte et al., 1997, 97; Levs et al., 2011; Culwick et al., 2020). As the substrate is likely rougher and bottom currents are higher at HSB than at LSB, the bottom water probably experiences 622 623 more intense mixing and turbulence at HSB. Finally, periodic supply of fresh phytoplankton 624 derived material during the spring bloom (Figure S8, Figure S9) increases the food availability 625 of passive suspension feeders living on the sponge grounds. In short, the stronger tidal currents at HSB enhance bottom water mixing which replenishes oxygen, dissolved organic matter, 626 POM, and (inorganic) nutrients in the benthic boundary layer, and thereby increases food 627 628 supply to benthic fauna (Davison et al., 2019; Hanz et al., 2021b, 2021a).

#### 629 4.3 Primary production and benthic-pelagic coupling

630 The Hudson Strait outflow water is known to increase nutrient concentrations in the surface waters on the northern Labrador Shelf (Kollmeyer et al., 1967; Sutcliffe et al., 1983; 631 Drinkwater and Harding, 2001). A thermal front, associated with the offshore branch of the 632 Labrador Current, is located along the 1,000 m isobath of the Labrador slope/shelf (Cyr and 633 634 Larouche, 2015). The increased nutrient supply support high primary productivity in an area extending from the Hudson Strait to the southern Labrador Shelf, bounded by the thermal front 635 636 associated with the 1,000 m isobath (Frajka-Williams et al., 2009; Frajka-Williams and Rhines, 2010; Cyr and Larouche, 2015). Our CTD profiles show elevated chl-a concentrations in the 637 CIL (~150 m depth), as was observed by Frajka-Williams et al., (2009). The fact that primary 638 639 production rates are comparable above the two lander station sites (Frajka-Williams and

Rhines, 2010), suggests that differences in primary production alone are insufficient to explain
the differences sponge biomass between regions. Furthermore, studies elsewhere in the
Canadian Arctic have shown that benthic biomass is explained not only by surface productivity
but also by local hydrodynamics and benthic-pelagic coupling (Thomson, 1982; Grebmeier
and Barry, 1991; Roy et al., 2014).

645 The lander fluorescence observations showed the arrival of relatively fresh phytodetritus at the 646 seafloor three months before the start of the phytoplankton bloom (Fuentes-Yaco et al., 2007; Frajka-Williams and Rhines, 2010). We suggest that this results from phytoplankton growth 647 that had already started in early March in the Hudson Strait outflow (Harrison et al., 2013). At 648 649 this time, the water column was still relatively cold and poorly stratified, allowing for relatively high export, which resulted in fluorescent material transported towards the seafloor at each 650 651 semidiurnal tidal cycle (Figure S9B). Sea ice retreat in mid-April relaxed light limitation and 652 further stimulated primary production (Carmack et al., 2004), explaining the fluorescent 653 material peaks at both landers at this time. In summer, there appears to be a decoupling between high surface primary production (Frajka-Williams and Rhines, 2010) and low chl-a 654 655 concentration on the seafloor (this study), likely due to enhanced stratification and intense zooplankton grazing (Rivkin et al., 1996; Turner, 2015). Strong tidal mixing, including a strong 656 657 neap-spring tidal cycle, at HSB could inhibit water column stratification for a longer period 658 than at LSB, thereby extending the period of fluorescent material deposition at the seafloor 659 (Sharples et al., 2006; Sharples, 2008; this study). Our findings suggest strong benthic-pelagic 660 coupling started weeks before the peak of the phytoplankton bloom, supplying fresh fluorescent material to the seafloor in spring for a period of weeks to months. Since the timing of 661 662 phytoplankton bloom for high-latitude seas is shifting to earlier in the year due to rising 663 temperatures and earlier sea-ice retreat (Edwards and Richardson, 2004; Wu et al., 2007; Hunter-Cevera et al., 2016), and since deep-sea sessile organisms, such as cold-water corals 664 665 and deep-sea sponges demonstrate seasonality in their phenology (Levs and Lauzon, 1998; 666 Maldonado, 2011; Maier et al., 2020a), the early arrival of phytoplankton-derived material could have consequences for their overall fitness and survival. Nevertheless, the effect of a 667 668 shift in spring bloom timing for benthic suspension feeders, including deep-sea sponges, 669 remains unknown.

670 Recent ABS measurements reveal a layer of increased 300 kHz backscatter along the northern 671 Labrador Shelf, indictive of high abundance of micronekton and macrozooplankton 672 (Chawarski et al., 2022). Earlier studies showed a high zooplankton biomass on the 673 Newfoundland Shelf from July onwards (Head et al., 2003, 2013). In our traps the highest flux 674 of unsaturated alcohols, a biomarker for zooplankton (specifically copepods; Dalsgaard et al., 675 2003), and the highest numbers of swimmers were in summer and autumn. During the spring bloom, trapped material at LSB had the highest relative amount of unsaturated alcohols while 676 677 at HSB the level of PUFAs, markers for phytoplankton derived-material, was highest (Dalsgaard et al., 2003). Furthermore, our observations suggest that the number of trapped 678 679 swimmers was higher at LSB than at HSB. These results are consistent with the hypothesis that zooplankton biomass is high over the northern Labrador Shelf (Saglek Bank) and that 680 681 zooplankton is transported by the southerly current along the Labrador Shelf together with the high phytoplankton biomass plume (Sutcliffe et al., 1983; Drinkwater and Harding, 2001). 682 683 Overall, there was a larger fraction of zooplankton marker lipids in trapped material at LSB, which implies that zooplankton play a more important role in benthic-pelagic coupling at LSB 684 685 than at HSB.

686 4.4 Organic matter cycling at the seafloor

Organic matter deposition was higher at HSB than at LSB. Overall, deposition was highest 687 during the winter months and consisted of more degraded material than during summer, 688 indicated by high C:N ratios, high  $\delta^{15}$ N, and low fluorescence. This increased deposition in 689 winter is likely resuspended material, when bottom current speeds were higher. The C:N ratio 690 691 of deposited matter was higher at LSB (~13) compared to HSB (~8), indicating the material 692 was more degraded at LSB. Hanz et al. (2021a, 2021b) also found higher mass and carbon fluxes during winter months and low carbon fluxes when the spring/summer phytoplankton 693 694 bloom arrived. They attributed this to the presence of more degraded and resuspended material 695 in winter. Data concerning mass fluxes from sponge grounds remain scarce, but the fluxes measured here (HSB 2.46  $\pm$  1.76 g m<sup>-2</sup> day<sup>-1</sup>, LSB: 1.43  $\pm$  0.93 g m<sup>-2</sup> day<sup>-1</sup>) were comparable 696 to those of a *Vazella pourtalesii* sponge ground on the Scotian Shelf  $(3.17 \pm 3.42 \text{ g m}^{-2} \text{ dav}^{-1};$ 697 Hanz et al., 2021a) but substantially higher than those of a sponge ground on the Arctic mid-698 Atlantic ridge  $(0.03 - 0.30 \text{ g m}^{-2} \text{ day}^{-1}$ ; Hanz et al., 2021b). Overall, our data suggest organic 699 700 matter deposition fluxes are higher at HSB compared to LSB, and that the organic matter is of higher quality. The organic carbon fluxes (HSB:  $3.07 \pm 1.91 \text{ mmol C} \text{ m}^{-2} \text{ d}^{-1}$ ; LSB:  $1.91 \pm 0.71$ 701 mmol C m<sup>-2</sup> d<sup>-1</sup>) reported in our study are considerably lower than those of a more shallow (150 702 - 250 m depth) V. pourtalesii sponge ground on the Scotian Shelf (8.3 mmol C m<sup>-2</sup> d<sup>-1</sup>; Hanz 703 704 et al., 2021a), but high compared to an Arctic mid-Atlantic ridge sponge ground (peak of 1.6 mmol C m<sup>-2</sup> d<sup>-1</sup>; Hanz et al., 2021b). The higher organic matter deposition rate and relative 705 706 fresher material at HSB compared to LSB are likely related to its shallower position on the shelf and the more dynamic water column. 707

708 No estimates of organic carbon utilization by the sponge grounds on the Northern Labrador Shelf were available for comparison with these sediment trap data at the time of writing. Here, 709 710 we estimate the organic matter requirements of the sponge grounds from published respiration 711 rates and biomass estimates obtained from bottom trawls using a depth stratified random 712 sampling design and images taken with a Remotely Operated Vehicle (ROV). Bottom-trawl estimates gave a biomass of 35 g WW sponge m<sup>-2</sup> at HSB and 0.01 g WW sponge m<sup>-2</sup> at LSB 713 714 (Lirette and Kenchington, pers. com.). Assuming a sponge respiration rate of 0.010 mmol O<sub>2</sub> g<sup>-1</sup> WW d<sup>-1</sup> (measured at 6 - 9 °C; Kutti et al., 2013; Leys et al., 2018; Bart et al., 2021), this 715 corresponds to a benthic respiration rate of 0.35 mmol O<sub>2</sub> m<sup>-2</sup> d<sup>-1</sup> at HSB and 0.0001 mmol O<sub>2</sub> 716 m<sup>-2</sup> d<sup>-1</sup> at LSB. Image analysis from ROV transects suggested higher biomass levels: 500 g 717 sponge WW m<sup>-2</sup> at HSB and 50 g sponge WW m<sup>-2</sup> at LSB (Wolff et al., 2020), equivalent to 718 benthic respiration rates of 5 mmol  $O_2 m^{-2} d^{-1}$  and 0.5 mmol  $O_2 m^{-2} d^{-1}$  for HSB and LSB, 719 respectively. The large difference in sponge biomass estimates between the trawl and ROV 720 721 methods is surprising, and we cautiously attribute this to: 1) the different spatial scales over 722 which both methods work combined with spatial heterogeneity within the area, 2) under-723 sampling of sponges by bottom trawling (Wassenberg et al., 2002), and 3) potential bias in 724 ROV imaging, as the trajectory of ROV transects is usually not randomized and potentially 725 biased to higher sponge cover areas. The ROV-based biomass and respiration are more in line, albeit on the lower end, with earlier observed sponge community benthic respiration values in 726 Norway ( $15 - 45 \text{ mmol } O_2 \text{ m}^{-2} \text{ d}^{-1}$ ; Kutti et al., 2013; Cathalot et al., 2015). As bottom trawling 727 data are the only sponge biomass estimates available on a shelf-wide scale, we consider the 728 729 trawl-based respiration rates to be the most representative for sponge respiration on the northern Labrador Shelf region, with the ROV-based respiration rates giving upper bounds. 730

#### 731 4.5 Food sources of benthic macrofauna

Although the sample size was limited, the stable isotope data revealed interesting patterns of 732 733 organic matter utilization by the benthic community. The gorgonian coral *P. resedueformis* is 734 found one trophic level (Fry, 2006) above the sediment trap material and therefore likely feeds 735 on sinking organic matter, confirming previous observations (Sherwood et al., 2005, 2008). Sponges can generally be classified into two groups based on their associated microbial fauna, 736 737 those with high microbial abundance (HMA) or those with low microbial abundance (LMA; Vacelet and Donadey, 1977). Geodia spp. can occur in high abundance and biomass on sponge 738 739 grounds (Kutti et al., 2013). These sponges are considered HMA (Radax et al., 2012) and feed mostly on dissolved organic matter with additional particulate sources such as bacterioplankton 740 (Bart et al., 2021). Many hexactinellidae that can form sponge grounds, for instance Vazella 741 742 pourtalessii and Aphrocallistes vastus, are considered LMA sponges and feed mostly on

bacterioplankton (Kahn et al., 2015). The high  $\delta^{15}$ N isotopic ratios for the sponges Asconema 743 spp.  $(12.6 \pm 0.3 \text{ \% } \delta^{15}\text{N})$  and *Mycale* spp.  $(13.1 \pm 0.4 \text{ \% } \delta^{15}\text{N})$ , has been observed previously 744 745 for LMA sponges (Iken et al., 2001; Polunin;, 2001; Kahn et al., 2018). Deep-sea LMA sponges typically have elevated  $\delta^{15}$ N values in the benthic food web (Kahn et al., 2018), a phenomenon 746 747 that is still poorly understood. Possible explanations could be selective feeding on <sup>15</sup>N enriched 748 bacteria (Wilkinson et al., 1984), feeding on resuspended benthic bacteria (Kahn et al., 2018), 749 or nitrogen (re)cycling within the sponge holobiont (Rooks et al., 2020; Hanz et al., 2022). Interestingly, the HMA massive sponge *Geodia* sp. has distinct  $\delta^{13}$ C and  $\delta^{15}$ N values, 750 751 indicating different feeding or metabolic strategies. Recent research on Geodia baretti has 752 indeed demonstrated that these sponges rely for a large part on DOM for their metabolic requirements (Bart et al., 2021; de Kluijver et al., 2021). In this study, *Geodia* spp.  $(8.2 \pm 0.2)$ 753  $\% \delta^{15}$ N) was one trophic level higher than oceanic DOM  $\delta^{15}$ N (~5 %; Benner et al., 2005; 754 Sigman et al., 2009) and  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup> (~5%; Sigman et al., 2009; Sherwood et al., 2021), limiting 755 our ability to distinguish between DOM and NO<sub>3</sub><sup>-</sup> (by i.e., denitrification; Hoffmann et al., 756 2009) as potential nitrogen sources. The  $\delta^{13}$ C value of *Geodia* spp. (-18.4 ± 0.17 ‰  $\delta^{13}$ C) is 757  $\pm 3.5\%$  higher than bottom water  $\delta^{13}$ C-DOC values on the Labrador Shelf (Barber et al., 2017), 758 i.e. more than four times higher than the expected 0.8%  $\delta^{13}$ C step per trophic level (Vander 759 Zanden and Rasmussen, 2001). Alternatively, Geodia spp. could capitalize on DIC via their 760 761 symbionts (de Kluijver et al., 2021), as recently observed in Arctic Geodia spp. assemblages 762 (Morganti et al., 2022) and other deep-sea sponges (van Duyl et al., 2020). Even limited chemoautotrophic assimilation of high  $\delta^{13}C\text{-DIC}~(\sim\!\!0~\%~\delta^{13}C)$  could explain the high  $\delta^{13}C$ 763 764 values of Geodia spp. These results indicate that passive suspension feeders benefit from high 765 tidal currents through an increased particulate organic matter flux (Shimeta and Jumars, 1991), 766 whereas sponges likely benefit from replenishment of nutrients, oxygen, and dissolved organic 767 matter (Schläppy et al., 2010).

#### 768 5 Conclusion

769 This study investigated the hydrodynamic- and environmental conditions at two contrasting 770 high- and low-biomass sponge grounds on the northern Labrador Shelf. The high-biomass sponge ground is in an area where three currents converge and there are strong tidal currents 771 772 throughout the year. This is also reflected in tidal periodicity of environmental conditions. The 773 high tidal currents increase the flux of available food resources to the benthic community. High 774 nutrient concentrations were found at the high-sponge-biomass site, which were associated 775 with the presence of Baffin Bay water and therefore related to large scale circulation patterns. 776 The Northern Labrador Shelf exhibits tight benthic-pelagic coupling during spring, and high

primary production alone seems to be a poor predictor for sponge biomass in this area. Intense
vertical mixing at the high-sponge-biomass site extends the period of benthic-pelagic coupling
by several months. High currents benefit the benthic community by increasing food availability
and replenishing nutrients, oxygen, and dissolved organic matter in bottom waters.

#### 781 6 Funding statement

This research was supported by the European Union's Horizon 2020 Research and Innovation 782 Programme under grant agreement nos. 678760 (ATLAS) and 818123 (iAtlantic). This output 783 784 reflects only the authors' view, and the European Union cannot be held responsible for any use that may be made of the information contained therein. Department of Fisheries and Oceans 785 786 contributions were funded through the departmental International Governance Strategy programme awarded to EK. DvO was supported by the Innovational Research Incentives 787 788 Scheme of the Netherlands Organisation for Scientific Research (NWO), respectively, under grant agreement 864.13.007. EdF was partly supported by ArcticNet Network of Centres of 789 790 Excellence, Glacier troughs as biodiversity and abundance hotspots in Arctic and subarctic 791 regions project, ArcticNet Phase V (Geoffroy et al.). The data presented herein were collected 792 by the Canadian research icebreaker CCGS Amundsen and made available by the Amundsen 793 Science program, which was supported by the Canada Foundation for Innovation and Natural 794 Sciences and Engineering Research Council of Canada. The views expressed in this publication do not necessarily represent the views of Amundsen Science or that of its partners. Ship-time 795 796 on the CCGS Amundsen was also funded by an NSERC ship-time grant (Edinger et al., grant nr.: RGPST-515528-2018), ArcticNet Network of Centers of Excellence Canada, and the 797 798 Department of Fisheries and Oceans Canada (DFO; Coté et al.). The funders had no role in 799 study design, data collection, and analysis, decision to publish, or preparation of the 800 manuscript.

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#### 802 7 Author statement

EDF: sample analysis, data analysis, and writing; IY: data collection, data analysis, and writing.
CM: conceptualization, data analysis and writing; JV: data collection and data analysis; FM:
conceptualization, sample analysis and data analysis; GD: conceptualization, data analysis;
EK, EH, IY, SWR, MR: conceptualization and site selection; SWR, MR, EK, BM, GT:
contribution and preparation of benthic landers; GW: conceptualization, sample analysis, data

analysis, and writing; SB: data collection and sample analysis; DvO: conceptualization, data
analysis, writing. All authors contributed to the article and approved the submitted version.

#### 810 8 Acknowledgements

811 We would like to thank the skilful crew and technicians on board CCGS Amundsen for their 812 support during the fieldwork. Specifically, we thank Dr. Paul Snelgrove (Memorial University of Newfoundland), Dr. David Cote (DFO) and Shawn Meredyk (Amundsen Science) for their 813 814 assistance in facilitating our field programme. Cam Lirette (DFO) assisted in preparing various data layers to assist in site selection. We would also like to thank Jan Peene for nutrient 815 816 analysis, Peter van Breugel and Jurian Brasser for help in measuring 817 macrofauna/POM/sediment trap stable isotopes, and Pascal Guillot for quality assurance of the 818 CTD profiles. Finally, we thank Kevin MacIsaac and Marc Ringuette for their help in 819 identifying the sediment trap swimmers.

#### 820 9 Data availability

- 821 Raw data and (some) processing scripts are available at
- 822 <u>https://doi.org/10.5281/zenodo.10571403</u>.

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### 1297 11 Supplementary material

#### 1298 11.1 Tables

station	instrument	date/period	latitude	longitude	depth
HSB_bl	benthic_lander	27-7-2018 to 2-7-2019	60.47	-61.29	410
LSB_bl	benthic_lander	27-7-2018 to 1-7-2019	59.38	-60.28	558
HSB_ctd1	CTD	2018-08-03 07:37:08	60.47	-59.26	2428
HSB_ctd2	CTD	2018-08-02 17:21:58	60.47	-60.38	1877
HSB_ctd3	CTD	2018-07-30 15:27:05	60.47	-61.30	391
HSB_ctd4	CTD	2018-07-30 07:31:07	60.46	-62.12	359
HSB_ctd5	CTD	2018-07-27 19:41:58	60.40	-62.90	289
LSB_ctd1	CTD	2018-07-29 04:30:19	59.53	-58.64	2563
LSB_ctd2	CTD	2018-07-28 23:25:52	59.48	-59.45	1938
LSB_ctd3	CTD	2018-07-28 09:52:11	59.38	-60.27	608
LSB_ctd4	CTD	2018-07-28 06:12:07	59.31	-61.02	192
LSB_ctd5	CTD	2018-07-28 03:10:24	59.22	-61.83	138

1299 Table S1: Overview of lander deployment and CTD casts

#### 1300

1301<br/>1302Table S2: Overview of rock dredge transects. HSB = high-sponge-biomass site, LSB = low-sponge-biomass site, (Coté et al.,<br/>2019).

Station Name	Start Lat	Start Long	End Lat	End Long	Logged bottom depth	Time at bottom	Length of cable	Max vessel speed	Comments
					(m)	(min)	out (m)	(knots)	
LSB_rd	59.38	-60.27	59.37	-60.29	552	10	1500	1	NA
HSB_rd	60.47	-61.28	60.48	-61.30	404	20	507	2	Small
									catch



Figure S1: slope direction or aspect estimation for HSB and LSB. A) map of study area with estimated slope aspects of 60° and 90° angle for HSB and LSB, respectively. Contour lines at 200, 400, and 1000 metre is shown. B) expanded detail on HSB shows angle of 60° is a good estimate. Contour lines at 400, 425, 475, 500 are shown. C) expanded detail on LSB site shows angle of 105° is better estimate. Contour lines at 450, 475, 500, 525, 550, 575, 600 metre depth are shown. Note the different colour scales for depth. Locations of lander is indicated by coloured dots, with HSB = blue, and LSB = brown/red.



1313<br/>1314<br/>1314Figure S2: slope angle and direction in north- to-south and west-to-east direction close to the high-sponge-biomass (HSB)<br/>and low-sponge-biomass (LSB) landers. The left column shows the depth around the landers (A, C, E, G), and the transect<br/>line for which the slope is calculated and plotted. The right column shows the slope along the black line from either north to<br/>south (B, F), or west to east (D, H), blue colours represent downhill angle and brown colours an uphill angle, z = depth in<br/>meters, d = distance from start transect (north or west) in meters. Landers are indicated by black dots in the left column, and<br/>coloured triangles in the right column. Note the different colour scales for plots in the left column and different y-axis scale<br/>for the plots in the right column.

Argofloat profile locations



1321<br/>1322Figure S3: Locations of Argo float profiles used for assessing the regional oceanography. Coloured squares indicate Argo<br/>float profiles, and black trigangles/dots the location of CTD profiles/benthic lander location.



1324Figure S4: CTD profiles with temperature (A), salinity (B), density (C), chlorophyll-a (D), Oxygen (E), Buoyancy frequency1325(F). LSB = Low-sponge-biomass transect, HSB = High-sponge-biomass transect. Buoyancy frequency is smoothed over 151326m for visibility, and the plot only shows top 650 m of the water column, as deeper waters have values close to zero.



1328Figure S5: nutrient profiles for the two transects over the complete depth. HSB = high-sponge-biomass, LSB = low-sponge-biomass.1329biomass.



1331Figure S6: Particulate organic carbon (POC) profiles for the two transects. HSB = high-sponge-biomass lander, LSB =1332low-sponge-biomass lander.





1334<br/>1335Figure S7: A) seasonal salinity signal, from Argo float data, of the water layer in which HSB lander is located. B) seasonal<br/>salinity signal of the water layer in which LSB is located.











Figure S9: progressive vector plots with chlorophyll-a as colour variable from 19 to 24 April 2019. With A) the highsponge-biomass (HSB) lander and b) the low-sponge-biomass (LSB) lander. Dotted lines represent the along slope direction at the respective sites. Note colour is in log-scale.



1344Figure S10: Spring Chlorophyll-a (A), bottom current speed (B), ice cover (C), during the spring bloom period (1 April-11345May, 2019), and ice cover for the whole deployment length (D).



1347<br/>1348Figure S11: Sediment trap lipid fluxes. A) Total lipid flux, B) unsaturated alcohol:total lipids ratio, C) poly-unsaturated fatty<br/>acid:total lipids ratio, D) sterol:total lipids ratio.



1351<br/>1352Figure S12: M2 tidal current ellipses in the Davis Strait case study area (OTIS inverse tidal model, hourly data, July/August<br/>2018).