THE PRIMARY METABOLITES OF SARDINE (SARDINA PILCHARDUS) IN THE MOROCCAN
NORTH ATLANTIC

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Abstract

Primary metabolites were determined in muscle, liver and gonads in sardines (Sardina pilchardus) captured in the Moroccan North Atlantic. It emerges from this study that in sardines, in the different seasons of the year, the lipid, protein and carbohydrate reserves are stored in the different organs. This important storage of metabolites in the gonads will undoubtedly be used for maturation; while the reserves stored in muscle and liver will be used for the production of energy necessary for catadromous migration. In addition, there are other biotic and abiotic factors that influence the use and storage of these metabolites such as the quality and quantity of foods and hydrological parameters.

Introduction:

In Morocco, the fishing sector plays an important role on the economic and social plan, its national production was 1,353,780 tons in 2014, which placed Morocco in the first rank of African production and in 17th place in the globally (FAO, 2016). From an economic point of view, this sector plays a very important role; during the last decade, its contribution to the GDP (Gross Domestic Product) is of the order of 2.3% on average. On the social level, the fishing industry generates more than 170,000 direct jobs and 500,000 indirect jobs. 80% of national production is destined for seafood processing industries and 20% for the local market for the consumption of fresh products (MPM, 2014)

This small pelagic species exhibits great seasonal variation in primary metabolites. This seasonal change induced by the seasonal fluctuation of gonad maturation and reproductive activity and somatic growth (Silva, 2006) and of the body condition of the sardine in the Moroccan North Atlantic. (Zwolinski, Stratoudakis, & Soares, 2001)

As in other pelagic species, primary metabolites play an important role in sardine nutrition for energy supply, as structural components of the body, and as essential membrane lipids (Hansen, 2008). Proteins for muscles and using sugars as a fuel for movement, however, the primary metabolites and its composition are associated with the life
cycle and depend on external factors, such as environmental conditions and diet (Bandarra et al., 2011; Watters, Iwamura, Ako, Et Deng, 2012). The primary metabolite composition is the net result of complex dynamic relationships between several factors, the details of which are not fully understood. The main factors are: food intake; oxidative catabolism rates; kinetics of metabolic reactions, where diet seems to play the most important role (Cejas et al., 2004; Hixson, 2014).

Catch and survey samples have been the most common source to provide biological data on Atlantic sardines for several studies.

The aim of this work is to evaluate the changes of the primary metabolites of sardine in the Moroccan North Atlantic in the different seasons.

**Material and Methods:-**

**Preparation of crude extracts:**

All the operations are carried out at 4 °C. Each tissue is cut into small pieces, then added with a buffer solution at pH 7.4 containing: 2 mM EDTA, 1 mM 2 β-mercaptoethanol, 1 mM PMSF, 0.25 M sucrose, 10 mM HEPES.

In order to burst the cells of the tissues, we performed a grinding by ultra-turrax type (Apolymix ®) then a pottering (maximum speed, 10 turns) using a potter (Elvehjem ®), and end sonication by a sonicator of the type (Bandelin electronic ®) followed by filtration on the biogas layers in order to eliminate cellular debris which has not been well ground. The collected filtrate is centrifuged at 11,200 g for 30 min at 4 °C. by type centrifugation (Hettich®). The supernatant obtained represents the cytoplasmic fraction which will be used for assays of the enzymatic activities. The pellet is resuspended in a small volume of the buffer, the latter is centrifuged at 100,000 g for 30 min at 4 °C and purified to obtain the mitochondrial fraction which will be used for the mitochondrial enzymatic assays.

**Protein determination:**

The proteins were quantified according to the technique of Bradford (1976) which is a sensitive, simple and rapid method. It is based on the use of the Bradford reagent (*) which contains coomassie blue G 250 which binds to the proteins giving a coloration ranging from brown for low concentrations to blue for high protein concentrations.

The volume of the samples 200 μl is added with 800 μl of Bradford's reagent. After 15 min of incubation at room temperature and in the dark, the absorbance of the color developed is measured at 595 nm using the spectrophotometer (3 Thermo electron corporation ®) and the unknown protein concentration is deduced by referring to the calibration curve produced using BSA (Bovine Serum Albumin).

**Electrophoresis under denaturing conditions (PAGE-SDS):**

The proteins are separated by vertical plate electrophoresis according to the technique of Laemmli (1970). It is carried out in a device for mini-gel (8 x 10 cm), the wedges used have a thickness of 1mm.

The use of a denaturing agent such as sodium dodecyl sulfate (SDS) causes the quaternary structure to dissociate into monomeric forms, to which it binds. Under these conditions, the separation of proteins on the gel takes place only according to their size, since their charge density is made negative overall due to the presence of SDS.

7.1. Preparation of gels

12% separation gel
1. Acrylamide 29.2% - Bisacrylamide 0.8% 4 ml
2. 1.5 M Tris-HCl pH 8.8 2.5 ml
3. SDS 10% 100 μl
4. Double-distilled water 3.33 ml
5. Ammonium persulfate 10% 50 μl
6. TEMED 20 μl
7. 4% concentration gel
8. Acrylamide 29.2% - Bisacrylamide 0.8% 0.675 ml
9. 0.5M Tris-HCl, pH 6.8 1.25 ml
10. SDS 10% 50 μl
11. Double-distilled water 3.05 ml
12. Ammonium persulfate 10% 30 µl
13. TEMED 20 µl
14. Migration buffer
15. It is made up of the following solutions:
16. 25 mM Tris-HCl pH 8.3
17. 320 mM glycine
18. SDS 0.1%

**Sample solubilization and migration:**
A volume containing 40 µg of proteins is dissolved in 10 µl of the loading buffer, the composition of which is as follows: 60 mM Tris-HCL (pH 6.8), SDS 1% (w / v), Glycerol 10% (v / v), Bromophenol blue 0.01% (w / v) and β-Mercaptoethanol 1% (v / v), then heat for 3 minutes at 100 °C, and loaded into the wells of the gel. Migration is carried out at room temperature with a constant voltage of 100V.

**Coloring and discoloration of gels:**
The gels are colored overnight with stirring at room temperature with a 0.25% solution of Coomassie blue, in a “methanol - distilled water - acetic acid” mixture with a ratio (4/5/1). The gels are then decolorized in the same mixture without the coomassie blue.

**Determination of total lipids:**
From each of the samples of sardine fillets, the total lipids were extracted by the method of Folch et al (1957). Five grams of sample are ground with the potter, 20 ml of methanol (MeOH) and 10 ml of chloroform (CHCl3) was added and vortexed for 2 minutes in a tube, 10 ml of (CHCl3) was added and stirred vigorously for 2 minutes. 10 ml of distilled water was added and the mixture was again vortexed for 2 minutes. The layers were separated by centrifugation at 448 g / 10 min. The lower layer was transferred to a vial with a Pasteur pipette. The method is repeated three times to exhaust the sample. After centrifugation, the CHCl3 phase was added to the first extract. Evaporation was carried out in a rotavapor at 45 °C. The weighing of the flask containing the lipid extract after evaporation of the solvent makes it possible to calculate the lipid content expressed in g per 100 g of sample, using the formula next:

\[ MG = \frac{(P2 - P1)}{Pe} \times 100 \]

P2: weight of the balloon containing the lipids.
P1: weight of the empty balloon.
Pe: test take.

**Gas chromatography coupled with mass spectrometer (GC / MS):**
The lipid extracts are saponified beforehand using KOH (0.5 N) and then methylated with a mixture of methanol-n-hexane according to the method of Nasopoulou et al. (2012). The methyl esters of the fatty acids are then separated, quantified and analyzed by gas chromatography coupled with a mass spectrometer (Chromatograph: Shimadzu CLASS VP (GC17A. Kyoto, Japan) on a capillary column 60 cm long and 250 μm in diameter. (Agilent, Santa Clara California, USA) The operating conditions for gas chromatography are as follows:
1. Injector and temperature sensor (220 °C and 225 °C) respectively.
2. The oven temperature has been programmed to increase from 45 °C to 240 °C (at the rate of 20 °C to 35 °C / minute).

1 µl aliquots were injected with bicyanopropil phenyl silicone as the stationary phase and hydrogen was used as the carrier gas. Peak fatty acids were identified by comparison with methyl retention time and quantification of alpha male acids was made by reference to an internal standard by palmitic acid (C16: 0).

**Determination of total sugars:**
The total sugars are determined according to the method of Dubois et al, (1956), the principle of which is based on the following reaction: the sample volume 100 µl is added with 200 µl of phenol and 1 ml of concentrated sulfuric acid causes, when hot, the departure of several water molecules from the ose. This dehydration is accompanied by the formation of a hydroxy-methylfurfural (HMF) in the case of hexose and of a furfural in the case of a pentose and which will give colored complexes (yellow-orange). The intensity of the coloring is proportional to the concentration of the ose. Optical density is measured at 490 nm using a spectrophotometer.
Resultants:
In this study, we investigated the monthly variations of the overall chemical composition of sardine: Sardina pilchardus in the Moroccan North Atlantic region. The present work, which concerns the variations of proteins, sugars and lipids in the different organs of the sardine (muscle, gonad and liver), in order to determine these modifications according to the months.

Determination of sugars:
The sugar content in the muscle, gonad and liver organs shows variation throughout the year as well as the variation is not very similar for the two sexes with a maximum value in February and a minimum value in June, knowing that the succession periods of sexual maturation and organ development even lead to significant fluctuations in the state of sugars.
**Protein determination:**
The monthly evolution of the sardine protein content is shown in figure 30. We observed an increase in this content from April to reach a maximum value in August for the male and female gonads, then we noted a slight increase in the protein content in the same period for the liver, at the end we observed a gradual increase for the muscle for both sexes.

![Graph showing protein content evolution](image)

**Evolution of the protein level of sardines**

**SDS-PAGE electrophoresis:**
Electrophoresis is the displacement of an ionized particle in a given electric field. The principle of this method is therefore based on the amphoteric nature of proteins. Ionization involves the acidic or basic end groups, amino residues that determine the net charge of the protein. The speed of protein migration inside the support (starch or polyacrylamide gel) depends on the net charge at a given pH, the molecular weight and the different interactions that the protein can have with its solvent.
Determination of total lipids:
The assays carried out on Sardina pilchardus during the aforementioned periods gave rise to the results shown in Table 13, these are the amounts of lipids in the various organs of the sardine reported in the fresh masses. During the period of our study, we note that the content of muscle lipids varies depending on the month with a maximum value of 0.05 g / g for males and for females 0.07 g / g of the mass of the fish. While the gonad lipid content is 0.03 g / g for males and 0.05 g / g for females. The lipid levels in the liver in males and females are the same 0.02 g / g.

| Month    | Male Muscle | Male Gonad | Male Liver | Female Muscle | Female Gonad | Female Liver |
|----------|-------------|------------|------------|---------------|--------------|--------------|
| January  | 0.01        | 0.03       | 0.01       | 0.02          | 0.03         | 0.01         |
| February | 0.03        | 0.03       | 0.01       | 0.04          | 0.04         | 0.02         |
| March    | 0.03        | 0.01       | 0          | 0.03          | 0.02         | 0            |
| April    | 0.01        | 0          | 0          | 0.02          | 0            | 0            |
| May      | 0.01        | 0          | 0          | 0.02          | 0            | 0            |
| June     | 0.03        | 0          | 0          | 0.03          | 0            | 0            |
| July     | 0.04        | 0.01       | 0          | 0.04          | 0.02         | 0            |
| August   | 0.05        | 0.01       | 0          | 0.06          | 0.01         | 0            |
| September| 0.05        | 0.02       | 0.01       | 0.07          | 0.05         | 0.01         |
| October  | 0.05        | 0.01       | 0.01       | 0.05          | 0.04         | 0.01         |
| November | 0.04        | 0.01       | 0.02       | 0.03          | 0.01         | 0.02         |
| December | 0.02        | 0.01       | 0.01       | 0.02          | 0.01         | 0.01         |

Quantity of lipids of sardine

Analysis of total fatty acids:
The fatty acids contained in the three organs were analyzed before performing the lipid class analysis. These results are reported in Table X. For all the analyzes, the majority fatty acid is palmitic acid.

The distribution in AGS, MUFA and PUFA of fatty acids in the different organs of the sardine is represented respectively 40.1%, 27.7% and 32.2% for the liver, 42.4%, 21.31% and 36.29% for muscle, 38.55%, 32.2% and 38.25% for gonads. The most important fatty acid families are respectively w3 about 22.5%, 28.85% and 30.2%, for w6 about 11.7%, 7.2% and 6.68%.

Distribution of fatty acids (%) within the three organs (liver, muscle and gonad) of the sardine
## Discussion:

The comparison of the three primary metabolites lipid, protein and carbohydrate of the different studied organs of the sardine that we have studied, allows noticing that at all levels there is variation over the whole year. This variation is well recorded during the breeding season than during the sexual rest period. This could be explained by the use of part of this primary metabolite for reproductive needs to which would be added the energy needs allowing the fish to overcome the winter season.

Lipid deposits increase steadily during ontogeny. For the liver is not involved in the transfer of lipid reserves necessary for vitellogenesis. This confirms the classification of S. pilchardus among fatty fish characterized by lipid accumulation in muscles and by subcutaneous fat deposition, unlike lean fish where lipids accumulate in the liver before passing to the gonads (Xu et al. 2000). In sardines, lipid storage takes place in the muscles, within or between muscle fibers (Yashodhara et al. 2009). Maximum adiposity values are recorded at the onset of maturation of the genitals, while they decrease in December and January when the gonad mass is maximum, thus confirming that S. pilchardus uses muscle and mesentery lipid reserves for develop gametes. The increase in fattening is due to the availability of food resulting from significant plankton production linked to the upwelling season (Makaoui et al. 2008). In fact, the sardine feeds intensively in summer and stores these energy reserves which will be allocated gradually to the maturation of the gonads and to reproduction. The adaptation condition of females is generally superior to that of males regardless of the study area. This indicates that the mass of females is greater than that of males for the same size (Bacha et al., 2007). Environmental conditions, in particular the temperature factor (Makaoui et al., 2005) as well as the quality and quantity of feed directly influence fattening. At the same time, the protein storage capacity is extremely limited. Fish do not have a reserve form of protein, such as glycogen for sugar or adipose tissue for lipids. If the amino acids that come from protein digestion are not used immediately by the body, they are converted into glucose for energy (Reynolds et al, 2003).

When the glycogen stock is depleted, such as during a bad weather period or a prolonged migration, and if the adipose tissue is poorly developed, the musculature therefore acts as a kind of reserve of amino acids.

The chemical composition of fish varies considerably from species and individual to individual depending on age, sex, environment and season. Changes in the chemical composition of fish are closely related to their diet, migratory

|          | Liver | Muscle | Gonad |
|----------|-------|--------|-------|
| C14:0    | 11.5  | 12.4   | 10.87 |
| C16:0    | 21.3  | 21.6   | 19.74 |
| C18:0    | 5.5   | 6.1    | 6.74  |
| C20:0    | 0.5   | 0.8    | 0.4   |
| Other    | 1.3   | 1.5    | 0.8   |
| **Saturated** | **40.1** | **42.4** | **38.55** |
| C16:1    | 9.2   | 10.43  | 10.73 |
| C18:1    | 10.9  | 8.95   | 8.82  |
| C20:1    | 3.4   | 0.16   | 1.45  |
| C22:1    | 3.4   | 1.4    | 2.2   |
| Other    | 0.8   | 0.37   | -     |
| **monounsaturated** | **27.7** | **21.31** | **23.2** |
| C16 : 2n4 | 1.4  | 0.4    | 0.82  |
| C18 : 2n6 | 6.8  | 6.08   | 5.89  |
| C18 : 3n3 | 0.6  | 0.24   | 1.23  |
| C18 : 4n3 | 1.1  | 0.4    | 0.1   |
| C20 : 4n3 | 1.5  | 0.96   | 1.34  |
| C20 : 5n3 | 6.5  | 10.95  | 11.34 |
| C22 : 5n3 | 1.1  | 1.1    | 0.7   |
| C22 : 6n3 | 13.2 | 16.16  | 16.83 |
| **polyunsaturated** | **32.2** | **36.29** | **38.25** |
| Σ Ω 3    | 22.5 | 28.85  | 30.2  |
| Σ Ω 6    | 11.7 | 7.2    | 8.68  |
| Ω 3/ Ω 6 | 1.92 | 4.01   | 3.48  |
movements and sexual changes in relation to spawning. Fish will have periods of starvation for natural or physiological reasons (such as migration and spawning) or because of external factors such as lack of food. Usually spawning, whether it occurs after a long migration or not, is very energy intensive. Fish that have energy reserves in the form of lipids will rely on these. Species that migrate for a long time, before reaching specific spawning sites, can use protein in addition to lipids for energy and therefore deplete their stores of both lipids and proteins. This leads to a general reduction in the physical condition of the fish. In addition, most species feed very little during their spawning migration and are therefore unable to obtain energy from food (Kacem, 2000).

One method, useful classified fish in lean species and fatty species, consists in considering as lean the fish which store lipids only in the liver and as fatty fish the fishes retaining the lipids in fat cells distributed in other tissues of the body. Typically lean species are ground fish such as cod and hake. Fat species include pelagic like sardines, sardinella, herring, and mackerel. Some species store lipids only in limited parts of their body tissue, or in smaller amounts than typically fatty species and are therefore called semi-fatty species, (mullet and shark) (Aymen et al 2010).

Our study shows the seasonal variation of fat deposition in sardines. We see that lipid levels in different tissues vary widely and this is consistent with the work of Shepherd et al, 2005. Lipid stores are typically used for large spawning migrations and gonad development.

**Conclusion:**
The comparison of the primary lipid, protein and carbohydrate metabolites in different muscle, liver and gonad organs of sardines in the Moroccan North Atlantic that we have studied, allows us to notice that at all levels there are fluctuations of the three metabolites in the different seasons of the year which is well noticed in the breeding period (February) than in the period of sexual rest (August). This could be explained by the use of a part of the metabolites for the needs of reproduction to which would be added the energy needs allowing the fish to overcome the winter season.

Furthermore, the results of this study show that the primary metabolites are stored and used unevenly in the sardine organs as needed.

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