
RICARDO SERRÃO SANTOS


The main histologic aspects of the reproductive organs of *Parablennius sanguinolentus parvicornis*, a rocky intertidal pool fish of the Azores with alternative life history tactics, are described and illustrated. Eight stages of oocyte development (plus atretic ones) are considered. The development of the ovaries is classified into 6 stages. Ovaries may contain, according to female size, from 4000 up to 41000 oocytes in the month which precedes the reproductive season. Gonad maturation of the males is assigned to 8 stages. Seasonal variations of gonad histology, of both males and females, are illustrated. Histology of the livers shows that they play an important role in the storage of energy. Lipidic reserves are used while the breeding season progresses.

SANTOS, RICARDO SERRÃO 1995. Anatomia e histologia dos caracteres sexuais secundários, gónadas e fígado do peixe que habita as poças rochosas do estrato entre-maris dos Açores. Esta espécie apresenta táticas alternativas de acasalamento. São considerados e descritos oito estádios de desenvolvimento dos óócitos (mais um estádio degenerativo). O desenvolvimento dos ovários é classificado em seis estádios. Os ovários, no mês anterior ao início da postura, contêm, conforme o tamanho das fêmeas, entre 4000 e 41000 óócitos em diferentes estádios de desenvolvimento. O processo de maturação das gónadas dos machos é classificado em oito estádios. São ilustradas, através de preparações histológicas, as variações sazonais do desenvolvimento das gónadas. A histologia hepática mostra que o fígado desempenha um importante papel como reservatório energético. As reservas lipídicas, acumuladas durante o Inverno e Primavera são gastas durante a curta estação de reprodução.

Ricardo Serrão Santos, Universidade dos Açores, Departamento de Oceanografia e Pescas, PT-9900 Horta, Açores, Portugal. - Unidade de Estudos Oceânicos IMAR, PT-9900 Horta, Açores, Portugal.

INTRODUCTION

The reproductive biology and phenology of several species of blennies and their close relatives of the north-eastern Atlantic and the Mediterranean and adjacent seas, have been studied from a variety of viewpoints (see GIBSON 1969, 1982; ALMADA & SANTOS 1995). Most of the studies were concentrated on populations from the British Isles, Mediterranean and Adriatic Seas.

Blennies, together with gobies, comprise the majority of the resident (sensu GIBSON 1969) fish species in the shallow water of these regions (GIBSON 1969, 1982; ZANDER 1972; ARRUDA
1990; Santos et al. 1994). They all lay demersal eggs and provide parental care to the developing embryos. Their general biology was reviewed at length by Gibson (1969, 1982, 1986).

In general, the blennies live more than two years and occasionally up to ten or thirteen years (e.g. Lipophrys pholis: Qasim 1956b; Dunne 1977). Most of them are iteroparous; sexual activity extending for two or more years. In temperate zones the reproductive season lasts for two or more months, with several spawnings (Qasim 1956a, 1956b; Shackley & King 1977a; Santos 1992). At lower latitudes females may spawn all the year round (Nursall 1977, 1981; Johannes 1978). Eggs are attached to the substratum by adhesive discs (e.g. Santos 1989) or threads.

The testes of blennies and gobies are unique among teleosts for possessing a special gland. In blennies the testicular gland has cells containing lipids (Patzner & Seiwald 1987c; Seiwald & Patzner 1987). The spermatids mature in the testicular gland where they receive nutrients (Lahnsteiner & Patzner 1990a, 1990b; Lahnsteiner et al. 1990). The testicular gland is also an important source of mucus secretion, which is added to the spermatozoa. At the end of the spawning season it is involved in the phagocytosis of remaining spermatozoa (Lahnsteiner & Patzner 1990b; Lahnsteiner et al. 1990).

There are often structures used to enhance behaviour associated with reproduction. Colour patterns of males frequently change during the reproductive phase (temporary dichromatism), with development of strikingly conspicuous colour marks (head masks) (Gibson 1969). Most species of blennies possess special structures in the head region such as tentacles and crests. Crests are usually present in breeding males, or, if present in both sexes, they are specially developed in males during the reproductive season (Qasim 1956a; Papaconstantinou 1979; Patzner et al. 1986; Patzner & Seiwald 1987b; Almada 1989). Male anal glands are distinctive secondary sexual characters in many species (Scartella cristata by: Smith 1974; Parablennius gattorugine: Kotrschal & Goldschmid 1983; P. pilicornis: Deniox 1984; P. ruber: Santos 1987; Salaria pavo: Patzner & Seiwald 1987a, 1987b, and see also Zander 1975). Their function is still obscure.

Parablennius sanguinolentus parvicorns (Valenciennes in Cuvier & Valenciennes 1836) is the dominant resident rock-pool intertidal species in the Azores (Santos et al. 1994). It is one of the few species of blennies (the other being Salaria pavo, Almada et al. 1994, 1995) which is known to have developed alternative mating tactics, in which the large males act as parentals, while the small males stay as satellites of the parental territories (Santos 1985; Santos & Almada 1988). The demography and growth, and the biometry of reproductive phenology of this species was studied in detail (Santos et al. 1995, 1996). Aggressions in the context of reproductive phenology, both in males and in females, were analysed by Santos & Nash (1996).

In this paper we provide a description of gonad development in the general context of the phenology of P. s. parvicorns, with a view to enhance detail to the biometric phenological analysis presented by Santos et al. (1996). The histology of the gonads and liver are examined in relation to the process of maturation and reproductive effort. From this the pathways of energy allocation and transfer can be elucidated. It is well known that some of the lipids deposited in the oocytes during maturation do not come directly from ingested food, but are transferred from lipid stores in the liver and muscles (Zahn 1959; Larson 1974; Htun-Han 1978; Crupkin et al., 1988). It has been shown in Lipophrys pholis (Shackley & King 1977a, 1978, 1979) that the yolk incorporates exogenous protein during oocyte development, which was synthesized in the liver.

In the present work other morphological and histological aspects associated with reproduction have also been considered, such as the development of the anal glands and testicular structures. Some of these subjects are poorly understood for many other blennies.
MATERIAL AND METHODS

1. Sampling

A total of 2,580 fishes were sampled monthly between January 1987 and December 1988 with and without the anaesthetic Chinaldin® (Merck) in rock pools at Feteira on the south coast of Faial Island, Azores. From these, the gonads of ca. 200 males and females, and the livers of ca. 100 males and females were prepared for histology. Total length and weight of the body, gonads and livers were obtained from recently dead fishes (always killed with an overdose of the anaesthetic Quinaldine). Width (diameter) of male anal glands (first and second transformed rays of anal fin - see Fig. 1) were measured with a caliper.

2. Preservation and preparation of material

Gonads and liver were preserved for histological studies and further examination. One or both of the gonads and the whole liver or a part of it, depending on its size, were preserved in aqueous Bouin's solution (COSTA & CHAVES 1943) for a period of between two and four days. Subsequently they were prepared for histological examination (following the standard histological series, adapted from COSTA & CHAVES 1943), by being stained with the eosine/haematoxiline method (after COSTA & CHAVES 1943).

Development of the gonads and the livers were described from histological preparations and relative weights. Diameters of oocytes at different stages as well as other structures (e.g. the thickness of adherent discs) were measured with an accuracy of ±1 μm. Diameter of fat vacuoles of the liver were measured in some histological preparations.

Sub-samples of the gonads of known weight (WSG) of 33 pre-spawning mature females were preserved in Gilson solution to enable separation of the oocytes for counting and measuring (KARTAS & QUIGNARD 1984). Since Gilson solution was observed to reduce egg diameter, 50 eggs, from two different females, were measured in fresh condition and remeasured after immersion. The mean reduction (mean ± standard deviation) of the egg diameter was 21 ± 5%.

Estimates of the total number of oocytes per female were made using a sub-sampling technique. All the oocytes were put into a small receptacle which was divided into 200 quadrats and twenty of these quadrats (NQs) were randomly selected. All the oocytes they contained (NEQs) were counted and measured. Oocytes were measured to ±1 μm accuracy using a Mitutoyor profile projector (PJ-100). Total number of oocytes in the sample (NES) was estimated as NES = NEQs/NQs x 200. Total number of oocytes per female (TFE) was estimated as TFE = GW/WSG x NES, where GW is gonad weight.

RESULTS

1. General morphology

1.1. Female reproductive organs

The female gonad is composed of two sausage-shaped ovaries, that extend along the body cavity in a dorsal position above the intestines and below the bladder (Fig. 2). Colouration varies from transparent to milky white in immature females to purple in ovaries with fully developed oocytes. Separate oviducts emerge from each gonad. These connect anterior to the single genital opening which is horseshoe shaped and posterior to the anus.

The maximum diameter of mature oocytes was 0.9 mm to 1.1 mm. In a single ovary there were oocytes in several stages of development. The adherent disc, a structure that enables the egg to stay fixed to a surface after spawning, was distinguishable during early egg development being fully developed by the end of oocyte maturation.
Fig. 1. A- Parental (above) and satellite (below) males of P. s. parvicornis. B- ventral internal view of the males (parental above, satellite below) to show the gonads and liver. C- Ventral external view of the males (parental above, satellite below) to show the urino-genital papilla, the anus and the anal glands.
1.2. Male reproductive organs and secondary sexual characteristics.

Males can be easily distinguished by external characters such as the shape and structure of genital papillae. The males have three openings on the urinogenital papillae, all posterior to the anus. Their arrangement is transverse to the body (Fig. 1). This is apparent very early being present just after metamorphosis. Small individuals need to be examined with a hand lens to confirm the shape of the genital papillae.

Larger males, in reproductive condition, have other obvious characters that make the distinction from females easy. They tend to be much darker, even black, during the territorial/parental phase. The profile of the head is more rounded and proeminent in males, probably due to the storage of fat tissues. Males also display two distinctive anal glands on the first and second anal fin rays.

Fig. 2. A- Female of Parablennius sanguinolentus parvicornis. B- Ventral external view of the female, showing the urino-genital papilla, the anus and the beginning of the anal fin. C- Ventral internal view of the female to show the liver and gonads.
Development of anal glands is ontogenetic and seasonal (Fig. 3). Small individuals have very small glands, which are difficult to distinguish (Fig. 1). They are more apparent during the summer due to a darker colouration. In bigger animals glands are highly reduced after the reproductive season but undergo rapid growth in May.

![Image](image_url)

**Fig. 3.A**- Monthly variations of the width diameter of the first anal gland of the males. **B**- Relationship of the width diameter of the first anal gland of the males and their total length, $r^2$ (May to August)=0.72; (September to April)=0.44.

The testes of *P. s. parvicornis* (Fig. 1) are similar to those of other Blenniidae composed of paired elongated bodies situated below the kidney and consisting of tubules of the unrestricted spermatogonial type (GRIER et al. 1980; GRIER 1981; PATZNER & SEIWALD 1987a). When mature their colour is milky white. The *vas deferens* emerge posteriorly from both testes.

Like in all other Blenniidae, the testes have other specialized structures. The most obvious is the testicular gland, which is situated ventrally in the testes. Its colour is milkish-rose. The *vas deferens* are connected with the testicular gland and conduct the sperm to the genital openings.

2. Development

2.1. Ovaries

The eight stages of development of the oocytes are described with detail in Table 1, and illustrated in Figure 4. The six stages of macroscopic gonad maturation, which are directly related with gonadosomatic indices (see SANTOS et al., in press) are described in Table 2. Oocyte stages compared with the ovary condition are shown in Table 3.

2.2. Testes and accessory organs

Male gonads of *P. s. sanguinolenrus* are composed of two main, and very distinctive components: the testis and the testicular gland (Fig. 1 and Fig. 5). The testicular gland is situated ventrally in relation to the testis. It has connections to the tubules and the *vas deferens* (see Fig. 5). It is composed of tubuli, which are separated by cell membranes. In ripe gonads, tubules are easily distinguishable in the testicular gland as filled with sperm (see Fig. 5). The *vas deferens* are located at the end of the testicular gland. Blood vessels and blood cells are easily distinguishable (see Fig. 5).

Both small (i.e. non-parental males, including satellites) and parental males have well developed gonads during the months of reproduction. Mature spermatozoa are present in the *vas deferens* before spawning in April and May). This indicates that male gametes mature before female gametes. After the spawning season, in September and October, mature spermatozoa still remain in the *vas deferens*. Six stages of gonad development were characterized (Table 4).
The size of the testis, relative to the testicular gland, was much bigger in small mature non-parenatal males (e.g. satellites) (mean±s.t.d. in May with n= 16: 6.80±1.1) than in older parental males (May, n= 14: 1.84±1.27). The testis shrink substantially after the reproductive season. The proportional relationship between testicular gland/testis clear distinct before and after the reproductive season (May, n= 33: 0.41±0.24 and in September, n= 14: 2.46±0.41).

3. Preliminary estimates of “fecundity”

The number of oocytes in the ovary varied from just under 4,000 in a female 8.50 cm total length, to over 41,000 in a female 17 cm total length (mean±s.d.= 13,420±5,117). In May, only a small proportion of oocytes belonged to the mature size classes. Being a multiple spawner, it is likely that only a small proportion of the oocytes in the ovary are released at any one time. Therefore, in a multiple spawning species such as this, absolute fecundity in terms of eggs released, is difficult to estimate (KARTAS & QUIGNARD, 1984). It was found that the number of eggs (F) was best correlated with total weight (TW) (F= 3606.4+399.176 TW, \( r^2 = 0.56, p< 0.001, n= 33 \)) rather than with gonad weight (GW) (F= 8102.63+3036.243 GW, \( r^2 = 0.32, p< 0.001, n= 33 \)) or total length (TL) (F= -18091.7+2549.504 TL, \( r^2 = 0.46; p< 0.001; n= 33 \)). The relationship between number of eggs and total length is best expressed by the function: F=63.096 TL^{2.129} (r^2=0.53, p<0.001).

<table>
<thead>
<tr>
<th>Table 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scale of oocyte development of <em>P. s. parvicornis</em>.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Stage</th>
<th>Name and Size</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Oogonia (up to 30 µm)</td>
<td>Present singly or in small nests in the germinal epithelium. Nucleus large. No nucleoli visible. Cytoplasm dark.</td>
</tr>
<tr>
<td>II</td>
<td>Primary oocyte (30 to 100 µm)</td>
<td>Oocytes already surrounded by a thin layer of follicular epithelium. Nucleus located in the middle of the cell and surrounded by nucleoli. Cytoplasm dark.</td>
</tr>
<tr>
<td>III</td>
<td>Oocytes between 100 to 150 µm.</td>
<td>Easily distinguishable in histological preparations since the cytoplasm is much lighter. Nucleus, located in the middle of the cell, surrounded by several nucleoli at the periphery. Development of the adherent disc recognizable at one pole of the oocyte forming a layer, 10 µm thick.</td>
</tr>
<tr>
<td>IV</td>
<td>Oocyte diameter about 230 µm</td>
<td>Cytoplasm appears much lighter and a few small vacuoles distinguishable. Yolk development begins. Nucleoli dispersed all over the nucleus. The thickness of the germinal chorion around 3 µm. Thickness of the adherent disc around 17 µm.</td>
</tr>
<tr>
<td>V</td>
<td>Oocyte diameter about 300 µm</td>
<td>Cytoplasm nearly filled with vacuoles, except for a ring circling the nucleus. Nucleoli spread all over the nucleus. The adherent disc is 27 µm thick.</td>
</tr>
<tr>
<td>VI</td>
<td>Oocyte diameter about 500 µm.</td>
<td>Oocytes characterized by the presence of yolk droplets associated with vacuoles.</td>
</tr>
<tr>
<td>VII</td>
<td>Oocyte diameter about 700 µm.</td>
<td>The cytoplasm completely filled with yolk granules. Nucleoli concentrated in the middle of the nucleus in most cases. Thickness of adherent disc now up to 30 microns, while that of the chorion is 10 µm.</td>
</tr>
<tr>
<td>VIII</td>
<td>Oocyte diameter up to 1mm.</td>
<td>Ripe oocytes ready to be released. Cytoplasm completely filled with large yolk platelets.</td>
</tr>
</tbody>
</table>
Fig. 4. Histology of female gonads: stages of oocyte development, from 1 to 7, are illustrated (see Table I for descriptive details). A- histology of a gonad from a female with 5.27 cm TL, from March. B- histology of a gonad from a female with 12.80 cm TL, from March. C- histology of a gonad from a female with 14.60 cm TL, from February. D- histology of a gonad from a female with 13.94 cm TL, from June. E and F- histology of a gonad from a female with 13.00 cm TL, from June (scale: 100 μm).
Table 2

Scale of ovary condition of *P. s. parvicornis*, with reference to oocyte development and gonadosomatic index.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Immature</td>
<td>Ovaries are very small. Weight less than 0.06g, representing 0.07 to 0.9 % of total body weight and 0.09 to 0.8% of the eviscerated weight. Ovaries form a thin transparent to yellowish ribbon. Oocytes mainly in stage I and II.</td>
</tr>
<tr>
<td>II</td>
<td>Maturing</td>
<td>Ovaries slightly swollen and creamy/yellowish. Oocytes now just visible macroscopically. Some ovaries do not complete development to mature stages. Weight not exceed 0.8g representing a maximum of 8% of total body weight. Oocytes present at stages of development I to V.</td>
</tr>
<tr>
<td>III</td>
<td>Ripening</td>
<td>Ovaries very large, occupy a large proportion of abdominal cavity. Oocytes easily visible macroscopically. Ovaries light yellow. Weight between 0.2g and 3.2g representing up to 8% of total body weight. Oocytes from stage I to VI.</td>
</tr>
<tr>
<td>IV</td>
<td>Ripe/pre-spawning</td>
<td>Ovaries distended and occupying almost the whole abdominal cavity. Oocytes easily visible macroscopically. Ovaries purplish. Weight may reach 6.5g representing up to 15% of total body weight and 18% of the eviscerated weight. Ripe/pre-spawning females found at the end of May and beginning of June. Ovaries have oocytes in all developmental stages.</td>
</tr>
<tr>
<td>V</td>
<td>Ripe/spawning</td>
<td>Ovaries may be less distended and less purple due to recent release of mature oocytes. Weight varies between 0.7g and 4.2g representing up to 10% of total body weight and 13% of eviscerated weight. Ripe/spawning ovaries found from June until August. All developmental stages of oocytes present.</td>
</tr>
<tr>
<td>VI</td>
<td>Post-spawning</td>
<td>Ovaries shrunken and coloured cream to dark-cream. Non-released eggs and atretic eggs can be seen macroscopically. Weight from 0.04g to 1g representing up to 2% of total body weight. Oocytes from stage I to stage III.</td>
</tr>
</tbody>
</table>

Table 3

Summary of oocyte stages and ovary condition in different months of the year

<table>
<thead>
<tr>
<th>Month</th>
<th>Oocytes stages</th>
<th>Ovary condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>January</td>
<td>I to III (+ atretic)</td>
<td>I and II</td>
</tr>
<tr>
<td>February</td>
<td>I to IV</td>
<td>I and II</td>
</tr>
<tr>
<td>March</td>
<td>I to IV</td>
<td>I and II</td>
</tr>
<tr>
<td>April</td>
<td>I to V</td>
<td>I and II</td>
</tr>
<tr>
<td>May</td>
<td>I to VI</td>
<td>I to III</td>
</tr>
<tr>
<td>June</td>
<td>I to VIII</td>
<td>I to V</td>
</tr>
<tr>
<td>July</td>
<td>I to VIII</td>
<td>I, II, III, V and VI</td>
</tr>
<tr>
<td>August</td>
<td>I to VIII</td>
<td>I, II and VI</td>
</tr>
<tr>
<td>September</td>
<td>I to II, VIII (+ atretic)</td>
<td>I, II and VI</td>
</tr>
<tr>
<td>October</td>
<td>I to III, VIII (+ atretic)</td>
<td>I, II and VI</td>
</tr>
<tr>
<td>November</td>
<td>I to III (+ atretic)</td>
<td>I and II</td>
</tr>
<tr>
<td>December</td>
<td>I to III (+ atretic)</td>
<td>I and II</td>
</tr>
</tbody>
</table>

4. Liver

4.1. General description

The liver is one of the largest organs located in the abdominal cavity. It is found just behind the transverse septum on the first half of the abdominal cavity (Fig. 2 and 3). Its colour may vary from dark cream to brown. As an organ with an important role in the storage of energy the liver exhibits considerable variation in its weight and structure that cannot be simply related with individual growth. These variations are strongly seasonal.

4.2. Histology

4.2.1. Seasonal variations in the structure of the female liver

In February the cells were filled with fat vacuoles of small and medium size (Fig. 6/A) in most of the individuals, but some of the bigger animals had large fat vacuoles (Fig. 6/B
Table 4

Scale of male gonad development in *P. s. parvicomis*, with reference to gonadosomatic index.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Maturation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Immature</td>
<td>Testes very small and transparent. Immature testes found in &quot;0&quot; individuals until January. Mean gonadosomatic index 0.14%</td>
</tr>
<tr>
<td>II</td>
<td>Maturing</td>
<td>Gonads much larger accounting from 0.5% to 3.4% of total body weight. Maturation begins in January/February. In fishes maturing for their second or third time testicular gland is consistent and cells look more compact than in younger fish. Testicular gland proportionally bigger than testis in large animals. In smaller males testicular gland only develops to a small size. From January until April the enlargement of the testis occurs. In April, the tubules may or may not be completely filled. Small cavities may be present in the centre. Not all stages of spermatogenesis will have been completed.</td>
</tr>
<tr>
<td>III</td>
<td>Ripe</td>
<td>By the end of May maturation appears to be complete. Gonad milky white in appearance and testicular gland slightly pink. Testes full of cells in different stages of spermatogenesis. Seminiferous tubules with spermatozoa distinct in the testicular gland. Spermatozoa found in <em>vas deferens</em>. Testes reach maximum size in relation to the testicular gland. Gonadosomatic indices reach their maximum value.</td>
</tr>
<tr>
<td>IV</td>
<td>Spawning</td>
<td>Spawning occurs in June, July and early August. The testis full of ripe spermatozoa. Cells in all other stages of development are present. Spermatozoa mainly seen in the testicular gland. In most cases tubuli empty in the middle indicating that spermatozoa have been released into the sperm ducts. Testicular gland showing a loose formation, with cavities, in August.</td>
</tr>
<tr>
<td>V</td>
<td>Post-spawning</td>
<td>Gonads much shrunken and greyish. Testicular gland prominent. The testis look like a fine band over the gland. Spermatozoa remaining in the testis and in the testicular gland. Condition maintained until November. Testicular gland proportionally wider than the testis. Gonadosomatic indices at their minimum in adults.</td>
</tr>
<tr>
<td>VI</td>
<td>Recovering</td>
<td>Recovery of the gonad initiated by December, when spermatogenesis occurs again.</td>
</tr>
</tbody>
</table>

and Table 5). In May, at the onset of the reproductive season, the livers were completely filled with fat vacuoles. These were large in bigger females (Fig. 6/C), and of intermediate size in smaller females (Fig. 6/D).

After the reproductive season, in August, the liver was much more compact and the fat vacuoles were rare, if present at all. The nucleus of the cells were much closer, and were very distinct (Fig. 6/E). The liver seemed to rest in this condition until October (Fig. 6/F). The storage of lipids appears to be reinitiated in November (Fig. 6/G), but some of the livers were still of the compact type at this time.

**4.2.2. Seasonal variations in the structure of the male liver**

The pattern of seasonal variations of the structure of the males liver seems very similar to that of the females. In March fat vacuoles fill the liver (Fig. 7/A). These fat vacuoles were very enlarged in May (Fig. 7/B) and still present in June (Fig. 7/C). In July the liver showed a compact structure (Fig. 7/D), which remained at least until October.

Table 5

A descriptive quantitative scale of liver structures for both males and females. (measurements in μm)

<table>
<thead>
<tr>
<th>diameter of lipid vacuoles</th>
<th>distance between nuclei</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>compact</td>
<td>---</td>
<td>4.58 ± 1.93</td>
</tr>
<tr>
<td>small fat vacuoles</td>
<td>≥ 5</td>
<td>6.44 ± 3.12</td>
</tr>
<tr>
<td>medium fat vacuoles</td>
<td>≥ 10</td>
<td>10.00 ± 1.67</td>
</tr>
<tr>
<td>large fat vacuoles</td>
<td>≥ 20</td>
<td>---</td>
</tr>
</tbody>
</table>
Fig. 6. Histology of female liver. A- female of 7.79 cm TL from February. B- female of 12.10 cm from February. C- female of 15.83 cm TL from March. D and E female of 11.75 cm TL from October. F female of 17.40 cm TL from August. G- female of 16.09 cm TL from October. H- female of 11.75 cm TL from November (fv) fat vacuoles (bc) blood cells (N) cell nucleus (scale 100 µm).
Fig. 7. Histology of male liver. A- male of 15.38 cm TL from March. B- male of 14.55 cm TL from May. C- male of 16.38 cm TL from June. D- male of 15.75 cm TL from July. E- male of 11.85 cm TL from August. F- male of 15.36 cm TL from September. G- male of 10.71 cm TL from October (f.v.) fat vacuoles (B) blood cell (bc) blood cells (N) cell nucleus (scale 100 μm).
DISCUSSION

Anal glands are distinctive male sexual traits of many blennid species (e.g., *Salaria pavo*, PATZNER & SEIWALD 1987b; *Parablennius ruber*, SANTOS 1987; and unpublished observations on *P. incognitus* and *Ophioblennius atlanticus* of the Azores). Their appearance and shape may be rather different even among species from the same genus (GOLDSCHMID et al. 1980), as for instance in *P. ruber* (SANTOS 1987) and *P. gattorugine* (KOTRSCHAL & GOLDSCHMID 1983) whose taxonomic distinction only recently was confirmed (ALMEIDA 1979; BATH 1982). The exact function of these organs is still being debated. It has been speculated that they produce pheromones (LOSEY 1969; LAUMEN et al. 1974), or that they produce antibiotic secretions that could be used by the male to protect the eggs against bacterial infections (QASIM 1956a; PETERSON 1984). The size of the glands is reduced out of the reproductive season. Since these glands are only fully developed in parental males during the reproductive season, and not in sneakers and satellites, it is a clear indication that they have a parental or epigamic function. This subject deserves to be further investigated.

Male gonads of blennioïds are unique among teleosts in possessing a testicular gland. It stores lipids and spermatozoa (LAHNSTEINER & PATZNER 1990d; LAHNSTEINER et al. 1990). LAHNSTEINER & PATZNER (1990c, 1990d) proposed that structural differentiation of the male gonads of blennies ("testis" and testicular gland), would have the advantage of shortening the duration of the spermatid cycle in the "testis", since final differentiation and consequent storage occurs in the testicular gland. Spawning in blennies may occur on successive days over a long period of time. The ability to accelerate the initial rate of spermatogenesis, associated with the possibility of spermatozoa storage would be an advantage, under these circumstances. The size and morphology of testicular glands of large and small mature males of *P. s. parvicornis* are very distinct, as in *Tripterygion tripterotus* and *T. delaisi* (JONGE et al. 1989). Testicular glands are clearly more developed in parental males. It is probable that this gland has other roles, besides the storage of spermatozoa. This subject is worth to be further investigation.

The histology of female gonads show that young females (size between 7.5 and 10 cm) only produce one batch of eggs during their first year as mature females. The oocytes of some of them reach only intermediate stages of maturation and are probably retained for the next spawning season. Major investment in reproduction begin when females reach their second year (SANTOS et al. 1996). Only a small proportion of the oocytes mature at a time, but this small proportion occupies the majority of the ovary volume. These are distributed by several spawning acts. In *Salaria pavo* more than 70% of the number of oocytes in the ovary are in the first stage of oogenesis (PATZNER 1985). Ripe oocytes occupy from 66.8% up to 75.3% of the ovary volume (PATZNER 1985). One out of 7 oocytes present in the ovary in the middle of the reproductive season will be retained for the next season. SHACKLEY & KING (1977a) found that each female produces eight different clutches of oocytes each breeding season. The production of multiple batches of eggs, reaching maturation and being spawned at different times, may present, at least, two types of advantages. One is related to female performance and management of resources, and the other to the survival of the embryos and the larvae. The first benefit is related to hydrodynamic performance. As MILLER (1979), following an hypothesis discussed by WILLIAMS (1959) pointed out: "a direct increase in gonadosomatic index raises problems of hydrodynamic performance and predator attraction, in addition to intrinsic levels of food intake. However, a high reproductive effort may be achieved by summation of repeat (partial, batch, heterochronal) spawning in which successive batches of eggs are produced during a lengthy period of reproduction." The other benefit is linked to the advantages of polyandry, on the one hand, and to temporal shifting of larvae in an unpredictable habitat, on the other. Distributing the eggs among several males,
reduces the risks of absolute loss of progeny due to wrong choice of the male. This is particularly relevant for a species in which survival of the embryos is directly dependent on the ability of the male to care for them. Distributing eggs for several spawnings during two months also assures that larvae will occur in the plankton at different times. This may contribute to reduced risks of total loss of progeny due to extreme unfavourable biotic and/or abiotic conditions during planktonic life. This is a good strategy in warm temperate and subtropical areas without marked seasonality of production such as the spring bloom, typical of cold temperate areas.

The dynamics of storage and depletion of resources in the liver is another important aspect of the phenology of *P. s. parvicornis*, which is clearly illustrated by histology. Liver growth is predominately linked with an increase in cell numbers (LOVE 1970), but it also undergoes changes that can be related to seasonal variations in storing and utilization of fat and glycogen by the individuals. It is known that the liver of the fish stores lipids and glycogen (LOVE 1970; PODROSCKO et al. 1985). Both males and females seem to rely on lipids stored in the liver after the winter during the reproductive season. It is known that energy stored in the liver is re-utilized in muscular activity (LOVE 1970), and effort of reproduction (e.g. in the process of ovary maturation). Females store lipids, which are later transferred to the ovary to be used by the developing embryo before it can feed (LOVE 1970). They transfer energy directly from feeding to the developing oocytes, but also rely on the fat they have stored in the liver. *P. s. parvicornis* with its well differentiated reproductive season shows the expected seasonal differences in liver size and histology, as well as structural differences related to age. Maturing females (sized from about 9 cm to 12 cm) store lipids in the liver from January until May. The compact structure of the livers of females sized between 7.5 cm and 10 cm during the months from January to March, and of smaller females until April shows that at these immature sizes do not store fat vacuoles (lipids) in the liver. It is probable that they invest directly in growth. Their investment in reproduction later on that year will be nil in the smaller size class. In May all females show abundant fat stores in the liver. The size of the fat vacuoles is less in small females than in larger ones. In June some of the big females have lost a great part of their liver fat reserves, certainly transferred to egg yolk, to be used by the embryos during their development. From August to December the liver has lost most of the fat reserves in all size classes. It is most probable that females will be investing directly in body restoration, maintenance and growth during this period. The synthesis of protein yolk is exogenous to the egg. It occurs in the liver from which it is transferred to the developing oocyte SHACKLEY & KING (1978, 1979). These observations confirm the conclusion that the liver is an organ with a role in the storage of energy which is spent during the reproductive season in this species.

ACKNOWLEDGEMENTS

The author acknowledge the field support given by Norberto Serpa. I am also grateful for the comments and/or help of Stephen J. Hawkins and Richard D. M. Nash (Port Erin Marine Laboratory), Robin Gibson (Scottish Marine Biological Association), Robert A. Patzner (University of Salzburg), Helen Rost Martins (DOP/UA) and two anonymous referees. The drawings included were made by Les Gallagher. This work was supported by JNICT (Ciência Program) and Secretaria Regional da Agricultura e Pescas (DRP), to whom I am grateful.

REFERENCES


(Pisces: Blenniidae) in an area where nest sites are very scarce. *Journal of Fish Biology* 45: 819-830.


LAHNSTEINER, F., U. RICHTARSKI, R.A. & PATZNER, 1990. Functions of the testicular gland in two blenniid fishes, *Salaria (=Blennius) pavo* and *Lipophrys (=Blennius) dalmatinus* (Blenniidae,


Accepted 17 April 1995.