Immunohistochemical localization of hepatopancreatic phospholipase A2 in Hexaplex Trunculus digestive cells

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Abstract

Background: Mammalian sPLA2-IB localization cell are well characterized. In contrast, much less is known about aquatic primitive ones. The aquatic world contains a wide variety of living species and, hence represents a great potential for discovering new lipolytic enzymes and the mode of digestion of lipid food.

Results: The marine snail digestive phospholipase A2 (mSDPLA2) has been previously purified from snail hepatopancreas. The specific polyclonal antibodies were prepared and used for immunohistochemical and immunofluorescence analysis in order to determine the cellular location of mSDPLA2. Our results showed essentially that mSDPLA2 was detected inside in specific vesicles tentatively named (mSDPLA2+) granules of the digestive cells. No immunolabelling was observed in secretory zymogene-like cells. This immunocytolocalization indicates that lipid digestion in the snail might occur in specific granules inside the digestive cells.

Conclusion: The cellular location of mSDPLA2 suggests that intracellular phospholipids digestion, like other food components digestion of snail diet, occurs in these digestive cells. The hepatopancreas of H. trunculus has been pointed out as the main region for digestion, absorption and storage of lipids.

Introduction

The Muricidae family of snails includes about 1,000 species, which represent a diverse and important component of marine communities [1]. The banded murex, Hexaplex trunculus (Linnaeus, 1758), is found in the Mediterranean sea and adjacent Atlantic ocean from the Portuguese coast, southward to Morocco and to the Madeira and Canary islands [2] and [3]. This species is a commercially important marine snail in the Mediterranean coasts.

The digestive gland or the hepatopancreas of gastropod molluscs like the marine snail H. trunculus is the key organ for metabolism. It is the main source of production of digestive enzymes, and it is involved in absorption of nutrients, food storage and excretion [4]. It combines many functions of the liver, pancreas, intestine and other organs in vertebrates [5]. For this reason, it has captivated scientists for more than 180 years [6] and [7] and it was considered as an excellent model for food digestion and cell secretion [8]. Its primary role is the synthesis and secretion of digestive enzymes, swallowing and final digestion of the ingested food and subsequent uptake of nutrients. Hepatopancreas is also implicated in storage and excretion of inorganic reserves, lipids and carbohydrate metabolites. Moreover, it is the primary metabolic center for the production of materials required for the temporally distinct events of molt and vitellogenesis [9].

Furthermore, the ultrastructure and functional aspects of several aquatic invertebrate hepatopancreases were studied. Crustaceans and mollusc hepatopancreases, similar to that of the marine snail, show mainly two specialized cell types in the digestive diverticula [10] and [11]. Secretory zymogene-like cells which are responsible for extracellular digestion, while digestive cells are involved in absorption, intracellular digestion and nutrients transport. Unfortunately, the digestive process is

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not thoroughly investigated in invertebrate and the digestive processes remain unclear.

We previously purified a new marin snail digestive phospholipase A₂ (mSDPLA₂) from the hepatopancreas of H. trunculus [12]. This mSDPLA₂ of 30 kDa, which contrasts with common 14 kDa-digestive PL₂, is of interest as it exhibits hemolytic properties and could be used as model to study digestive and cytotoxicity mechanisms. Zarai et al [12] have shown that the potential mSDPLA₂ activity was measured, in vitro, in presence of bile salts like NaTDC or NaDC. This result confirms that mSDPLA₂ presents a high interaction power which allows it to bind to its substrate even in presence of bile salts.

Many studies have described the morphology as well as the digestive system of snails. The hepatopancreas and the salivary glands of the snails are the main sources of digestive enzymes. To our knowledge, the presence of bile salts in the molluscs digestive glands has never been described. As recently shown by Amara et al (2010) [13] no bile salts are detected in the land snail digestive gland. One can say that, in vivo, the digestive system of sail contains probably some molecules (bile salts-like) which allow the snail phospholipase to hydrolyse efficiently its substrate.

To investigate the digestive function of mSDPLA₂ in invertebrate, and to elucidate its digestive mechanism in hepatopancreas, we performed immunohistochemistry and immunofluorescence analysis using rabbit anti-mSDPLA₂ whole serum. Our current studies demonstrate that marine snail mSDPLA₂ has both intracellularly and extracellularly food digestion role. The digestive processes by mSDPLA₂ were accurately followed by the use of the anti-mSDPLA₂ polyclonal antibodies. A digestive model in hepatopancreas is therefore formulated.

Results

General digestive gland description

The hepatopancreas of molluscs is a large digestive gland formed by a vast number of blind ending tubules; the digestive diverticula. This organ is involved in several functions including the extracellular and intracellular food digestion, lipids, glycogen and minerals storage. It is also the main site of nutrient absorption and plays a major role in detoxification [14-17].

H. trunculus, like other molluscs, has an acini digestive glands. The lumen of acinous gland communicates by ducts with stomach lumen. The epithelium lining the tubules of the glands consists chiefly of tall columnar cells; their apical surface is typically covered by microvilli (Figure 1).

The digestive diverticula consist of an epithelium with a single layer of cells, separated from the surrounding connective tissue and muscle cells by a basal lamina. In several molluscs these epithelia consist of the digestive and basophilic cells [18-21]. However, in gastropods other cell types have been reported in addition to these two cell types [22-24]. Digestive cells are the most abundant cell type in the digestive diverticula of H. trunculus.

Molluscs digestive cells are mainly characterized by the presence of a large number of heterolysosomes, in which food digestion is completed. These columnar shaped cells are the most abundant in the hepatopancreas and their apical surface is covered by microvilli. Lipid droplets and glycogen granules are usually present in the cytoplasm of digestive cells. The basophilic cells, also called secretory or crypt cells, are typical the main protein secreting cells. These pyramidal-shaped cells contain large amounts of rough endoplasmic reticulum, a well-developed Golgi complex and accumulate secretion granules. They seem to be responsible for the secretion of digestive enzymes, which undertake the extracellular food digestion [18-20,25,23]. In mollusc hepatopancreas, we suppose that duct cells can be involved in lipid storage and nutrient absorption [26]. Moreover, some results suggest that these cells may play a role in digestion [25].

Immunodetection of mSDPLA₂ in the crude hepatopancreas homogenate

Supernatant of the marine snail hepatopancreas homogenate containing 100 μg of total proteins was subjected to SDS-PAGE analysis followed by immunoblotting. Anti-mSDPLA₂ polyclonal antibodies were found to react with a single band of 30 kDa corresponding to the mSDPLA₂ present in the crude extract. No other protein bands were recognized by anti-mSDPLA₂, suggesting a good specificity of these mSDPLA₂ antibodies (Figure 2). Based on the specificity towards mSDPLA₂, polyclonal antibodies were used for immunocytolocalization of mSDPLA₂ in the hepatopancreas sections.

Immunocytolocalization of mSDPLA₂

Sections of hepatopancreas were immunostained against mSDPLA₂ as described above (Figure 4). Only digestive cells displayed a positive labelling for the presence of mSDPLA₂. Conversely, secretory zymogene-like cells were not immunostained. In the control experiments without anti-mSDPLA₂ antibodies, no labeling was observed (Figure 3). Interestingly, we noticed that only few intracellular granules of the digestive cells were immunoreactive. These granules containing mSDPLA₂, tentatively named mSDPLA₂⁺ granules, which were irregular in shape and did not have a specific location in the digestive diverticula (Figure 4 and 5). To further confirm the presence of mSDPLA₂ and the specificity of the mSDPLA₂ polyclonal antibodies, we performed a laser
capture microdissection targeting stained cells of hepatopancreas tissue sections. Selected cells were dissociated and protein extracts were analysed by western blot. Western blotting revealed a strong broad band at around 30 kDa, corresponding to the mSDPLA₂ (Data not showed).

**Discussion**

The ultrastructure of the hepatopancreas was described in some Opisthobranchs [27-30,24], but such studies were never carried out on Neogastropoda, which includes the genus *Hexaplex*. To enlarge our knowledge about these marine molluscs, the structure and function...
of the digestive gland of the gastropod mollusc, H. trunculus, was investigated through electron microscopy analyses and immunohistochemistry.

The digestive gland is composed of two main cell types, the “digestive” cells and the “secretory” cells. The digestive cells appear to be concerned with the absorption and digestion of nutrients, while the secretory cells produce digestive enzymes and calcareous concretions. Undifferentiated cells are scattered between these two cell types.

In previous studies, we have purified and biochemically characterized an original marine snail digestive phospholipase A₂ from the hepatopancreas [12]. Here, we report the ultrastructural study and some cytological aspects of the digestive cells. We performed immunocytochemical and immunofluorescence analysis to specify the tissular and subcellular location of this digestive enzyme. We showed that mSDPLA₂ is located in the digestive cells while no labeling was observed in the secretory zymogene-like cells.

From the granular labeling observed in the digestive cells, mSDPLA₂ appears as an intracellular enzyme involved in the intracellular food digestion process as described for other invertebrates [31]. In mammals, classical digestive enzymes as pancreatic enzymes were detected in intracellular zymogene granules in pancreatic acinar cells. However they act in the lumen of the gastrointestinal tract which supposes a secretory process from acinar cell [32]. The presence of mSDPLA₂ in the digestive cells is not a definitive argument for involving it in phospholipid digestion mechanism located inside these cells. Invertebrate digestive cells can take up partially digested food by endocytosis through the microvilli. It was shown that, using this process, digestive cells of a mollusc Sepia officinalis absorb 80% of a radiolabeled food source [33]. According to Boucaud-Camou and Yim [34], the pinocytotic vesicles fuse together to form heterogeneous phagosomes known as heterophagosomes (Figure 6E). When they combine with primary lysosomes containing the intracellular digestive enzymes, they form secondary lysosomes or heterolysosomes, where the intracellular digestion takes place. We noted that the mSDPLA₂ immunoreactive (mSDPLA₂+) granules belonging to the digestive cells were irregular in shape and size. Small mSDPLA₂+ granules of about 2 μm might be primary lysosomes or lysosomes-like vesicles. However, larger mSDPLA₂+ granules of about 7 μm were most likely secondary lysosomes or heterolysosomes. That can result from the fusion of lysosome-like vesicles and phagosomes (Figure 4). The larger dense granules lacking mSDPLA₂ take a large part of digestive cells are thought to be phagosomes. Furthermore, similar vesicles were previously observed in digestive cells of many invertebrates [35,36]. The number and size of such vesicles can be related to the digestion stage.

The secretory zymogene-like cells must be responsible for secretion of digestive enzymes. They undertake an initial and rapid extracellular food hydrolysis in the diverticula lumen. Then, the partially digested products are likely to be absorbed by pinocytosis (Figure 6E) and stored in
the digestive cell where they will be slowly hydrolyzed by intracellular enzymes for energy generation.

The anatomy and the histology of the molluscs digestive system have been previously studied. However, the knowledge of the ultrastructure and the physiology of the *H. trunculus* digestive system remain unknown (poor). In this histologic study, the ultrastructure of digestive cells of *H. trunculus* was investigated by electron microscopy transmission and immunofluorescence.

In semi-thin sections of *H. trunculus* hepatopancreas, many digestive were observed in the epithelium of the digestive diverticula. Digestive cells exhibited numerous heterolysosomes sometime fused together or with phagosomes (Figure 6A and 6B). The apical surface of digestive cells was covered with microvilli, almost could reach 10 μm in length (Figure 6C). We describe a second cell types for which a single large vacuole occupying almost entirely the cell is observed, while the cytoplasm is reduced to a very thin peripheral layer (Figure 6C).

In digestive cells, the apical zone contained many endocytic vesicles with electron dense materials, in addition to vacuoles with a clear background, usually containing low amounts of dense material. A large part of the cytoplasm was filled with heterolysosomes organelles with variable length and contents. This heterogeneity reflects different stages of intracellular digestion.

In molluscs, the extracellular digestion products found in the lumen of digestive diverticula are collected by digestive cells for further breakdown in heterolysosomes. In some species digestive cells can collect relatively large

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**Figure 4** Immunolabelling of the digestive gland (A and B) Overall view of a sections of the digestive diverticula after incubation with the solution containing pAbs anti-mSDPLA2 (1:100) and then hybridized with appropriate secondary biotinylated-antibodies (1:200) counter-stained with eosine and hematoxylin. (C and D) The section of digestive diverticula showing the occurrence of mSDPLA2 in the lumen of digestive cells, mSDPLA2 localize in the around cells. (E and F) enlarged view of the digestive cells (400×).
food particles by phagocytosis [37], but in others only dissolved substances are captured by small endocytic vesicles for extracellular digestion [38,39].

For *H. trunculus*, many endocytic vesicles filled with electron dense materials were observed in the apical region of *H. trunculus* digestive cells. Probably, these vesicles contained extracellular digestion products that would be transferred to heterolysosomes of digestive cells, to complete the digestive process. In some digestive cells the number of endocytic vesicles was very high, indicating a very intense endocytic activity (Figure 6E). Conversely, some other cells had only few vesicles. That suggests the existence of different digestive steps at the cellular level, with low endocytic activity. The Golgi stacks with dilated cisternae containing dense substances were detected in digestive cells of *H. trunculus*. In the hepatopancreas of *H. trunculus*, digestive cells can also be distinguished by size and electron-density of vesicles in their cytoplasm.

Small and electron dense vesicles are associated with secretory functions, whereas large and electron-lucid vesicles are associated with the absorption functions. Furthermore, the presence of lipid droplets and glycogen granules in the cytoplasm of digestive cells suggests their involvement in the metabolism of lipids (Figure 6G).

The thorough involvement of the hepatopancreas of *H. trunculus* in secretion, digestion and, metabolism absorption was evidenced by the decondensed aspect of nuclear chromatin, presence of rough endoplasmic reticulum, Golgi complex region, lysosomes, vesicles and cytoplasmic inclusions in the digestive cells. The hepatopancreas of *H. trunculus* has been pointed out as the main region for digestion and absorption. The cytoplasm of digestive cells contain granules similar to hemozoin, vesicles with protein content and lipid droplets, indicating the possible function in digestion, absorption, and storage of lipids.

### Method and materials

**Animals dissection and tissue collection**

The marine snails *H. trunculus* were collected from the sea cost of Sfax, Tunisia (Figure 1). They were kept on ice until use in the laboratory. The carapace and the internal organs were separated and the hepatopancreases were immediately collected and stored at -80°C.

**Preparation of hepatopancreas extracts**

Extraction buffer (50 mM Tris-HCl, pH 8.5) was added to the hepatopancreas sample in the proportion of 5 ml per 1 g of fresh tissue and stirred for 45 min at room temperature. Then, 2 mM benzamidine was added to the mixture and centrifuged for 45 min at room temperature. The clear supernatant was collected for protein analysis and phospholipase activity measurement.

**Protein concentration and glycosylation analysis**

Protein concentration was measured spectrometrically according to the Bradford method [40], using bovine serum album as a standard. The presence of glycan chains in the pure mSDPLA2 was checked by anthrone-sulfuric acid method using glucose as a standard [41].

**Production of anti-mSDPLA2 polyclonal antibodies**

Rabbits were injected subcutaneously and intra-muscularly every 3 weeks with 0.5 mg of purified mSDPLA2. The first injection included complete Freund’s adjuvant, and the last two injections contained incomplete adjuvant. Anti-mSDPLA2 serum was collected and its immunoreactivity was verified.

**SDS-PAGE and immunoblotting Analysis**

Analytical polyacrylamide gel electrophoresis of proteins in the presence of sodium dodecyl sulfate was performed as described by Laemmli [42]. The specificity of anti-mSDPLA2 polyclonal antibodies was confirmed by protein blotting. Proteins from SDS-PAGE were blotted onto nitrocellulose membranes. Membranes were then rinsed three times with PBS (10 mM phosphate, 150 mM NaCl, pH 7.2), and saturated with 3% powder milk in PBS for 1 h at room temperature. Thereafter, membranes were incubated for 1 h at room temperature with anti-mSDPLA2 polyclonal antibodies diluted at 1:1000 in PBS containing 0.05% tween-20 (PBS-T). Afterward, they were washed three times with PBS-T and incubated for 1 h at room temperature with a 1:2000 dilution of...
Figure 6 Electron micrographs of the digestive cells. (A, B) the apical region of the cell contains a brush border of microvilli and numerous clumps of α-glucogen granules. (C) digestive adjacent cells. Note some microtubules (arrow) and a cilia root (arrowhead). (D) The cytoplasm of digestive cells contain granules similar to hemozoin, vesicles with protein content and lipid droplets, indicating the possible function in digestion, absorption, and storage of lipids. (E) The pinocytotic vesicles fuse together to form heterogeneous phagosomes known as heterophagosomes. (F) A large residual body is about to extruded into lumen at the end of the process. (G) The vacuola become bigger and distributed through the whole cell.
alkaline phosphatase-conjugated anti-rabbit immunoglobulin (Sigma). After washing as above, membranes were incubated with 0.3 mg/ml of nitroblue tetrazolium chloride (Sigma), 0.2 mg/ml of 5-bromo-4-chloro-3 indolyolphosphate (Sigma) and 0.2 mg/ml of MgCl₂ to reveal the specific immunoreactivity.

Histological, immunochemical and immunofluorescence studies

Immunohistochemistry

Fragments of tissues, with a size of around 1 mm³, were embedded in Optimal Cutting Temperature (OCT) and cryopreserved in isopentane chilled on liquid nitrogen. Sections with a thickness of 4 μm were cut. Slides were saturated with 5% of normal goat serum in PBS with 0.05% Triton X-100 for 15 minutes and subsequently hybridized with primary anti-mSDPLA₂ polyclonal antibodies diluted at 1:200 in Dako Diluent (S3022) for 2 h at 4°C. Slides were rinsed twice and hybridized with (1:200) secondary anti-rabbit biotinylated antibodies diluted at 1:200 for two hours at room temperature.

Immunofluorescence

Tissue sections were fixed in PBS with 4% paraformaldehyde, permeabilized with 0.5% Triton X-100, saturated in 5% normal goat serum in PBS-T and hybridized with anti-mSDPLA₂ polyclonal antibodies diluted at 1:100 in Dako Diluent (S3022) overnight at 4°C. After two washes, cells were hybridized for 2 h with a chromoTM-488-conjugated secondary antibody from rabbit (Abcam, Cambridge, UK) diluted at 1:200 in PBS. The fluorescence analyses were performed with BX41 Olympus upright microscope and pictures were taken with an Olympus C-5060 digital camera. Control experiments were carried out in absence of anti-mSDPLA₂ antibody.

Electron microscopic study

Fragments of tissues, with a size of about 1 mm³ were fixed in 2.5% glutaraldehyde for 30 minutes, rinsed in 0.1 M cacodylate buffer and post-fixed for one hour with 1% OsO₄. Tissues were dehydrated in graded concentrations of alcohol, and then in propylene oxide. Thereafter, tissues were embedded in epoxy resin for 24 h at 60°C. Finally, ultrathin sections were cut and observed with a Joel, Jem-1010 Electron Microscope.

Conclusion

In conclusion, the marine snail digestive phospholipase A₂ is an enzyme localized in the digestive cells of the snail hepatopancreas. Immunocytolocalization study shows that only one irregular-shaped vesicle type; mSDPLA₂⁺ present in the digestive cells of the snail hepatopancreas was found to contain mSDPLA₂. The enzyme appears to be homogenously distributed in these vesicles. This cellular location suggests that intracellular phospholipids digestion, like other food components digestion of snail diet, occurs in these digestive cells.

Abbreviations

mSDPLA₂: marine snail digestive phospholipase A₂; SDS-PAGE: sodium dodecyl sulfate-polyacrylamide gel electrophoresis; ESA: bovine serum albumin; pAbs: polyclonal antibodies; PBS: phosphate buffer saline; DAPI: Di Aminido Phenyl Indol; OCT: Optimal Cutting temperature.

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Authors’ contributions

ZZ carried out all the studies, analyzed the data and drafted the manuscript. NB, AK and LM helped with the analysis, discussion of the data and correction of the manuscript. SB helped with the correction of the manuscript. TR helped with the discussion of the data. YG and HM participated in the study design and helped to draft the manuscript. All authors have read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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