Determinacy of fecundity and oocyte atresia in sole (Solea solea) from the Channel, the North Sea and the Irish Sea

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Abstract

Changes in the morphology of the ovary are described during maturation and it is established that it is a homogeneous structure for sampling purposes at maturity stages four and five. Several aspects of vitellogenic oocyte growth and recruitment were examined as criteria to ascertain whether the annual fecundity can be determined in Solea solea just prior to spawning. The first criterion was whether a hiatus develops in the size frequency distribution between the previtellogenic and vitellogenic oocytes just prior to spawning. We were unable to demonstrate this feature in sole from division VIIId (N = 5) or IVc (N = 5) but it was present in two out of five fish from division VIIia and all fish examined from division VIIia (N = 5) and IVb west (N = 5). The hiatus occurred between 175 and 250 μm (cohort size 25 μm) in the size frequency distribution and measured between 25 and 75 μm. Oocyte size frequency distribution from mid spawning and late spawning-spent sole from division IVc showed a hiatus in 9 out of 10 fish which became wider as the residual annual potential fecundity declined. Next the rate of recruitment and growth of vitellogenic oocytes was studied in sole sampled at monthly intervals (July 1993 to February 1994) from division VIIId in relation to the start and duration of the annual spawning season. The growth rate of the leading oocyte cohort was $2.73 \times 10^{-5}$ mm$^3$ per day on 20 September and increased to $3.91 \times 10^{-5}$ at a mean diameter of 839 μm by 10 February just prior to spawning. However, the observed maximum growth rate, even when adjusted for the positive influence of temperature, allowed little scope for previtellogenic oocytes to complete maturation during a spawning period of 60 days. The recruitment of vitellogenic oocytes was 3013 oocytes per day on 3 August and declined rapidly as spawning approached. In the three months prior to spawning no significant increase was found in the annual potential fecundity. An analysis of a sample of spawning fish from IVc demonstrated that 88% had less than 50% of their predicted potential fecundity remaining in the ovary. The mean batch fecundity was 8400 (SE ± 13363) and in conjunction with existing data on spawning frequency and duration gave a realised fecundity close to the potential fecundity. It is concluded that for all practical purposes the sole can be regarded as having a determinate fecundity in all the areas studied.

The regulation of potential fecundity by follicular atresia was assessed in pre-spawning and spawning sole. In ICES areas IV and VII the prevalence of atretic oocytes in pre-spawning fish varied between 0.04 and 0.69 and the relative intensity varied between 0.017 and 0.076. During spawning in division IVc the prevalence of atresia varied between 0.45 and 0.57 and the relative intensity between 0.022 and 0.058. If we assume a turnover rate of 9 days the loss from the potential annual fecundity, during spawning, in division IVc would be 12.4%.

Keywords: Determinacy, atresia, oocytes, fecundity, spawning season, maturity, vitellogenesis, Solea solea.
INTRODUCTION

The southerly extent of the distribution of sole is along the Mediterranean coast and as far south as Senegal (30° north) on the eastern Atlantic coast (Whitehead et al., 1986). The northern limit is bounded by the western Baltic Sea, Trondheim Fjord on the Norway coast (60° north) and the Faroe Islands. The stocks around the British Isles are thus towards the northern limit of the range of this species. Rijnsdorp et al. (1992) and Horwood (1993) have recently reviewed the biology of the sole and the fishery. An asynchronous pattern of oocyte development, and the release of mature oocytes in a series of batches during the spawning season, has been described by Ramos (1983) and Urban (1991).

The assessment of sole stocks for management purposes is presently done by virtual population analysis (Pope and Shepherd, 1988) using catch-at-age data from commercial vessels. An alternative method can be used, in which the total number of eggs in the plankton is estimated using a research vessel survey and divided by the number of eggs produced per female. This method can only be used if the total annual fecundity is present and can be counted prior to spawning. If this is the case, the fecundity is termed determinate. If, however, some previtellogenic oocytes commence vitellogenesis and develop to ovulation for some period during spawning, then the fecundity measured before spawning does not represent the true fecundity and is termed indeterminate. In the latter case, the daily egg production method can be used for assessment (Parker, 1980). In the past, it has been assumed that the sole has a determinate fecundity. The criterion for his assumption has been the presence of a gap or hiatus in the size frequency distribution of oocytes between the previtellogenic oocytes which mature in subsequent years and the maturing (vitellogenic) oocytes which mature and are spawned in the current year. Horwood and Greer Walker (1990) showed that a hiatus developed prior to spawning in sole from the Bristol Channel (ICES division VIIIf) and concluded that the fecundity of sole from that area was determinate. However, Urban (1991) found no hiatus in the size frequency distribution of oocytes from German Bight sole (ICES division IVb east) and concluded that the fecundity was therefore indeterminate. During a discussion of these differences between stocks, Urban (1991) pointed out that the two stocks may have different...
spawning strategies. It is also apparent that the sole from the German Bight were small fish, possibly in their first spawning season, and caught at the end of the spawning season (late June). Under these circumstances they may not be considered representative of the whole spawning population.

In the present work we have studied changes in ovary morphology in relation to maturity stage to verify sampling procedures. Primarily we set out to establish whether or not sole stocks from the Channel (ICES divisions VIIId and VIId), the North Sea (ICES divisions IVb west, IVb east and IVc) and the Irish Sea (ICES division VIIa) have a determinate fecundity. To this end three lines of evidence were examined:

1. the development of a gap or hiatus in the size frequency distribution between previtellogenic and vitellogenic oocytes prior to and during spawning;
2. the rate of recruitment and growth of vitellogenic oocytes in relation to the length of the spawning season;
3. changes in the number of vitellogenic oocytes in the ovary during spawning.

Finally as an adjunct to the main objective we have quantified atresia (the resorption of oocytes) in sole from all the ICES areas mentioned above prior to spawning (maturity stage 4) and in ICES division IVc during spawning.

### MATERIALS AND METHODS

#### Collection of fish

Collections of female sole, detailed in table 1 and figure 1, were made between 1984 and 1994. Samples collected from division IVc during 1984 were used to study gross ovarian morphology. During 1991, female sole were collected from several places in ICES areas IV and VII to give a comprehensive geographic coverage in one year. Fish were collected at a time just before spawning, which was January in division VIIc, February in VIId and March in area IV (De Clerck, 1974) to study hiatus development and oocyte atresia. Spawning fish were also collected in area IV to continue the study of hiatus development, and estimate oocyte atresia and batch fecundity. Finally, samples were collected during the period from July 1993 to February 1994 from ICES division VIIId to measure the rates of increase in fecundity and growth in size of the leading cohort of vitellogenic oocytes cohort prior to spawning. In 1991, stratified length samples were generally taken from the catch at sea and the ovaries dissected out and preserved immediately after the fish were caught. However, some of the fish sampled in division VIIe were selected after landing but from a part of the catch made 24 h before the return to port. From July 1993 until January 1994 10 recovering-maturing fish per month were selected.

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**Table 1.** Details of samples collected in ICES areas IV and VII between 1984 and 1994.

<table>
<thead>
<tr>
<th>ICES division</th>
<th>Method of capture</th>
<th>Date</th>
<th>No. of fish</th>
<th>Length (cm)</th>
<th>Maturity stage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>range</td>
<td>mean (SD)</td>
</tr>
<tr>
<td>IVb west</td>
<td>Beam trawl</td>
<td>19 March 1991</td>
<td>45</td>
<td>24-46</td>
<td>33 (5)</td>
</tr>
<tr>
<td>(Flamborough)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IVb east</td>
<td>Beam trawl</td>
<td>15 March 1991</td>
<td>40</td>
<td>26-48</td>
<td>36 (6)</td>
</tr>
<tr>
<td>(German Bight)</td>
<td>27 May-16 June 1991</td>
<td>69</td>
<td></td>
<td>28-36</td>
<td>32 (2)</td>
</tr>
<tr>
<td>IVc west</td>
<td>Beam trawl</td>
<td>26 April-5 June 1984</td>
<td>10</td>
<td>29-40</td>
<td>34 (4)</td>
</tr>
<tr>
<td>(Southern Bight)</td>
<td></td>
<td>1 March 1991</td>
<td>55</td>
<td>24-46</td>
<td>34 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>15-16 May 1991</td>
<td>93</td>
<td>26-41</td>
<td>32 (2)</td>
</tr>
<tr>
<td>VIIa west</td>
<td>Beam trawl</td>
<td>21-25 March 1991</td>
<td>29</td>
<td>28-42</td>
<td>34 (4)</td>
</tr>
<tr>
<td>(Irish Sea)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VIIId west</td>
<td>Beam trawl</td>
<td>4-7 Feb. 1991</td>
<td>49</td>
<td>27-43</td>
<td>35 (3)</td>
</tr>
<tr>
<td>(Eastern Channel)</td>
<td></td>
<td>19 July 1993</td>
<td>10</td>
<td>34-38</td>
<td>35 (1)</td>
</tr>
<tr>
<td></td>
<td>Trammel net</td>
<td>4-8 Aug. 1993</td>
<td>10</td>
<td>34-42</td>
<td>37 (2)</td>
</tr>
<tr>
<td></td>
<td>Beam trawl</td>
<td>20-21 Sept. 1993</td>
<td>10</td>
<td>32-38</td>
<td>34 (2)</td>
</tr>
<tr>
<td></td>
<td>Trammel net</td>
<td>10-11 Oct. 1993</td>
<td>10</td>
<td>33-42</td>
<td>38 (3)</td>
</tr>
<tr>
<td></td>
<td>Otter trawl</td>
<td>20 Nov. 1993</td>
<td>10</td>
<td>33-49</td>
<td>38 (5)</td>
</tr>
<tr>
<td></td>
<td>Beam trawl</td>
<td>8 Dec. 1993</td>
<td>10</td>
<td>33-42</td>
<td>38 (3)</td>
</tr>
<tr>
<td></td>
<td>Beam trawl</td>
<td>11 Jan. 1994</td>
<td>10</td>
<td>33-43</td>
<td>37 (3)</td>
</tr>
<tr>
<td></td>
<td>Beam trawl</td>
<td>10 Feb. 1994</td>
<td>35</td>
<td>30-40</td>
<td>37 (2)</td>
</tr>
<tr>
<td>VIIe west</td>
<td>Beam trawl</td>
<td>27 Jan. 1991</td>
<td>33</td>
<td>30-51</td>
<td>39 (5)</td>
</tr>
</tbody>
</table>

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(Ramsay, 1993) randomly from fish caught 24 hours before landing by commercial fishing boats working in division VIIa. A stratified length sample was collected at sea in February 1994 of which the first 10 mature fish comprised the final monthly sample. The total fish length was measured to the nearest centimetre. Ovaries collected were assigned to one of seven maturity stages table 2. Ovary weights were estimated prior to preservation in the 1993-1994 samples.

**Ovary morphology**

Two freshly-caught whole fish (table 1-1984) were clamped down flat on a metal plate and dipped into isopentane cooled to $-150^\circ$C by liquid nitrogen on board ship. After several days storage, at about $-100^\circ$C in the vapour phase of a liquid nitrogen refrigerator, the ovaries and surrounding tissue and were cut free from the body and mounted in a block of 2% (w/v) sodium carboxy-methyl cellulose. They were transferred to a deep freeze at $-20^\circ$C for transport and sectioning. Sections of 25 $\mu$m were cut on a whole body cryostat (Bright LKB 2250) at the Huntingdon Research Centre, Cambridge. Sections were collected on adhesive cellulose acetate tape and stained with Harris' haematoxylin and eosin.

**Homogeneity of the ovary**

Female sole were caught close to Lowestoft between 26 April and 5 June 1984 (table 1, fig. 1). Eight pairs of ovaries (maturity stages 4, 5 and 6) were preserved in 3.6% buffered formaldehyde and returned to the laboratory. A cross-section of tissue was cut out from the middle of the eye-side ovary and from the anterior end of the blind-side ovary (two sites that might be expected to show maximum diversity in terms of oocyte developmental stages). These sections were prepared for histological analysis as described below. The density (the number of oocytes per unit area) of vitellogenic oocytes from the two samples

Table 2. — Description of the ovary maturity stages used in the collection. Maturity stages 5 and 6 are shown in figure 2A-C.

<table>
<thead>
<tr>
<th>Maturity stage</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Immature:</td>
<td>Ovaries are small, extending no more than 5 cm into the body cavity; translucent in appearance; no oocytes visible.</td>
</tr>
<tr>
<td>2 Resting:</td>
<td>Ovaries cream coloured, extending further into the body cavity. Lumen filled with fluid. No oocytes visible.</td>
</tr>
<tr>
<td>3 Early developing:</td>
<td>Ovaries extend to fill half the body cavity. Yellow in colour. Blood supply well developed. Some developing oocyte visible.</td>
</tr>
<tr>
<td>4 Late developing:</td>
<td>Ovaries fill the body cavity and body is distended. Yellow in colour. Lumen and granular oocytes visible.</td>
</tr>
<tr>
<td>5 Ripe:</td>
<td>Ovary swollen and the body distended. Translucent hyaline oocytes can be seen inter-spaced with opaque granular oocytes. Oocytes can be extruded under pressure.</td>
</tr>
<tr>
<td>6 Running:</td>
<td>Ovary full becoming flaccid as spawning progresses. Lumen filled with ovulated hyaline oocytes which can be extruded under gentle pressure. They are often visible through the transparent ovary wall.</td>
</tr>
<tr>
<td>7 Spent:</td>
<td>Ovaries reduced in size and flaccid. Cream coloured. Areas of red tissue indicate atretic oocytes. Lumen filled with fluid and resorbing or atretic oocytes.</td>
</tr>
</tbody>
</table>
sites was estimated in the following manner. The vitellogenic oocytes were divided into three easily-identifiable developmental stages: cortical alveoli, yolk granule and yolk fusion stages (Guraya, 1986). These stages were counted within a square grid; oocytes that impinged on two of the four sides were ignored but counted when they impinged on the other two sides. A number of grid counts (about 10) were made in such a way that the ovary section was traversed.

Determimacy of fecundity and oocyte atresia

Histological methods

The nomenclature and identification of oocyte development stages that we have adopted were described by (Ramos, 1983) in Solea solea and Yamamoto and Yamazaki (1961) in Carassius auratus. Ovaries for the study of determinacy and atresia were prepared for analysis in the following manner. They were fixed in 3.6% formaldehyde buffered to pH 7.0 with 0.1M sodium phosphate for a period of 36 h. After fixation, they were stored in 70% industrial methylated spirit. The ovary volume was determined using a displacement method (Scherle, 1970) and transverse sections 0.5 cm thick were dissected out from the midpoint of each ovary using a scalpel and placed in a histological cassette. Following dehydration in ethyl alcohol and embedding in methacrylate resin, sections were cut at 5 μm using a motorised microtome situated in a refrigerated cabinet at -12°C. Sections were stained with periodic acid Schiff’s (PAS) and Mallory’s trichrome to identify which oocyte development stages were present in the ovary. Oocytes which contained cortical alveoli were included with yolk granule stage oocytes in the fecundity estimates (Khoo, 1979).

Assessment of the presence of a hiatus in the oocyte size frequency distribution

The presence or absence of a hiatus in the oocyte size frequency distribution was established using the method of Foucher and Beamish (1980). Two measurements of diameter were made, using a visual image display system (VIDS 5, Synoptics Ltd, Cambridge), from each oocyte where a nucleus was visible in the cross section, and the mean taken. It was calculated that the maximum error at 200 μm was 0.89 x true size and 0.97 x true size at 800 μm. This error was calculated from the possible range of diameter measurements of the outer circle (the cell wall) when the inner concentric circle (the nucleus) still appears in the cross section. All oocytes > 100 μm were measured and classified as either vitellogenic or previtellogenic until up to 50 vitellogenic oocytes had been measured from each ovary. Oocytes which had commenced final oocyte maturation (yolk coalescence) were excluded. The observed oocyte size frequency measured in histological section was corrected to the actual distribution as described by Greer Walker et al. (1994).

Estimation of the numbers of vitellogenic and atretic oocytes and the growth rate of vitellogenic oocytes

A stereological technique (Emerson et al., 1990) was used to count the numbers of vitellogenic and atretic oocytes. Five point and profile counts were made from each of the two ovaries using a Weibel grid (M162) such that the grids traversed the ovary section. Up to 50 vitellogenic oocytes were measured to estimate both the k distribution coefficient and maximum oocyte size found in the ovary. It is possible that both the relatively small sample size, and measuring oocytes where part of the nucleus was visible in the cross section, could contribute to an underestimate of maximum oocyte size. We therefore compared our results of maximum oocyte size from sections (method 1) with measurements of whole oocytes from the same ovaries which were preserved in formaldehyde (method 2). In the latter case 20 larger oocytes were selected from the eye side ovary and measured using VIDS 5. The maximum oocyte diameters found in 20 fish (table 1, VIIId 20-21 September, 10-11 Oct. 1993) using the two methods were compared by a linear regression (method 1 = α + β method 2). The results were as follows.

Method 2 = Method 1 x 0.856 - 0.619

R² = 0.95 p < 0.001

The intercept was not significantly different from zero p > 0.97 and method 1 underestimated the maximum diameter in formaldehyde preserved oocytes by 14%.

The length range of sole available on the market to provide monthly samples for division VIIId in 1993-94 (table 1) was often very restricted. To enable us to compare the development of fecundity in different sized fish during the time series the observed fecundity at length L was corrected to a standard length of 35 cm using equation 2:

Corrected fecundity = Fecobs/Fecpred x Fec35 (2)

Fecobs = observed fecundity of fish of length L cm
Fecpred and Fec35 = predicted annual potential fecundity of fish of L cm and of 35 cm derived from the 1994 fecundity length equation in table 3.

Estimating the progress through spawning

The relationship between the potential annual fecundity, that is, the total number of vitellogenic oocytes calculated prior to spawning, and length in ICES areas IV and VII is described in Anonymous (1992) and is summarised in table 3. The difference between the predicted annual potential fecundity and the measured fecundity enabled us to estimate the progress through spawning providing, of course, that we can accept that the sole has a determinate fecundity.
The relationship between the potential annual fecundity (defined as the number of oocytes at or beyond the cortical alveoli stage of development immediately before spawning) and length of soles from ICES areas IV and VII. All collections were made during 1991 except in VIIId where collections were made in 1991 and 1994. The regression parameters fitting the model \( \ln \text{fecundity} = b + a \ln \text{length} \) are given with the predicted fecundity at a length of 35 cm.

<table>
<thead>
<tr>
<th>ICES division</th>
<th>Date</th>
<th>N</th>
<th>a</th>
<th>b</th>
<th>( R^2 )</th>
<th>Predicted fecundity at 35 cm (thousands)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IVb east</td>
<td>15 March</td>
<td>40</td>
<td>3.7407</td>
<td>-0.2986</td>
<td>0.90</td>
<td>443</td>
</tr>
<tr>
<td>IVc</td>
<td>1 March</td>
<td>55</td>
<td>3.8065</td>
<td>-0.5617</td>
<td>0.88</td>
<td>430</td>
</tr>
<tr>
<td>IVb west</td>
<td>19 March</td>
<td>45</td>
<td>4.2851</td>
<td>-2.7495</td>
<td>0.88</td>
<td>264</td>
</tr>
<tr>
<td>VIIa</td>
<td>21-25 March</td>
<td>29</td>
<td>3.9531</td>
<td>-1.3122</td>
<td>0.72</td>
<td>342</td>
</tr>
<tr>
<td>VIIId</td>
<td>4-7 Feb. 1991</td>
<td>49</td>
<td>4.1699</td>
<td>-2.1270</td>
<td>0.76</td>
<td>327</td>
</tr>
<tr>
<td>VIIe</td>
<td>27 Jan.</td>
<td>33</td>
<td>3.3365</td>
<td>-6.3090</td>
<td>0.74</td>
<td>258</td>
</tr>
<tr>
<td>VIIId</td>
<td>10 Feb. 1994</td>
<td>35</td>
<td>4.1702</td>
<td>-2.2147</td>
<td>0.56</td>
<td>300</td>
</tr>
</tbody>
</table>

**Batch fecundity**

The batch fecundity was estimated using the hydrated oocyte method (Hunter and Goldberg, 1980). Fish were collected (table 1) from ICES division IVc between 15 and 16 May 1991 (N = 93) and IVb east between 27 May and 16 June 1991 (N = 69). All the ovaries from maturity stage 5 fish were screened histologically and those with fresh post-ovulatory follicles were rejected as, in these cases, ovulation had clearly commenced. Fresh follicles were identified by their cord-like appearance. Degenerating follicles from previous batches were recognised because of their cell hypertrophy and their more compact appearance. A full description is given by Hunter and Macewicz (1985a). A total of 38 fish measuring between 28 and 36 cm were selected after screening. The number of hydrated oocytes, present in each of two weighed sub-samples per fish, was counted using an adapted Zeiss microfiche reader (magnification × 6.5) and subsequently raised to the weight of the whole ovary.

**Identification of atresia**

The nomenclature and general characteristics used to identify atretic stages follow those of Bretschneider and Duyvene de Wit (1947) and Lambert (1970). Estimation of the number of atretic oocytes which still retained some characteristics of cortical alveolic or yolk granule oocytes was based on the first part of the alpha stage. The truncated alpha stage was more suitable for stereological estimation because it is the end of this stage, the structure’s boundary becomes diffuse and poorly defined. The following criterion were used:

**Cortical alveoli stage (fig. 3A and B)**

The first visible signs of alpha atresia are a folded and pitted zona pellucida, increased vacuolation, a disorganised ooplasm and intense PAS staining. At the end of this stage the cell is reduced in size, highly vacuolated and the zona pellucida having migrated towards the centre of the cell has all but disappeared. The follicle is hypertrophied.

**Yolk granule stage (fig. 3C, D, E and F)**

The first visible signs of alpha atresia occur in the zona pellucida which shrinks, thickens and takes on a wavy outline. Fissures occur in the zona radiata interna and in the early stages yolk products from the ooplasm form vesicles inside the zona radiata externa which remains intact for a while. These vesicles discharge their products through breaks in the zona radiata externa towards the follicle. As atresia progresses, these fissures enlarge and larger granules pass through. The zona pellucida shrinks towards the centre of the oocyte and at the end of the alpha stage the cell is much reduced in size, no yolk remains, the follicle is hypertrophied, prominent vacuoles are present and only remnants of the zona pellucida remain. This definition for the end of the alpha atretic stage implies a somewhat shorter period of developmental time than that described for the northern anchovy (Engraulis mordax) by Hunter and Macewicz (1985a). In the latter case, the end of the alpha stage was taken as being when the resorption of the oocyte including yolk, cytoplasm and the zona pellucida was complete.

The terminology used to describe oocyte atresia is as follow:

- **Prevalence.** The proportion of female fish with alpha atretic oocytes.
- **Intensity.** The number of alpha atretic oocytes found in ovaries from an individual fish.
- **Relative intensity.** The number of alpha atretic oocytes divided by the predicted potential annual fecundity calculated from fish length (table 3). It should be noted that estimates of potential fecundity do not include atretic oocytes.

**RESULTS**

**Ovary morphology**

In the ripe and running fish (fig. 2A and B) the eye-side ovary is approximately twice the size of
Fecundity and oocyte atresia in *Solea solea*

Table 4. – A comparison of the density (number per unit area) of cortical alveoli, yolk granule and yolk fusion stage oocytes in the eye-side and blind-side ovaries of the sole at different maturity stages using the chi-squared test.

<table>
<thead>
<tr>
<th>Fish no.</th>
<th>Ovary: eye-side (A)</th>
<th>Maturity stage</th>
<th>Total oocyte counts</th>
<th>Chi-squared test</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>blind-side (B)</td>
<td></td>
<td>Cortical alveoli</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>stage</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Yolk granule stage</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Yolk fusion stage</td>
<td></td>
</tr>
<tr>
<td></td>
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<td></td>
</tr>
<tr>
<td>1</td>
<td>A</td>
<td>4</td>
<td>7</td>
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</tr>
<tr>
<td></td>
<td>B</td>
<td></td>
<td>3</td>
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</tr>
<tr>
<td>2</td>
<td>A</td>
<td>4</td>
<td>8</td>
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</tr>
<tr>
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<td>B</td>
<td></td>
<td>6</td>
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<td>3</td>
<td>A</td>
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<tr>
<td>5</td>
<td>A</td>
<td>6</td>
<td>13</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td></td>
<td>5</td>
<td>27</td>
</tr>
<tr>
<td>6</td>
<td>A</td>
<td>6</td>
<td>4</td>
<td>47</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td></td>
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</tr>
<tr>
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<td>A</td>
<td>6</td>
<td>5</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td></td>
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<td>40</td>
</tr>
<tr>
<td>8</td>
<td>A</td>
<td>6</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td></td>
<td>1</td>
<td>34</td>
</tr>
</tbody>
</table>

the blind-side ovary and in both cases the cross-sectional area increases anteriorly to a point where they form a common oviduct which leads into the cloaca posterior to the operculum (fig. 2C). Prior to a spawning, hydrated eggs appear to occur at random throughout the cross-section but following ovulation they are concentrated in the lumen which expands as far as the ovary wall and appears as a transparent window along the length of the ovary.

Homogeneity of the ovary

The frequencies of cortical alveoli, yolk granule and yolk fusion stage oocytes in sections from the mid-region of eye-side ovary and the anterior region of the blind-side ovary are compared using a chi-squared test in table 4. The heterogeneity for all fish is shown to be significant and this analysis is consequently carried out for individual fish. No difference appears between samples within maturity stage 4 and within stage 5 or when stages 4 and 5 are combined. However, two of the four maturity stage 6 fish show significant differences. These results show that the two sample sites contain the same densities of vitellogenic oocytes at maturity stages 4 and 5 but that caution has to be exercised when sampling fish after ovulation but immediately prior to the spawning of a batch of eggs. Clearly, the anterior portion of the ovary is not representative of the rest of the ovary at this time and sampling of fish at this particular developmental stage should be avoided.

Determinacy of fecundity

Oocyte size frequency distribution

The numbers of previtellogenic and vitellogenic oocytes, in 25 \( \mu m \) class intervals, ranging from 100-250 \( \mu m \) in pre-spawning sole from three representative divisions (VIIId, IVb west and IVb east) are shown in table 5 (groups 1-3). The number of previtellogenic oocytes in each class interval shows a similar pattern in all three divisions with 1.0% of those counted greater than 175 \( \mu m \). However, the number of vitellogenic oocytes in the different class intervals from the various divisions are considerably different. In the five fish from division IVb west there are only 6 vitellogenic oocytes found in the range 150 to 250 \( \mu m \) compared to 53 in VIIId and 103 from division IVb east. Clearly, the size range 100-250 \( \mu m \) is the part of the oocyte size frequency distribution where we should look for a hiatus in prespawning fish to satisfy our first line of evidence. Also shown in table 5 (groups 4 and 5) are some data from mid-spawning and late spawning-spent fish collected in division IVc. This data together with the data from IVb east, which is a geographically adjacent sample with a similar fecundity (table 3), forms a trend (groups 3, 4 and 5 in table 5) representing maturity stages from just before spawning (stage 4) to spent (stage 7). As spawning progresses there are plainly fewer vitellogenic oocytes present between 100 and 250 \( \mu m \) which would be expected if a hiatus was developing in the size frequency distribution.

Oocyte size frequency distributions of individual sole (length 31-50 cm) with maturity stage 4 ovaries
from divisions VIIa, VIId, VIIe, IVb west, and IVc are shown in figure 4. In none of the ovaries examined were post-ovulatory follicles present and the maximum vitellogenic oocyte diameter was 750 µm or more in all the divisions, indicating that the sole were close to but not yet spawning. An inspection of the histograms (fig. 4) reveals some regional differences. A hiatus in the oocyte size frequency distribution between 100 and 250 µm is not present in any of the sole from divisions VIId and IVc but in division VIIe 25 µm hiatuses are found in two of the five fish (fig. 4, row 3c-200-225 and d-175-200 µm). In divisions VIIa and IVb west all the sole show a hiatus in the oocyte size frequency distribution between 175 and 250 µm. In most cases it is 25 µm in size but on two occasions (VIIa, e, 175-225 µm and IVb west, a, 175-250 µm) it is larger. Also shown in figure 4 are size frequency distributions from spawning fish (50% or more of the predicted annual potential fecundity remaining) and late spawning-spent fish (< 10% of the predicted annual potential fecundity remaining) collected from division IVc. This series of distributions from IVc make an interesting comparison. We found no hiatuses in the pre-spawning sole but four (fig. 4 row 6 a, c-e) of the five sole caught in mid spawning have hiatuses in the oocyte size distributions, one of 25 µm (c), two of 50 µm (a and d) and one of 75 µm (e). In the sample labelled as late spawning-spent, two sole (fig. 4 row 7, c and d) have no vitellogenic oocytes > 250 µm remaining while the three other sole have a reduced distribution of vitellogenic oocytes with hiatuses of between 175 (a), 275 (e) and 300 µm (b).

We can conclude that sole from divisions IVc and VIIId show no hiatuses in the oocyte size frequency distribution but two out of five do in VIIe and all five sole in VIIa and IVb west. However, these hiatuses are only of the order of 25 or 50 µm and may not therefore constitute a true hiatus. The series of samples from IVc (pre-spawning, spawning and spent) however shows that the hiatus develops during spawning. In the near-spent sole a well developed hiatus is present and in the fully spent fish no vitellogenic oocytes remain.

**Rate of recruitment and growth of vitellogenic oocytes in relation to the length of the spawning season**

The samples collected from division VIIId between July 1993 and February 1994 are summarised in table 1. Little change in ovary weight and condition (ovary weight/length³ × 1000) was seen between July and September (fig. 5A, B) but this coincided with a steady increase in the proportion of fish (July 30% August 50%, and September 90%) which have commenced ovarian recrudescence (fig. 5C). Ovarian recrudescence started with the appearance of cortical alveoli stage oocytes but no oocytes with yolk granules were visible. Those fish without cortical alveoli stage oocytes all contain lipid vesicle stage previtellogenic oocytes except one fish in the July sample which only had perinuclear stage previtellogenic oocytes. In several fish there were residual signs of the previous annual spawning in the form of advanced gamma atresia. Between September and October the proportion of fish with yolk granules increased from 20 to 90% respectively marking the start of exogenous vitellogenesis and associated increase in ovary weight.
Fecundity and oocyte atresia in *Solea solea*

Figure 3. - Histological sections (5 μm) of atretic oocytes embedded in methacrylate resin and stained with periodic acid–Shiff’s and Mallory’s trichrome. The scale bar is 50 μm. A, an oocyte containing cortical showing alveoli signs of early alpha atresia, note pieces of the zona radiata interna in the cytoplasm; B, an oocyte containing cortical alveoli showing signs of late alpha atresia; C, a yolk granular oocyte showing signs of early alpha atresia, note the breaks in the zona pellucida; D, a yolk granular oocyte showing signs of late alpha atresia; E, the zona pellucida of a yolk granular oocyte showing alpha atresia. Note the formation of breaks in the zona pellucida and the fact that the zona radiata externa remains intact at this stage; F, a later stage than (E) showing the zona radiata externa ruptured and the contents of the oocyte being discharged towards the follicle. *Zp* = zona pellucida, *ZRI* = zona radiata interna, *ZRE* = zona radiata externa, *F* = follicle, *V* = vesicle containing mainly yolk, *Y* = yolk granule.

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Figure 4. - The oocyte size frequency distribution found in each of thirty five sole (L = fish length cm), shown in seven rows (ICES divisions VIIa, VIIb, VIIc, IVb west, IVc pre-spawning and mid and late spawning fish in IVc respectively) of five fish (a-e) collected during 1991. Each distribution was calculated from an oocyte sample size dependent on the number of oocytes > 100 μm encountered with a nucleus visible until 100, or all, vitellogenic oocytes present in the two ovary sections, had been measured. The smallest size class of oocytes displayed is 150-175 μm and where the frequency exceeds the Y-axis scale in any class the value is shown underlined above.
and condition in the following months. Spawning in the population was not detected in January, either after macroscopic examination of the commercial catch on the market, or in our random sample. However, by February, one stage 5 fish was found at sea whilst collecting the fecundity sample and a small proportion (2%) of the fecundity sample was rejected because of the presence of hydrating oocytes or post ovulatory follicles in the ovary sections.

The growth rate of oocytes comprising the annual fecundity was estimated from September until February, when most fish had commenced ovarian recrudescence, and derived by differentiating the following model fitted to the data in figure 5D:

\[ y = ax^2 + b \]  

where \( a = 1.36 \times 10^{-5} \) (SE ± 1.1 \times 10^{-6}), \( b = 0.030 \) (SE ± 0.011), \( y \) = volume of the leading oocyte cohort, and \( x \) is in days (\( R^2 = 0.742 \) N = 60 \( p < 0.0001 \)). The oocyte growth rate \( Y_g \) (mm\(^3\) per day) was thus:

\[ Y_g = 2ax \]  

Table 5. - The numbers of previtellogenic and vitellogenic oocytes in the size range 100-250 \( \mu \)m from prespawning sole collected from ICES divisions IVb west (1), VIIId (2) and IVb east (3). Also included are sole from ICES division IVc that have 50% (4) and less than 10% (5) of their predicted potential annual fecundity remaining. In each case the results from five soles are combined.

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

(1) Division IVb west (\( N = 5 \), collected 19 March 1971

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

(2) Division VIIId (\( N = 5 \), collected 4-7 February 1971

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

(3) Division IVb east (\( N = 5 \), collected 15 March 1971

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

(4) Division IVc (\( N = 5 \), collected 15 May-15 June 1971

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

(5) Division IVc (\( N = 5 \), collected 15 May-15 June 1971

<table>
<thead>
<tr>
<th>Oocyte diameter (( \mu )m)</th>
<th>Previtellogenic oocytes</th>
<th>Vitellogenic oocytes</th>
<th>Total number of oocytes</th>
<th>Previtellogenic oocytes (%)</th>
<th>Vitellogenic oocytes (%)</th>
</tr>
</thead>
</table>

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and increased from $2.73 \times 10^5$ on 20 September (day 1) proportionally with the number of days elapsed. A period of 143 days from September 20 until February 10 was required for oocytes to grow from 386 $\mu$m to 839 $\mu$m the latter value being close to, but just below the final maturation size.

Two approaches were used to study the recruitment of vitellogenic oocytes to the potential annual fecundity. In the first the rate of recruitment was measured over the whole period from August to February whilst in the second changes in fecundity were studied in the final three months from November to February prior to spawning. In the first approach we used fecundity data (fig. 5E) normalised to a standard length of 35 cm using equation 2. We then fitted a polynomial model to the data.

$$y = ax^2 + bx + c$$  (5)

where $a = -8.32$ (SE = 2.26), $b = 3030$ (SE = 440), $c = 8888$ (SE = 19628), $p < 0.0001$, $N = 70$, were $y$ is the fecundity at 35 cm and $x$ is in days starting on the 3 August when half the population had commenced ovarian recrudescence. The daily rate of production of vitellogenic oocytes can be derived from differentiating the polynomial equation (5) thus:

$$y = 2ax + b$$  (6)

and shows maximum production on the August 3 (3013 oocytes day$^{-1}$) rapidly falling to just below zero by February 10.

Approach 2 focussed on the change in fecundity between November 20 and February 10 and ANCOVA was used to fit the following model with day and log length as covariates:

$$\ln(\text{fecundity}) = a + b \times \ln(\text{length}) + c \times \text{(days)}$$  (7)

where $a = -0.038$ (SE = 1.589), $b = 3.509$ (SE = 0.440), $c = 0.002$ (SE = 0.001), $N = 40$, $p < 0.0001$. If the mean length (37.5 cm) of the four samples is substituted into equation 7 then

$$\ln(\text{fecundity}) = 12.680 + 0.002 \times \text{day}$$

predicts a mean increase of 0.18% per day but this is not significantly different from zero ($p > 0.1$).

There are two further samples (table 1 divisions IVc March 1 and IVb east March 15 1991) collected in the central southern North sea which were amenable to the same analysis. These samples were from locations 200 km and 14 days apart and almost certainly from the same spawning stock with a very similar potential annual predicted fecundity at length (table 3). The parameters of the equation 7 were estimated as $a = -0.477$ (SE = 0.500), $b = 3.782$ (SE = 0.142), $c = 0.002$ (SE = 0.003), $N = 95$, $p < 0.0001$. Substituting the mean fish length (35.1 cm) into the equation predicts an increase in fecundity of 0.2% per day but again this was not significantly different from zero ($p > 0.5$).

Figure 5. - Measurements of ovarian recrudescence in eight samples of 10 female fish collected at monthly intervals commencing 19 July 1993 and finishing 10 February 1994 just prior to spawning. (A) Mean ovary weight + one standard deviation. (B) Mean ovary condition (ovary weight/length$^3 \times 1000$) + one standard deviation. (C) Maximum recorded vitellogenic oocyte diameter. Between July and September a number of fish, recorded just above the X axis, had no oocytes containing cortical alveoli. (D) The growth in volume of the largest oocyte, containing either cortical alveoli or yolk granules, found in the monthly samples of fish. A model $ax^2 + b$ describing the growth in oocyte volume from 20 September is fitted where $a = 1.36 \times 10^{-5}$, $b = 3.0 \times 10^{-2}$ and $x$ is days. (E) The increase in fecundity from 3 August found in the monthly samples of fish.
Changes in the number of vitellogenic oocytes in the ovary during spawning in sole from ICES division IVc

In order to study the change in the numbers of vitellogenic oocytes during the spawning season, two samples were compared (table 1), the first collected prior to the spawning season (1 March 1991, \( N = 55 \)) and the second around peak spawning (15/16 May 1991, \( N = 93 \)). The ratio of the observed to the expected fecundity (table 3) at length in the first sample was 1.00, SD ± 0.25 and in the second sample it was 0.27, SD ± 0.20, demonstrating a reduction in the number of oocytes present. We then examined the sample collected towards the end of the spawning season in more detail (fig. 6). 88% of the sample retained half or less of their predicted annual potential fecundity which included 22% near or completely spent fish the latter having no cortical alveoli or yolk granule stage oocytes.

Batch fecundity

Estimates of batch fecundity from 38 fish (fig. 7) varied between 1000 and 20000 oocytes, but there is no significant relationship between batch fecundity and fish length (\( R^2 = 0.12 \)). The mean batch fecundity was 8400 (SE ± 1363) oocytes at a mean fish length of 31.6 cm. The majority (88%) of fish were 4 years old and contained 0.333 (SE ± 0.005) of their predicted annual fecundity.

Figure 6. – Analysis of a sample of sole (\( N = 93 \)) collected from ICES division IVc on 15/16 May 1991 showing the proportion of the potential annual fecundity remaining in the ovary.

Estimation of atresia

Levels of atresia prior to spawning

The prevalence and relative intensity of atresia in fecundity samples collected prior to spawning in areas IV and VII are shown in table 6. The prevalence varied between the lowest value (0.044) found in IVb west and the highest (0.690) in VIIa. Intermediate values were found in IVc (0.182), IVb east (0.175), VIIId (0.082) and VIIe (0.060).

In ovaries with atresia the relative intensity was low. The highest value was recorded in sole from division VIIa (0.076) with values from other divisions varying between 0.017 and 0.060.

Levels of atresia during spawning

If we can again accept that the sole has a determinate fecundity then progress through spawning can be measured by the decline in standing stock of vitellogenic oocytes during spawning. On this basis, the spawning period can be divided into three parts based on the portion of the predicted annual potential fecundity remaining in the ovary: pre-spawning to 50% spawned, 51 to 75% spawned, and 76-99% spawned. The results from division IVc are shown in table 7 with the 3 completely spent fish appended. The prevalence of atresia increased slightly as spawning progressed from 0.45 in the first half, through 0.52 in the middle to 0.57 in the final stages of spawning but declining to zero in the spent group. However, the relative intensity of atresia rises from 0.035 in the first period to a peak of 0.058 during the second portion and declines to 0.022 during the final portion. A one way ANOVA was used to
Table 6. – The number of fish with atresia in fecundity samples of pre-spawning female sole collected during 1991 from ICES areas IV and VII. The prevalence of atresia is shown together with the geometric mean of the relative intensity of atresia in each division.

<table>
<thead>
<tr>
<th>ICES division</th>
<th>No. of fish sampled</th>
<th>No. of fish with atresia</th>
<th>Prevalence</th>
<th>Relative intensity (± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IVb east</td>
<td>40</td>
<td>7</td>
<td>0.175</td>
<td>0.0178 (± 0.0072)</td>
</tr>
<tr>
<td>IVb west</td>
<td>45</td>
<td>2</td>
<td>0.044</td>
<td>0.0228 (± 0.0171)</td>
</tr>
<tr>
<td>IVc</td>
<td>55</td>
<td>10</td>
<td>0.182</td>
<td>0.0214 (± 0.0072)</td>
</tr>
<tr>
<td>VIIa</td>
<td>29</td>
<td>20</td>
<td>0.690</td>
<td>0.0764 (± 0.0181)</td>
</tr>
<tr>
<td>VIIId</td>
<td>49</td>
<td>4</td>
<td>0.082</td>
<td>0.0285 (± 0.0151)</td>
</tr>
<tr>
<td>VIIle</td>
<td>33</td>
<td>2</td>
<td>0.060</td>
<td>0.0601 (± 0.449)</td>
</tr>
</tbody>
</table>

Table 7. – The number of fish with atresia in relation to the progress through spawning in sole from division IVc (15-16 May 1991). A spawning season of 60 days is divided into three parts (30, 15, 15 days) proportional to the reduction in the annual predicted potential fecundity. The mean loss from the predicted annual potential has been calculated as: Prevalence x geometric mean of the relative atresia intensity x spawning duration/duration of early alpha atresia (9 days). No early alpha atresia remained in the spent fish.

<table>
<thead>
<tr>
<th>Progress through spawning (%)</th>
<th>Duration of spawning (days)</th>
<th>No. of fish sampled</th>
<th>No. of fish with atresia</th>
<th>Prevalence</th>
<th>Geometric mean of the relative intensity of atresia (± SE)</th>
<th>Mean loss from potential (weeks) over period</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-50</td>
<td>30</td>
<td>11</td>
<td>5</td>
<td>0.45</td>
<td>0.035 (± 0.018)</td>
<td>5.25</td>
</tr>
<tr>
<td>51-75</td>
<td>15</td>
<td>42</td>
<td>22</td>
<td>0.52</td>
<td>0.058 (± 0.014)</td>
<td>5.03</td>
</tr>
<tr>
<td>76-99</td>
<td>15</td>
<td>37</td>
<td>21</td>
<td>0.57</td>
<td>0.022 (± 0.006)</td>
<td>2.11</td>
</tr>
<tr>
<td>&gt; 99 (Spent)</td>
<td>3</td>
<td>3</td>
<td>0</td>
<td>0.00</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>60</td>
<td>93</td>
<td>48</td>
<td></td>
<td></td>
<td>12.39</td>
</tr>
</tbody>
</table>

The size frequency distribution of alpha atretic oocytes during spawning

There are some obvious problems associated with measuring the size of atretic oocytes. The outline of atretic oocytes is often irregular and the nucleus degenerates progressively, it is therefore difficult to measure their size precisely. This may account for wider size frequency distributions than is usually the case with groups of healthy oocytes, also some shrinkage will inevitably have occurred. The problem was reduced by measuring only early alpha stage atretic oocytes. The size frequency distribution of early alpha atretic oocytes from division IVc collected between 15 May and 16 May 1991 is shown in figure 8. Nineteen fish (length 33.3 cm, SD ± 3.2) were selected where > 0.015% of relative atresia was present in the ovaries and 10 atretic oocytes were measured in each fish. The mean fecundity of the sample was 142,263, SD ± 88,926 oocytes and this constitutes 0.37, SD ± 0.018 of the predicted annual potential fecundity. The size frequency distribution was between 100 and 525 µm centred around 250 µm.

DISCUSSION

Traditional methods of estimating fecundity employ volumetric (Burd and Howlett, 1974) or gravimetric (Simpson, 1951) techniques. In these methods, the oocytes from both ovaries are released from the connective tissue by chemical and mechanical means.
and a number of subsamples taken and subsequently the result is raised to the whole ovary by volume or weight. However, when stereological methods are used the number of oocytes in the ovaries is calculated from only one or two sample sites. Rosenboom (1985) has shown that the eye-side ovary of the sole can be up to twice the size of the blind-side ovary and this feature has raised doubts as to whether, from the point of view of sampling procedure, the two ovaries have the same densities of oocytes. A comparison (Emerson et al., 1990) using a stereological method to count the oocytes in one ovary and a volumetric method and an automated particle counter to count the oocytes in the other ovary of the same sole showed no significant differences in density between either the three methods or the two ovaries.

Horwood (1993) approached this problem in a different way taking samples alternatively from the eye-side and blind-side of fish. An analysis of covariance of fecundity with length showed that results based upon the blind-side ovaries were 18% greater than those using the eye-side ovaries. This difference although relatively large was not significant. Here we have examined the density of three vitellogenic oocyte development stages from the two ovaries and find no significant difference in oocyte density in maturity stage 4 and 5 ovaries. The current practice of taking sections from the mid-point of both ovaries would therefore seem to provide a representative sample. However, our results also show that during production of a batch of eggs when the hydrated oocytes are concentrated in the lumen and at the anterior end of the ovary close to the cloaca (fig. 2) a section in that area would not be representative of the whole ovary.

The present sampling programme was conducted over a wide temporal and spatial range (fig. 1) and the question arises as to how isolated or discrete were the populations in the various sampling areas. However, tagging data reported in Anonymous (1989) concluded that "when a sole has spawned on a ground it remains associated with it subject only to seasonal movements". Nil or low values of interchange were reported between the spawning groups between the north and south coasts of the eastern English channel.

As regards the hiatus in the oocyte size frequency distributions it has been shown to exist in sole from division VIIb by Horwood and Greer Walker (1990). These sole were assessed as being prior to spawning using the absence of hydrated oocytes as a criterion but were not histologically screened. We have collected pre-spawning sole over a much wider area and found evidence for a small hiatus in the adjoining division VIIa and IVb west (see also Deniel, 1981) but this was not so for every division (table 5 and fig. 4). However, in sole from IVc, where there was no suggestion of a hiatus in pre-spawning fish, a large hiatus develops during spawning. We suggest that the degree of development in a hiatus prior to spawning may be related to area differences in fecundity which are independent of length (Anon, 1992). The geographic differences in relative fecundity, reported above, also appear to be inversely related to the egg size found in plankton samples collected in each ICES division (Anonymous, 1992; Rijnsdorp and Vingerhoed, in press). Our results therefore confirm those of Urban (1991) in that there is no clear hiatus in the size frequency distribution of oocytes in sole prior to spawning from division IVb east and IVc. The results from the other divisions are more difficult to interpret but there does seem to be consistent evidence for the development of a hiatus either closer to or during spawning. However, we believe that other criterion for determinacy should be considered where the species in question has an indistinct hiatus which develops more clearly during spawning as is the case with the sole.

The second line of evidence we consider for a determinate fecundity is related to the rate of growth and recruitment of vitellogenic oocytes in relation to the spawning season. Oocytes take 143 days to grow from 386 to 839 μm (fig. 5D) which is much longer than the estimated spawning duration of 60 days (Anonymous, 1992), but their daily growth rate increased rapidly reaching 3.91 × 10⁻³ mm² per day on February 10. The question is; what growth rate would be realised by an oocyte commencing vitellogenesis in February, or later, and would it be fast enough to produce a viable egg within the estimated 60 day spawning duration? The observed increase in oocyte growth rate must be dependent on the increase in surface area of the zona pellucida and follicle capillary bed which parallels the change in

Figure 8. – The size frequency distribution of alpha atretic oocytes (N = 191) of Solea solea (N = 10 per fish) caught in ICES division IVc during 15-16 May 1991. Fish were selected if the relative intensity of atresia found in their ovaries was > 0.075%.

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oocyte volume. We must also consider the effect of seasonal temperature changes which increased 3.7°C from 5.0°C on February 10 to a mean of 8.7°C during the March-April spawning period (Directorate of Fisheries Research Lowestoft sea surface temperature database). Several cellular processes associated with vitellogenesis are influenced by the metabolic rate (Moomsen and Walsh, 1988), and therefore the oocyte growth rate should be adjusted for changes in temperature using the \( Q_{10} \) equation (Schmidt-Niesen, 1975; Kjesbu, 1994).

\[
R_2 = R_1 Q_{10}^{(T_2-T_1)/10}
\]  

(8)

where \( Q_{10} \) is 2.3, \( R_1 \) is the rate at temperature \( T_1 \), and \( R_2 \) is the rate temperature at \( T_2 \). If we assume a 60 day spawning duration, at a mean temperature of 8.7°C, then a mean daily growth rate of \( 5.08 \times 10^{-3} \text{ mm per day} \) would be required for an vitellogenic oocyte to grow from 200 \( \mu \text{m} \) to maturity at 839 \( \mu \text{m} \). Applying a temperature correction amounting to 3.7°C in equation 8 the revised growth rate of \( 3.74 \times 10^{-3} \) would be equivalent to that measured at 137 days (equation 4) which was found in oocytes of 817 \( \mu \text{m} \) using equation 3. It would appear therefore that the maximum growth rate found would only just allow the smallest oocyte comprising the potential annual fecundity in February to reach maturity. Thus the asynchronous pattern of oocyte development in the sole originates from both vitellogenic oocyte recruitment, which takes place over several weeks (fig. 5E), and the acceleration of oocyte growth rate during development.

The fecundity recruitment rate of 3013 oocytes per day measured on August 3 is probably an underestimate because the sample comprises some individuals which have still to start the recruitment process. We observed a steady increase in the proportion of fish commencing ovarian recrudescence from July 19 to September 20. In early August 50% of the population had not commenced production of cortical alveoli oocytes and they would suppress the observed fecundity recruitment rate. However, in the three months prior to spawning, from November to February, we found no evidence that sole from division VIIId produced any significant increase in fecundity. A similar result was obtained in divisions IVb east and IVc. The small but not significant increase in fecundity with time, which was detected, may stem from fish which are the last individuals in the population to commence vitellogenesis. These individuals can be identified by the relatively small size of oocytes comprising the leading oocyte size cohort (Kjesbu, 1994). We conclude that the oocyte recruitment process appears to have finished well before spawning commences. Although this does not prove that recruitment of vitellogenic oocytes does not occur during spawning when taken together with the information we have on oocyte growth rates and the developing hiatus it would seem highly unlikely they could mature during a 60 day spawning period.

In our third line of evidence we consider the spawning process. We observed a marked decline in the number of vitellogenic oocytes in the ovary during spawning (fig. 6). This sample taken on the 15th May was probably taken from a population which we estimate had been spawning for 5 to 6 weeks based on data in Anonymous, 1992.

A comparison with other species which have been classified as having an indeterminate annual fecundity is also useful in assessment of the spawning strategy of Solea solea. A study of the gonadosomatic index has shown that the closely related sand sole, S. lasecarius and S. impar, from western Brittany have estimated spawning seasons of 135 and 105 days respectively (Deniel et al., 1989). These two species show a continuous recruitment of oocytes into vitellogenesis during the spawning season and therefore an indeterminate fecundity. The continued maturation of these oocytes is presumably possible because of the longer spawning season and warmer water temperatures. More extreme examples of fish with indeterminate fecundities are the northern anchovy (Engraulis mordax) and the queenfish (Seriphus politus) which have spawning seasons of 150 days (Lasker and Smith, 1977) and 180 days (De Martini and Fontaine, 1981) respectively. In both cases the number of eggs spawned is more than an order of magnitude greater than the standing stock of oocytes present prior to spawning (De Martini and Fontaine, 1981). However, sole from division IVb east (and possible IVc also) appear to develop a hiatus after the beginning of spawning and we have suggested that this late developing hiatus could be associated with the larger potential fecundities found in these areas possibly because the more fecund fish produce vitellogenic oocytes over a longer time period, and closer to spawning, such that the hiatus develops later. Alternatively, it is possible that as the winter temperatures (Jones and Jeffs, 1991) and therefore the oocyte growth rates are higher in the West of the study area that the hiatus in the oocyte size frequency distribution may develop earlier. In this respect, it would be interesting to compare the oocyte growth rates in the different areas; it would seem unlikely that the sole would have determinate and indeterminate fecundities in adjacent stocks.

It is possible in division IVb east to compare the potential annual fecundity prior to spawning with the realised fecundity calculated from the product of the batch fecundity, the batch frequency and the duration of the spawning season. The mean batch fecundity (8400, SE \( \pm 1363 \) ) found was to be independent of length and therefore atypical to data published on other teleosts. (Hunter and Goldberg, 1980; Urban, 1991; Watson et al., 1992). A similar result (mean batch fecundity 7600 SE \( \pm 1035 \) and independent of length) was reported by Urban (1991). We conclude like Urban (1991) that the low variability of batch fecundity with length maybe attributable to the narrow size range of small fish which were also caught towards the
end of their annual spawning season. Devauchelle et al. (1987) measured the batch size in captive sole (Solea solea) varying both the temperature and the day length. The results showed that batch size was highly variable (1 500-138 000 eggs per batch) and in some circumstances much larger than that recorded in wild populations. There is little information available on batch frequency. Houghton et al. (1985) reported that three captive sole in the same tank in the laboratory spawned on 22 nights over a period of 47 nights and on this basis a frequency of two days was adopted which is consistent with the incidence of fresh postovulatory follicles in sole from the German Bight (Anonymous, 1992). Le Bec (1983) has suggested that sole from the Gulf of Gascony produce between 8 and 12 batches of oocytes during the spawning season on a weekly basis but there are no direct measurements to support this conclusion. As regards the duration of spawning, Horwood (1993) used a duration of 40 days for sole from division VIII as being compatible with the available evidence from that area. However, cumulative frequency plots of maturity stages 5 and 6 from the Dutch market sampling programme (1970-79) indicate a spawning duration of 60 days in the Southern Bight and the German Bight (Anonymous, 1992) and this was thought to be more relevant on a geographical basis to the present investigation. The realised annual fecundity based on the product of batch size, spawning frequency and duration is 252 000 which is + 94% and - 30% respectively around the predicted potential annual fecundity (178 000-480 000) for the sampled length range (table 3). The predicted potential fecundity (which does not include losses through atresia, table 7) for most of the population is thus above or about the same as the batch size based estimate and these results are consistent with the fact that the fecundity of sole from division IVb east is determined prior to spawning. We therefore reach the same conclusion as Deniel et al. (1989) that for all practical purposes the sole should be regarded as having a determinate fecundity over the geographic range studied.

Follicular atresia has been described in a wide range of species (Guraya, 1986) but little attention has been paid to studying atresia as a method of regulating fecundity. In order to quantify oocyte losses through atresia it is necessary to estimate the rate of turnover, bearing in mind that small numbers with a rapid rate of turnover found at any instant in a preserved ovary, could significantly alter our estimate of the number of oocytes resorbed and that the process is temperature dependent. Turnover rates have been estimated as 7-9 days in anchovy (Hunter and Maciewicz, 1985b), 10 days in cod (Gadus morhua) (Kjesbu et al., 1991) and 7.5 days in mackerel (Scomber scombrus) (Anonymous, 1993) at ambient temperatures. We adopted a stage duration of 9 days as used by Horwood (1993) but the truncated stage may have a shorter duration and this is the subject of current research.

We surveyed the extent of atresia in pre-spawning (table 6) and spawning sole populations table 7. The prevalence of atresia in pre-spawning fish (maturity stage 4) in divisions IVc-e, VIIId + e ranged from 0.04-0.18 (N = 222), and in division VIIa 0.69 (N = 29). The value for prevalence in division VIIa is obviously high and this is also the case when the values for relative intensity from the various divisions are compared (division VIIa = 0.076, area IV and VII range of 0.017-0.06). These values compare with a prevalence of 0.10 and relative intensity of 0.01 (N = 50) in division VIIIf during 1990 (Horwood, 1993). A comparison of atresia in spawning fish caught in division IVc in 1991 and VIIIf in 1990 strongly suggests area and or annual differences in the degree of fecundity regulation by atresia. A prevalence in IVc of 0.45, 0.52, and 0.57 depending on spawning progress compared with 0.50 and 0.90 in maturity stages 5 and 6 sole respectively from VIIIf. The relative atresia in sole from division IVc appeared to be much less abundant with values of 0.035, 0.058, and 0.022, depending on spawning progress, compared to 0.069 in maturity stage 5-6 sole reported by Horwood (1993) for VIIIf in 1990. We found a similar trend in the relative intensity of atresia as Greer Walker et al. (1994) in mackerel (Scomber scomber L.) which suggests that this may have a more general role in the regulation of fecundity and spawning strategy in some species that have a determinate fecundity. The anchovy, in contrast, which has an indeterminate spawning strategy shows maximum levels of atresia towards the end of spawning (Hunter and Maciewicz, 1985b).

Our measurements of alpha atretic oocytes in spawning fish clearly show a narrow size frequency distribution with a peak around 250 μm (fig. 8) when compared with the oocytes comprising the residual potential fecundity (fig. 4 division IVc spawning). These smaller oocytes may become atretic as a consequence of the relatively slow growth rate in the smallest (200-350 μm) cortical alveoli stage oocytes. It is unlikely that they could maintain a growth rate equivalent to an oocyte of 817 μm diameter to reach maturity during the 60 days spawning season and there maybe insufficient supplies of yolk present (Kjesbu et al., 1991). Whatever the cause of their demise their removal would contribute to the formation of an increasingly large hiatus in the oocyte size frequency distribution between previtellogenic and vitellogenic oocytes.

Our results can be compared with those of Horwood (1993) from the Bristol Channel (division VIIIf) who reported the predicted annual potential fecundity would be reduced by 30.5% due to atresia. In the case of mackerel (Anonymous, 1993), it was calculated that the reduction would be 8.8%. An extreme instance of fecundity regulation through atresia was found in tank-held cod which accumulated a potential annual fecundity that is more than twice that of wild cod but
36% of the eggs were lost through atresia (Kjesbu et al., 1991).

Some major differences in the values of atresia are therefore evident as the ovary passes from maturity stage 4 to 7 which may differ between geographic areas within one year, and possibly between years. Clearly, high levels of atresia in the ovaries would seem to be an inefficient use of resources and it is likely that atresia plays some part in regulating the potential fecundity. Anthropic effects on ovarian recrudescence and occurrence of atresia have been reported in specific cases (Brule, 1987; Johnson et al., 1992), and this too may interact with the natural regulation.

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