



Climate warming enhances polar cod recruitment, at least transiently



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ABSTRACT

Polar cod (*Boreogadus saida*) is the dominant forage fish in Arctic seas and the main prey of the ringed seal (*Pusa hispida*), the beluga (*Delphinapterus leucas*) and several seabird species. Changes in the abundance of polar cod will have cascading effects on arctic marine ecosystems. We tested the hypothesis that an earlier sea ice breakup and warmer sea surface temperatures (SST) in spring-summer result in the higher recruitment of juvenile polar cod in late summer. The density (number m⁻²) and biomass (mg m⁻²) of age-0 polar cod in August and September, estimated by hydroacoustics over 9 years in 9 areas of the Canadian Arctic, were negatively correlated to ice breakup week and positively correlated to SST. The timing of the ice breakup was the main determinant of recruitment, with mean juvenile biomass in September up to 11 times greater for early breakup (late May) than for late breakup (early September). Early ice breakup in spring increased juvenile biomass in August and September by allowing the survival of larvae hatched in winter and spring. Since 1979, ice breakup has occurred earlier by as much as 9.3 days per decade in some areas. We thus forecast a transient increase in polar cod biomass over the first part of the present century. Thereafter, the relaxation of extreme climatic conditions in Arctic seas should harbingers the replacement of the hyper-specialized polar cod by subarctic and boreal forage fish.

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1. Introduction

Sea surface temperature (SST) has increased in most Arctic seas over the last century, with warming intensifying since 1995 (Steele et al., 2008). Arctic sea ice extent (Stroeve et al., 2012) and volume (Laxon et al., 2013) have declined during all seasons, and ice breakup has occurred progressively earlier (Stroeve et al., 2012). Among the biological consequences of climate warming, an increase in primary production (Tremblay and Gagnon, 2009), a shift towards smaller phytoplankton and zooplankton species (Hop et al., 2006; Li et al., 2009), the development of an Atlantic-type autumn bloom (Ardyna et al., 2014), perturbations of the phenology of key species (Ji et al., 2013; Søreide et al., 2010), and the poleward expansion of boreal species (e.g. Falardeau et al., in press;

Fossheim et al., 2015; Lenoir et al., 2011; Perry et al., 2005; Renaud et al., 2012) already impact arctic marine ecosystems.

The Arctic pelagic food web is relatively simple and dominated by a few species (Welch et al., 1992). The intermediate trophic level between zooplankton and top predators is occupied almost exclusively by the small polar cod *Boreogadus saida*, a fish with large biomasses and high trophic connectivity (Fortier et al., 2015; Kortsch et al., 2015). Changes in its abundance in response to the ongoing warming of Arctic seas could have profound effects on arctic marine ecosystems and the services they provide to society, especially Inuit communities (Fortier et al., 2015; Tynan and DeMaster, 1997).

In teleosts, interannual fluctuations in the survival of the eggs, larvae, and early juveniles determine the number of fish that recruit to the adult population in a given year (Browman, 2014; Houde, 2008). Hence, any factor influencing early survival may affect adult population size. The hatching season of polar cod extends from January to July, and higher temperatures have been linked to enhanced growth and survival of the larval stages (Bouchard and Fortier, 2008, 2011; Fortier et al., 2006). A large pre-winter size has been suggested as an important survival advantage (Bouchard and Fortier, 2008, 2011). Summer hatchers

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experience higher temperatures and more abundant food that result in faster growth. However, winter hatchers benefit from a longer growing season and reach pre-winter weights 100–150 times those of summer hatchers (Bouchard and Fortier, 2011). An early sea ice breakup should favour the survival of early hatchers and the recruitment of large juvenile polar cod in late summer. The density and biomass of juveniles (age-0) are predictors of fish recruitment (e.g. Laurel et al., 2016a; Sogard, 1997). In polar cod, these variables can be assessed using hydroacoustics validated by ichthyoplankton net collections (Geoffroy et al., 2016).

In this study, we first test the hypothesis that an early sea ice breakup and the associated high spring-summer SST promote the recruitment of juvenile polar cod. We then explore whether higher recruitment is primarily caused by higher survival of early-hatched larvae or by faster growth over the summer. Hydroacoustic estimates of the density and biomass of age-0 polar cod from August to October validated by net sampling were correlated to satellite-based estimates of sea ice breakup dates and mean SST in spring-summer. Juvenile density and biomass in late summer-early fall were calculated in different years over a 9-year period for 9 sea ice areas of the Canadian Arctic as defined by the Canadian Ice Service. The hatch date of age-0 polar cod collected by nets was determined by otolith analyses. Based on trends in ice breakup date since 1979, we project the pre-winter size and biomass of age-0 polar cod in August 2050 and 2100.

2. Materials and methods

2.1. Study area

From 2005 to 2015, the research icebreaker CCGS *Amundsen* and the trawler F/V *Frosti* conducted acoustic-trawl surveys in Canadian Arctic seas between early August and late October (Table 1, Fig. 1). Conductivity-Temperature-Depth (CTD) profiles from the *Amundsen* (SBE-911 plus[®]) and the *Frosti* (SBE-25 and SBE-19 plusV2[®]) were used to determine sound speed in water c_w (Mackenzie, 1981) and the frequency-dependent coefficient of sound absorption α_a (Francois and Garrison, 1982) for acoustic calculations.

2.2. Remote sensing data

The annual date of the ice breakup since 1979 was determined for 9 areas defined by the Canadian Ice Service (www.ec.gc.ca/glaces-ice). These included M'Clintock Channel, Peel Sound, Larsen-Victoria Sounds and Coronation-Maud in the Kitikmeot; West Baffin Bay and North-West Baffin Bay in Baffin Bay; and Mackenzie Shelf, Amundsen Gulf and Amundsen Gulf Mouth in the Beaufort Sea (Fig. 1). For each area and year, the week during which ice concentration fell below 50% was used as the date of

the ice breakup (Scott and Marshall, 2010). In M'Clintock Channel, the ice cover remained >50% through the summer in 21 years. In these cases, week 41 (latest week of ice breakup in this area + 1) was used. For each area and year, mean SST from 1 May to 31 July (corresponding to the main growth season of larval polar cod) was calculated using Level 3 daytime 11 μm wave band Aqua MODIS remote sensing data at a 4 km² resolution (<https://oceancolor.gsfc.nasa.gov/cgi/13>).

2.3. Ichthyoplankton sampling

Ichthyoplankton nets were deployed to identify the juvenile fish assemblage and validate the acoustic signals from the epipelagic layer (Fig. 1). A double square net bearing two square-conical nets (1-m² aperture, 500- μm and 750- μm meshes) and a rectangular midwater trawl (8-m² aperture, 1600- μm mesh) were deployed from the *Amundsen*. A bongo net (0.25-m² aperture, 500- μm mesh) and an Enzenhofer and Hume midwater trawl (9-m² aperture; 0.63-cm mesh) were used on the *Frosti*. Ichthyoplankton nets were cast from the surface to depths varying from 10 to ≤ 100 m, corresponding to the epipelagic layer where age-0 polar cod typically make up >90% of the fish assemblage (Geoffroy et al., 2016). The standard length (SL) of up to 50 gadids (i.e. possibly both polar and ice cods – see below) per station was measured fresh before preservation in 95% ethanol. For each year, a monthly mean SL was calculated for each area. Values from the *Amundsen* and the *Frosti* for a given year and area were pooled.

2.4. Otolith analysis

The lapillar otoliths of a stratified subset of 962 polar cod from all areas and years were extracted and polished (see Bouchard and Fortier, 2008 for further details). Daily growth rings were enumerated under a microscope coupled to an image analyser to determine the age and hatch date (HD: date of capture – age at capture) of each fish. Based on the size of the otolith nucleus (Bouchard et al., 2013), 34 *Arctogadus glacialis* were detected and discarded from the analysis. To compare SL at age between years and areas, SL on 6 August (the earliest date a polar cod was captured over the 2005–2015 period) was back-calculated from increment width using the biological intercept method (Campana and Jones, 1992).

2.5. Hydroacoustics

On both the *Amundsen* and the *Frosti*, hydroacoustic data were continuously recorded with a Simrad EK60[®] split-beam echosounder with hull-mounted transducers at 38 kHz and 120 kHz (both with nominal beam angle of 7°). The ping rate varied from ~ 1 to 2 s depending on maximum depth, and pulse duration was set to 1024 μs . Power was 2 kW at 38 kHz and 500 W (2006–2011) or

Table 1

Annual ichthyoplankton net and hydroacoustic sampling periods (date range) for each region. No hydroacoustic data were recorded in 2005.

Year	Beaufort Sea			Kitikmeot			Baffin Bay		
	Aug	Sep	Oct	Aug	Sep	Oct	Aug	Sep	Oct
2005				27–30			16–22		
2006					23–27			6–20	
2007						11–13			1–4
2008								9–23	
2010	14–31		1–8	8–14					
2011		1–31	1–4	7–12		3–8			15–22
2012	6–31								
2013	1–31						9–29		
2014	2–31	1–25		10–16			1–7		
2015				15–20	20–23		4–12		5–19

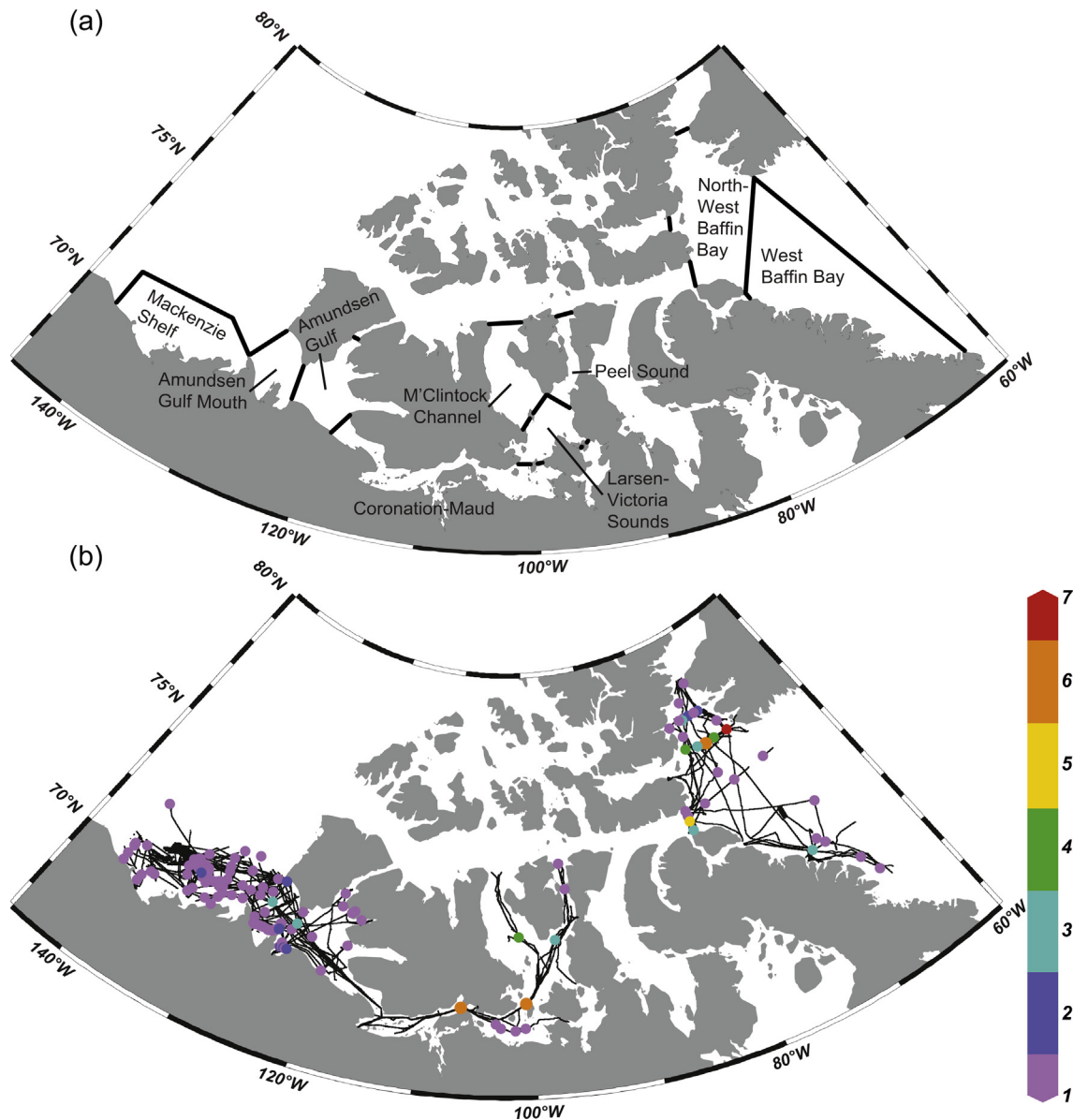


Fig. 1. (a) Boundaries of the 9 Canadian Ice Service areas. (b) Hydroacoustic and net sampling. Black lines represent all hydroacoustic transects from 2006 to 2015. Coloured dots are the locations of net sampling with colour coding the number of times (colour scale) each location was sampled from 2005 to 2015. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

250 W (2012–2015) at 120 kHz. Transducers were calibrated annually using the standard sphere method (Demer et al., 2015). The echograms were edited with Echoview® (versions 5 and 6) to correct for bottom detection and to remove signals distorted by ice breaking noise, the lowering of scientific instruments, and interferences from other acoustic sources. Time-varied gain (TVG) profiles were added based on the mean c_w (sound speed) and α_a (frequency-dependent coefficient of sound absorption) calculated from the closest CTD cast. A time-varied threshold ($TVT = 20 \log R + 2\alpha_a R - 140$, where R is the range from the transducer) was also added to the volume backscattering strength echograms to compensate for noise amplification at depth by the TVG (e.g. Geoffroy et al., 2016). The difference in mean volume backscattering strength ($\Delta MVBS$) at 38 and 120 kHz between corresponding echo-integration cells 0.25 nautical mile long by 3 m deep allowed discriminating pelagic fish from zooplankton ($\Delta MVBS_{120-38}$ ranging from -10 to 5 dB re $1 \text{ m}^2 \text{ m}^{-3}$, Benoit et al., 2014; Geoffroy et al., 2016). Outliers and echo-integration cells with a volume backscattering strength $S_v > -40$ dB re $1 \text{ m}^2 \text{ m}^{-3}$

m^{-3} were also discarded to avoid the inclusion of remaining bottom and artefact noise signals.

For each area and year, the monthly mean polar cod SL measured from net collections was transformed into mean target strength ($TS_N = 14.33 \log_{10}(\text{mean SL}) - 65.13$; Geoffroy et al., 2016). TS_N (dB re 1 m^2) and the nautical area backscattering coefficient (NASC, in m^2 nautical mile $^{-2}$) were then used to estimate the density of polar cod (Parker-Stetter et al., 2009) integrated between 13.5 m (shallowest effective sampling depth of the transducers due to their position on the Amundsen's hull and exclusion of near-field effect) and 100 m depth (or 1 m above the bottom in areas <100 m deep). We assumed that the SL of juvenile fish captured by nets from the surface to 100 m was representative of the fish producing the acoustic signals in the 13.5–100 m depth layer. Integrated density (D in number m^{-2}) was then transformed into biomass (B in mg m^{-2}) based on the monthly mean weight estimated from net samples (Benoit et al., 2008; Simmonds and MacLennan, 2005) and the weight-length regression for polar cod ($\text{Weight} = 0.0055 \text{ SL}^{3.19}$; Geoffroy et al., 2016).

The fine-scale vertical distribution of young polar cod is poorly documented, but 59% of larvae and juveniles 5–35 mm captured from early June to early August by a multinet sampler between the surface and 140 m were found in the top 10 m of the water column (Bouchard et al., 2016). Hence, by excluding the top 13.5 m of the water column, our acoustical estimates of age-0 polar cod abundance and biomass are certainly underestimates. However, the bias is assumed constant across years and areas. Hereafter, the 13.5–100 m depth interval is defined as the epipelagic layer.

2.6. Statistics

Correlations between environmental variables and the density or biomass of epipelagic age-0 polar cod were tested with the non-parametric Spearman rank correlation coefficient. Linear regressions of ice breakup date on year were calculated to predict breakup date and juvenile biomass in 2050 and 2100. A two-way ANOVA was performed to test the effect of hatch date and spring-summer SST on SL on 6 August. All statistical analyses were conducted with R version 1.0.136.

3. Results

3.1. Ice breakup, sea surface temperatures, and the survival of age-0 polar cod

Over the period 1979–2016, ice breakup week occurred progressively earlier in all areas (Spearman rank correlation, $P \leq 0.05$). The linear regression of breakup week on year was statistically significant for all areas except North-West Baffin Bay and Amundsen Gulf (Table 2). The rate of hastening of the breakup varied from 2.8 d decade⁻¹ in the Coronation-Maud area to 9.3 d decade⁻¹ in the Mackenzie Shelf area (Table 2).

From 2005 to 2015, mean SST in spring-summer (1 May to 31 July) increased significantly (Spearman rank correlation, $P \leq 0.0001$) with an earlier ice breakup (Fig. 2). For all areas, mean SST reached ~2–5 °C when ice breakup occurred before week 25 (19–25 June) and remained around 0 °C when ice broke up after week 31 (end of July).

The density (number m⁻²) of age-0 polar cod in the epipelagic layer (13.5–100 m) in August and September decreased exponentially with later breakup week and increased exponentially with increasing spring-summer SST (Fig. 3; Spearman rank correlation, $P = 0.04$). No significant correlation was found in October (Fig. 3).

The standard length on 6 August of age-0 polar cod sampled by net increased exponentially with earlier ice breakup and warmer SST (Fig. 4a and b). Mean SL on 6 August ranged from 7.6 mm in Larsen-Victoria Sounds in 2005 to 30.7 mm in Mackenzie Shelf in 2014, corresponding respectively to 2.3 mg and 197 mg wet weight, an 85-fold difference.

SL on 6 August was strongly influenced by hatch date (HD; Fig. 4c) and, to a lesser extent, by spring-summer SST (two-way

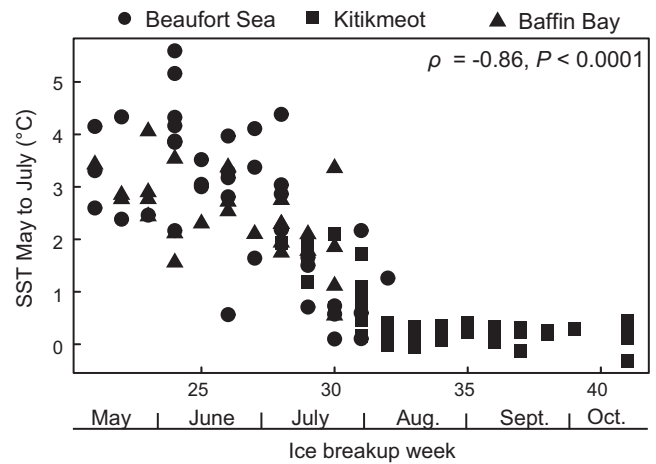


Fig. 2. Mean SST in spring-summer (from 1 May to 31 July) in relation to ice breakup week (IBW) from 2005 to 2015 for all areas in the Beaufort Sea, the Kitikmeot, and Baffin Bay. ρ is the Spearman rank correlation coefficient.

ANOVA, $F_{HD} = 6126$, $P < 0.0001$; $F_{SST} = 1156$, $P < 0.0001$). There was no significant interaction between HD and SST ($P = 0.73$).

The biomass of age-0 polar cod in the epipelagic layer in August and September increased exponentially with earlier breakup and increasing spring-summer SST (Fig. 5). Ice breakup week explained 63% of biomass variations in August and 65% in September (linear regressions, $P \leq 0.002$). Epipelagic juvenile biomass in September, obtained from the linear regression, ranged from 38 mg m⁻² for the latest sea ice breakup observed on week 36 (early September) to 420 mg m⁻² for the earliest observed breakup on week 21 (mid May), an 11-fold difference. There was no indication that the relationships between biomass and ice breakup week or SST differed among the Beaufort Sea, Kitikmeot, and Baffin Bay regions (Fig. 5). No significant correlation was found in October.

3.2. Forecasting age-0 polar cod biomass in 2050 and 2100

The regressions of ice breakup week on years for different areas of the Canadian Arctic (Table 2) predict that breakup will occur 1–5 weeks earlier than nowadays in 2050 and 4–11 weeks earlier in 2100 (Table 3). Depending on the area, the projected average standard length (weight) of epipelagic juveniles on 6 August will be 15–30 mm (19–176 mg) in 2050 and 20–42 mm (49–519 mg) in 2100. By comparison, observed mean SL (weight) on 6 August from 2005 to 2015 varied from 11 to 25 mm (7–102 mg) among all areas.

According to our results, earlier ice breakup will also increase the biomass of age-0 polar cod in the epipelagic layer in late summer. Depending on the area, an earlier ice breakup would increase the biomass of epipelagic juveniles in August by a median value of 2.7 folds (0.9–18) by 2050 and 9.7 folds (3–131) by 2100 (Table 3).

Table 2

Regression of ice breakup week (IBW) on year (Y) for 1979–2016 ($n = 38$ years) by sea ice area, and corresponding hastening of the breakup in days per decade.

Area	Regression	r^2	P	Days per decade
W Baffin Bay	IBW = $-0.0742Y + 177.77$	0.35	<0.0001	5.2
NW Baffin Bay	IBW = $-0.0551Y + 134.58$	0.08	0.08	3.9
M'Clintock Channel	IBW = $-0.1265Y + 291.40$	0.23	0.002	8.9
Larsen-Victoria Sounds	IBW = $-0.1010Y + 236.92$	0.25	0.001	7.1
Peel Sound	IBW = $-0.0646Y + 162.75$	0.10	0.05	4.5
Coronation-Maud	IBW = $-0.0407Y + 111.73$	0.17	0.011	2.8
Amundsen Gulf	IBW = $-0.0743Y + 176.28$	0.05	0.16	5.2
Amundsen Gulf Mouth	IBW = $-0.1131Y + 252.21$	0.14	0.02	7.9
Mackenzie Shelf	IBW = $-0.1327Y + 293.47$	0.14	0.02	9.3

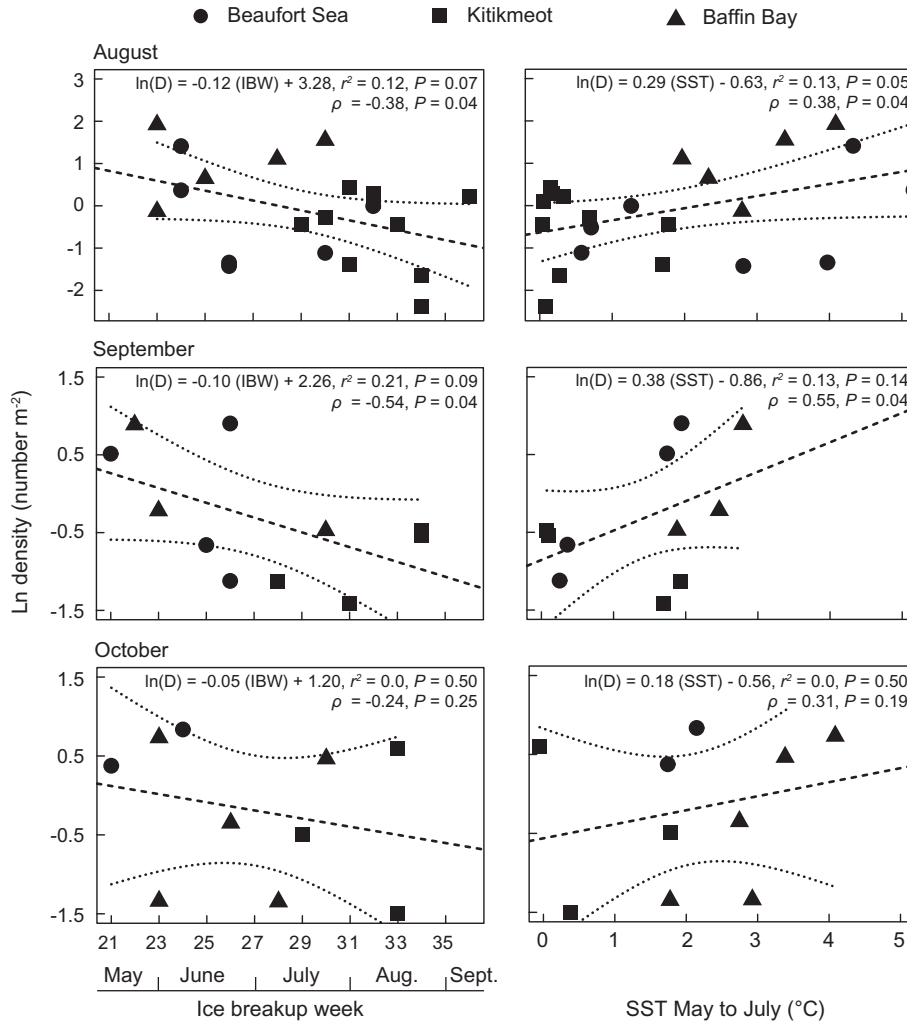


Fig. 3. Density (D) of age-0 epipelagic polar cod in relation to ice breakup week (IBW, left panels) and mean SST in spring-summer (right panels). Dashed lines are the linear regressions and dotted lines the 95% confidence intervals. ρ is the Spearman rank correlation.

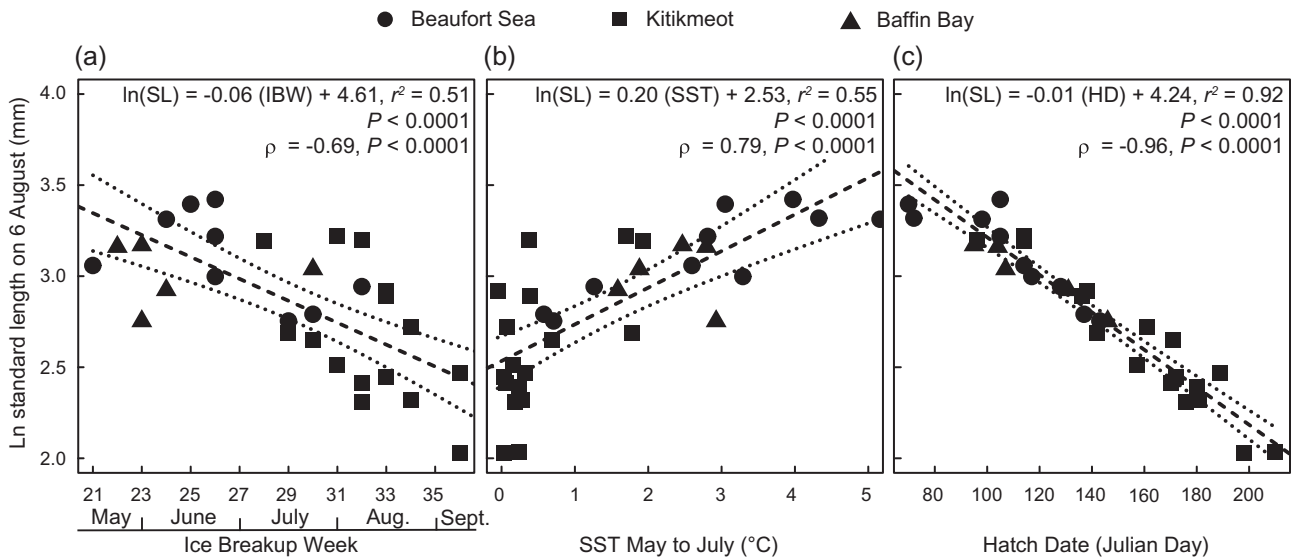


Fig. 4. Standard length (SL) of age-0 polar cod on 6 August in relation to (a) ice breakup week (IBW), (b) mean spring-summer SST, and (c) hatch date (HD). Dashed lines are the linear regressions and dotted lines the 95% confidence intervals. ρ is the Spearman rank correlation.

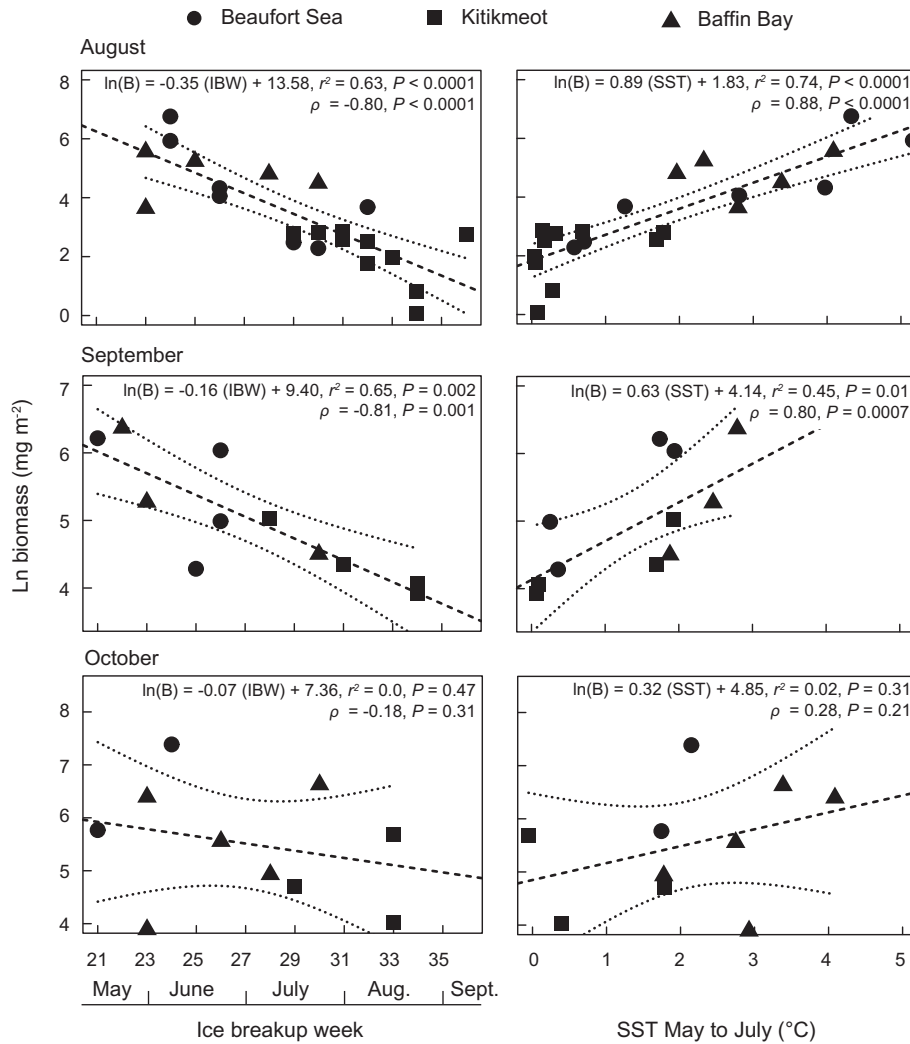


Fig. 5. Biomass (B) of age-0 epipelagic polar cod in relation to ice breakup week (IBW, left panels) and mean spring-summer SST (right panels). Dashed lines are the linear regressions and dotted lines the 95% confidence intervals. ρ is the Spearman rank correlation.

Table 3
Mean ice breakup week (IBW) observed from 2005 to 2015 and projected ice breakup week for 2050 and 2100 based on the equations in Table 2 for different areas of the Canadian Arctic; corresponding observed and projected standard length (mean SL) and weight (mean W) on 6 August, and epipelagic juvenile biomass (mean B) in August. Projected SL from $\ln(\text{SL}) = -0.06(\text{IBW}) + 4.61$ (Fig. 4a), transformed to weight according to $W = 0.0055 \text{SL}^{3.19}$ (Geoffroy et al., 2016); projected biomass in August from $\ln(B) = -0.35(\text{IBW}) + 13.58$ (Fig. 5).

Area	Week of ice breakup			SL on 6 August (mm)			Weight on 6 August (mg)			Biomass in August (mg m^{-2})		
	2005–2015	2050	2100	2005–2015	2050	2100	2005–2015	2050	2100	2005–2015	2050	2100
W Baffin Bay	28	26	22	21.1	21.6	26.9	59.4	63.7	129.5	112	99	363
NW Baffin Bay	23	22	19	20.6	27.5	32.4	55.4	138.0	233.7	172	408	1068
M'Clintock Channel	36	32	26	15.8	14.7	21.4	23.8	18.6	62.6	–	10	96
Larsen-Victoria Sounds	34	30	25	10.6	16.7	22.7	6.7	28.5	74.8	5	23	133
Peel Sound	33	30	27	11.1	16.3	19.8	7.7	26.1	48.5	12	19	60
Coronation-Maud	30	28	26	20.6	18.4	20.8	54.8	38.4	56.8	13	39	80
Amundsen Gulf	27	24	20	23.1	23.9	29.8	79.0	88.1	179.3	10	180	659
Amundsen Gulf Mouth	25	20	15	20.7	29.6	41.6	55.8	176.1	519.1	35	636	4595
Mackenzie Shelf	26	21	15	25.0	27.8	41.4	101.6	142.8	509.2	338	434	4437

4. Discussion

4.1. The mechanism linking polar cod recruitment to ice breakup timing

A central paradigm of fisheries science is that the size of an annual cohort of fish (the strength of a year class) is determined by survival during the early larval life from hatching to metamor-

phosis into juvenile (e.g. Houde, 2008). The match/mismatch hypothesis proposes that early growth and survival depend on the synchronized development of the larvae and their plankton food (Cushing and Horwood, 1994). Somewhat at odds with this hypothesis, the hatching season of polar cod extends from as early as January to the first week of July, with most larvae hatching under the ice in April and May when light, temperature, and prey availability are low – an apparent evolutionary aberration

(Bouchard and Fortier, 2008, 2011; Fortier et al., 2006). It has been suggested that the selection force driving some polar cod to hatch under the ice in winter and early spring is the survival advantage provided by the large pre-winter size resulting from early hatching (Bouchard and Fortier, 2008, 2011; Fortier et al., 2006). In juvenile fish, a large size is associated with better winter survival through increased lipid content, predator avoidance, resistance to starvation, and physiological tolerance (e.g. Hunt et al., 2011; Sogard, 1997). In polar cod, the largest pre-winter sizes are achieved by larvae hatched in mid-winter in areas where river plumes provide a freshwater thermal refuge against sub-zero temperatures (Bouchard and Fortier, 2011). Hatching beyond mid-July, when temperature, light and food availability are at their maximum, is selected against because the pre-winter size achieved would be too small to ensure survival over the subsequent winter.

Our results are consistent with this notion that maximizing pre-winter size rather than ensuring a match between larval growth and the maximum production of plankton food is the driving force behind the intriguing reproductive strategy of polar cod. Given an extended hatching season from January to early July, and notwithstanding suboptimal survival conditions in winter and spring relative to summer (Fortier et al., 2006), the survival of early hatchers is clearly enhanced in years and areas with an early ice breakup relative to late breakups, resulting in more abundant and heavier epipelagic juveniles in late summer (August and September). The resulting differences in size were spectacular, with mean weight on 6 August ranging from 2.3 mg in Larsen-Victoria Sounds in 2005 (ice breakup in week 36 or early September) to 197 mg on the Mackenzie Shelf in 2014 (ice breakup in week 26 or end of June). Hence, despite suboptimal conditions for the survival of first-feeding larvae in winter and spring relative to summer (Fortier et al., 2006), winter-spring hatching provides the potential for the population of juveniles to achieve large biomasses in the fall and, presumably, strong recruitment after the first winter. Given the central importance of polar cod in the pelagic food web of Arctic seas, poor juvenile recruitment in years of late breakup is expected to negatively impact polar cod predators. Consistent with this, poor reproduction in ringed seals has been associated with years of heavy ice conditions in the mid-1970s and mid-1980s in the Beaufort Sea (Stirling, 2002).

Among areas and years, an earlier ice breakup was correlated with warmer SST in spring-summer (Galbraith and Larouche, 2011; Wood et al., 2013). The early growth of polar cod is positively correlated to SST (Bouchard and Fortier, 2011) and survival is low at near-zero temperatures (Fortier et al., 2006). In the present study, the length achieved on 6 August was correlated to SST, but more strongly so to the mean hatch date of the fish collected in an area and a given year (Fig. 4b and c). Hatch date explained 5.3 times more variability in length on 6 August than did SST (ratio of the F statistics in the two-way ANOVA). Hence, early ice breakup favours a large pre-winter size primarily by allowing the survival of early hatchers and to a lesser extent by providing warmer temperatures for growth over the summer. Polar cod larvae and epipelagic juveniles are visual predators that feed primarily on calanoid copepod nauplii (Bouchard et al., 2016; Michaud et al., 1996). We suspect that, in addition to higher SST, advanced blooms of ice algae and phytoplankton (e.g. Kahru et al., 2011), the associated hastening of the production of copepod nauplii (e.g. Daase et al., 2013; Fortier et al., 1995; Ringuette et al., 2002), and a greater availability of light (e.g. Bouchard and Fortier, 2008; Ponton and Fortier, 1992; Varpe et al., 2015) were the proximal causes of the improved survival of early hatchers in years of early ice breakup.

Interestingly, the biomass of the epipelagic juveniles in August and September was more strongly correlated to ice breakup week or to spring-summer SST (Fig. 5) than were either density (Fig. 3) or

size on 6 August (Fig. 4a and b). This suggests that biomass, which reflects both the number of fish surviving and the weight each survivor achieves, is the best integrator of the effect of the environment on the combined growth and survival of young epipelagic polar cod and on their ultimate recruitment.

The weakening in October of the slope of the relationship between biomass and ice breakup week or SST is not unexpected. Polar cod juveniles leave the epipelagic layer to descend to overwintering depths when they reach 35–55 mm in length (Benoit et al., 2014; Geoffroy et al., 2016). In years and areas with early ice breakup, juveniles will attain this threshold length early, and most will have left the epipelagic layer by October resulting in relatively low epipelagic biomasses in that month. In years and areas of late breakup, small size will delay migration to depth resulting in relatively high biomass in the top 100 m in October. Hence the importance of conducting surveys at the appropriate period (August and September in this case) when assessing the recruitment of juvenile polar cod.

4.2. Polar cod and climate warming: a likely bust beyond the initial boom

Age-2+ polar cod are mostly mesopelagic, forming dense aggregations in deep embayments and along the shelf slope (Benoit et al., 2008, 2014; Geoffroy et al., 2016, 2011). The epipelagic larvae and age-0 juveniles develop in the top 100 m surface layer over the spring-summer (e.g. Bouchard and Fortier, 2011; Bouchard et al., 2016; Geoffroy et al., 2016). By early fall, age-0 juveniles descend to intermediate depths to join the deep adult population (Geoffroy et al., 2016) or colonize the ice pack as age-1 (David et al., 2016). Hence, it is primarily the epipelagic age-0 and the sympagic age-1 fish that are exposed to shifts in the sea ice and SST regimes induced by climate warming.

As in other Arctic regions (Stroeve et al., 2012), trends towards an earlier ice breakup have been observed over the period 1979–2016 in all nine areas of the Canadian Arctic studied here. Based on our results, this hastening of sea ice breakup forced by global warming will increase the biomass of age-0 polar cod juveniles by a median value of 2.7-fold (0.9–18) by 2050 and 9.7-fold (3–131) by 2100 across the different areas. Because interannual differences in mortality occur primarily during the larval stages, the density and biomass of juveniles at the end of their first growing season are valid proxies of gadid recruitment (Laurel et al., 2016a). Hence, the ongoing warming of the lower atmosphere will likely increase the abundance of polar cod in the Canadian High Arctic, at least as long as the observed relationships between ice breakup timing and juvenile biomass hold. Similar trends can be expected in other High Arctic regions, but not necessarily in sub-arctic zones of the distribution of the species, where any hastening of an already early ice breakup could be detrimental to the early survival of polar cod, and likely would favour competing boreal forage fish.

Our optimistic predictions for future polar cod recruitment assume that these relationships will persist through the century, despite the expected major shifts in sea ice and SST regimes. Statistical relationships linking fish recruitment to some environmental variable often fail when new observations are added that fall outside the range of environmental conditions covered by the initial data set (Frank, 1997; Leggett and Frank, 2008; Myers, 1998). As climate warming unfolds, the reduction of the ice cover is expected to ultimately impact the cryopelagic polar cod negatively (e.g. Kohlbach et al., 2017; Moline et al., 2008; Tynan and DeMaster, 1997). The question is how and when will the presently beneficial relaxation of the extreme arctic conditions start to negatively impact the reproduction of the highly specialized polar cod? In the laboratory, the eggs of polar cod develop normally in the tem-

perature range -1.0 to 3.5 °C, but not beyond ≥ 5 °C (Drost et al., 2016; Kent et al., 2016). Age-0 polar cod grow optimally at 9.0 °C (Laurel et al., 2017, 2016b). The winter ice cover of Canadian Arctic seas is likely to continue to form over the present century (Stroeve et al., 2012), providing the frigid conditions needed for normal egg development. As well, mean spring-summer SST in the surface are presently in the range 0 – 5 °C (Fig. 2), which leaves much room for positive effects from warming before the optimal developmental temperature for age-0 polar cod (9 °C) is exceeded. Hence, increasing SST in spring and summer may have little immediate negative effect on polar cod recruitment.

As discussed above, a progressively earlier ice breakup will improve juvenile recruitment in the fall by increasing the survival of early hatchers, at least initially. There may exist a threshold beyond which an ever-earlier ice breakup becomes detrimental to the survival of polar cod early hatchers. For instance, ice algae start growing when light levels at the ice-water interface reach $\sim 7 \mu\text{E m}^{-2} \text{s}^{-1}$ (Gosselin et al., 1985), typically in April at the latitudes studied here. If ice breakup eventually regresses to April, ice algal production would no longer occur for lack of an ice substrate. A reduced ice algal production might disrupt the reproduction of copepods such as *Pseudocalanus* spp. and *Calanus glacialis* (Tourangeau and Runge, 1991), the nauplii of which are the main prey of polar cod larvae at first feeding (Bouchard et al., 2016). Similarly, juvenile polar cod rely on the nauplii and copepodites of the lipid-rich *Calanus glacialis* and *Calanus hyperboreus* to sustain growth (e.g. Bouchard et al., 2016). Warmer waters could trigger a replacement of these species by smaller, relatively lipid-poor species such as the subarctic *Calanus finmarchicus* (Kjellerup et al., 2012; Slagstad et al., 2011). For instance, Hunt et al. (2011) report that large numbers of age-0 walleye pollock (*Gadus chalcogrammus*) survive to summer in years of extremely early ice breakup in the Bering Sea. However, extremely early ice breakups also reduce the production of large, lipid-rich, crustacean prey, which results in low condition of age-0 pollock, increased vulnerability to predation as fish switch from zooplankton prey to age-0 pollock, and poor recruitment to age-1. Finally, an early ice melt could shrink the habitat of some of the age-1 and age-2 polar cod that recruit to the ice pack of the Arctic Ocean (David et al., 2016).

However, the ongoing migration to northern waters of subarctic species in response to changing conditions (e.g. Berge et al., 2015; Fossheim et al., 2015; Kortsch et al., 2015) could eventually be the main cause of the demise of the hyper-specialized polar cod, with all the expected cascading impacts on the unique pelagic ecosystems and biodiversity of Arctic seas (e.g. Moline et al., 2008; Tynan and DeMaster, 1997). Already, indications are that generalist forage fish such as the capelin (*Mallotus villosus*) and the sand lance (*Ammodytes* spp.) are outcompeting polar cod in Hudson Bay (Provencher et al., 2012). Potential predators of polar cod, such as the Atlantic cod (*Gadus morhua*), haddock (*Melanogrammus aeglefinus*) and Atlantic mackerel (*Scomber scombrus*), are thriving in Svalbard waters and the Greenland Sea (Berge et al., 2015), and other piscivorous species, such as redfish (*Sebastes* spp.), recently became ubiquitous in the area (M. Geoffroy, unpublished data). In coming decades or by mid-century, competition and predation from invading subarctic species will likely curb the ongoing positive effect of a progressively earlier ice breakup on the survival of young polar cod.

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