Oceanographic flow regime and fish recruitment: reversed circulation in the North Sea coincides with unusual strong sandeel recruitment

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**INTRODUCTION**


The lack of successful applications of environmental predictors (e.g. physical and biological) to date may be partly due to the strong emphasis placed on the stock–recruitment (S-R) relationship (Ricker 1954), which assumes that recruitment variability (i.e. where
recruitment is understood as the number of young-of-the-year individuals at the life phase transition from larvae to juveniles; ICES 2018) is primarily explained by the biomass of the reproductively active segment of a stock, also termed the spawning stock biomass, and thus any environmental influence must therefore be secondary (Myers 1995, Myers & Barrowman 1996). However, modelled S-R relationships for individual stocks are often weakly supported by the data, due to high inter-annual variability in recruitment, in combination with relatively short time-series (Fogarty et al. 1991, Olsen et al. 2011, Szuwalski et al. 2015). Established correlations are therefore often quite weak, and tend to decompose or change over time (Myers 1998, Skern-Mauritzen et al. 2016).

The perception that recruitment is a function of the biological and physical oceanography dates back a century (Hjort 1914, 1926). Corollaries from this work have since given rise to hypotheses, such as the ‘match–mismatch’ hypothesis (Cushing 1975, 1990), which build on the realization that biological and physical processes might be complementary and emphasize the importance of matches in time and space between the early life stages and the availability of food and the ‘right’ physical conditions. In a recent meta-analysis, Vert-pre et al. (2013) found that although some stocks showed strong S-R relationships, net population increases for the majority of stocks were driven by environmental regimes.

Examples of significant correlations between recruitment and planktonic food resources are common in the literature (e.g. Beaugrand et al. 2003, van Deurs et al. 2009, Burrow et al. 2011, Kristiansen et al. 2011). The fate of fish larvae and food sources are to a large extent influenced by the same oceanographic features, and yet the local oceanographic processes are often neglected in these studies or replaced by higher-level climate indicators such as average temperature or teleconnection indices like the North Atlantic Oscillation (NAO, an index of the predominant air circulation patterns over the North Atlantic) (Arnott & Ruxton 2002, Solow 2002).

The purpose of the present study was to explore the potential for using advanced oceanographic model output in short-term forecasts of fish stock production (i.e. recruitment), which we investigated using a well-studied North Sea case. The ecologically and economically important lesser sandeel Ammodytes marinus stock on the Dogger Bank (DB) in the central North Sea is an example of a fish stock in which recruitment dynamics drive most of the population variability and catch opportunities. This stock is a typical ‘boom-and-bust’ type of stock (Schwartzlose et al. 1999), characterized by dramatic fluctuations and occasional unusually strong year-classes that sustain a large fishery in the following years. Hence, the ability to predict the coming of an exceptionally strong year-class is of major importance (de Moor et al. 2011). The recruitment of sandeel has been linked to both food abundance and climate indicators, such as sea surface temperature (SST) and the NAO (Arnott & Ruxton 2002, van Deurs et al. 2009, Régnier et al. 2017, Lindegren et al. 2018). With respect to food availability, in particular the temporal match–mismatch between occurrence of first-feeding larvae and the abundance of the 2 largest copepod species in the North Sea, Calanus finmarchicus and C. helgolandicus, has been stressed (Arnott & Ruxton 2002, van Deurs et al. 2009, Régnier et al. 2017). Simulation studies of hydrodynamic larval drift and circulation processes that affect larval growth have also been carried out (Christensen et al. 2007, Gurkan et al. 2012), but studies on the connection between local oceanographic events and inter-annual variability in sandeel recruitment do not exist.

The general flow regime of the North Sea can be described as a counterclockwise circulation produced by prevailing southwesterly winds, which occasionally reverses with easterlies. This occasional shift from the predominant counterclockwise circulation to other regimes can include a large-scale gyre flowing in the opposite direction over the whole North Sea or multiple smaller-scale gyres with alternating flow directions and even temporary circulation halts (Kauker & von Storch 2000, Sündermann 2003, Sündermann & Pohlmann 2011). Changes in the physical setting affect fish production, especially for forage fish that display boom-and-bust dynamics (Fréon et al. 2005). For example, upwelling events driving the production of Pacific sardine stocks off the coast of Central America (Rykaczewski & Checkley 2008) are strongly linked to certain types of ocean circulation patterns. Another example is the Norwegian spring spawning herring, where the physical conditions of the Norwegian Coastal Current determine survival and growth of the larvae as they drift toward the nursery grounds in the Barents Sea (Skagseth et al. 2015).

During a preliminary investigation, it was observed that the largest sandeel recruitment event on record coincided with a circulation shift in the northwest Atlantic during the winter of 1996, first mentioned by Greene & Pershing (2003) and later coupled to possible circulation reversals in relation to settlement dynamics of 2 species of flatfish in the North Sea (Lacroix et al. 2013, Tiessen et al. 2014). While many
studies have flagged hydrodynamics as playing various roles for sandeel recruitment involving different biophysical levels and larval life stages (Bernsten et al. 1994, Proctor et al. 1998, Gallego et al. 2004, Christensen et al. 2007, Gurkan et al. 2013), they all agree on the notion that at least the earliest life stages, or first-feeding larvae, should be considered as passive tracers whose fate is dictated by the ocean currents. Although sandeel larvae develop active swimming abilities before they metamorphose at a length of ~40 mm, it is assumed that they generally follow current surface flow until settlement (Christensen et al. 2008). The drifting phase lasts from 1–3 mo (i.e. February to April) (Wright & Bailey 1996, Conway et al. 1997, Jensen et al. 2003).

We therefore hypothesized that unusual strong year-classes of DB sandeel are associated with certain types of flow regimes during the early parts of the drifting period when first-feeding larvae are most abundant. Flow regimes were quantified using original output from an oceanographic model (HIROMB-BOOS), and the period of first feeding was examined from hatch dates estimated using larval otoliths. Subsequently, the oceanographic output was subjected to a cluster analysis to tease apart different types of flow regimes, before relating these to sandeel recruitment. To test the usefulness of flow regimes in short-term forecasting of strong year-classes, we used a probabilistic approach. This approach, also known as the risk-ratio analysis, is widely used in risk assessments (Wilson & Crouch 1987), but rarely applied in fisheries science (but see Mantzouni & MacKenzie 2010 and Skagseth et al. 2015). Nevertheless, it may be suitable for detecting mechanisms associated with infrequent extreme year-classes, as seen for our case stock. We applied this approach to the flow regime data that resulted from the cluster analysis, as well as to other environmental variables previously suggested to be influencing sandeel recruitment. This comparative approach not only allowed us to rank the performance of the different recruitment predictors, but it also provided a basis for discussing possible causal mechanisms.

**MATERIALS AND METHODS**

**Oceanographic data**

We inspected 3 papers (i.e. Kauker & von Storch 2000, Sündermann 2003, Sündermann & Pohlmann 2011) in order to identify an oceanographic time-series of flow regimes that captured the essentials of the Central North Sea water circulation system as well as the transport of water over the DB, where the most important sandeel fishing grounds are located. Based on these studies, we settled on a transect that followed the 50 m depth contour, along the northern edge of the DB, stretching from the centre of the North Sea to 60 nautical miles (n miles) from the UK coastline (0.0° E, 55.5° N to 4.0° E 56.0° N), the latter to avoid the influence of the Scottish coastal current (i.e. Brown et al. 1999) (Fig. 1). Based on the assumption that first-feeding larvae are passive drifters distributed in the water column (e.g. Christensen et al. 2007), hourly values of the net flow of surface water (0–15 m depth) were then derived by summing up 3D water fluxes from an oceanographic model (described in more detail below). Otolith-based estimates of hatch dates in the Shetland sandeel population, early-stage larvae captured over the DB by the Continuous Plankton Recorder (CPR), and peak larval survival estimated by individual-based modelling (Wright & Bailey 1996, Gurkan et al. 2013, Lynam et al. 2013) seem to indicate that February and March covers the most important period for first-feeding larvae; thus we chose to focus on these months in the data analysis (this assumption was tested later in the present study; see ‘Estimation of hatch dates’, below).

The oceanographic data were provided by the Danish Meteorological Institute (DMI) and are a validated hindcast (1990–2013) produced by the North Sea-Baltic Ocean Model (HBM Model) in the operational setup used by DMI (http://ocean.dmi.dk/models/hbm.uk.php). The model is a 3D ocean circulation model, where the vertical flow assumes hydrostatic balance and incompressibility of seawater, and horizontal transport is modelled using the Boussinesq approximations (Smagorinsky 1963, Canuto et al. 2002, Berg & Poulsen 2012). The horizontal resolution (grid spacing) was 6 n miles, which is a standard resolution used by DMI for coarse-scale applications, and has previously been used to model fish larval drift in the North Sea and water fluxes between the North Sea and the Baltic Sea (e.g. She et al. 2007, Gurkan et al. 2013, Ulrich et al. 2017). The main consequence of this relatively coarse grid spacing is the potential risk of failing to capture the full effect of small-scale eddies and the risk of exaggerating the wind effect. However, in the present study, we were only interested in the trends in net flow averaged across 300 km wide transects (i.e. the geographical scale of interest is 27 times larger than the model resolution), where small-scale processes are likely to be averaged out as fluctuations along the transect. The model was forced by atmospheric data from a numer-
ical weather prediction model, High Resolution Local Area Model (HIRLAM, http://hirlam.org/), with 10 m wind fields, sea level pressure, temperature, humidity, and cloud cover. At open model boundaries between Scotland and Norway, and in the English Channel, we used tide surges from a barotropic model of the North Atlantic (Dick et al. 2001). All other model variables at open boundaries were set to monthly climatological values. The freshwater daily river runoff data come from the operational hydrological model HBV run by the Swedish Meteorological Hydrological Institute (Bergström 1992) and observations made by German Bundesamt für Seeschifffahrt und Hydrographie. The hindcast period (1990–2013) was limited by the overlap of applied forcing data sets.

In the present study, we expressed the net flow of water across the transect as the signed (i.e. ±) accumulated length (km) of water that has moved through the transect since time 0, averaged across the surface of the transect (where ‘+’ indicates water movement from north to south and ‘−’ indicates the opposite movement, from south to north). Time 0 was either 1 February or 1 March, and the final dataset consisted of hourly values for all of February and March, respectively. To get from length of water to the volume of water, we must multiply by the dimensions of the transect surface area, which was 300 km (transect length) times 0.015 km (transect depth), and the hourly average transport rate can be derived from the volume at time $h$ minus the volume at time $h-1$. However, in relation to the present study, the length of water was found to be the most useful unit, as it offers a direct approximation of the scale of potential horizontal transportation of passively drifting biological particles (such as fish larvae or copepods).
Identification of flow regimes using principal component analysis and hierarchical clustering

To identify distinct annual and monthly flow regimes, we carried out a principal component analysis (PCA) on hourly values of the accumulating water transport across the selected transect. PCA is a useful statistical signal-processing technique to reduce the multi-dimensionality of datasets for pattern recognition. The PCA algorithm constructs principal components (PCs) obtained as linear combinations of the original variables, where the first PC is allowed to have the largest possible variation in the data, and the second PC is estimated under the assumption of being orthogonal to the first PC and to have the largest possible variance, and so forth for other additional PCs that might be important (see Abdi & Williams 2010 for details). In order to choose the correct number of PCs to be kept in the analysis, we used a scree test (Cattell 1966), which is a graphical detection of a bend in the distribution of the successive variance explained.

Data from February and March were treated in 2 separate PCAs, which assigned each year in the study period to clusters, each of which represented distinct hydrodynamic patterns (hereafter referred to as flow regimes). The Euclidean distances between years were computed from the projection of years on the PCs, and subsequently a hierarchical clustering analysis was conducted. We used the Ward linkage method (Ward 1963), which has a minimum variance criterion that minimizes the total within-cluster variance. For each step, we identified a pair of clusters which leads to a minimum increase in total within-cluster variance when merged. The PCA and the hierarchical clustering analysis were implemented in the R-packages ‘PTA-k’ (Leibovici 2010) and ‘ade4’ (Chessel et al. 2004).

Sandeel recruitment

As a measure of sandeel recruitment on the DB, we used the number of young sandeel in July (age 0) as estimated by the International Council for the Exploration of the Sea (ICES) for Management Area 1 (ICES 2016a). These recruitment estimates are produced by a stochastic multispecies stock assessment model (Lewy & Vinther 2004) and are essentially back-calculated in the model from the number of 1 yr old sandeel estimated by the model in the following year using the natural mortality for age 0 sandeel in the second half year (ICES 2016a). Prior to the update of recruitment time-series in March 2017, input to the stock assessment model used by ICES was changed. We therefore included recruitment time-series from before and after the 2017 update (ICES 2016b, 2018).

Estimation of hatch dates

To identify and confirm the important months (i.e. February/March; Gurkan et al. 2013, Lynam et al. 2013) of first-feeding larvae in the DB area, we derived hatch dates from sandeel otoliths taken from larvae caught on the DB in 2006, 2008, and 2009. All larvae samples were collected between 17 and 22 April. The larvae were caught with a 1 m Methot-Isaac-Kidd (MIK) net, then frozen at sea and later thawed and preserved in ethanol. Three MIK samples were randomly selected from each year (9 samples in total). From each sample, 20 larvae were sub-sampled using a stratified sampling procedure to cover the full length distribution. Otoliths were dissected from the larvae under polarized light, imbedded in thermoplastic cement, and then photographed under a microscope. Daily increment widths were measured from the central primordium to the otolith edge, but not all increments were visible. Hence, the age in days was determined by dividing the distance from yolk sac absorption to the otolith edge by the mean daily increment width and then adding 12 d to account for the yolk sac period (Wright 1993, Wright & Bailey 1996). The distance along the axis of fastest growth from the primary primordium to the point of yolk sac absorption was assumed to be 20 µm (Wright 1993). A linear age−length relation derived from the subsampled otoliths was then used to convert the length distribution in the original MIK sample to a hatch date distribution. Lastly, each hatch date distribution was corrected for natural mortality. Daily mortality rates ranging between 0.042 and 0.07 (Fujiwara et al. 1990, Régnier et al. 2017) have previously been assumed, and we therefore investigated the sensitivity of mortality. In order to make our results directly comparable to those of Wright & Bailey (1996), we adopted a mortality rate of 0.07, but we also included high (0.12) and low (0.01) mortality rates by simply increasing mortality by ~100%.

Quantifying and comparing the performance of different predictors of recruitment

Previous studies have reported positive relationships between sandeel recruitment and Calanus
copepods and negative relationships with SST and the NAO (Arnott & Ruxton 2002, van Deurs et al. 2009, Régnier et al. 2017, Lindegren et al. 2018). In the present study, we wanted to re-investigate all of these recruitment predictors together with our new oceanographic predictor. Information about the abundance of *C. finmarchicus* and *C. helgolandicus* were acquired from the CPR programme. These data were provided by the Sir Alister Hardy Foundation for Ocean Science (SAHFOS), and came as monthly indices calculated specifically for Management Area 1 that covers the DB (ICES 2016b). The NAO was downloaded as station-based monthly values from the Hurrell data base: https://climatedataguide.ucar.edu/climate-data/hurrel-north-atlantic-oscillation-nao-index-station-based. SST was downloaded from the Hadley Centre observational database as monthly values in a 1 × 1° grid. For the present study, we used the average SST for an area defined as 1−3° E and 54−55° N. Only data from February and March were included in the analysis to match the period of first feeding and the oceanographic data (see explanation above). One exception to this rule was NAO, where we used an early (mean NAO for the period from December to February) and a late winter index (mean NAO for the period from January to March) instead of monthly values.

In order to compare the different candidate recruitment predictors, we computed the risk-ratio (RR) as in Mantzouni & MacKenzie (2010):

\[
RR = \frac{G / (g + G)}{C / (c + C)}
\]

where *G* is the number of years where favourable environmental conditions coincided with good recruitment (i.e. unusually strong year-classes), *g* is the number of years where favourable conditions did not coincide with good recruitment, *C* is the number of years where unfavourable conditions coincided with good recruitment, and *c* is the number of years where unfavourable conditions did not coincide with good recruitment. In cases where *G* or *C* was 0, we replaced 0 with 1/24, since we have 24 observations in total.

Since the intention was to forecast the most extreme recruitment events rather than the stock trends, good recruitment was defined as recruitment that exceeded the mean of the time-series plus 1 SD. This approach suited its purpose and captured the distinct spikes in the recruitment time series, which are of particular importance in relation to managing boom and bust stocks (de Moor et al. 2011). Based on the previously mentioned reports of a positive relationship between sandeel recruitment and *Calanus* copepods and negative relationships for SST and NAO, we defined favourable conditions in relation to each of those predictors as values either above or below the time-series mean ± 1 SD. RR > 1 indicates that the probability of observing good recruitment when the environmental condition is expected to be favourable is higher than the probability of observing good recruitment when the environmental condition is unfavourable. Hence, high values of RR were considered indicative of high performance in relation to recruitment forecasting. We also applied the same RR approach to derive RR values for the opposite question; i.e. the relative probability of seeing less than good recruitment when environmental conditions are favourable (i.e. \( RR = [c/(C+c)]/[g/(G+g)] \)). In this test, low RR values were considered indicative of high performance. To distinguish between the 2 different applications of the RR equations, we used RR₁ and RR₂ to label the former and latter application, respectively. Finally, we obtained p-values by applying the chi-squared test of independence to a 2 × 2 contingency table

\[
\begin{bmatrix}
G & g \\
C & c 
\end{bmatrix}
\]

The p-values refer to the null hypothesis that the 2 categorical variables (environmental condition and recruitment) are independent.

**RESULTS**

The net flow of water across the selected transect (analysed for February and March) went from being almost consistently positive (i.e. running in a north−south direction) or negative (i.e. running in a south−north direction), through highly variable, alternating between positive and negative, to nearly stagnant (Fig. 2A,C). The classification of clusters was mainly driven by the overall direction of the net flow (positive or negative), but the temporal development of the trajectories was also important. Division of years into clusters was based on 2 PCs (PC1 and PC2, Fig. A1 in the Appendix, together explaining 93.0 and 94.5% of the total variation in February and March, respectively (Fig. 2B,D). These clusters captured certain types of flows: (1) unidirectional flow and relatively large negatively or positively signed net flow (advection), (2) bidirectional flows associated with negatively or positively signed low net flow (retention), (3) relatively weak or stagnant flow that suddenly shifts to a negatively or positively signed directional flow of higher net flow (shifting flow), and...
(4) relatively weak flow alternating between positive and negative flow centered around 0 (stagnant).

PC1 explained most of the variation (76.9 and 84.8% for February and March, respectively) and discriminated among the types of flows listed above, making a distinction between years of predominantly positive net flow and years of predominantly negative net flow, which represented 2 overarching clusters in both February and March (Fig. 3) and are hereafter referred to as negative and positive flow regimes. However, each flow trajectory was also associated with a temporal behaviour over the transect, for example whether a flow trajectory had a flow with a directional motion, either a continuous or oscillating motion over the transect, or a motion with close to 0 flow in any direction. This temporal behaviour was captured by the second PC2, explaining 16.1 and 9.7% of the variation in February and March, respectively (Fig. 2). The years also had correlations with total contributions to each PC that supported this (Figs. A2 & A3). The PCA identified 5 different types of flows in February, but only 4 in March, where the analysis failed to distinguish between advection and retention (Figs. 2 & 3).

The year 1994 differed from all other years (in February), as it went from almost stagnant to the highest continuous negative flow observed; thus it is essentially an intermediate between the ‘stagnant’ and ‘retention/advection’ flow types, which is also why the analysis places this year in its own sub-cluster labelled ‘shifting flow’ (Figs. 2 & 3, green). Another outlier was 1991 (March), characterized by an ex-
tremely stagnant flow and net transport close to 0 (Figs. 2 & 3, purple).

Also noteworthy was the decreasing trend in net flow in February, where all years (except 2000) assigned to the cluster of positive ‘retention’ flows appeared after 2002. Likewise, only 1 year after 2002 was assigned to the cluster of positive ‘advection’ flows (2012, the weakest among the advection years). Hence, it appears that a shift in the winter oceanography in February from an advection-dominated system to a retention-dominated system occurred around 2002 (Fig. 3A).

Good recruitment (i.e. recruitment > mean + 1 SD) was observed in 1994, 1996, 2001, and 2009 (Fig. 4). In February, all 4 years with good recruitment were assigned to the same overarching cluster consisting of negative flow regimes. However, only 1996 and 2001 were assigned to negative flow regimes in both February and March (Fig. 3). Furthermore, 3 out of the 4 good recruitment years were assigned to the same sub-cluster of negative ‘retention’ flows. Only 1994 was assigned differently and placed (as the only year in the February time-series) to a sub-cluster labelled ‘shifting flow’ (the flow phenomenon of 1994 is described in more detail above, this subsection).

According to the analysis of daily increments in larval otoliths, hatching was mainly confined to March. The earliest hatching occurred on Day 53 (22 February) and the last hatching on Day 83 (24 March). Mean time of hatching also varied between years (by ca. 10 d). Hatching in February was only observed in 2009, whereas in 2006 and 2008 all hatching took place in March.
place in March (Fig. 5). The sensitivity analyses showed that mortality did not have a substantial effect on the hatching distributions. A given percentage hatching was reached earlier with increased mortality, but the maximum difference between the low and high mortality scenario was 5 d (Fig. 5).

In the first RR test (RR1), the flow regime in February performed better than all other recruitment predictors. In fact, the RR1 values computed for the flow regime in February suggested that the probability of observing the combination of good recruitment and a negative flow regime was 243 times higher than the probability of observing the combination of good recruitment and a positive flow regime. In support of this result, the chi-squared test revealed a significant relationship between the distribution of recruitment and the distribution of flow regime in February (i.e. rejecting the null hypothesis of independence: χ² = 7.91, p = 0.005; Table 1). In comparison, the corresponding RR1 for the negative flow regimes in March and *Calanus finmarchicus* in February was only 5.00.

Table 1. Risk ratios (RR) and chi-squared test of independence for various sandeel recruitment predictors. *G*, *g*, *C*, and *c* correspond to the values entered into the RR formulas (RR1 and RR2) (see ‘Materials and methods: Quantifying and comparing ...’ for more details). Note that high performance of a predictor is indicated by large RR1 values and small RR2 values. Chi-squared statistics and p-values refer to the null-hypothesis that the 2 categorical variables (environmental condition and recruitment) are independent. The tests were performed on 2 × 2 contingency tables.

<table>
<thead>
<tr>
<th>Environmental predictor</th>
<th><em>G</em></th>
<th><em>g</em></th>
<th><em>C</em></th>
<th><em>c</em></th>
<th>RR1</th>
<th>RR2</th>
<th>Chi-squared</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative flow regime (February)</td>
<td>4</td>
<td>3</td>
<td>0</td>
<td>17</td>
<td>243.43</td>
<td>0.43</td>
<td>7.91</td>
<td>0.005</td>
</tr>
<tr>
<td>Negative flow regime (March)</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>18</td>
<td>5.00</td>
<td>0.55</td>
<td>1.50</td>
<td>0.221</td>
</tr>
<tr>
<td>High <em>Calanus helgolandicus</em> abundance (February)</td>
<td>0</td>
<td>2</td>
<td>4</td>
<td>18</td>
<td>0.11</td>
<td>1.20</td>
<td>0.00</td>
<td>1.000</td>
</tr>
<tr>
<td>High <em>C. helgolandicus</em> abundance (March)</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>18</td>
<td>2.33</td>
<td>0.78</td>
<td>0.00</td>
<td>1.000</td>
</tr>
<tr>
<td>High <em>C. finmarchicus</em> abundance (February)</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>20</td>
<td>10.78</td>
<td>0.02</td>
<td>5.35</td>
<td>0.021</td>
</tr>
<tr>
<td>High <em>C. finmarchicus</em> abundance (March)</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>19</td>
<td>3.67</td>
<td>0.58</td>
<td>0.11</td>
<td>0.741</td>
</tr>
<tr>
<td>Low NAO (February)</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>18</td>
<td>2.33</td>
<td>0.78</td>
<td>0.00</td>
<td>1.000</td>
</tr>
<tr>
<td>Low NAO (March)</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>17</td>
<td>0.67</td>
<td>0.67</td>
<td>0.81</td>
<td>0.369</td>
</tr>
<tr>
<td>Low SST (February)</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>18</td>
<td>2.33</td>
<td>0.78</td>
<td>0.00</td>
<td>1.000</td>
</tr>
<tr>
<td>Low SST (March)</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>18</td>
<td>2.33</td>
<td>0.78</td>
<td>0.00</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Fig. 4. Sandeel recruitment. Two recruitment time-series are presented: ICES (2016a) (solid line) and ICES (2016b) (black dashed line) (see ‘Materials and methods’ for details). Good recruitment years (unusual recruitment spikes extending beyond the dashed line) are highlighted with red bullet points (ICES 2016a) or green bullet points (ICES 2016b). Red and green dashed lines: corresponding mean (±1 SD).

Fig. 5. Cumulative hatch-day frequency. Hatch dates were estimated from larval length distribution and the age−length relationship found by otolith daily increment analysis for sampled years: 2006 (red), 2008 (green), and 2009 (blue). Hatch frequencies were corrected for cumulative mortality assuming different mortality rates of 0.01 (circles), 0.07 (triangles) and 0.14 d⁻¹ (squares). DOY: day of the year.

Table 1. Risk ratios (RR) and chi-squared test of independence for various sandeel recruitment predictors. *G*, *g*, *C*, and *c* correspond to the values entered into the RR formulas (RR1 and RR2) (see ‘Materials and methods: Quantifying and comparing …’ for more details). Note that high performance of a predictor is indicated by large RR1 values and small RR2 values. Chi-squared statistics and p-values refer to the null-hypothesis that the 2 categorical variables (environmental condition and recruitment) are independent. The tests were performed on 2 × 2 contingency tables. NAO: North Atlantic Oscillation, SST: sea surface temperature.
and 10.78, respectively (ranking third and second best), and the distributions of recruitment and flow regime were independent (i.e. the null-hypothesis of independence could not be rejected). In the second RR test (RR2), *C. finmarchicus* in February performed best (RR2 = 0.02, $\chi^2 = 5.35$, $p = 0.021$; Table 1). For example, the relative probability of observing less than good recruitment in combination with a high abundance of *C. finmarchicus* was very low. However, it should be noted that only 2 years qualified as high *C. finmarchicus* abundance years (1994 and 2009). Time-series of *C. helgolandicus*, *C. finmarchicus*, NAO, and SST are presented in Fig. 6.

**DISCUSSION**

**Circulation patterns in the North Sea**

The common circulation in the North Sea is counterclockwise, produced by prevailing south-westerly winds (Sündermann 2003, Sündermann & Pohlmann 2011), but occasionally the flow direction is reversed (i.e. clockwise circulation). According to reports by Greene & Pershing (2003) and Tiessen et al. (2014), such an event took place in 1996. In the present study, the year 1996 was assigned to the overarching cluster encapsulating negative flow regimes in both February
and March and the ‘retention’ sub-cluster in February (in March the PCA could not distinguish between ‘advection’ and ‘retention’ flow types). Considering February in isolation, 7 out of the 24 years analysed were allocated to the cluster of negative flow regimes. However, 2 of these years were classified as ‘stagnant’ or ‘shifting flow’ and not ‘retention’ (in the sub-clusters) and thus whether all 7 years correspond to a reversed (clockwise) circulation on a regional scale (as in 1996) is not clear from the present results.

The schematic illustration by Sündermann & Pohlmann (2011) of the overall circulation in the North Sea suggested 5 possible patterns, of which Kauker & von Storch (2000) provided the most detailed description presented by a 15 yr simulation (1979–1993) using the European Centre of Medium-Range Weather Forecast reanalysis and an ocean circulation model. Here the dominant regime displayed a circulation pattern of large gyres circulating either in a counterclockwise or clockwise direction resulting from northeasterly and southwesterly winds, respectively. In line with our findings, clockwise rotations, represented by negative associated flow regimes in our data, are relatively unusual, taking place only 15% of the time (and in 22% of the trajectories analysed in our study). In comparison, clockwise directions occur 30% of the time. The ‘stagnant flow’ observed in our study was also identified by Kauker & von Storch (2000) and, as in our study, these were also relatively rare (10% in their study and 6% in our study). Kauker & von Storch (2000) also pointed out that it was rare for regimes to last for longer periods, and only 20% persisted for >5 d. Some of the flows that they characterized as stagnant may therefore be comparable to what we refer to as ‘shifting flow’ (which we observed in 6% of the trajectories analysed). They also described a fourth intermediate type of circulation, which occurred 45% of the time, and the full nature of which we may not have captured within our study design. In this type, northwesterly or southerly winds produced bipolar circulation structures, shifting between southern and northern parts of the North Sea, which might be translated to either ‘shifting flow’ or ‘retention’ in our case.

Another remarkable pattern emerging from the present analysis was that during the first half of the time-series (1990 to ca. 2002), flow regimes of the ‘advection’ type dominated in February, whereas in the latter half of the time-series, flow regimes of the ‘retention’ type prevailed. Such an indication of an oceanographic regime shift is striking, in particular because a series of other studies have documented recent biological regime shifts in the North Sea, such as shifts in the zooplankton species composition and reduced growth and recruitment of planktivorous fish, including sandeel (e.g. Reid et al. 2001, Beaugrand 2004, Clausen et al. 2018, Lindgren et al. 2018). However, these shifts typically occurred around 1990 (plus/minus several years), which represents the beginning of the time-series analysed in the present study, and therefore prevented us from aligning the findings from previous regime-shift studies with our data.

**Flow regimes in relation to sandeel recruitment and first feeding**

All 4 good recruitment years (i.e. unusually strong year-classes, see definition of ‘good’ in ‘Materials and methods’) coincided with negative flow regimes in February (only 2 in March), suggesting that the flow regimes in February, rather than in March, determine the fate of sandeel larvae on the DB. In the present study, otolith-based hatch dates were mainly confined to March, which corroborates similar studies from nearby sandeel populations (Shetland Islands and Firth of Forth), showing peak hatching in the same month (Wright & Bailey 1996, Régnier et al. 2017). In contrast, the CPR survey also captured substantial numbers of sandeel larvae on the DB in February (Lynam et al. 2013), indicating that the otolith-based approach may be biased toward relatively late hatch dates. With this in mind, a closer inspection of the hatch date distributions revealed a heavy tail into February in 2009, which happened to coincide with a negative flow regime and the second largest recruitment event in the time-series. In fact, Wright & Bailey (1996) also observed a similar extended tail in the 1990 hatch data distributions, which coincided with a large year class (Poloczanska et al. 2004). Hence, we suggest that negative flow regimes in February facilitate better first-feeding conditions and survival of the earliest larvae, creating the extended tails that we observed in the otolith-based hatch date distributions. This view is in line with Arnott & Ruxton (2002), suggesting a strong link between year-class strength and survival of early larval stages in February (and not March). Unfortunately, we only had otoliths from 3 years in our possession. A more complete comparison of the timing of first feeding and recruitment was therefore not possible. A comparison of the oceanographic conditions within the first-feeding period each year (instead of fixed windows, February and March) would also have been useful.
Utility of flow regime and other environmental descriptors in probabilistic forecasts

On several occasions, *Calanus finmarchicus* has been highlighted as an important indicator of sandeel recruitment (Arnott & Ruxton 2002, van Deurs et al. 2009, Lindegren et al. 2018). This link is based on correlations between *C. finmarchicus* abundance in February and sandeel recruitment, and has been explained by the potential naupliar production in February, when the overall zooplankton concentration is at its lowest (Arnott & Ruxton 2002). However, the present study revealed that, while the 2 best *C. finmarchicus* years coincided with good recruitment years, other recruitment spikes occurred in very poor *C. finmarchicus* years (i.e. 1996). In comparison, recruitment never occurred outside a negative flow regime in February. These findings suggest that sandeel larvae do not rely entirely on *C. finmarchicus* for survival, as previously hypothesized. Instead, it may be that sandeel recruitment and the retention of *C. finmarchicus* near the DB are both favoured by a negative associated flow regime. Planque & Batten (2000) explained the very low number of *C. finmarchicus* recorded by the CPR in the North Sea in 1996 by the fact that there was an unusually small *C. finmarchicus* population in the same year in the North Atlantic. However, it is counterintuitive that distinct *C. finmarchicus* spikes occur in negative flow regimes, given the consensus hypothesis that *C. finmarchicus* drift in a south-easterly direction into the North Sea in the first quarter of the year through the Orkney-Shetland channel and the Norwegian trench (Gallego et al. 1999, Madden et al. 1999). One plausible explanation could simply be that the surface water flow in the central North Sea is largely disconnected from the deep inflow events (below 150 m) responsible for transporting *C. finmarchicus* into the North Sea (Reid et al. 2003, Jónasdóttir & Koski 2011). Reid et al. (2003) also showed that the deep water inflow (<150 m) is negatively correlated to the inflow in the upper water column (0–150 m). Hence, the correlation between surface flow on top of the DB and deep inflow events is likely to be weak. Recently, Régnier et al. (2017) presented *C. helgolandicus* as a more important copepod species in relation to the year-class strength of sandeel west of the DB. This is with good reason, as *C. helgolandicus* has become the dominant *Calanus* species in the North Sea in recent decades, presumably due to climate change (Fromentin & Planque 1996, Hinder et al. 2014). However, in relation to the DB, we found no matches between years with large numbers of *C. helgolandicus* and years with good sandeel recruitment.

Why did SST and NAO coincide poorly with recruitment in the present study, when previous studies have shown significant relationships in time-series correlations (Arnott & Ruxton 2002, Lindegren et al. 2018)? The explanation regarding SST may already have been provided by Lindegren et al. (2018), who described how SST defines the recruitment trends, but not the extreme recruitment events. Therefore, while SST could potentially be used to inform managers about productivity regimes (Claussen et al. 2018, Lindegren et al. 2018), it has little value in relation to short-term forecasts of fishing opportunities, in particular for stocks displaying boom-and-bust dynamics (Stocker & Hilborn 1981, de Moor et al. 2011). NAO has been linked to *C. finmarchicus* regimes on a decadal time scale (Planque & Taylor 1998, Heath et al. 1999), and some degree of correlation between sandeel recruitment and winter NAO has also been reported (Arnott & Ruxton 2002). NAO influences the North Sea circulation usually with a positive relationship, where stronger westerly winds amplify the general cyclonic circulation during NAO+ phases. Nonetheless, positive phases also seem to favour wind conditions that could lead to circulation reversals on weekly and monthly scales (Mathis et al. 2015), which limits the chance of finding any robust correlations between NAO and local circulation patterns at the DB. It is therefore not surprising to find that NAO is a poor predictor of the inter-annual variation in sandeel recruitment.

As mentioned above, all good recruitment events coincided with a negative flow regime in February. In contrast, when a positive flow regime is observed in February, the prediction is that the recruitment will always be moderate to low. However, 3 out of the 7 years (1991, 2007, and 2010) assigned to the cluster of negative flow regimes in February were not good recruitment years. In this respect, it is important to mention that 1991 and 2007 represent peaks in the recruitment time-series, although they did not classify as ‘good recruitment’ years under the criteria set in our study. Furthermore, although 1991 had a flow trajectory almost identical to that of other good recruitment years, it also showed a unique signature of total stagnation in March, which could explain why good recruitment failed to take place in 1991 despite suitable conditions in February (i.e. assuming that stagnation is suboptimal for larval growth and survival). The year 2007 was characterized by a low spawning biomass (less than the time-series average minus 1 SD), suggesting some influence from the S-R relationship. Strong density-dependent regulation induced by 1 yr old fish in the population has been
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A recruitment hypothesis

While the consensus in the literature regarding sandeels is that they appear to have strong links with specific prey types, coupling the ‘match–mismatch’ hypothesis (Cushing 1969, 1975, 1990) to mainly the relative large Calanus copepods advected into the North Sea (e.g. Arnott & Ruxton 2002, Régnier et al. 2017), we suggest considering alternative hypotheses, such as the ‘weak link/strong filter’ hypothesis proposed by Runge (1988), or the ‘Stable Ocean’ and ‘Optimal Environmental Window’ theories (Lasker 1988, Curly & Roy 1989). Applying these to the sandeel case would imply that the link to Calanus copepods advected into the North Sea (e.g. Arnott & Ruxton 2002, Régnier et al. 2017), we suggest considering alternative hypotheses, such as the ‘weak link/strong filter’ hypothesis proposed in former studies (e.g. van Deurs et al. 2009, Lindegren et al. 2018). Numbers of age-1 fish were about average or below in 1991 and 2007. However, in 2010 the second highest number of age-1 fish in the study period was recorded, potentially explaining the poor recruitment in this year (time-series of spawning stock biomass and numbers of age-1 fish are available in ICES 2016b).

If favourable oceanographic winter (February) conditions occur often enough, early hatching may even be favoured, as it extends the duration of the growth period and increases the probability (relative to larvae hatching in March or April) that the critical threshold size is reached before winter (van Deurs et al. 2011). Studies on the yolk sac period in sandeel may support to this view. The yolk sac provides nutrition for sandeel larvae during the first ~12 d after hatching (Wright 1993), which helps them cope with low food availability for as much as 2 wk (Buckley et al. 1984, Régnier et al. 2017). At the same time, all winter-hatching sandeels are apparently capable of feeding prior to yolk sac absorption (Yamashita & Aoyama 1985), enabling advantageous utilization of an extended growth period whenever oceanographic conditions are already favourable in February.

The remaining question is: What does ‘favourable oceanographic conditions’ mean? In the search for direct causal relationships, we suggest that larval retention and advection processes on the DB are important. The flow regimes associated with unusually strong year-classes are characterized by either bidirectional flow or relatively limited net flow in a northerly direction in February (in contrast to a relatively strong net flow in a southerly direction, which prevails in most years). We hypothesize that this flow regime promotes dominance of southern water masses (e.g. from the Southern Bight, German Bight, and Wadden Sea), where primary production is highest in February (Joint & Pomroy 1993, Moll 1998). At the same time, the relatively limited net flow over the DB ensures larval retention and concentrates microzooplankton and phytoplankton for the early hatching larvae to feed on (Malzahn et al. 2007, Malzahn & Boersma 2009). These speculations are in line with earlier model simulations identifying the optimal sandeel hatching time to be February in the southeastern North Sea (Gurkan et al. 2013) and the DB to be a location where retention dominates over advection, facilitating perfect conditions for accumulating food and retaining sandeel larvae near suitable settlement habitats (Christensen et al. 2008).

Despite our efforts to identify and discuss possible causal relationships, the underlying mechanisms remain unclear, and the data presented here were not designed to support strong conclusions about anything other than the striking concurrence between unusually strong sandeel year-classes and negative flow regimes. We therefore encourage further research, pursuing the hypotheses and ideas developed in this study.
Conclusions and prospects of using oceanographic data to forecast stock productivity

Here we have demonstrated, for the first time, how internal North Sea circulation patterns can be used to predict fish recruitment in the North Sea. We found that good sandeel recruitment occurred only when negative flow regimes prevailed in February (and is most likely to occur in the ‘retention’ type of negative flow regimes). It is also plausible that these negative flow regimes can be translated into reversals of the predominant North Sea circulation pattern (i.e. changed from clockwise to counterclockwise), although this may be a rather oversimplified description of complex oceanographic processes, requiring further validation studies using, for example, North-Sea-wide 2D flow fields. Moreover, other candidate drivers, such as chlorophyll (Bakun 2006), might provide further insight into the mechanistic understanding behind these large year-classes. We also found indications of a previously unnoticed oceanographic regime shift in

Fig. 7. ‘Recruitment window’ hypothesis. Vertical grey arrows signify the order of events in the flowchart. The recruitment window represents the critical period where hatching and first feeding larvae are present. Within this window, it is the sequence of oceanographic events (i.e. flow regimes) that determines growth and survival of the larvae (i.e. flow regimes define the foraging arena and/or larval retention processes). For example, a given sequence can be comprised of both optimal conditions (labelled ‘A’) and poor conditions (‘B’), where poor conditions lead to dramatically impaired survival (i.e. ≤10% of the survival experienced when conditions are optimal; see Houde 2008 regarding the scaling of larval mortality). In the flow chart, the different types of flow regimes found in the present study are used to exemplify optimal and poor conditions (illustrated with arrows and labelled A & B). Note that in stagnant flow regimes (flow regime on the far right) it is unclear from the present results whether this translates into optimal or poor conditions for the larvae. In the end, total recruitment is approximately proportional to the ratio between optimal and poor conditions within the recruitment window.
the North Sea taking place around 2002, which also warrants further investigation. Lastly, we recognise that a relatively coarse resolution in ocean models (such as in the model used here) can lead to an underestimation of the volume transport, making it easier for wind forcing to affect surface flow (Påtsch et al. 2017). Hence, we recommend that future studies use a higher horizontal model resolution.

The use of oceanographic data in recruitment forecasts is still very limited (e.g. Myers 1998, Payne et al. 2017), despite the fact that oceanographic and meteorological data have proven useful in understanding recruitment dynamics of some of the largest fish stocks in the world. For example, there is a link between productivity of Pacific sardine and wind-driven upwelling systems off the coast of Central America and coupling between certain physical conditions of the Norwegian Coastal Current and the survival and growth of Norwegian spring spawning herring larvae drifting toward the nursery grounds in the Barents Sea (Rybakowski & Checkley 2008, Skagseth et al. 2015). In this study, we have demonstrated that information about local flow regimes outperforms other previously considered recruitment predictors with respect to forecasting the most extreme recruitment events. Our approach differs from many other time-series studies of fish recruitment by being probabilistic (i.e. the RR approach) and focussing on the anomalies (i.e. the most extreme recruitment events) rather than on the average trends. This approach may be useful in regions with highly variable oceanographic conditions. Lastly, we propose a hypothesis for sandeel recruitment and its relation to physical oceanography, which we hope will inspire future research (Fig. 7).

While our study has quantified a probabilistic link between flow conditions and recruitment, there are additional considerations which must be addressed before such a relationship can be applied in stock forecasting. These include verification of the proposed recruitment hypothesis and a demonstration that such a forecast will effectively improve the stock advice with respect to forecast accuracy and stakeholder needs.

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Appendix. Additional information on the PCA

Fig. A1. Scree plots of eigenvalues ordered from largest to the smallest; (A) February, (B) March. The number of principal components (PCs) is determined at the point beyond which the remaining eigenvalues are all relatively small and of comparable size. Black bars signify PCs retained for PCAs.

Fig. A2. Correlation circles (A, February; B, March) showing correlations between years and principal components, PC1 (i.e. Dim1) and PC2 (i.e. Dim2), where correlated years are grouped together. The distance between arrows and the origin measures the quality of the years on the factor map. Variables that are away from the origin are well represented on the factor map.
Fig. A3. Total contribution of years to principal components PC1 and PC2 in February and March. Contributions of variables in accounting for the variability in a given PC are expressed in percentage (%). The red dashed line on the graph indicates the expected average contribution.