## Histological, histochemical and immunohistochemical study on the growing oocytes of the abyssal teleost *Hoplostethus mediterraneus* (V)

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Abstract: The oocytes of the abyssal Teleost, *Hoplostethus mediterraneus* were studied. Four stages of growth were observed and the oocytes of all the stages were surrounded by follicular cells and had several nucleoli in the nucleus. In the oocytes of the II° stage, vacuoles without contents, in oocytes of the III° stage several vacuoles with a basophilic contents and small yolk globules were identified. General and basic proteins, ribonucleoproteins, acid proteoglycans with -COOH groups were recognized in the cytoplasm, in the nucleoli of oocytes in the II° stage and in the vacuolar contents of oocytes in the III° stage. In the follicular cells, in the pellucid zone, in the yolk globules, from their beginning, glycoproteins were present. Positivity, for all lectins used, was revealed in the follicular cells and in the four stages of oocytes growth.  $\alpha$ -D-glucose and  $\alpha$ -D-mannose binding sites were in the pellucid zone and in the initial yolk globules. In the lather galactose and  $\beta$ -N-acetyl glucosamine were present too. nNOS and VIP immunopositivity revealed at the periphery of the cytoplasm and at network of nerve fibres between oocytes, suggests NO is involved in a mechanism of regulation of the gametogenesis and of the spawning.

Key words: Vitellogenesis - Teleost - Hoplostethus mediterraneus - Oocytes - Glycoconjugates - nNOS - VIP - Nerve fibres

### Introduction

In the oocytes of a lot of Invertebrates and Vertebrates the different staining affinities among the initial, intermediate and definitive yolk globules were correlated with the variations in the chemical constitution of the nucleoli during vitellogenesis [1,2].

Only in *Mictophum punctatum* the yolk globules had the same chemical constitution, though different staining affinities, [3].

The different GAG and endogen agglutinins (lectins) found in the cortical granules and in the jelly coat of the oocytes in some species of *Teleostes* [4,5] were considered as substances preventing the polyspermia.

In the oocytes of different species of teleosts sugar specific sites were found in the follicular cells [6] as well as in the pellucid zone [7]. The phylogeny and functional roles of nNOS and VIP, present in different tissues (epidermic, tegumental connective, musculature of swim-bladder, gills, lung) of Invertebrates and Vertebrates are still unclear. However recent studies indicate involvement of these bioactive compounds in the non adrenergic-non cholinergic (NANC) regulatory mechanism [8-10].

A recent immunohistochemical investigation on the mantle integument of the mussel *Mytilus galloprovincialis* [10], in order to detect, in both sexes, the presence of nNOS and VIP, resulted positive in cortex of sperm follicles and nNOS positive sperm only in the male, in the female positivity in the nerve cells bodies and fibres and negativity in the maturating oocytes.

The aim of this study was the investigation on the oocytes of abyssal teleost *Hoplostethus mediterraneus* in order to ascertain the presence of bioactive molecules too [9-13] and to confirm their role in the reproductive function.

#### Materials and methods

**Specimen samples.** Specimens of *Hoplostethus mediterraneus* caught in the Strait of Messina (Italy) were sacrificed and female gonads were isolated. They were fixed in Bouin and in 10% for-

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malin in sea water, for the morphological and structural study and in 4% paraformaldehyde buffered with 0.01M sodium phosphate (PBS), pH=7.4 at 4°C for the histochemical and immunohistochemical study, then they were embedded in paraplast. Serial sections (5  $\mu$ m tick) were stained with Mallory mod. Ignesti (Carazzi hemalum, 1% acid fuchsin, Mallory's solution) and Galgano I (Mayer's hemalum, 0.1% acidic fuchsin, 1% phoshomolybdic acid, Mallory's solution).

**Proteins.** Following histochemical methods demonstrating proteins were used: bromophenol blue method, ninhydrin- Schiff, chloramines T-Schiff, performic acid-Schiff, Morel and Sisley, Millon, Adams reactions, diazoreaction and Sakaguchi reaction.

**Glycoproteins.** Following histochemical methods demonstrating proteins were used: PAS reaction according to Mc Manus, dimedone- PAS reaction. Acid proteoglycans AB (pH=1.0); AB (pH=2.5). Distinction between glycoproteins and acidic proteoglycans; AB pH=1.0-PAS; AB pH=2.5 PAS.

**Distinction between acid proteoglycans with different acidities.** AB - C.E.C. (critical electrolyte concentration): 0.05% AB, inhibition of alcianophilia at graded molarities of 0.5 - 1 M MgCl<sub>2</sub> in acetic buffer (pH=5.8) weak methylation-AB (pH=2.5); strong methylation-AB (pH=2.5); weak methylation-saponification-AB (pH=2.5). Acid hydrolysis in 0.1 M acetate buffer (pH=2.5) for 4h at 60°C followed by AB (pH=2.5), also without saponification.

**Methods of enzymatic digestion.**  $\alpha$ -amylase - PAS; AB-neuraminidase (pH=2.5), from Clostridium perfringens with and without saponification, testicular hyaluronidase [14] (Table 1).

**Specific sugar residues.** Biotinylated lectins and avidin-biotinperoxidase method (ABC) was employed. The biotinylated lectins used (Vector Laboratories, Burningame, California, USA) their origin, specificity and inhibiting sugars are reported in Table 2. Incubating the sections with specific sugar performed controls.

**Indirect immunoperoxidase method.** Section of ovary were incubated overnight at 4°C in humid chamber with polyclonal rabbit anti-human brain nNOS antibody diluted 1:200 (nNOS type I Code 606-259-1550; Biomol, Hamburg, Germany) or polyclonal antihuman VIP antibodies (Biomeda, Foster City CA, USA). Afterwards, sections were washed in PBS and incubated for 2 h with a goat anti-rabbit IgG-peroxidase conjugate (1:100; Sigma, Munich, Germany). Peroxidase activity was visualized by incubation of the sections for 5-12 min at room temperature in a solution of 0.015% 3,3'-diaminobenzidine in 0.01 M tris buffer (pH=7.6) containing 0.005%  $H_2O_2$ . Negative controls included omission of the primary antibody or its substitution with non-immune rabbit serum.

**NADPH diaphorase.** To demonstrate NADH diaphorase activity, pieces of tissue were fixed for 2 h at room temp in a solution of 4% paraformaldehyde immediately after dissection. Afterwards, the pieces of tissue were incubated for 5 min in 0.1 M PBS containing 0.25% Triton X-100, and subsequently for 60 min at 37°C in a solution containing 3 mg/ml NADPH (Sigma, St Luis MO, USA) and 0.5 mM nitroblue tetrazolium (Sigma). Then, the tissue were washed thoroughly, dehydrated and embedded in paraplast and sectioned (5  $\mu$ m thick). Control incubations were carried out using pieces of tissue incubated in media lacking NADPH.

#### Results

The structural examination on the ovarian isolated in the different months of a year from specimens of *Hoplostethus mediterraneus*, revealed four stages of

 
 Table 1. Conventional histochemical methods for visualization and identification of glycoconjugates (GCs).

Procedures	GCs revealed	
PAS	Glycoproteins	
AB pH=1.0	Proteoglycans with sulphate groups	
AB pH=2.5	Proteoglycans with carboxylic groups	
AB pH=2.5 PAS	Proteoglycans with carboxylic groups and glycoproteins	
Neuraminidase enzymatic digestion	Sialic acid	
AB (C.E.C.)	Sialic acid	

growth [15,16]. The oocytes in the I° stage, about 20 000 of medium  $\mu$ m volume, had a homogeneous cytoplasm and were surrounded by a single layer of follicular cells. In nucleus there were up to six nucleoli, of which one was spherical and greater in size, the spherical others were smaller arranged along the nuclear membrane. By Mallory's modification Ignesti and Galgano I structural methods, the cytoplasm was strongly basophylic and the nucleoli were red stained (Fig. 1).

The oocytes in II° stage, about 180 000 µm of medium volume, were larger and characterized by two layers of follicular cells arranged along the oocyte's membrane, in orders to form an epithelium at first monostratified and then pluristratified. Furthermore, they are characterized by the presence in the nucleus of a greater number of nucleoli, adhering to the membrane mainly scattered of round shape, with changing dimensions and strongly red stained with the above mentioned methods.

The oocytes in III° stage, about 3 000 000  $\mu$ m of medium volume, were surrounded by several layers of follicular cells, of which, one thicker and thicker, with evident streaking, constituted the radiate or pellucid zone, strong red stained by Mallory.

In the cytoplasm of these oocytes there were also vacuoles with basophilic contents arranged in several layers (Fig. 2). In the cortical zone between the radiate zone and the vacuoles there were yolk globules extending from the periphery to the centre, at first they were red stained then turning from red into blue (Fig. 3). The nucleoli, of which one or two seemed more conspicuous, partly kept the red stain.

The oocytes of IV° stage, about 20 000 000  $\mu$ m of medium volume, were surrounded by one layer of follicular cells, that in the most advanced vitellogenesis became shorter, cubic-shaped, surrounding the pellucid zone. In the oocyte's cytoplasm, the number of vacuoles decreased till to disappearing at the end of the vitellogenesis, at the same time, the yolk globules

Lectin	Origin	Carbohydrate specificity	Inhibitory sugar
ConA	Canavalia ensiformis	α-D-mannose α-D-glucose	D-Mann
RCA	Ricinus communis	β–D-galactosyl	Lactose; D-galactose
PNA	Arachis hypogaea	D-galactose	D-gal
SBA	Glycine max	N-acetyl-D- Galactosamine>D-galactose	D-galNAC
UEA I	Ulex europaeus	α-L-fucose	L-fucose
WGA	Triticum vulgaris	N-acetyl-D-galactosamine> Sialic acid	D-GlcNAc

Table 2. Lectins employed and their major binding specificities.

increased in number and in size gradually showing a blue stain. The nucleus became smaller in size. The nucleoli appeared red stained. The pellucid zone changed stain from red to blue (Fig. 3).

The basophilia in the cytoplasm of oocytes of I° and II° stages was confirmed by positivity for general and basic proteins (bromophenol blue with and without mercury dichloride methods). The pironinophilia in the cytoplasm is due to ribonucleoproteins. AB pH=1.0 and AB pH=2.5 methods revealed the presence of acid proteoglycan with -SH and -COOH groups respectively. The negative result to PAS reaction sec. Mac Manus showed absence of glycoproteins. In the follicular cells, glycoproteins, (Fig. 4), acid proteoglycan with -SH and -COOH groups. In the nucleoli general and basic and ribonucleoproteins were present.

The contents of vacuoles, where present, was positive for general and basic proteins, glycoproteins (Fig. 4) without glycogen, acid proteoglycan with -SH and -COOH groups coexisting with glycoproteins (Fig. 5). Follicular cells of oocytes of I° and II° stage and the II° stage oocyte's cytoplasm were positive after use of the RCA, ConA, PNA, SBA, WGA, UEA I lectins. The follicular cells, the cytoplasm and vacuolar contents of oocytes II° in advanced stage, were positive for  $\alpha$ -D-mannose and  $\alpha$ -D glucose (ConA) and for galactose (RCA) (Fig. 6).

The oocytes of III° stages showed a gradual decrease of basophilia, pironinophilia and positivity for the proteins. The latter substances were present in the yolk globules, in the nucleoli of oocytes and in the cytoplasm of the follicular cells. In the oocytes cytoplasm glycoproteins, without glycogen, acid proteoglycans with -SH and -COOH groups were present.

AB (pH=1.0)-PAS and AB (pH=2.5)-PAS reactions revealed glycoproteins and acid proteoglycans with -SH and -COOH groups (Fig. 7) both in the cytoplasm and in the vacuolar contents. Only glycoproteins were present in the yolk globules, pellucid zone and follicular cells (Fig. 7). The acid proteoglycans were absent in the yolk globules, in the pellucid zone, in the nucleoli.

Use of all lectins always revealed positivity of binding sites for follicular cells. In most of the initial yolk globules the galactose (RCA) was present (Fig. 8). WGA revealed  $\beta$ -N-acetyl-glucosamine in the smaller yolk globules and in the follicular cells (Fig. 9).

In the oocytes of IV° stage there was no correspondence between changes in stain and changes in the chemical constitution. In fact in the yolk globules there were proteins and glycoproteins. By using ConA and WGA (Fig. 10) positivity was shown only in the periphery. The use of the other lectins simply revealed negativity.

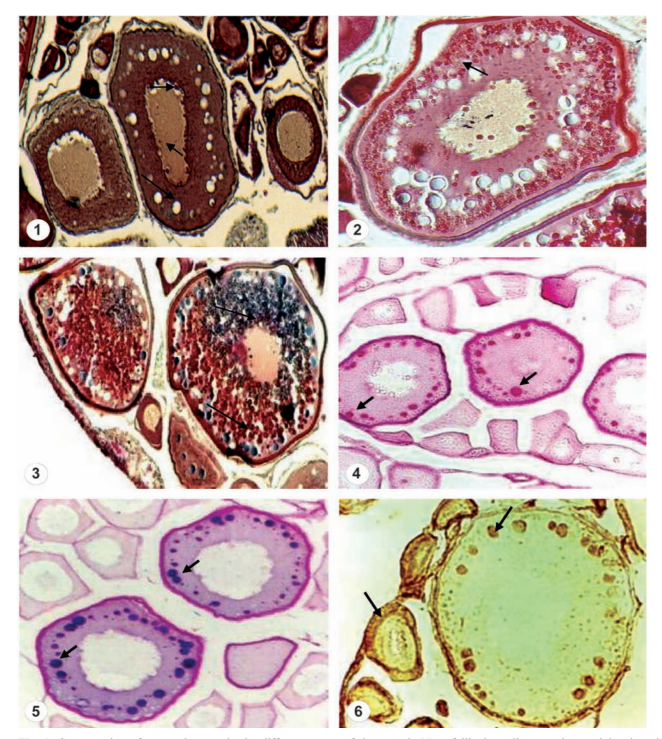
After immunohistochemistry analysis the cytoplasm of vitellogenic oocytes and of follicular cells was nNOS (Fig. 10) and VIP positive nerve fibres nNOS positive were found adjacent to maturating oocytes (Fig. 11).

Incubations in the absence of primary antibodies or their substitution by non-immune sera resulted in nerve fibres and follicular cells, were negative for nNOS and VIP. The final reaction product of NADPH diaphorase activity was present in nerve fibres and follicular cells. Distribution patterns of NADPH diaphorase were similar to the patterns of nNOS positive staining.

#### Discussion

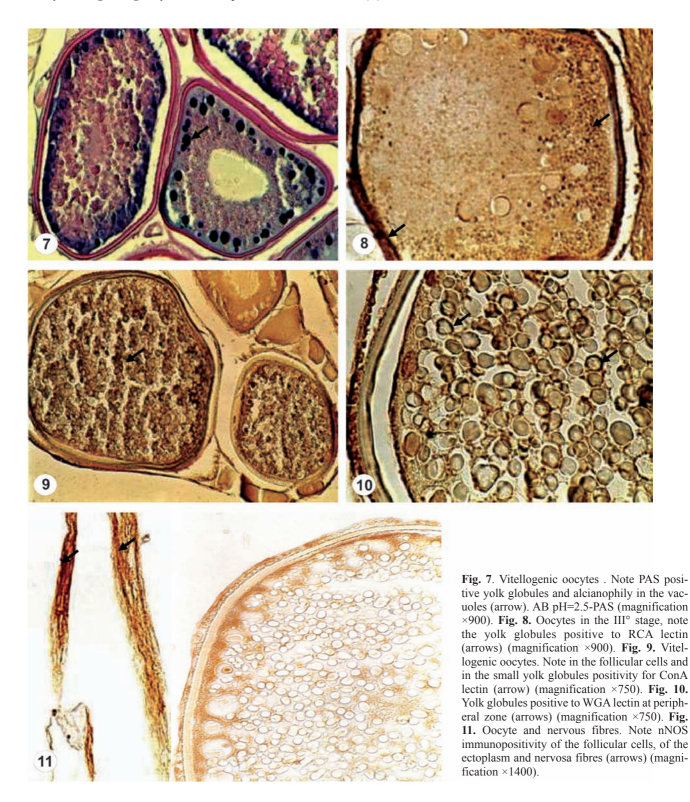
The results obtained in *Hoplostethus mediterraneus* were revealed both analogies as well as differences to ward the other species studied, in particular with *Coelorhinchus coelorhynchus* abyssal teleost.

According to other authors [2,17-19], in oocytes of III° stage of *Hoplostethus mediterraneus* there were several layers of follicular cells and a pellucid zone with evident streaking. The yolk globules, found in *Coelorhinchus* appeared from the beginning of the



**Fig. 1.** Cross section of ovary. Oocytes in the different stages of the growth. Note follicular cells, vacuoles, peripheral nucleoli (arrows). Mallory's triple stain (magnification  $\times$ 750). **Fig. 2**. Vitellogenic oocytes. Note the yolk globules red stained (arrow). Mallory's triple stain (magnification  $\times$ 750). **Fig. 3**. Previtellogenic and vitellogenic oocytes. Note yolk globules blue stained (arrows). Galgano I stain (magnification  $\times$ 900). **Fig. 4**. Oocytes in different states of the growth. Note PAS positivity of the follicular cells and of the vacuoles (arrows). PAS sec. Mc Manus reaction (magnification  $\times$ 750). **Fig. 5**. Oocytes of the I° and the II° stages of the growth. The blue- purple stain in the vacuoles shows the coexistence of the glycoproteins and acid proteoglycans with -COOH groups (arrows). AB pH=2.5- PAS (magnification  $\times$ 750). **Fig. 6**. Oocytes in the II° stage. Note the follicular cells and vacuoles positive to RCA lectin (arrows) (magnification  $\times$ 900).

third stage of growth preceded by vacuologenesis. The contents of the vacuoles, were at first without contents, and later basophilic for the presence of proteins and acid proteoglycans with -SH and -COOH groups. The yolk globules, first red stained by Mallory's method, during the vitellogenesis, changed stain from red to



blue, their chemical constitution remaining however unchanged.

The absence of the contents in the vacuoles from the beginning, was according to was found in *Coelorhinchus coelorhynchus* [19] and in *Channa* [20], but not according to Mictophum punctatum [3], an abyssal species, in whose oocytes, the vacuoles had a basophilic contents constituted by glycoproteins, that is by the same substances found in the yolk globules from the beginning of the vitellogenesis. These results confirmed what supposed for *Coelorhinchus*, and *Channa*, *i.e.* it was a delayed condensation.

Nucleolar apparatus in oocytes of IV° stage showed small variations in chemical constitution and in stain-

ing affinities; no correspondence with staining variations of yolk globules. Therefore *Hoplostethus mediterraneus* was different from abyssal species *Coelorhinchus coelorhynchus*, [21] and from pelagic species Coris julis [22] and Gobius cobitis [2], in fact in the nucleoli of these species there occurred a coincidence between variations in staining affinities and chemical constitution.

The presence of follicular epithelium structurally different in the different stages of growth and the invariability of chemical constitution of yolk globules permitted to state that also in this examined species, the substances responsible to constitute yolk globules are exogenous, probably synthesized in liver [22-24], carried by blood to the follicular epithelium and at last in the oocyte.

The positivity revealed after use of all lectins is different by *Coelorhinchus*, in which the follicular cells were positive for some lectins only.

Similar results in *Hoplostethus* and in *Coelorhinchus* with regard to positivity for the WGA and SBA were found both in the yolk globules and in the periphery of globules at the end vitellogenesis. The hypothesis was that in *Hoplostethus* other groups in the molecular structure might mask the inreactivity of specific sugar radicals for all lectins used.

VIP and nNOS immunoreactivity elevated at the periphery of the cytoplasm of the oocytes and in the follicular cells, and the evidence of network of nerve fibres between oocytes during vitellogenesis, as found in other species [8,10] suggested that the presence of NO and its synergism with VIP is a challenging subject to study due to wide variety of biological actions of NO and VIP on regulation of gametogenesis and spawning of both invertebrates and vertebrates.

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