Liver energy, atresia and oocyte stage influence fecundity regulation in Northeast Arctic cod

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ABSTRACT: Marine ecosystems are changing; global warming-induced increases in water temperatures and fishing have caused truncated age structures and small sizes at maturity in many stocks. This may affect both populations’ total reproductive output and the link between population demography and recruitment, yet detailed information on fecundity regulation is generally lacking for marine fishes. We therefore examined associations between liver energy, oocyte stage, leading cohort oocyte size \( (LC_{20}) \), atresia and fecundity for the Northeast Arctic cod \((Gadus morhua)\) from 2006 to 2008 in a comprehensive field and laboratory study. The relationship between the relative liver size (hepatosomatic index, HSI) and specific liver energy content was best described by an asymptotic curve, increasing rapidly at first, then levelling off at HSI > 6%. \( LC_{20} \) increased towards the spawning season, but was also positively associated with total length. At present there is thus a tendency towards larger NEAC females spawning earlier. The incidence of atresia was highest during the advanced yolk granule stage. Only 1% of females that reached an \( LC_{20} > 300 \mu m \) absorbed all oocytes and thereby aborted spawning. Potential fecundity showed a parabolic relationship with \( LC_{20} \), peaking around 614 \( \mu m \), i.e. approximately on February 1st, and was positively associated with weight, liver energy and, presently, age. In summary, NEAC females that start vitellogenesis will likely spawn. Atresia and fecundity down-regulation appears only to become pronounced close to spawning. Finally, the size-dependent spawning time, which appears to have emerged in the stock recently, may alter the link between population demography and recruitment.

KEY WORDS: \( Gadus morhua \) · Fecundity regulation · Atresia · Liver energy · Size-dependent spawning time
Marine ecosystems are also changing in ways that may affect allocation patterns. Fishing has severely depleted commercial stocks (Hutchings & Reynolds 2004), and common denominators of contemporary stocks are a truncated age structure and smaller size at maturity compared to historic data (Trippel et al. 1997, Marteinsdottir & Thorarinson 1998, Murawski et al. 2001). Although the importance of an in-depth understanding of the biological processes affecting maturation and thereby indirectly recruitment is acknowledged, essential information is lacking for most marine species.

In terms of fecundity styles, fish can be broadly distinguished into 2 categories: indeterminate and determinate spawners (Hunter et al. 1992, Murua & Sabarido-Rey 2003). Indeterminate spawners recruit new oocytes throughout the spawning period, whereas determinate spawners recruit a finite number of oocytes to the maturing pool prior to spawning. For fish employing the latter strategy, such as the batch-spawning Atlantic cod Gadus morhua L., only a down-regulation of potential fecundity is possible after the recruitment of oocytes is completed. This occurs through the process of atresia, i.e. the reabsorption of vitellogenic oocytes (Woodhead & Woodhead 1965, Kjesbu et al. 1991, Øskarsson et al. 2002).

The Northeast Arctic cod (NEAC) is now the largest remaining cod stock in the world (ICES 2008). Although NEAC has been the object of close scientific scrutiny, detailed studies on individual fecundity, energy reserves, energy flow or atresia are scarce. Using field samples, Thorsen et al. (2006) found that potential fecundity was positively correlated with female length and condition and decreased due to atresia in the final weeks prior to spawning. This study only included data from late in the maturation cycle (mid-February to mid-March) and was limited to data on weight and length. Thus, this type of sampling design cannot describe allocation patterns in the earlier part of the maturation cycle or supposed peak fecundity (Kjesbu 2009). Neither did this analysis incorporate liver energy as a proxy for fecundity, which is closely linked to egg production in NEAC (Marshall et al. 1998, 1999). Recently, Skjøraasen et al. (2009) examined mechanisms regulating oocyte growth, atresia and spawning omission of NEAC in a laboratory experiment from early August to late January. It was found that females whose leading cohort oocyte size advanced beyond 300 µm spawned and that spawning females started oocyte maturation from September to early November, whereas the transition from the cortical alveoli to the yolk granule stage (Tyler & Sumpter 1996) occurred around 400 µm. However, there is a clear need to validate these laboratory results with detailed field investigations.

We therefore present data on NEAC collected in a comprehensive field and laboratory study, where individual data on weight, length and age were combined with detailed analyses of fecundity, oocyte stages, the incidence of atresia and liver energy content. This allowed us to investigate allocation and oocyte development patterns on a fine scale hitherto not possible.

**MATERIAL AND METHODS**

**History of fish.** Female NEAC were collected using commercial or research vessels from autumn 2006 to spring 2008 (Fig. 1, Table 1). To examine potential spawners we specifically targeted cod >59 cm in total length (Skjøraasen et al. 2009). Aboard the vessels all cod were measured for total length, 1 cm below, i.e. a female of 60.5 cm will be given the length 60 cm, and whole body, liver, gonad and gutted weight (1 g below). A subsample of the gonad was also stored in neutral buffered formaldehyde (3.6%) for later laboratory analyses of oocyte sizes, fecundity and histology. Otoliths were collected for ageing and stock classification (Stransky et al. 2008). In 2007 to 2008, the livers of 164 female cod, which also included samples of non-maturing fish, were frozen at −20°C immediately after collection for subsequent chemical analyses of liver energy content (see below). Further information of the sampling period, samples obtained and specific analyses performed is given in Table 1.

**Proximate analysis of liver energy content.** The liver was partly thawed and 3 tissue pieces of approximately 1 g each were excised and analysed for dry weight content after one day at 105°C followed by calculation of mean and coefficient of variance (CV) (CV = 100 × SD × mean−1) for each fish. Likewise, total fat content (wet weight based) of the same triplicates was found gravimetrically with ethyl acetate as extraction solvent (Lauter & Trams 1962). In both cases the CV was typically around 1 to 2% and maximally 5%.

**Oocyte measurements.** All samples were subjected to digital image analyses (Thorsen & Kjesbu 2001). This method uses the contrast between vitellogenic and previtellogenic oocytes in relation to the set background to specifically select and measure the diameter of vitellogenic oocytes. Only females that could be (1) measured in the automated analyses, i.e. possessed vitellogenic oocytes, and (2) classified as NEAC based on their otolith shape (Rollefson 1934), were included for any further analyses, which left a total sample size of 379 cod (Table 1). For each measured sample, the size of 200 developing oocytes was reported. From these data the average size of the leading cohort oocytes (LC20) (n = 20, average of the largest 10% of the oocytes) was calculated, since this is indicative of
the time to start of spawning (Kjesbu 1994). Potential fecundity was calculated as:

\[ F_p = 2.139 \times 10^{11} \times OD^{-2.7} \times OW \]  

where \( F_p \) is potential fecundity, \( OD \) is average vitellogenic oocyte diameter, estimated by the image analyser, and \( OW \) is ovary weight (g) (Thorsen & Kjesbu 2001).

**Histological analyses.** The ovarian samples were prepared for histology using standard protocols for resin embedding (Technovit® 7100) producing 4 μm sections stained with 2% toluidine blue in 1% sodium tetraborate buffer. Under the light microscope, the presence or absence of atresia was noted. If yolk granules (YG) were found, around 150 YG (vitellogenic) oocytes were examined for atresia. A similar type of estimation was done for the cortical alveoli (CA) stage. Atretic cells were classified as either in the alpha-stage, containing yolk, or the beta-stage, without yolk (see Hunter & Maciewicz 1985). Females were then grouped under 1 of the following 3 categories (1) no atretic oocytes, (2) some atretic oocytes present, (3) all oocytes atretic, i.e. the females were aborting spawning. This ‘rough’ estimation of atresia was considered adequate for the scope of the present study, as the tracking of fecundity should implicitly reflect the actual level of atresia (Kjesbu 2009).

**Data analyses. Liver energy:** As noted above, our chemical analysis was limited to total dry weight content, i.e. the sum of lipids, proteins and glycogen, and lipid dry weight content. Based on the results of Lambert & Dutil (1997) we assumed the glycogen content to increase linearly from 0 to 5% of the total liver wet weight in the HSI (hepatosomatic index; \( HSI = 100 \times \frac{\text{Liver weight}}{\text{Total body weight} - \text{Gonad weight}} \)) range 0 to 6. Hence, for HSI values >6 we assumed the glycogen content to be 5% of the liver wet weight. The remaining non-lipid wet weight was considered to be proteins. A priori an exponential increase to a maximum value in lipid content with HSI was expected (e.g. Lambert & Dutil 1997). To calculate the specific liver energy content (SLEC, in kJ g\(^{-1}\)) of liver tissue at different HSI we used the following equivalences of SLEC (y) for different compounds: glycogen, 17.1 kJ g\(^{-1}\); protein, 23.6 kJ g\(^{-1}\); and lipids, 39.5 kJ g\(^{-1}\) (Kleiber 1975), against HSI (x). For the latter regression analyses we first examined if the assumption of normally distributed errors and constant variance was met. If not, we then tested for the potential influence of outliers by using Cook’s distance, leverage and standardized residuals (Venables & Ripley 1999).
Finally, we aimed to establish a biologically realistic statistical model for the explanatory variables of potential fecundity and liver energy content. It is known that potential fecundity is affected by female size and some metric of relative liver energy content, i.e. $\text{HSI} \times \text{SLEC}$, was estimated with a stepwise backwards multiple regression. Time of year was coded in the following way: the earliest date for which vitellogenic oocytes were found was given the value 0, e.g. September 3, and later passed since this date, e.g. October 1 is allocated the value 1. 

**RESULTS**

**Liver energy**

The estimated liver protein content varied between 0 and 20% of the liver wet weight. The corresponding measured fat content varied between 30 and 80% and was positively associated with HSI (Fig. 2a, $F_{1,163} = 162.6$, p < 0.0001, $R^2$-adj = 0.49). This relationship and the one between SLEC and HSI was best described by an asymptotic power curve (Fig. 2b, $F_{1,163} = 247.8$, p < 0.0001, $R^2$-adj = 0.60). When including all data, the assumption of normally distributed errors was not met. Five outliers were identified; each had a standardized residual >2.5 (Fig. 2b). After removal of these outliers all assumptions were met. The equation given in Fig. 2b was used to calculate the specific and total energy content for all individual fish referred to below.
Gonadosomatic index (GSI) and \( LC_{20} \) increased as the spawning season approached (Fig. 3a,b). In the latter case, the stepwise backwards multiple regression yielded a significant effect of both time of year \((B = 2.59, p < 0.0001, \text{i.e. leading cohort oocyte size increased by about } 26 \, \mu m \text{ in } 10 \, d \text{, Table 2})\) and female length \((B = 1.40, p < 0.0001, \text{Table 2})\). For comparison, on February 1, a 70 cm female’s expected \( LC_{20} \) was 610 \( \mu m \), whereas a 90 cm female’s \( LC_{20} \) was 638 \( \mu m \), i.e. a 20 cm size difference in length equated to 10 to 11 d difference in the predicted onset of spawning. There was no effect on \( LC_{20} \) of either \( HSI \times SLEC \) or residual condition. Time of year and length explained 72% of the variation in \( LC_{20} \).

The main transition from the CA to the YG stage occurred around 400 \( \mu m \) (Fig. 4). In the population, the CA stage dominated in September and October, then decreased sharply in November and by January the CA stage was virtually non-existent (Fig. 5).

Atresia

The prevalence of atresia was low early in the sampling period, but increased more or less linearly during maturation, and in February 34% of all sampled females possessed atretic oocytes (Fig. 5). Females with atretic oocytes had lower liver energy content in
November ($F_{1,46} = 7.95$, $p < 0.01$), but not significantly so between December to February (Table 3). Residual condition was not associated with the incidence of atresia in either of these months. Even so, values for both our metric of liver energy content and residual condition tended to be consistently lower in females with atretic oocytes (Table 3). The ANOVAs could not be performed for other months due to insufficient samples. Only 2 (1.01%) of the 198 fish examined for intensity of atresia were deemed to undergo complete reabsorption and, thereby, spawning omission.

**Energy content, fecundity and growth**

The total liver energy content (TLEC) was significantly negatively correlated with LC$_{20}$ (Fig. 6), and positively correlated with gutted weight and residual condition (Table 4). There was also a significant effect of season (Table 4) as NEAC had higher liver energy content in 2007–2008.

Fecundity varied greatly from a few hundred thousand to more than 15 million vitellogenic oocytes in the most fecund females (Fig. 7a). The most parsimonious model for fecundity included LC$_{20}$, (LC$_{20}$)$^2$, gutted weight, age and relative liver energy content, which accounted for 78% of the observed variation in fecundity (Fig. 7b, Table 4). In addition to gutted weight fecundity was also clearly positively associated with both relative liver energy and age. For comparison, at an LC$_{20}$ of 600 μm, a 6 yr old 5 kg female, with an HSI of 10 had an estimated fecundity 13.0% higher than a female of the same age and gutted weight with an HSI of 2, i.e. an estimated 3.07 million compared to
2.67 million vitellogenic oocytes (Fig. 7b). On the other hand, an 8 yr old female, with a gutted weight of 5 kg and a HSI of 10 produced an estimated 14.4% more oocytes than the 6 yr old female with the same HSI, i.e. 3.52 million compared to 3.07 million oocytes (Fig. 7b). The LC20 at which maximum fecundity occurred was estimated to be 614 μm (95% CI 566 to 662 μm), i.e. around February 1 for a 70 cm cod. We found no differences between 6, 7 and 8 yr old NEAC in body length growth (all p-values > 0.07, data not shown).

DISCUSSION

The present study sheds light on several important aspects of oocyte recruitment and fecundity regulation in the determinate batch-spawner, the NEAC. The dedicated, exploratory analyses on fecundity, atresia and liver energy content provides fine-scaled data that allows us to make inferences about life-history traits difficult to target.

Oocyte recruitment and energy flow

In teleost fish the main yolk protein, vitellogenin, is synthesized from proteins and lipids (Tyler & Sumpter 1996). In lean fish such as cod, the main storage organ for lipids is the liver (Takama et al. 1985, Kjesbu et al. 1991, Lambert & Dutil 1997). Due to its direct link to oocyte recruitment and growth, liver energy has therefore been suggested as a proxy for egg production (Marshall et al. 1999, 2000). Liver energy content is not solely a function of liver size, as water content varies substantially with relative liver size (Lambert & Dutil 1997). It is therefore necessary to establish robust relationships between liver energy content and easily measurable metrics of condition. Lambert & Dutil (1997) did so for the Gulf of St. Lawrence cod and their relationships have since been applied to other cod stocks (e.g. Skjæraasen et al. 2006), including NEAC (Marshall et al. 1999). The present study indicates that the specific energy content is somewhat higher in NEAC compared to Gulf of St. Lawrence cod for a given liver size, although the liver energy content shows the same general relationship with the hepatosomatic index (HSI). For comparison, the specific energy content of a liver with an HSI of 3 was estimated to be 21.0 kJ g⁻¹ for NEAC compared to 17.8 kJ g⁻¹ for Gulf of St. Lawrence cod. Our estimated asymptotic maximum

Table 4. Gadus morhua. Estimated values and significance levels of all parameters included in models for total liver energy content (TLEC) and potential fecundity (Fp). GWeight: Gutted weight; HSI × SLEC: see definition in Table 1

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Explanatory variable</th>
<th>Estimate</th>
<th>t</th>
<th>p</th>
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</thead>
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<tr>
<td>TLEC (R²-adj = 0.60)</td>
<td>Intercept</td>
<td>1114</td>
<td>1.00</td>
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<tr>
<td></td>
<td>(LC20)²</td>
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<td>−4.50</td>
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<td>GWeight</td>
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<td>21.13</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>Residual condition</td>
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<td>3.37</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Season</td>
<td>2328</td>
<td>4.38</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Fp (R²-adj = 0.78)</td>
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<tr>
<td></td>
<td>LC20</td>
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<tr>
<td></td>
<td>(LC20)²</td>
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<td>Age</td>
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<td>3.97</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>HSI × SLEC</td>
<td>1607</td>
<td>2.38</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>

Fig. 7. Gadus morhua. Relationship between potential fecundity and leading cohort oocyte size (LC20). (a) Total variation in potential fecundity plotted against LC20. (b) Predicted fecundity as a function of LC20 size for 3 'model' females, demonstrating the effects of differing liver energy and age calculated from the parameters given in Table 4. The 3 'model' females have a gutted weight of 5 kg each. (····) 6 yr old female, hepatosomatic index (HSI) of 2; (−−−) 6 yr old female, HSI of 10; (——) 8 yr old female, HSI of 10.
value of 28.6 kJ g\(^{-1}\) is also 3.7 kJ g\(^{-1}\) higher. Even so, the similarity of the response curves between these 2 studies (Fig. 2, Lambert & Dutil 1997) makes it unlikely that the results of past studies, applying the relationship of Lambert & Dutil (1997) to other stocks, would have been significantly altered by employing the present relationship instead. In both studies a lot of variation remained unexplained. The decrease in energy content with LC\(_{20}\) is in accordance with the utilization of liver energy for yolk production and reduced appetite during maturation (Skjæraasen et al. 2004). Our model predicts similar energy usage regardless of fish size. This is arguably not biologically realistic given that larger females produce more eggs and conversely should utilize more energy. It may be that larger females are able to replenish the energy used for oocyte development through feeding to a greater extent than smaller females, but such size-specific feeding has yet to be supported statistically (Michalsen et al. 2008).

Successful reproduction in marine pelagic spawners includes larvae finding sufficient food. NEAC larvae are connected to the spring *Calanus finmarchicus* nauplii peak production, i.e. what became known as the critical period (Hjort 1914) or match-mismatch hypothesis (Cushing 1974, 1975). We found that LC\(_{20}\) increased as the spawning season approached, yielding a positive relationship with time of year, but also with female length. LC\(_{20}\) is indicative of the start of spawning (Kjesbu 1994) and highly fecund, large females will thus to a larger extent be present during the early part of the spawning season. This could have positive consequences for successful reproduction and recruitment due to an increased temporal overlap with the zooplankton peak for the first-feeding cod larvae (Solemtdal 1997). However, this would probably be only one contributory factor to recruitment variability since many other factors can affect the survival of offspring from early life history to maturity and joining the adult population. More interestingly, no such link between female length and spawning time was found for NEAC in the 1980s (Kjesbu 1994). Ongoing research indicates that elevated water temperatures may create such a coupling between female length and spawning time due to different physiological responses at a large and small body size in agreement with general thermal tolerance theory (see Kjesbu et al. 2010 and references therein). Our observations tentatively agree with this as there has been a consistent trend towards higher temperatures in the water masses occupied by NEAC over the last decades (Ottersen et al. 2006). At present the length-based separation in spawning time is small, but, if temperature related, it may become more pronounced with expected water warming (IPCC 2007). This is an important topic for future research.

### Fecundity regulation

Fecundity showed a parabolic relationship with leading cohort size clearly illustrating the recently introduced concept of ‘down-regulation’ in the final months prior to spawning (Kennedy et al. 2009). Down-regulation likely occurs to some extent in all ovaries (Kjesbu 2009), but is accelerated in fish in poor condition (e.g. Kjesbu et al. 1991, Kennedy et al. 2007). We only found partial support for the latter explanation in the present study. Although both residual condition and liver energy content was generally lower in females with compared to females without atresia, this difference was only significant for liver energy content in November (Table 3). It may be that atresia in NEAC reflects temporal changes in energy reserves rather than energy reserves per se as has been shown in place *Pleuronectes platessa* L. (Kennedy et al. 2008). However, the estimated down-regulation (Fig. 7b) suggests that all females show atresia at some point in the later part of the maturity cycle. Thus, absence of atresia in several females is a likely indicator of low overall, but not necessarily zero, atresia levels in these females or, alternatively, that our sample was obtained before or after the atretic window of that particular fish. Interestingly, our estimated peak fecundity, LC\(_{20}\) = 614 μm, occurs quite late. This suggests that (1) the window for oocyte recruitment remains open until the LC\(_{20}\) are well into the YG stages, but perhaps more importantly that (2) atresia becomes pronounced only close to spawning. The latter is also supported by monthly examination of the frequency of females with atretic oocytes. Notably, complete reabsorption of oocytes only occurred in 1.0% of the maturing females. Thus it seems that with little exception, females that truly start vitellogenesis, i.e. LC\(_{20}\) > 300 μm, will spawn. This latter result, and the transition from the CA to the YG stage around 400 μm, closely mimics the results obtained on NEAC in the laboratory (Skjæraasen et al. 2009). Based on the sharp reduction in the frequency of females whose LC\(_{20}\) was at the CA stage from October to November, i.e. females that have fairly recently started maturation, it would seem that females that are to spawn in the coming spring join the maturing fraction before the end of November. This is in accordance with the suggested onset of maturation from previous studies on NEAC (Kjesbu 1994, 2010). Female fecundity was also positively associated with female size, i.e. weight, and our metric of liver energy content. In our field study we have not considered the effect of feeding, logically improving the energetic status of each female and thereby its liver energy content, complicating straightforward interpretations of the present data. It is well known from experiments on cod that there is an inverse relationship between body and gonad...
growth during vitellogenesis (Taranger et al. 2006). In light of the earlier mentioned generally reduced appetite at advanced maturity, the fecundity reduction at LC_{20} > 620 μm is likely due to insufficient energy resources to fuel the on-going oocytic yolk uptake in these long-migratory fish.

The present data indicate that older females produced more eggs than younger ones of similar size. Increased reproductive investment theoretically comes at the expense of growth (Roff 2002) and likely survival, as poor condition increases post-spawning mortality in gadoids (Dutil & Lambert 2000, Lambert & Dutil 2000). Thus, from a life-history perspective, younger females would be expected to invest less in fecundity and more in growth than older ones, whose future reproductive success is expected to be low (Houston & McNamara 1999, Clark & Mangel 2000, Roff 2002). We clearly find support for the former at present, but, arguably, not the latter as there were no growth differences between females 6 to 8 yr old. However, NEAC females of 6 to 8 yr are not particularly ‘old’ and it may be that a growth related trade-off would have emerged if we had had more samples from older females. Moreover, it may be that, if present for NEAC, the trade-off is more related to storage and thereby survival than growth per se. For the North Sea cod, similar patterns of age-dependent reproductive investment after controlling for fish size have been attributed to fisheries-induced genetic change, given that such patterns were not observed in the same stock in the late 1960’s (Yoneda & Wright 2004). Investigation of the IMR (Institute of Marine Research) time series of fecundity data collected at Andenes (69° 20′ N, 16° 06′ E) indicates that this age-dependent investment has been seen intermittently in the last decades for NEAC (O. S. Kjesbu unpubl.). Arguably, an age-dependent response to environmental conditions is therefore a more likely explanation for our results, but this could also be due to insufficient sample sizes in some years. Consequently, identification of the causes behind the presently observed allocation pattern should be further tested.

CONCLUSIONS

The results highlight a number of important processes for oocyte recruitment and fecundity regulation; females that reach a LC_{20} > 300 μm will, almost without exception, spawn. Atresia and down-regulation only becomes pronounced during the final period prior to spawning for maturing females. NEAC females now tend to be temporally structured in spawning time according to length, which may affect the link between population demography and recruitment. There were also indications of an age effect on fecundity, not reported earlier. Finally, we have established a liver energy model for the NEAC and used this successfully to predict fecundity and we have detailed the phenomenon of fecundity down-regulation. This was achieved by a comprehensive sampling programme on adult female cod in the Barents Sea covering all stages of oocyte growth, instead of the traditional programme limited to prespawners only.

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