1 Recruitment of shrimp (*Pandalus borealis*) in the Barents Sea related to spawning stock

2 and environment

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4

5 Abstract

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The shrimp spawn in autumn, and the females carry their eggs as out roe until spring when the 7 larvae hatch. Within a period of 2 months the shrimp larvae settle to the bottom. It has been 8 9 claimed that the year class strength probably is determined during the larval phase. Today's assessment and forecast of the shrimp stock productivity and potential fishing yields is weak. 10 This is partly due to poor knowledge on population dynamics from hatching until the shrimp 11 are caught in the fishery at the age of 3 or 4 years. We therefore here identify the most 12 important abiotic and biotic factors that effect recruitment in addition to spawning stock 13 biomass. Since 1995 a net attached to the underbelly of the survey trawl used at the annual 14 cruise in the Barents Sea has caught juvenile shrimp. The abundance of settled shrimp larvae 15 varies in time and space. The recruitment to the fishery has been quite stable with the 16 17 exception of the 1996 year-class which was observed as 1 year old but has not been registered 18 since. The temporal pattern of the three youngest year-classes are studied in relation to abiotic factors such as sea temperature, ice index and North Atlantic Oscillation, as well as biotic 19 factors such as spawning stock biomass and presence of copepods, euphausiids and predating 20 cod. Recruitment indices and factors identified by Spearmann correlation to be significantly 21 correlated with recruitment were used as input in a principal component analysis (PCA) and a 22 generalized additive model (GAM) was applied. Abundance of 1 year old shrimp is positively 23 correlated to spawning stock biomass the previous year and temperature the last winter and 24 negatively correlated with the number of 1 year old cod. Two year old shrimp show 25 significant correlation with temperature while there is a strong negative correlation with 26 euphausiids. Three year old shrimp are significantly correlated with the number of 2 year old 27

28	shrimp the previous year but negatively correlated to temperature at sampling time. This is
29	probably due to less overlap with the main predator cod when cold. Ricker functions indicate
30	an increased density dependent mortality with age. When predicting the recruitment of shrimp
31	to the fishery the spawning stock biomass, the abundance of cod and euphausiids, as well as
32	the temperature should be included.
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34	Key words: Pandalus borealis, recruitment, Northeast Atlantic, Barents Sea, Spawning stock,
35	environmental factors
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- 43 **1.** Introduction
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The shrimp (*Pandalus borealis*) is a protandric hermaphrodite changing sex from male to female at an age of 4 to 7 years in the Northeast Atlantic (Nilssen and Hopkins, 1992). The shrimp spawn in autumn and the eggs are carried as out roe by the females (ovigerous females) until spring when the larvae hatch. Within a period of 2–3 months the shrimp larvae pass through six developmental stages after which they settle to the bottom as post larvae (Shumway et al., 1985; Ouellet et al. 1990; Bergström, 2000).

It is of major importance for the shrimp stock assessment to get information on the strengths 51 of the recruiting year classes as early as possible. Today's assessment forecast of the shrimp 52 stock productivity and potential fishing yields is weak, partly due to the lack of knowledge on 53 the population dynamics from hatching until the shrimp are caught in the fishery. According 54 to Shumway et al. (1985) the year class strength of shrimp is probably largely established 55 during the pelagic larval stage. In the Barents Sea shrimp larvae are transported 0-300 km 56 during the larval phase (Pedersen et al., 2003). It is assumed that the transport processes 57 58 influence the recruitment both directly as advectional losses of larvae and indirectly through temperature, food availability and predator-prey interactions (Apollino et al., 1986; Lysy and 59 Dvinia, 1991; Clarke et al., 1991; Ouellet et al., 1995). Ouellet et al. (2007) found that 60 survival of the pelagic shrimp larvae was dependent on warming rate and the depth of the 61 mixed water layer. The object of this study is to define relevant recruitment indices and 62 identify the environmental factors determining recruitment success. Annual variation in 63 recruitment and larval survival are related to abiotic factors; temperature, ice coverage, the 64 North Atlantic Oscillation (NAO), and to biotic factors; spawning stock biomass (SSB) 65 defined as biomass of ovigerous females the previous year, abundance of zooplankton, 66 euphausiids and capelin (Mallotus villosus). We will also study the effect of the main predator 67

68	cod (Gadus morhua) as predation has been shown to overrule the positive effect of favourable
69	temperature conditions on shrimp survival (Wieland et al., 2007).

- 71 2. Material and Methods
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73 2.1. Study area
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The Barents Sea is a shelf sea with an average depth of 230 m. The circulation is dominated 75 76 by the Norwegian Atlantic Current that enters the Barents Sea through the Bear Island Trench. Some of the Atlantic Water flow eastward parallel to the coast towards Novaya 77 Zemlya while another part flows northeast wards and into the Hopen Deep (Fig.1). The 78 79 relative strength of these two branches varies with the atmospheric fields (Ingvaldsen et al., 2003). South of the Atlantic inflow, the Norwegian Coastal Current continues along the 80 81 Finnmark and Kola coast. In the northern and eastern parts of the Barents Sea, Arctic water flow southwest wards near the surface. The Atlantic inflow continues towards northeast 82 83 below this layer. The Barents Sea has several bank areas with associated anticyclonic 84 circulation. The Atlantic inflow to the Barents Sea shows considerable inter-annual variability (Ådlandsvik and Loeng, 1991). Fig. 1 85

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The major water masses in the investigated area are Atlantic Water with salinity over 35‰, and the colder and fresher Arctic Water. These water masses are separated by the Polar Front. In the western Barents Sea the position of the front is relatively stable, although it seems to be pushed northwards during warm climatic periods (Loeng, 1991; Ingvaldsen, 2005). In the eastern part the position of the front has large seasonal, as well as year to year variations. Ice conditions also show large seasonal and year to year variations. In the winter the ice can cover most of the northern Barents Sea, while in the summer the whole sea may be ice-free.

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95 2.2. Abiotic factors

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97	The climate of the Barents Sea shows substantial variations, and the temperature fluctuations
98	at the Vardø-N section along 31°13'E (Fig. 1) gives a good representation of the climate
99	variability in the central Barents Sea (Ingvaldsen et al., 2003) and the position of the Polar
100	Front in the Hopen Deep (Skjoldal et al., 1987). The mean temperature in the section,
101	between 50–200 m depth, was calculated from 72°15'N to 74°15'N. The section is sampled 4
102	times a year (January, March, June-July and August-September). In addition to January and
103	March temperatures (best available measure of winter temperature) the annual mean
104	temperature was used in the correlation analysis.
105	
106	From a temperature atlas where all observations from August-October each year have been
107	interpolated to a regular grid, a section along the Hopen Deep between $74^{\circ}N$ and $76^{\circ}30'N$ at
108	50 to 200 m was defined to provide a time series of temperature in the northernmost parts of
109	the Barents Sea where shrimp is most abundant (Fig. 1). Temperatures from
110	August-September each year were used instead of annual means, because by including winter
111	temperatures the section could not extend as far north. As the temperature level for the rest of
112	the year is determined by the winter temperature, the annual variation is clear also in the
113	summer temperature (Ottersen et al., 2000). Bottom temperatures in the Hopen Deep (Fig. 1
114	Area E) were extracted from a similar temperature atlas and mean temperatures were
115	calculated. The Russian section defined along the Kola meridian at 33°30' E (Bochkov 1982
116	and Tereshchenko, 1996) gives a good indication of the temperature variation in the southern
117	Barents Sea (Ingvaldsen et al., 2003). The annual mean temperature for the upper 200 m from
118	70°30'N to 72°30' N, were calculated and used in the correlation analysis.
119	

The North Atlantic Oscillation (NAO) is one of the most prominent and recurrent patterns of
seasonal and long-term atmospheric variability in the North Atlantic Ocean and is there fore

122	used in this study. The NAO index from NCAR (Hurrell, 1995) is a much used, but crude
123	indicator of the south-westerly winds in the Norwegian and Barents Sea, and has significant
124	effect on the Barents Sea temperatures (Ottersen et al., 2003).

The ice index is an integrated value for the ice coverage between 25°E and 45°E in the
Barents Sea and is a good indicator of the climate conditions. A definition of the index is
given by Ådlandsvik and Loeng (1991), and values are updated annually by the Institute of
Marine Research (Skogen et al., 2007). Note that a low index corresponds to heavy ice
conditions.

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132 2.3. Biotic factors

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134 Since 1987, the zooplankton abundance has been monitored on annual surveys during the Norwegian-Russian 0-group and capelin surveys that are carried out in autumn. The 135 zooplankton biomass in the Barents Sea is dominated by Calanus finmarchicus, and the 136 137 biomass estimate is based on net-hauls from bottom to surface (Dalpadado et al., 2003; Skogen et al., 2007). Euphausiids (Thysanoessa inermis, T. raschii, T. longicaudata and 138 Meganyctiphanes norvegica) have been monitored in the Barents Sea since 1952 at demersal 139 fish surveys conducted by Russian scientists at PINRO. A net is attached to the survey trawl 140 and sampled in the autumn and winter period when euphausiids are concentrated close to the 141 142 bottom and show no vertical migration (Drobysheva et al., 2003). 143 The number of 1 year old capelin sampled in August was included in the correlation analysis 144

since there may be a relationship between the shrimp and the capelin directly by capelin

146 feeding on shrimp juveniles or indirectly by both being prey for the cod (Skogen et al., 2007).

147 The Arctic Fisheries Working Group report (Anonymous, 2006) lists the number of cod in

- age group 3, whereas Bjarte Bogstad (IMR) provided data on age groups 1 and 2. The number 148 of cod in each year class is the output of the VPA. 149
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Shrimp spawning stock and recruitment 151 2.4.

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Annual shrimp surveys have been conducted by Norway in the Barents Sea every spring in 153 the period 1982–2004. Between 100 and 200 stations have been towed with 3 knots¹ for one 154 hour, 30 minutes or, since 1992 for 20 minutes. Shrimp are sexed and length measured on 155 board the vessel. The spawning stock biomass is presented as the weight (ton) of reproducing 156 females (egg carrying females and females with hatched eggs) per square nautical mile² (nm⁻ 157 ²). Aschan and Sunnanå (1997) described in detail the procedures for sampling, stratification, 158 length measurements and sex determination. 159

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As the survey trawl (Campelen 1800) only caches large shrimp (carapace length (CL) >15 161 mm equals total length > 6 cm) efficiently, sampling techniques for sampling the smallest 162 shrimp have been tested and evaluated (Larsen et al., 1993; Aschan et al., 2000). In 1995 a 163 fine meshed (8 mm) juvenile bag with a 1 m² opening was attached to the lower trawl belly 164 one meter in front of the junction to the cod end (Nilssen et al., 1986; Aschan and Sunnanå, 165 1997). Because the juvenile bag samples both 1 and 2 year old shrimp escaping the trawl 166 167 trough the belly and thereby does not require additional sampling time, it was chosen as the main sampling tool for small shrimp. Indices for 3 years old and also for 2 year old shrimp 168 were produced from the cod end sample. 169

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The samples were weighted to the shrimp catch at each station and the strata size in a swept 171 area calculation before calculating the average density by sub-area (A-F, Fig. 1). Number of 172 shrimp in each 1 mm interval was calculated for the years 1989–2004 and for the belly bag 173

 $^{^{1}}$ 1 knot = 1 nautical mile/h = 1.852 km/h = 0.514 ms-1 2 1 square nautical mile = (1.852 km)² = 3.4299 km²

174	for the years 1995–2004. The young shrimp (< 15 mm) are divided into age groups; age 1: CL
175	<9 mm, age 2: 9 mm \leq CL< 12 mm and age 3: 12 mm \leq CL< 15 mm, according to the length
176	at age key generated from the Hopen Deep (Aschan, 2001). This length at age key is based on
177	data collected since 1990 and is not necessarily applicable on shrimp collected in the 1980's.
178	However, when describing the long time spatial recruitment pattern the index of 2 year old
179	shrimp, defined as above, is presented by area for year class 1980 to 2002.
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181	2.5. Numerical analysis
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183	Regression analysis between 2 year old shrimp caught in the juvenile bag and same age
184	shrimp caught in the cod end was run to identify if the latter could be used as a recruitment
185	index and thereby provide a longer time series.
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187	The spawning stock recruitment relationship was studied by fitting linear regression and the
188	Ricker (1954) spawning stock recruitment model to recruitment indices of age 1 (1994-2003
189	year class), age 2 (1989–2002 year class) and age 3 shrimp (1986–2001 year class) and
190	spawning stock biomass. The formula for the Ricker model is:
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192	$R = a \cdot SSB \cdot e^{(-b \cdot SSB)}$
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194	where R is the recruitment, SSB is our measure of the spawning stock biomass in the year of
195	recruitment, and a and b are constants.
196	
197	We used two criteria for determining the fit of the models, the adjusted R^2 where a larger
198	value indicates a better fit, and the akaike information criterion (AIC) where smaller value
199	indicate a better more parsimonious model (Quinn and Keough 2002).

As the shrimp recruitment is likely to depend on the spawning stock biomass a survival index was established for age 1 and age 2 as recruitment per unit of SSB which produced these recruits. For age 3 the survival index is the number of age 3 recruits per age 2 recruits the previous year.

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A correlation analysis between recruitment measures, recruitment indices and survival indices 206 (age 1, age 2 and age 3), and abiotic and biotic parameters was run using Spearman's rho 207 statistic to estimate a rank-based measure of association. The rank Spearman correlation 208 coefficient (rho) is calculated after the variables have separately been transformed to ranks 209 and is a conservative measure on monotonic relationships between the variables. The 210 Spearman correlation is robust and has been recommended when data do not necessarily 211 come from a bivariate normal distribution (Quinn and Keough 2002). The populations in 212 nature, in this study temperature, species abundance etc., seldom have a normal distribution, 213 214 and one can not assume that the data are normally distributed. When using our few data Shapiro-Wilkins normality test indicated normality for some variables (annual mean 215 temperature Vardø P = 0.02, Capelin P = 0.008, Cod 1 P = 0.05, Cod 2 P = 0.02, Age 2 P = 216 0.03 and Survivals = Age 3/Age 2 P = 0.01), but normality plots showed that the outermost 217 points did not follow the normal distribution. 218

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All variables were tested for autocorrelation for a lag of one, two and three years. Where autocorrelation was significant ($P \le 0.05$), we corrected for it by adjusting the degrees of freedom used in the Spearman's rho statistics. This adjustment was done in accordance with Pyper and Peterman (1998):

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$$\frac{1}{N^*} = \frac{1}{N} + \frac{2}{N} \sum_{j=1}^{N/5} r_{xx}(j) r_{yy}(j)$$

227 where N^* is the effective number of degrees of freedom on the time series X and Y, N is the sample size and $r_{xx}(j)$ and $r_{yy}(j)$ are the autocorrelations of X and Y at lag j. Following the 228 recommendation by Pyper and Peterman (1998) a maximum of N/5 lags were included in the 229 calculation of effective number of degrees of freedom. Therefore two lags where included for 230 1 year old shrimp and three lags for 2 and 3 year old shrimp. All statistical analyses in this 231 study were run in R 2.5.0 (R Development Core Team, 2007). 232 233 Relevant abiotic and biotic factors identified by the Spearman correlation analysis as 234 235 significant, were used as input when running a principal component analysis (PCA). A generalized additive model (GAM) in the function "ordisurf" in R fits surfaces of density of 236 recruits (age 1, age 2 and age 3) to the respective principal component ordinations (PC1 and 237 PC2). The generalized additive model (GAM) is a generalized linear model (GLM), in which 238 the linear predictor is replaced by a user specified sum of smooth functions of the covariates 239 plus a conventional parametric component of the linear predictor. The function "gam" uses 240 thinplate splines in two dimensions, and automatically selects the degree of smoothing by 241 generalized cross validation (Oksanen, 2007). 242 243

244 **3. Results**

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246 3.1. Abiotic factors

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From 1985 to 1989 there were in general low temperatures in the Barents Sea, while the period 1990–1995 was characterized by warm conditions (Fig. 2). In 1996 a sudden temperature drop occurred and it stayed cold until 1998. Since then temperatures have been high, with the exception of 2003 which was close to the long-term mean. Prior to 2000–2001 the mean temperature, the January and the March temperatures in the Vardø-N section, the NAO index and the ice index varied much in the same manner indicating warm conditions

with small amounts of ice during high NAO, but after 2000–2001 there is not a good
correlation between the NAO and other parameters. The ice index follows in general the
variations in mean temperature, but with a lag of 1–3 years. An exception was in 2003, when
there were large amounts of ice despite the temperatures not being correspondingly low. The
reason was a late onset of melting in spring 2003. In the Hopen Deep the ice edge was close
to 75°N through the entire winter, which is close to normal, but the ice edge reached 76°N
about a month later than in 2002.

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It was cold in the Hopen section in 1987, 1996 and 2003, while the warmest years were 1989, 1999 and 2004 (Fig. 2). The bottom temperature in the Hopen Deep (area E) shows much of the same variability as the mean temperature in the Hopen section (50–200 m). However, some differences are evident, and the most pronounced difference is the strong increase in bottom temperature in 1987–1992, followed by a rapid decrease in 1993. In 1996 and 2003 the bottom temperatures were low, but the decrease was not as extreme as observed at 50–200 m depth.

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The temperature along the Hopen section is significantly correlated with the bottom temperature in the Hopen Deep, with the Vardø annual mean temperature and with the NAO (Spearmann correlation rho = 0.68, 0.70 and 0.85 respectively). The temperature in Hopen and Kola sections has the same variability, but there are large deviations between the time series after the year of 1999 and therefore there is no correlation (rho -0.40). The very low temperatures measured in Hopen in the years 1996 and 2003 were not observed in Kola.

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278 3.2. Biotic factors

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280	The zooplankton and euphausiid biomass increases in the 90's and stay above the long-term
281	mean until 2004. However, both variables show great variation between years (Fig. 3). Cod
282	has very good recruitment in the period 1991–1998. Capelin has a negative correlation to cod
283	of age $1-4$ (rho: -0.44 to -0.63) while zooplankton has a positive correlation with cod age 1, 2
284	and 3 (rho: 0.66-0.69). This is the consequence of low densities of cod giving rise to high
285	numbers of capelin predating on copepods. Fig. 3
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287 3.3. Shrimp spawning stock and recruitment

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The recruitment indices given for age 2 shrimp caught in the cod end of the survey trawl and in the juvenile bag attached to the belly of the trawl are well correlated ($R^2 = 0.80$) (Fig. 4). We therefore used the longer time series for number of age 2 shrimp caught in the cod end in the correlation analysis. Fig. 4

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294 The biomass of spawning females, recruitment index for age 2 and age 3 shrimp may vary by a factor of 4–5, while the recruitment index for 1 year old shrimp vary by a factor of 10. This 295 296 indirectly indicate that the greatest mortality occur between age 1 and 2. Recruitment indices for 2 and 3 year old shrimp and spawning stock biomass sampled in the cod end each year 297 since 1989 reveals high spawning stock and good recruitment of age 2 shrimp in the early and 298 the late 90's (Fig. 5). The recruitment indices for 1 year old shrimp sampled in the belly bag 299 since 1995 show strong year-classes in 1999 and 2000. Both spawning stock biomass and 300 recruitment of all age classes are on a historically low level in 2004. Fig. 5 301 302

303 The recruitment of shrimp varies between areas and years (Fig 6). The Hopen Deep (E) is the most important area for recruitment with the highest density of recruits. The mean density of 304 age 2 shrimp in the Tiddly Bank (B) and Thor Iversen Bank (C) is less than 30% of the 305 density in the Hopen Deep (E). Off Finnmark (A), in the Bear Island Trench (D) and off Bear 306 Island (F) the density of age 2 shrimp is even lower with a mean density less then 15 % of the 307 mean density in the Hopen Deep. Except for the period 1994–1997, the number of recruits in 308 the Hopen Deep (E) has a strong covariation with the bottom temperature in the Hopen Deep 309 (Fig. 2 and Fig. 5). The poor covariation in 1994–1997 is probably due to the high number of 310 311 1 and two year cod in that period (Fig. 3), and possibly due to unusual inter-annual variability in the temperature. Fig. 6 312

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The spawning stock recruitment relationship fits both to the linear model and the Ricker model, but F–statistics give the linear model the best fit for all age groups (P<0.001) (Fig. 7 and Table 1). The AIC indicates that the stock recruitment model fit gets weaker for each age group. That is due to the biotic and abiotic factors affecting the survival rate of young shrimp over time. Fig. 7

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Some of the variables (SSB, Age 2, Cod 1) turned out to have a significant ($P \le 0.05$) 320 autocorrelation, but only at lag 1. The correlation between Age 1 and Cod 1 (-0.6) and Age 2 321 322 and SSB (-0.46) turned out to be insignificant when degrees of freedom were adjusted for (Table 2). However, survival (Age 1/SSB) and Cod 1 still have a significant negative 323 correlation. All abiotic and biotic factors presented were included in the correlation analysis, 324 but only variables giving significant correlation (P < 0.05) were included in Table 2. Thereby 325 the Kola temperature, ice index, zooplankton and cod age 3 are excluded. The correlation 326 analysis reveals that for age 1 shrimp the most important positive factor, after spawning stock 327 biomass the previous year, is the temperature in January and March the same year. The 328 presence of cod age 1 and 2 the previous year is important negative factors (Table 2.). Age 1 329

330 shrimp is positively correlated with capelin but negatively, although not significantly correlated (rho = -0.45) with zooplankton. Age 2 shrimp is not significantly correlated with 331 spawning stock biomass (rho = 0.46), is significantly correlated with temperatures (Vardø and 332 Hopen sections and Hopen bottom) and NAO the previous year, but has a significant strong 333 negative correlation with euphausiids (rho = -0.8). Age 3 shrimp is significantly correlated 334 with age 2 shrimp the previous year and negatively correlated to the temperature at the 335 sampling time (Vardø in March). Simultaneously, the survival of 3 year old shrimp, as 336 previously seen for age 1 shrimp, seem to be good when the young capelin thrives. 337 338

The principal components 1 and 2 contribute 92%, 79% and 70% to the correlation between 339 factors selected for the PCA ordination to be fitted to the distribution of age 1, 2 and 3 shrimp 340 (Table 3). The generalized additive model has the best fit to the age 1 and age 2 shrimp while 341 the deviance explained by the model for age 3 is only 16% (Table 3). The density of shrimp 342 of age 1 and 2 is projected on the respective PCA ordination plots (Fig. 8). In the PCA plot 343 the isolines show how the recruitment, of age 1 and age 2 change along the 3 most important 344 factors. The PCA for age 1 and age 2 visualizes how the recruitment is very dependent on 345 SSB and temperature. While the presence of young cod and euphausiids reduces shrimp 346 recruitment. Fig. 8 347

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349 **4. Discussion**

When evaluating the effect of environmental factors on shrimp recruitment, the ambient temperature is the most obvious factor to study. The NAO and the ice index were included as they may represent the large scale climate variability of the Barents Sea. The NAO influences on the Barents Sea in several ways. It may have an effect through changing the wind field, thereby changing the position of the Polar Front and the ice edge, and by increasing the northward flow of Atlantic Water in the Barents Sea. This response is direct with no lag as it is related to the wind conditions at the time. The NAO has also an indirect effect as it often

cause changes to the temperature of the Atlantic Water in the Norwegian Sea which is 357 subsequently advected into the Barents Sea. Due to this a lag between the NAO and the 358 Barents Sea temperatures has been identified by several authors (e.g. Furevik, 2001). When 359 using the ocean temperatures this advected, lagged signal is already in the time series, and 360 there will therefore be no additional information available by lagging NAO and shrimp. 361 The influence of the NAO is strongest when it is in a well defined positive or negative phase 362 with several high- or low-index years following. Prior to 2001 the NAO index had well 363 defined positive or negative phases, while since 2001 it has been low and irregular (Fig. 2). 364 365 This is probably the cause for the poor correlation between the NAO and the other parameters after 2001. The age 1 and 3 shrimp show no correlation with NAO while the age 2 shrimp is 366 significantly correlated with this index (Table 2). Age 2 shrimp also show a significant 367 correlation to the Vardø and Hopen temperatures and as the NAO and these temperatures are 368 significantly correlated (0.88 for NAO and Hopen), we consider this a temperature effect. 369

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The best recruitment index for age 1 shrimp is received by using the juvenile bag. However, 371 the time series is short and because age 2 shrimp caught in the cod end is correlated to the 372 373 shrimp caught in the juvenile bag and provides a longer time series they are considered the best recruitment measure (Fig. 4). Due to high and variable natural mortality from age 1 to 374 age 3, age 1 shrimp is not the best recruitment indicator to be used for fishery prognosis. The 375 index of age 2 shrimp is significantly correlated to available abiotic and biotic factors whereas 376 this correlation is weaker for the index of age 3 shrimp (Table 2 and Table 3). We consider 377 age 2 shrimp to be the best indicator of recruitment as has also been concluded for shrimp 378 stocks off West Greenland, Iceland and elsewhere(Wieland, 2004; Skuladottir, 1990). 379 380

The index of 3 year old shrimp from the cod end is hard to predict as the effect of several abiotic and biotic factors co-occur over time and the correlation to these factors get weak. A better correlation might have been received if the environmental factors (e.g. temperature and

cod) would have been integrated over the last three years before running the correlation
analysis. However, the index of age 3 shrimp gives an indication on the recruitment to the
shrimp fishery the next year, and may be used as a recruitment index in future assessment
work. A problem using this index is however, that a fraction of this age group is caught in the
commercial trawl and may have been exposed to fishing mortality.

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One year old shrimp are significantly correlated to spawning stock biomass, and the relation 390 is almost linear (Table 1, Fig. 7). Even if the Cushings match/mismatch hypothesis, effective 391 392 for the larval phase, may explain some of the inter annual variability in shrimp recruitment (Ouellet et al., 2007), the winter temperature and cod predation seem to be of essential 393 importance as they together with the SBB stand for more than 97 % of the deviance explained 394 by the GAM function (Fig. 8, Table 3). Crustaceans (krill, amphipods and shrimp) seem to be 395 the dominant prey of 1 year old cod, composing up to 40-80% of their diet. In some years 396 Pandalus borealis made up for 30-37% of the Total Fullness Index (Dalpadado and Bogstad, 397 2004). According to Dalpadado and Bogstad (2004) the cod reduces its crustaceans consume 398 at age 2 and moves to fish prey at age 3, consequently the negative correlation with shrimp 399 400 recruits gets weaker with shrimp and cod age (Table 2).

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The positive correlation between shrimp and the winter temperature may be a direct effect of 402 403 decreasing natural mortality, as temperatures below -1 °C are known to result in reduced abundance and temperatures below -1.6 °C result in extinction of shrimp (Smidt, 1981; 404 Wieland 2005). However, the positive correlation between one year old shrimp and 405 temperature and capelin may be related to food competition and overlap between shrimp and 406 capelin as also shrimp feed on Calanus finmarchicus (CI-CVI) (Harvey and Morrier 2003). 407 During winter the capelin is distributed according to the position of the Polar Front, so during 408 cold winters the capelin is concentrated in the Hopen Deep while during warm winters it is 409 spread over a larger area (Gjøsæter, 2008). In warm years, the distribution of capelin in 410

411 summer has a more northerly distribution, north and east of the Hopen Deep, than in cold years. Consequently there is less overlap between shrimp and capelin in warm years than in 412 cold years both during summer and winter. Although capelin by far is controlling the 413 abundance of copepods (Dalpadado and Skjoldahl 1996; Gjøsæter et al. 2000), less overlap in 414 warm years leaves more C. *finmarchicus* available for the shrimp in the Hopen Deep. This 415 may explain the negative, although not significant, correlation (-0.45) between one year old 416 shrimp and zooplankton. Dalpadado et al. (2003) found correlation coefficients 0.57–0.72 417 between temperature and capelin biomass. The positive response of one year old shrimp and 418 419 capelin to high temperatures may thereby be the result of less overlap and consequently less food competition. 420

421

The 2 year old shrimp show a positive response to temperature (Table 2). This relationship is 422 common when studying fish recruitment in the Barents Sea. Abundance estimates of 0-group 423 cod exhibit a close relationship with sea temperature variability at the Kola section (Nilssen et 424 al., 1994). Covariability in early growth and year class strength of cod, haddock and herring is 425 explained by their common positive response to temperature (Ottersen and Loeng, 2000). The 426 striking negative correlation with euphausiid biomass may have several explanations. The 427 dominating euphausiids in the Barents Sea are *Meganyctiphanes norwegica*, a boreal species 428 found in Atlantic water, and *Thysanoessa inermis*, *T. raschii* and *T. longicaudata* having an 429 430 arctic boreal distribution (Drobysheva et al., 2003). M. norwegica reaches the length of 45 mm and is primarily a carnivore aiming for copepod prey, including overwintering Calanus 431 *spp.* (Dalpadado et al., 2003; Kaartvedt et al., 2002). *Thysanoessa ssp.* is smaller (25–35 mm) 432 and has been classified as herbivores although T. raschii seems to switch to detritus feeding in 433 winter (Hopkins et al., 1989, Drobysheva et al., 2003; Pedersen et al., 2008). The age 2 434 shrimp and *M. norwegica* have the same size and are likely to compete for the same food 435 source, as juvenile shrimp are known to be active feeders and obtain most of their food, 436 euphausiids and copepods, from the macro plankton (Berenboim, 1981; Wienberg, 1980; 437

Wienberg 1982). In winter shrimp and *M. norwegica* may crop on overwintering copepods 438 that may be a scarce food source in deep water also in the Barents Sea (Kaartvedt et al. 2002). 439 As indicated before the mortality in the second year seems to be high, especially in years after 440 good recruitment. The reduced food supply may be a limiting factor not only when 441 euphausiids are abundant, but also when density of 2 year old shrimp is high. This increase in 442 density-dependent mortality with age is illustrated by the change in the shape of the Ricker 443 curve with age (Fig. 7). This is probably caused by intra-specific competition due to 444 limitations in available prey in combination with predation. 445

446

The recruitment indices are influenced by variable natural mortality rates and catchability 447 (Hannah, 1993). Although the spawning stock number was high in 1996 to 1999 the 448 recruitment of age 2 shrimp did not show an increase in the same manner as seen previously, 449 probably as a result of increased natural mortality due to cod predation (Fig. 3). Cold 450 temperature conditions in the Hopen Deep as observed in 1987, 1996 and 2003 (Fig. 2) 451 increases the natural mortality of 1 and 2 year old shrimp either directly, or indirectly as the 452 habitat overlap with cod probably increases as female shrimp and larvae move along 453 temperature gradients (Rasmussen et al., 2000), and get a more southern distribution as they 454 escape the cold water. Pedersen et al. (2003) showed by a transport model that high numbers 455 of larvae settled in the area around Bear Island, that also serves as nursery area for cod 456 (Skogen et al., 2007). The young cod feed on the young shrimp causing high natural mortality 457 and low abundance of age 2 shrimp off Bear Island (Fig 6). 458

459

The absence of the 1996 year class in the survey as 3 year olds in year 1999 is thereby caused by several co-occurring factors, the low spawning stock number and the distribution of the spawning females in 1996, temperature conditions and cod consumption. Similar environmental conditions may explain the low abundance of the 1987 and the 2001 year class at age 3 (Fig. 5). Wieland (2005) concluded that a moderate increase in temperature from 1°C

to 3°C above a lower threshold of optimal range in the northern regions off West Greenland
extended the favourable distribution area for shrimp and recruitment of age 2 shrimp. A
similar temperature reduction was observed in the in the Hopen Deep in the northern Barents
Sea in 1987, 1996 and 2003 (Fig. 2). As seen here for shrimp, studies on cod recruitment in
the Barents Sea indicate that the North East Arctic cod is also affected by temperature through
a variety of pathways (Hjermann et al., 2007)

471

The spawning stock biomass seem to have a significant effect on the number of age 1 and age 2 shrimp but this correlation is not detectable for age 3 shrimp, probably because the effect of temperature, predation by 1 and 2 year cod and euphausiids governs the development of the year classes (Table 1, Fig. 7). So, even if the female biomass may vary with a factor of 5 between years, a low number of female shrimp does not alone cause low recruitment.

477

Even if the larval face is important (Quellet et al., 2007), the year class strength does not
seem to be established during the larval stage as has been claimed by Shumway et al. (1985).
The density of 1 year old shrimp is directly dependent on SSB the previous year and is further
influenced by cod predation after the larvae settled.

482

Environmental fluctuations rather than changes in spawning female biomass are the primary 483 484 causes of shrimp stock fluctuations (Hancock, 1973; Garcia 1983). Therefore approaches including an environmental factor e.g. natural variability, temperature have been suggested 485 and three dimensional interpretations have been proposed (Garcia, 1983; Hannah, 1999). This 486 is supported by our study where the ambient factor temperature, and the biotic factors 487 spawning stock biomass, cod age 1 and 2 as predators and euphausiids as competitors all 488 affect the recruitment significantly. When modelling the recruitment of shrimp in the Barents 489 Sea the spawning stock number, the annual mean Vardø temperature, the number of 1 and 2 490 year cod, and the euphausiid abundance the previous year should be included. 491

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664 Figure text:

665

Fig.1. A schematic description of the circulation of the Barents Sea. Arrows show the current
of Atlantic water (dark grey) and Arctic water (light gray) (after Aure et al., 2000). Main
survey areas are East Finnmark (A), Tiddly Bank (B), Thor Iversen Bank (C), Bear Island
Trench (D), Hopen Deep (E) and Bear Island (F). The Hopen, the Vardø-North and the Kola
section (stippled lines) are shown.
Fig. 2. Annual mean, January and March temperature in the Vardø section (50–200 m),

summer temperature (August–September) in the Hopen section (50–200 m) and Kola section

annual mean (0–200 m), bottom temperature in the Hopen Deep (area E in Fig. 1), NAO

winter index and ice index. A low ice index corresponds to heavy ice conditions and viceversa.

677

Fig. 3. Zooplankton biomass, index of euphausiid abundance (updated from Drobysheva etal., 2003) and number of capelin, cod age 1, age 2 and age 3.

680

Fig. 4. Two year old shrimp caught in the juvenile bag attached to the underbelly of the
survey trawl plotted against 2 year old shrimp caught in the cod end, 1995–2004.

683

Fig. 5. Recruitment indices for 2 and 3 year old shrimp (standardized to the mean) and biomass of spawning females (SSB) sampled in the cod end each year. Recruitment indices for 1 year old shrimp (standardized to the mean) sampled in the belly bag since 1995. The indices represent the whole Barents Sea.

688

Fig. 6. Number of recruits as 2 year old shrimp by each sub area (A–E) in Fig. 1.

691	Fig. 7.	Number of	recruits at ag	e 1 (1994–20	003 year	class), at	t age 2 ((1989–	2002 ye	ear cla	ass)
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- and at age 3 (1986–2001 year class) plotted against spawning stock biomass for
- 693 corresponding year classes. Linear regression and the Ricker stock-recruitment model y = a *
- $x \exp(-b x)$ are fitted, and the parameter values received are given in Table 1.

695

- ⁶⁹⁶ Fig. 8. Spline surfaces of shrimp at age 1 (No. ^x 100 nm⁻²) upper panel and at age 2 (No. ^x 1000
- ⁶⁹⁷ nm⁻²) bottom panel, fitted to principal component ordinations of selected environmental
- 698 factors. The years plotted refer to the year-class in both panels.

Table 1. Parameter values (*a* and *b*) with respective standard error for linear regression

through origin and the Ricker curve. Adjusted R–squared, the degrees of freedom and,

702 P-value for F-test and AIC values are given.

	Age	e 1	Age	2	Age 3		
	Linear	Ricker	Linear Ricker		Linear	Ricker	
а	4.29	1.65	5.71	4.22	22.43	74.61	
st.							
error	0.67	1.36	0.77	2.57	2.99	23.69	
b		-0.35		-0.09		0.39	
st.							
error		0.22		0.17		0.12	
R ²							
(adj.)	0.80	0.78	0.79	0.74	0.77	0.76	
d.f.	9	12	13	12	15	14	
Р	<0.001	ns	<0.001	ns	<0.001	<0.01	
AIC	64	105	101	103	136	153	

704

706	Table 2. Spearmann rank correlation between shrimp at age 1, 2 and 3, and survival (Age
707	1/SSB, Age 2/SSB and Age 3/Age 2) and explanatory abiotic (January and March
708	temperature Vardø section the same year, mean annual temperature Vardø section, the Hopen
709	section and bottom temperature from the Hopen Deep, and the the NAO the previous year)
710	and biotic factors (capelin, euphausiids and cod age 1 and 2). SSB is lagged to the year of
711	recruitment of each age group. Correlation factor (rho) is given only for significantly
712	correlating factors (P< 0.05). Values given in parenthesis turned out not to be significant
713	when correction for autocorrelation was conducted.

	Age 1	Age1/SSB	Age 2	Age 2/SSB	Age 3	Age 3/Age 2
Sampling years	1995-2004		1991–2004		1989–2004	
Age 2 shrimp					0.43	
SSB	0.90		(-0.46)			
Temp. Vardø			0.75	0.55		
Temp. Vardø J.	0.77	0.90				
Temp. Vardø M.		0.70			-0.51	
Temp. Hopen			0.62	0.51		
Temp. Hopen b.			0.55			
NAO			0.58	0.48		
Capelin	0.63					0.49
Euphausiids			-0.81	-0.52		
Cod 1 year	(-0.60)	-0.63				
Cod 2 years		-0.62				

Table 3. Contribution of eigenvalues for PC1, PC2 and PC3 received from the Principal

717 Component Analysis and adjusted R^2 values and deviance explained from the Generalized

718 Additive model.

719

	Age 1 shrimp	Age 2 shrimp	Age 3 shrimp
Principal Component Analysis:			
Contribution to correlation			
PC1	70.8 %	63.6 %	47.0 %
PC2	21.1 %	16.2 %	24.1 %
PC3	5.7 %	14.3 %	22.3 %
Generalized Additive Model :			
R^2 (adj.)	0.92	0.56	0.04
Deviance explained	97.6%	62.6%	16.6%