Spermiogenesis particularities of a sperm storage species: *Helicolenus dactylopterus* (Teleostei: Scorpaenidae)

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SUMMARY: A study of the spermiogenesis and spermatozoa of *Helicolenus dactylopterus* was conducted. Females of this species have the capacity to store sperm within their ovaries, and male gametes have a considerable cytoplasmic mass surrounding their heads to survive the long period of intraovarian sperm storage. Our observations show that early spermatids are round-shaped cells and have a spherical nucleus with diffuse chromatin. The nuclear volume decreases as a result of progressive chromatin condensation during spermiogenesis, causing the nucleus to take on a U-shape. Flagellar insertion is not central to the nucleus but consistently occurs at an oblique angle towards one side of it. The flagellum is inserted into the nuclear fossa, without subsequent nuclear rotation. In mature spermatozoa, the flagellum is adjacent to the nucleus. A comparison of the spermatozoa in the testicular lobules and those in the intraovarian storage structures suggests that the increase in volume of the cytoplasmic mass may occur in the posterior region of the testis, in the testicular duct. Spermatozoa enter the ovary in groups that reach the ovarian lumen and are surrounded by the ovarian epithelium for storage in sperm storage crypts.

Keywords: *Helicolenus dactylopterus*, spermiogenesis, ultrastructure, sperm storage, spermatozoa.

INTRODUCTION

The bluemouth, *Helicolenus dactylopterus* (Dela-Roche, 1809), is a benthic species which inhabits the seabed between 200 and 1000 m depth (Whitehead *et al.*, 1986). It is regularly caught in the Mediterranean Sea and Atlantic Ocean, the target of both semi-industrial and artisanal fishing. The information on bluemouth reproduction includes three studies from the Azores (Isidro, 1989; Estácio *et al.*, 2001; Mendonça *et al.*, 2001).
et al., 2006), several studies from the Atlantic (Kelly et al., 1999; Sequeira et al., 2003; White et al., 1998), and many studies from the Mediterranean Sea (Allain and Lorane, 2000; Muñoz et al., 1999; Muñoz et al., 2000; Muñoz et al., 2002a,b,c).

*H. dactylopterus* belongs to the family Scorpaenidae, which is especially interesting from the reproductive point of view because scorpænid modes of reproduction are extremely varied. The group contains families which lay individual pelagic eggs (Anoplopomatidae, Congiopodidae, Hoplichthyidae and Triglidae) and others in which the spawn consists of sticky clusters of eggs (Agonidae, Cottidae, Cyclopteridae and Hexagrammidae). As far as is known, most scorpænids are basically oviparous species that produce pelagic egg masses enclosed in a gelatinous matrix. There are also viviparous genera such as *Sebastes* and *Sebastodes* (Washington et al., 1984). In addition, there are some, such as *Helicolenus*, that have particularities in their patterns of reproduction which are considered typical of viviparous species. In the case of *H. dactylopterus*, sperm storage is intraovarian and can last for ten months, an extremely long period for a teleost (Muñoz et al., 1999). Spermatozoa are stored within cryptal structures situated at the base of the interlamellar spaces. Once oocytes are ripe, spermatozoa are released from the storage structure, where they are maintained in viable condition throughout the storage period, into the ovarian lumen, where fertilization occurs (Muñoz et al., 1999).

The species under study is zygoparous (Muñoz et al., 2002a), a term referring to an oviparous condition in which fertilized ova (i.e. zygotes) are retained within the female reproductive tract for short periods of time before being released into the marine environment (Wourms et al., 1988).

In previous studies conducted by our research team, we have analyzed this storage process ultrastructurally and histochemically (Muñoz et al., 2000; 2002b) and have observed that the bluemouth’s spermatozoa show a special particularity: while male germinal cells within testicular lobules show the typical morphology of internal fertilizing species (Muñoz et al., 2002a), when we observe them within the male spermatid ducts and within the ovaries, these spermatozoa have a cytoplasmic mass surrounding their midpiece. This mass gradually decreases in volume as the spawning period approaches (Vila, 2004).

Given this peculiarity and the other special traits detected in relation to the reproduction of this species, in this study we aim to describe the transformation that male sexual cells undergo during spermiogenesis until spermatozoa are formed and released into the lumen of the testicular lobule. A further objective is the study of subsequent modifications in the testis related to the appearance of the cytoplasmic mass in the male gamete. This occurs in the spermatid ducts before sperm are introduced into the female’s body by the male’s urogenital papilla.

**MATERIALS AND METHODS**

Ten male specimens of *Helicolenus dactylopterus* were caught off the Costa Brava in the northwestern Mediterranean between March 2004 and February 2005, at a rate of approximately one per month, to study the fine structure of spermiogenesis. The spermatid duct samples were obtained from seven specimens caught between December 2005 and February 2006. Several portions of recently killed specimens were fixed in a mixture of 2.5% glutaraldehyde and 2% paraformaldehyde in a 0.1 M cacodylate buffer. After fixation at 4°C for 2 h, samples were washed several times with 0.1 M cacodylate buffer. Samples were then postfixed in 1% osmium tetroxide at 4°C for 1 h in cacodylate buffer. After washing, they were dehydrated through a graded alcohol series and finally embedded in Spurr’s mixture. Sections were examined in a Zeiss EM-910 transmission electron microscope, and images were analogically processed on Kodak 4489 electron microscopy film.

**RESULTS**

Very early in the spermatid stage, a pair of centrioles migrates to the plasma membrane (Fig. 1A). Concurrently, there is also a movement of mitochondria to this zone. They will surround the cytoplasmic mass of the flagellum. The centriolar complex is subsequently anchored to the plasma membrane and gives rise to the flagellum. In this complex, the proximal centriole is oblique to the distal centriole, and the distal centriole, which is anchored by radial fibrils to the plasma membrane, differentiates into the basal body. The cytoplasmic space is then formed by movement of the centriolar complex towards the nucleus, giving rise to the flagellar membrane and the initial segment of the flagellum, as noted by Mattei (1970).

At the same time, there is an evident reduction in nuclear volume as a result of a considerable amount of progressive chromatin condensation. This produces changes in the appearance of the nucleus, as chromatin is organized into clusters (Fig. 1B). This chromatin agglomeration is not uniform, beginning at the point where the centriolar complex is inserted and resulting in the nucleus taking on a U-shape (Fig. 1C). Nevertheless, in the centriolar complex zone there is always a less condensed chromatin mass. This transformation leaves a large space in the nucleus since the nuclear membrane has not been reduced; a vesiculation process would consequently be necessary to eliminate nuclear membrane debris. Some isolated double membrane-bound vesicles that are detected near the nucleus (Fig. 1C) appear to have been pinched off from the nuclear envelope. In this manner, the nuclear membrane will perfectly delimit the condensed chromatin.

The centriolar complex disposition undergoes changes between the initial spermatids and the late spermatozoa (Fig. 1D, E). In the initial stages of
Fig. 1. – A, very early spermatid, showing the centriolar complex in the spermatid plasma membrane (cc= centriolar complex; cm= cytoplasm membrane; n= nucleus ); B, late spermatid with a nucleus that contains highly condensed, thin fibres of chromatin (m= mitochondria; n, nucleus; *, nuclear fossa; arrow, cytoplasmic channel); C, “U-shaped” nucleus chromatin in a late spermatid, showing different examples of condensed chromatin depending on the site of the nucleus (cc, cytoplasm channel; hcc, high condensed chromatin; lcc, low condensed chromatin; m, mitochondria; arrow, centriolar complex); D, distal and proximal centrioles within the nuclear fossa are not perfectly orthogonal (clu, clusters of chromatin; dc, distal centriole; n, nucleus; pc, proximal centriole); E, high magnification of distal and proximal centrioles of a mature spermatozoon (cc, cytoplasm channel; dc, distal centriole; n, nucleus; pc, proximal centriole); F, mature spermatozoa, showing the lateral nuclear fossa and the nucleus lobule that protects it (f, flagellum; h, head; l, nucleus lobule; m, mitochondria; nf, nuclear fossa). Scale bars: A-C, F, 1 μm; D, 0.5 μm; E, 0.2 μm.
Fig. 2. – A, the flagellar axis is at an oblique angle to the spermatid nucleus (f, flagellar axis; ch, condensed chromatin; m, mitochondria; n, nucleus); B, spermatid midpiece, showing mitochondria with regularly organized cristae (f, flagellum; m, mitochondria); C, the spermatozoa inside testicular lobules do not exhibit cytoplasmic masses around their heads (f, flagella; h, heads); D, *H. dactylopterus* spermatozoa are of the complex type of anacrosomal introsperrm, typical of internal fertilizing species (cm, cytoplasmic mass; h, spermatozoa’s head; f, flagellum; m, mitochondria; t, sperm tails). The arrows show the lateral projections of the flagellum; E, spermatozoon midpiece, in which we can clearly see a cytoplasmic channel (c, centriole; cc, cytoplasm channel; f, flagellum; m, mitochondria; v, vesicles); F, undulating surface of spermatozoon flagellum (f, flagellum; pm, plasma membrane). Scale bars: A, 1 μm; B, 0.25 μm; C, 2.5 μm; D, 2 μm; E-F, 0.5 μm.
Fig. 3. – A, spermatozoa in the posterior part of the testis showing cytoplasmic masses around their heads (cm, cytoplasmic mass; h, sperm head); B, overview of the posterior part of the testis, in which epithelial cells of the sperm duct exhibit cytoplasmic projections (ec, epithelial cells; sz, spermatozoa; szt, spermatozoa’s tails); C, detail of the sperm duct epithelium releasing exocytosis vesicles (ev, exocytosis vesicles), D and E, efferent ducts during the spawning period, showing masses of sperm. Note the examples of alignment of heads with heads and tails with tails. Haematoxylin-Eosin-Mallory (szg, groups of spermatozoa). Scale bars: A, 2.5 μm; B-C, 1 μm; D, 200 μm; E, 100 μm.
spermiogenesis, the distal and proximal centrioles are oblique to each other. As spermiogenesis advances, they become more and more orthogonal, although in the final spermatooza the two centrioles are still not perfectly perpendicular, but form an angle smaller than 90°. As a result of the initial lateral association of the centrioles in the mature spermatooza, the nuclear fossa is situated laterally (Fig. 1F).

The flagellar insertion is not central to the nucleus but is at one side of the latter and oblique to it (Fig. 2A).

As the spermiogenesis process advances, the mitochondria that are concentrated around the flagellum surround its initial segment, giving rise to the midpiece of the future spermatooza. In the spermatids, mitochondria show tubular and regular cristae (Fig. 2B). At the end of spermiogenesis, the residual spermatid cytoplasm is phagocytized by Sertoli cells (Fig. 2C), which delimit the cysts. Thereafter, mature spermatooza are released into the lumen of the seminiferous lobe.

_H. dactylopterus_ spermatooza belong to the complex type of anacrosomal intromission (Muñoz et al., 2002a) defined by Jamieson and Leung (1991) and typical of species with internal fertilization (Fig. 2D). The spermatooza heads do not show any trace of an anacrosomal vesicle, and exhibit an elongated “pear-shaped” nucleus surrounded by a narrow strip of cytoplasm without organelles. The midpiece has four mitochondrial layers, each containing approximately ten mitochondria. A well-developed cytoplasm channel is evident (Fig. 2E). The flagellum shows the typical 9 + 2 axonemal structure, with nine doublets of microtubules and a central pair, all of which are surrounded by a slightly undulated flagellar membrane (Fig. 2F). Two lateral projections are evident in the medial and distal part of the flagellum (see Fig. 2D).

While newly-formed spermatooza are still in the testicular lobules, no particular cytoplasmic extensions are observed in them (see Fig. 2C). However, the spermatooza located in the posterior part of the testis, at the beginning of the spermiogenetic duct, have a cytoplasmic mass on one side of their nuclei and midpieces (Fig. 3A) with some small vacuoles in it. Furthermore, the epithelium of this area of the spermiogenetic duct exhibits numerous projections and free vesicles that penetrate the area in which spermatooza are waiting to be released through the spermiogenetic duct into the female’s body (Fig. 3B, C). Mature spermatooza are first released towards the lobular lumen and then accumulate at the spermiogenetic duct (Fig. 3D, E). They are organized in groups, are finally inseminated in this state into the female’s body, penetrate the ovaries and are stored until oocyte maturation.

DISCUSSION

According to Mattei (1970), the process of spermiogenesis in teleosts may result in two basic spermatoozon types (type I and type II), with an intermediate type between them. This classification is based on whether or not nuclear rotation occurs during spermiogenesis. When rotation does not occur, the flagellum becomes perpendicular and medial to the nucleus, and spermatoozoa are defined as type I. Without nuclear rotation, the flagellum remains parallel to the nucleus, and spermatoozoa are defined as type II. However, nuclear rotation may be incomplete, and in this case the flagellum is eccentric to the nucleus and spermatoozoa are of an intermediate type between I and II. Recent studies of Pimelodidae (Quagio-Grassiotto and Oliveira, in press) and Heptapteridae (Quagio-Grassiotto et al., 2005) show a medio-flagellar development, in which there is no nuclear rotation. These characteristics have not been found in other fish species until now, so Quagio-Grassiotto et al. (2005) postulated a new type of spermiogenesis, which they called type III.

Spermiogenesis in the bluemouth shows an oblique insertion of the flagellum relative to the nucleus but there is no nuclear rotation. Chromatin condensation begins where the centrioles are inserted, giving rise to mature spermatooza with perfectly aligned heads and tails and thus coinciding with type III spermiogenesis.

The final spermatooza keep some fraction of cytoplasm on one side of their heads when they are inseminated into the female’s body. This could suggest that the cytoplasm may contain nutrients that nourish stored spermatooza during the period of intraovarian storage. Electron microscope observations have shown that during this long storage period these cytoplasmic masses undergo a considerable decrease in volume, with the result that final spermatooza have a more hydrodynamic shape, thereby facilitating ova fertilization in the ovary. Comparing spermatooza inside the lobular testis with those inside the ovaries we have noted that the increase in volume of the spermatooza cytoplasmic masses occurs in the posterior part of the testis, where the spermiogenetic duct begins. This hypothesis is reinforced by the observations we have made in this zone of numerous vesicles and projections of the epithelium towards the area in which spermatooza are released before being inseminated into the female.

Most tissue cells are polarized and exhibit two different membrane domains to which different types of secretory vesicles are targeted. Epithelial cells from the posterior part of the bluemouth testis show a characteristic apical domain that is typically found facing the lumen, with specialized structures such as cilia or microvilli, and a basolateral domain that comprises the rest of the cell. The package of secretory vesicles is located in Golgi apparatus. The last phase of the secretion route of a vesicle is the release of the product it contains by means of exocytosis (Alberts et al., 1996). This may therefore be the purpose of the cellular projections towards the lumen referred to above, allowing spermatooza to absorb nutrients and store them so that they can survive the long intraovarian storage period.

At first glance, it is surprising to find this cytoplasmic mass on one side of the sperm heads, because it gives sperm cells a low hydrodynamic profile. How-
ever, sperm cells are organized into bundles within the testis and are also subsequently released in this agglomerated pattern into the female’s genital tract. The fact that they enter the ovary in bundles means that a hydrodynamic shape is not so important. Thus, spermatozoa enter the ovary in organized groups that reach the ovarian lumen and are surrounded by the ovarian epithelium to be stored in sperm storage crypts (Vila, 2007). Previous electron microscope observations (Vila, 2004, 2007) showed that during the ten-month period of intraovarian sperm storage these cytoplasmic masses decrease in size as the spawning period approaches.

When spermatozoa are released into the ovarian lumen to fertilize ovulated ova, they have a more hydrodynamic shape which better equips them to penetrate ripe oocytes. The lack of specialized structures to store spermatozoa has been linked to the short storage period. For example, in Alcichthys alcicornis spermatozoa float freely within the ovarian cavity (Koya et al., 1997). The spermatozoa of this species show an elongated nucleus and midpiece. The flagellum is joined to the posterior end of the nucleus, where many small mitochondria surround the axoneme (Koya et al., 2002). As in Helicolenus dactylopterus, ten mitochondrial sections were found in one transversal section of Alcichthys alcicornis.

Sperm can be maintained with varying degrees of specialization. In the viviparous embiotocid Cymatogaster aggregata, spermatozoa remain inside a sort of sperm pocket, where their heads are inserted into the ovarian epithelium for several months between copulation and fertilization (Gardiner, 1978). In Sebastes taczanowskii, the male gametes adhere to the ovarious lamella epithelium or are wrapped in its microvilli (Takemura et al., 1987; Hayashi, 1990). There is a some diversity in the way species store sperm within the ovaries, but all exhibit a sperm morphology suitable for travelling through the female’s genital tract.

The observed elongation of the sperm nucleus may be related to sperm transport and storage in internally fertilizing species (Jamieson, 1987) because the streamlined nucleus may aid the passage of spermatozoa through the narrow areas and viscous fluids of the female reproductive tract (Gardiner, 1978), or help form sperm bundles (Atwood and Chia, 1974). Elongation of the sperm nucleus may facilitate the side-by-side alignment of spermatozoa moving through the testicular ducts (Burns et al., 1995).

In fact, after the bluemouth mates, the sperm penetrates deep into the ovary, reaching the interlamellar space. Therefore, the elongated midpiece is an energy source and is necessary for sperm movement. This energy is supplied by the mitochondria, which respire and consume the endogenous substrate of the midpiece (Baccetti and Afzelius, 1976).

The process of spermiogenesis in Helicolenus dactylopterus is thus characterized by a number of particular ultrastructural features related to the sperm cytoplasmic mass that are not found in any other fish species. This could possibly be related to the fact that the spermatozoa can be stored within the female’s ovaries for up to ten months.

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