A Comparative study of hematological and blood chemistry of Indian and Italian Grey Mullet (Mugil cephalus Linneaus 1758)

Fazio Francesco¹, P. Satheeshkumar²*, D. Senthil Kumar³, Faggio Caterina⁴ and Piccione Giuseppe¹

Abstract

**Background:** Blood hematological and serum biochemistry parameters are often used to assess the health status and as stress indicators in fishes. In the present study was undertaken to comparative study of hematological and blood chemistry of Indian and Italian Grey Mullet (Mugil cephalus) to establish baseline values. Fifteen sexually immature and disease-free Indian wild fish (26.25 ± 0.23 cm total length, 347.55±14.27 g weight) and 15 Italian fish (31.53 ± 1.08 cm total length, 416.5 ± 14.56 g weight) were examined.

**Results:** Statistical analysis revealed that differences in hematological and biochemical parameters between two region mullet fish were significant (P<0.01). The RBC/ WBC level increased due to the decrease in WBC during the comparative study.

**Conclusions:** The study showed that the environmental conditions significantly impacted the status of the fish. It is suggested that these physiological parameters can be conveniently employed as health monitoring tools in fish culture practices. Biochemical parameters also are indicative of the habits of fishes, and can be used for confirming the maturity and monitoring any changes in the quality of waters.

**Background**

Mullets have worldwide distribution and inhabit tropical and temperate waters and few spend their lives in freshwater [1]. They are grouped taxonomically in the family Mugilidae includes 17 genera and 72 species in the world [2]. Mullet (Mugil cephalus) are Perciform species, which feeds mainly on zooplankton, benthic organisms and detritus, and was chosen because it possesses several characteristics required in an estuarine sentinel species, such as the extreme salinity tolerance [3]. Understanding the haematological characteristics is an important tool that can be used as an effective and sensitive index to monitor physiological and pathological changes in fishes. Normal ranges for various blood parameters in fish have been established by different investigators in fish physiology and pathology [4,5]. Hematological and biochemical parameters are being used as indicators in the measurement of health conditions and toxicological symptoms of organisms [6]. While providing information about the health status of organisms, these parameters may also indicate abnormal environmental conditions [7]. Information about the existence, status and degree of possible sickness in organisms can be rapidly obtained by with use of hematological and biochemical parameters. One of the difficulties in assessing the state of health of natural fish population has been the paucity of reliable references of the normal condition [5,8]. Although fish haematology continues to offer the potential of a valuable tool, progress in establishing normal range values for blood parameters has been sparse and literature in this area is isolated and often incomplete [5]. Despite advances in fish medicine in recent years, interpretation of fish haematology is often troubled by a lack of meaningful reference values and the bewildering diversity of fish species [9]. Only a few normal values for a small number of haematological parameters have been established for some teleosts, but these values range widely due to the lack of standardized collecting and measuring techniques. Haematological studies help in understanding the relationship of blood characteristics to the habitat and adaptability of the species to the environment. A multitude of intrinsic and extrinsic factors cause normal and abnormal variations in haematologic data [9] such as species and strain [10], temperature [10,11], age [12], stress [14], photoperiod [15], nutritional state [12,16], the cycle of sexual maturity, health condition [17], and water quality. The main objective of this present study was to comparative investigation on haematological and biochemical parameters on M. cephalus, captured in two different habitat Faro lake (Italy) and Vellar estuary (India). These data can be used to monitor the physiological status of individual species to assess in wild and in aquaculture.

**Material and methods**

**Study area**

**Faro Lake**

Cape Peloro is a brackish system located in the north-eastern corner of Sicily (Lat 38°15’57” N; Long 15°37’50” E). It consists of two basins, Ganzirri and Faro, communicating with the Tyrrhenian Sea by English canal and connected to each other by Margi canal [18].

**Vellar Estuary**

Healthy marine teleost fishes were collected from Vellar estuary (Lat. 11°29 N and Long 79°46 E), Parangipettai of southeast coast of Tamil Nadu, India. The estuary is seasonally bar-built and semi-diurnal type flows eastwards and empties into the Bay of Bengal at Parangipettai on southeast coast of India, carrying the wastes from the adjacent agriculture lands and industries in addition to domestic municipal and distillery effluents.

© 2012 SatheeshKumar et al; licensee Herbert Publications Ltd. This is an Open Access article distributed under the terms of Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0). This permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
Sampling and analytical methods

In this present study 30 *M. cephalus* were investigated in two different habitat Faro Lake (Italy) and Vellar estuary (India). They were divided into two equal groups on the basis of the site of collection. 15 fish were caught in Faro Lake – Sicily (group A) and 15 were caught in Vellar estuary (India) (group B). All fish were caught with bottom-set nets and blood samples were immediately collected. All fish samples were collected from 2007 to 2008. At the end of blood sampling on all subject weight and length were recorded (Table 1). On the basis of their weight and length all fish were considered sexually mature and with age between two and four years [19]. Water sampling was carried out in the same date of fish sampling, in three stations points of Faro Lake and Vellar Estuary. The three stations on the each location were selected randomly and the distances among them were about 3 meter to 1.5 Km. In Faro lake, water samples collected by Niskin bottle (General Oceanics, Inc.-Miami, Florida) for sampling and a multiparametric probe YSI 85 System for temperature, salinity, dissolved oxygen (DO) and pH. In Vellar estuary from the collection site, the water quality dissolved oxygen was estimated by Winkler’s methods [20], salinity by an Erma hand refractometer (Tokyo). The pH and water temperature were measured by using pH tester pen (Japan) and thermometer, respectively.

Blood samples were collected by caudal vein/ direct heart puncture by using a sterile plastic syringe (2.5 mL) and transferred into 2 different tubes, one (Miniplast 0.5 ml, LP Italiana Spa, Milano) containing EDTA (1.26 mg/ 0.6 mL) as an anticoagulant agent and the other without EDTA. The blood samples were collected in EDTA tubes were used for the determination of haematological profile. Heparin sodium (1%) was used as an anticoagulant [21]. The collected blood samples were immediately subjected to hematological analysis. Evaluation of the haemogram involves the determination of the Red Blood Count (RBC), Haematocrit (Hct), Hemoglobin concentration (Hgb), White Blood Cell Count (WBC), Mean Corpuscular Volume (MCV), Mean Corpuscular Haemoglobin (MCH), and Mean Corpuscular Haemoglobin Concentration (MCHC).

The bloods were diluted with appropriate diluting fluids for RBC and WBC counts and were determined using improved Neubauer haemocytometer and calculated [22,23]. Replicated counts were made for each blood samples to minimise the error. Hematocrit was determined by microhematocrit centrifugation. Microcapillary tubes were filled, plugged with clay, and centrifuged at 19,000g for 5 minutes. Measure the length of the columns containing packed red cells, and packed red cells plus supernatant. The calculation of hematocrit is as follows: (packed red cells/packed red cells plus supernatant)/100%. Haemoglobin concentration was measured with Hb test kit (Roach GmbH Mannheim, Germany) using the cyanmethemoglobin method [24] and Sahili Haemoglobinometer. Mean corpuscular volume (MCV), mean corpuscular haemoglobin (MCH) and mean corpuscular haemoglobin concentrations (MCHC) were calculated indirectly by the above direct parameters values using standard formulas.

For the assessment of glucose on whole blood, a portable blood glucose analyzer (ACCU-Check Active, Roche Diagnostics GmbH, Mannheim, Germany). On serum samples obtained from blood samples without EDTA by centrifugation for 10 min at 3000 rpm, total protein, cholesterol and urea were determined by means of commercial kits (BioSystem, Barcelona, Spain) using on UV spectrophotometer (Slim, SEAC, Florence, Italy). In Indian mullet, Biochemical estimation of blood glucose, protein, cholesterol, and urea were determined by following standard methods [25,26,27,28].

Statistical Analysis

Differences in haematological parameters between the two study stations marine teleost fish were statistically analyzed by Unpaired T-test. Mean and standard error (SEM) were calculated for each parameter. All these statistical analyses were performed using the statistical software Prism v. 4.00 (Graphpad Software Ltdt., USA, 2003).

Results

Table 1 shows the physico chemical parameters of both sampling stations Faro Lake (Italy) and Vellar Estuary (India). The temperature of Indian station was higher than Italian with a statistical p value of (p<0.05).The other parameters such as pH, salinity and DO were high in Italian station than India and with p value (p< 0.1, p< 0.001 and p< 0.001) respectively. Table 2 indicates the biometric parameters of Grey Mullet (*M. cephalus*) collected in both stations. 15 sexually immature and disease-free Indian wild fish (26.25 ± 0.23 cm total length, 347.55±14.27 g weight) and 15 Italian fish (31.53 ± 1.08 cm total length, 416.5 ± 14.56 g weight) were examined. Among these Faro lake fishes were with high mean length (31.53±1.08cms) and weight

| Site of collection | Length (cm) | Weight (g) |
|--------------------|-------------|------------|
|                    | Mean±SEM    | Min | Max | CV (%) | Mean±SEM | Min | Max | CV (%) |
| Faro Lake (Italy)  | 31.3±1.08   | 19.50 | 36  | 13.28  | 416.5±14.36 | 340 | 510 | 13.54 |
| (n=15)             |             |      |     |        |           |     |     |      |
| Vellar estuary (India) | 26.25±0.23* | 25.00 | 28.1| 3.40   | 347.55±14.27* | 235 | 410 | 15.90 |
| (n=15)             |             |      |     |        |           |     |     |      |

Significance: *Vs Faro Lake (Italy)

| Parameters | Faro Lake (Italy) | Vellar estuary (India) | P value |
|------------|-------------------|------------------------|---------|
| T (°C)     | 25.30±0.29        | 31.83±2                | <0.05   |
| pH         | 8.11±0.013        | 7.8±0.24               | <0.01   |
| Salinity (ppt) | 33.43±0.48   | 26.16±1.43            | <0.001  |
| DO mg l-1  | 5.89±0.033        | 4.48±0.30              | <0.01   |

T=Temperature; DO=Dissolved Oxygen
(416.50±14.56 gms). Both the biometric parameters were significant with Vellar estuary samples. The haematological parameters of *M. cephalus* of Faro lake (Group A) and Vellar estuary (Group B) were expressed in Table 3. RBC, RBC/WBC ratio and Hct % were all higher in Group A than B with a P value of 0.0001. Remaining parameters such as WBC, MCH, MCHC and MCV were elevated in Group B than A with p value of 0.0001 and 0.1 respectively. The biochemical parameters detected in the serum of both group fishes were shown in Table 4. Among the organic constituents proteins, glucose, urea and cholesterol were found higher in Group A and are statistically significant (p<0.0001 and p<0.2 in urea).

**Discussion**

The physiological condition in fishes required for their selection as broodfish can be determined by hematological studies. The blood constituents in teleost fishes are influenced by factors like temperature ecological habitat, food selection and mode of life. Therefore, it is difficult to establish any normal values for the class as a whole. But, if data are collected for different species as well as within species under different conditions some normal ranges of values can be arrived at, which can form a valuable diagnostic aid in fisheries [29]. Studies of blood parameters had proven to be a valuable approach for analysing the health status of fish and help in understanding the relationship of blood characteristics to the habitat and adaptability of the species to the environment [30,31]. The ranges of normal values of the key biochemical parameters are still undefined for same species living in different habitat. In the aquatic habitat, the fish homeostatic system is continuously affected by the changes of the level of salinity, temperature, pH, oxygen concentration [32]. The physiological response to environmental variations such as salinity, temperature and DO fluctuations in aquatic system has been investigated in fresh and marine water species [33,34]. In our study significant differences were observed in some haematological and biochemical parameters between groups A and B from Italian and Indian sampling sites respectively (Table 3 and 4). In particular our results showed that RBC values, Hct and RBC/WBC ratio were higher in group A compared to group B while Hgb, WBC, MCH and MCHC values were lower in group A compared to group B. As fish are very susceptible to environmental physical and chemical changes which may be reflected in their blood components [34], the significant differences in some haematological and biochemical parameters in the two groups of *M. cephalus* could be attributed to different habitat conditions. In fact, as shown in Table 2 significant differences in physico-chemical factors of water, investigated in the two sampling sites were observed. In particular salinity and DO values of Faro Lake were higher than those of Vellar estuary, while temperature was lower in the Italian site compared to Indian site. Salinity variation between the two sampling sites could justify the higher Hct values in group A compared to group B in according with a study on rainbow trout [35].

Our results showed that when water temperature increased and salinity and DO decrease RBC count decreased and Hb increased. It has been observed that blood parameters such as Hct, Hb and RBC count are related to environmental factors such as water temperature and salinity. Moreover, the relationship between haemoglobin and oxygen shows adaptations not only to environmental conditions but also to metabolic requirements, both of which govern oxygen availability and transport to tissue [36]. These adaptations may involve quantitative changes in total Hb content, or qualitative changes in Hb-oxygen-binding properties, and may appear both at the inter- and intra- specific level [37].

The differences in WBC count in the two groups of *M. cephalus* studied here could be the result of different body size of fish as there was statistical difference in biometric data of the two fish groups as it is known that differences in WBC may be attributed to many factors, both biotic (such as age, season, maturity, pathogens) and abiotic (including water temperature, pH, dissolved oxygen content) and in particular to stress [38]. There is an inverse relationship between haemoglobin and oxygen shows adaptations not only to environmental conditions but also to metabolic requirements, both of which govern oxygen availability and transport to tissue [36]. These adaptations may involve quantitative changes in total Hb content, or qualitative changes in Hb-oxygen-binding properties, and may appear both at the inter- and intra- specific level [37].

The differences in WBC count in the two groups of *M. cephalus* studied here could be the result of different body size of fish as there was statistical difference in biometric data of the two fish groups as it is known that differences in WBC may be attributed to many factors, both biotic (such as age, season, maturity, pathogens) and abiotic (including water temperature, pH, dissolved oxygen content) and in particular to stress [38]. There is an inverse relationship between haemoglobin and oxygen shows adaptations not only to environmental conditions but also to metabolic requirements, both of which govern oxygen availability and transport to tissue [36]. These adaptations may involve quantitative changes in total Hb content, or qualitative changes in Hb-oxygen-binding properties, and may appear both at the inter- and intra- specific level [37].

The differences in WBC count in the two groups of *M. cephalus* studied here could be the result of different body size of fish as there was statistical difference in biometric data of the two fish groups as it is known that differences in WBC may be attributed to many factors, both biotic (such as age, season, maturity, pathogens) and abiotic (including water temperature, pH, dissolved oxygen content) and in particular to stress [38]. There is an inverse relationship between haemoglobin and oxygen shows adaptations not only to environmental conditions but also to metabolic requirements, both of which govern oxygen availability and transport to tissue [36]. These adaptations may involve quantitative changes in total Hb content, or qualitative changes in Hb-oxygen-binding properties, and may appear both at the inter- and intra- specific level [37].

| Haematological parameters | Faro Lake (Italy) (group A) | Vellar estuary (group B) | P value |
|---------------------------|-----------------------------|--------------------------|---------|
| RBC (x106 μ/ L)           | Min: 2.21 Max: 4.47 Mean ± SEM: 3.53±0.16 | Min: 2.7 Max: 4.47 Mean ± SEM: 2.51±0.026 | P<0.0001 |
| WBC (x103 μ/ L)           | Min: 16.30 Max: 21.00 Mean ± SEM: 18.30±0.40 | Min: 27.56 Max: 29.21 Mean ± SEM: 28.14±0.13 | P<0.0001 |
| Hgb (g/dL)                | Min: 5.70 Max: 13.30 Mean ± SEM: 10.65±0.60 | Min: 37.03±0.21 Max: 38.5 Mean ± SEM: 35.4 | P<0.0001 |
| RBC/WBC (%)               | Min: 1.07 Max: 2.39 Mean ± SEM: 1.94±0.08 | Min: 0.076 Max: 0.088 Mean ± SEM: 0.82±0.04 | P<0.0001 |
| Hct (%)                   | Min: 21.0 Max: 50.0 Mean ± SEM: 39.60±2.10 | Min: 27.56 Max: 29.21 Mean ± SEM: 28.06±0.11 | P<0.0001 |
| MCV (fl)                  | Min: 95.02 Max: 138.6 Mean ± SEM: 111.5±2.45 | Min: 10.71 Max: 12.3 Mean ± SEM: 115.6±1.48 | P<0.1 |
| MCH (pg)                  | Min: 25.44 Max: 34.50 Mean ± SEM: 29.83±0.60 | Min: 140 Max: 156 Mean ± SEM: 148±1.17 | P<0.0001 |
| MCHC (g/ dL)              | Min: 21.09 Max: 31.43 Mean ± SEM: 26.88±0.62 | Min: 117 Max: 135 Mean ± SEM: 129.2±1.20 | P<0.0001 |

| Parameters | Faro lake (group A) | Vellar estuary (group B) | P value |
|------------|---------------------|--------------------------|---------|
| Total proteins (mg/dL) | 2.05±0.16 | 3.58±0.041 | P<0.0001 |
| Glucose (mg/ dL) | 70.80±1.51 | 84.05±1.06 | P<0.0001 |
| Cholesterol (mg/dL) | 175.7±2.79 | 195.4±2.88 | P<0.0001 |
| Urea (mg/dL) | 5.44±0.96 | 6.58±0.10 | P<0.2 |

Table 3. Mean values ± SEM of haematological parameters obtained into two experimental groups (abbreviations are explained in the text).

Table 4. Mean values ± SEM of biochemical parameters obtained into two experimental groups.
of MCH and MCHC values as effect of different habitat conditions are controversial. In our study high salinity and temperature lead to a lower MCH and MCHC values while [39] in pike perch noted an increase of these parameters with an increased salinity level.

Significant variations in some blood biochemical parameters between group A and B were observed in this study. In particular, our results showed lower concentrations of total proteins, glucose and cholesterol in group A than group B. These parameters are considered to be major indices of the health status of teleosts. Increased concentrations of total proteins can be caused by structural liver alterations reducing aminotransferase activity, with concurrent reduced deamination capacity and impaired control of fluid balance [40]. Elevated levels of cholesterol indicate disorders of lipid and lipoprotein metabolism, especially liver disease [41]. However the differences in blood biochemical parameters found in the two experimental groups of this study were attributed to the different habitat, in fact it is known that the ranges of serum biochemistry can be influenced by many biotic and abiotic factors such as water temperature, seasonal pattern, food, age and sex of the fish [36].

Conclusions
The results of this study provide the knowledge of the characteristics of haematological and biochemical parameters of M. cephalus from two different habitats and show that many physico-chemical factors of water influence the ranges of haematology and of serum biochemistry of fish within same species suggesting that blood parameters may therefore be a value in monitoring the effects of habitat changes on fish biology and fish culture practices.

Competing interest
The Authors declare that they have no competing interest.

Author information
1Department of Experimental Science and Applied Biotechnology. Laboratory of Veterinary Chronophiology. Faculty of Veterinary Medicine. Polo Universitario Annunziata, University of Messina. 98168, Messina, Italy.
2Kandaswamy Kandar Arts and Science College, Department of Zoology Paramathi-Velur 638 181, India.
3Department of Life Science “M. Malpighi” Faculty of Science MM.FF.NN. University of Messina. Viale Ferdinando Stagno d’Alcontres 31, 98166 S. Agata-Messina Italy.

Article history
Received: 21-Apr-2012 Accepted: 8-May-2012 Published: 9-May-2012

References
1. Nelson JS: Fishes of the World. John Wiley and Sons, Inc. 4th Edition. Edited by Hoboken p. New Jersey. 601. | Book
2. Turan C, Gürlek M, Ergüden D, Yägloğlu D, Oztürka B: Systematic status of nine Mullet species (Mugilidae) in the Mediterranean Sea. Turk J Fish Aquat Sci 2011; 11:315-321. | Article
3. Ferreira M, Moradas-Ferreira P, Reis-Henriques MA: Oxidative stress biomarkers in two resident species, mullet (Mugil cephalus) and flounder (Platichthys flesus), from a polluted site in River Douro Estuary, Portugal. Aquat Toxicol 2005; 71(1):39-48. | Article | Pubmed
4. Darvish Bastami K, Haji Moradiou L, Mohamadi Zaragabadi A, Salehi Mir SV, Shakiba MM; Measurement of some haematological characteristics of the wild carp. Comp Clin Patho 2009; 18(3):321-323. | Article
5. Satheeshkumar, P, Ananthan, G, Senthil Kumar, D, Jagadeesan, L; Haematology and biochemical parameters of different feeding behaviour of teleost fishes from Vellari estuary, India. Comp Clin Patho 2011. | Article
6. Rao JV: Biochemical alterations in euryhaline fish, Oreochromis mossambicus exposed to sub-lethal concentrations of an organophosphorus insecticide, monocrotophos. Chemosphere 2006; 65(10):1814-20. | Article | Pubmed
7. Elahee KB, Bhagwant S: Haematological and gill histopathological parameters of three tropical fish species from a polluted lagoon on the west coast of Mauritius. Ecotox Environ Saf 2007; 68:361-371. | Article | Pubmed
8. Kori-Siakpere O, Ake JEG, Idoge E: Haematological characteristics of the African snakehead, Parachanna obscura. Afr J Biotech 2005; 4: 527-530. | Article
9. Claus TM, Alistair DM, Arnold DJE; Hematological disorders of fish. Vet Clin N Am Exot Anim Pract 2008;11:445-462. | Article | Pubmed
10. Langston AL, Hoare R, Stefansson M, Fitzgerald R, Wergeland H, Mulcahy M; The effect of temperature on non-specific defence parameters of three strains of juvenile Atlantic halibut (Hippoglossus hippoglossus L.). Fish Shellfish Immun 2002; 12: 61-76. | Pubmed
11. Magill AH, Sayer MDJ.; The ejeect of reduced temperature and salinity on the blood physiology of juvenile Atlantic cod. J Fish Biol 2004;64:1193-1205.
12. Svetina A, Matasins Z, Tofant A, Vucemilo M, Fijan N; Haematology and some blood chemical parameters of young carp till the age of three years. Acta Vet Hung 2002; 50:459-467. | Article | Pubmed
13. Hofer R, Stoll M, Romani N, Koch F, Sordyl H; Seasonal changes in blood cells of Artic char (Salvelinus alpinus L.) from a high mountain lake. Aquat Sci 2000; 62:308-319. | Article
14. Cnani N, Timm N, Avidar Y, Ron M, Hulata G; Comparative study of biochemical parameters in response to stress in O. aureus, O. mossambicus and two strains of O. niloticus. Aquaculture Res 2004; 35:1434-1440. | Article
15. Leonardi MO, Klemppu AE: Artificial photoperiod influence on the immune system of juvenile rainbow trout (Oncorhynchus mykiss) in the Southern Hemisphere. Aquaculture 2003; 221: 581-591. | Article
16. Lim C, Klesius PH; Influence of feed deprivation on hematocrit, macrophage chemotaxis, and resistance to Edwardsiella ictaluri challenge of channel catfish. Aquat Anim Health 2003;15:13-20. | Article
17. Rey Vázquez G, Guerrero GA; Characterization of blood cells and hematological parameters in Chichosoma dimerus. Tissue Cell 2007;39:151-160. | Article | Pubmed
18. Mazzola A, Bergamasco A, Calvo S, Caruso G, Chemello R, Colombo F, Giaconne G, Gianguzza P, Guglielmo L, Leopardi M, Raggio S, Sarà G, Signa G, Tomassello A; Sicilian transitional waters: current status and future development. Chem Ecol 2010; 26:267-283. | Article
19. McDonough CJ, Roumillat WA, Wenner CA; Sexual differentiation and gonad development in striped mullet (Mugil cephalus L.) from South Carolina estuaries. Fish Bull 2005;103:601-619. | Article
20. Strickland JDH, Parsons TR (1972) A practical hand-book of seawater analysis. p. 1-310. Fisheries Research Board of Canada, Canada.
21. Svoboda M, Kourh J, Hamackova J, Kalap P, Savina L, Svobodova Z, Vykusova B; Biochemical profile of blood plasma of tench Tinca tinca during pre and post spawning period. Acta Vet Brno
2001;70:259–268. | Article
22. Blaxhall PC, Daisley KW: Routine haematological methods for use with fish blood. Fish Biol (1973) 5:771–781. | Article
23. Shah SL, Altingdag A: Alteration of immunological parameters of tench (Tinca tinca) after acute and chronic exposure to lethal and sublethal treatments with mercury, cadmium and lead. Tur J Vetand Anim Sci 2005;29:163–1618. | Article
24. Larsen HN, Snieszko SF: Comparison of various methods of determination of haemoglobin in trout blood. Prog Fish Cult 1961;23:8-17. | Article
25. Folin O, Wu H: Span diagnostic kits. J Biol Chem 1920; 9:341–367.
26. Annino JS: Clinical chemistry principles and procedures. Little, Brown and Company. 4th edition. Edited by Giese, Roger W. 1976. p. 230–257 Boston. | Book
27. Henry RJ: Clinical chemistry, principles and techniques. Harper and Row, New York. 1968. p. 664–666.
28. Fawcette JK, Scott JE: Practical clinical biochemistry. 4th edition. Arnold Harold Varley, India. 1960. p. 119–122.
29. Goel KA, Mishra BP, Gupta K, Wadhwa S: A comparative haematological study of a few freshwater teleosts. Ind J fish 1984; 3:108–112. | Article
30. Fazio F, Faggio C, Marafioti S, Torre A, Sanfilippo M, Piccione G: Comparative study of haematological profile on Gobius niger in two different habitat sites: Faro Lake and Tyrrhenian Sea. Cah Biol Mar 2012;53:213–219. | Article
31. Bahmami M, Kazemi R, Donskaya P: A comparative study of some haematological features in young reared sturgeons (Acipenser persicus and Huso huso). Fish Physiol Biochem 2001;24:135–140. | Article
32. Imsland AK, Gustausson A, Gunnarsson S, Foss A, Arnason I, Jonsion AF, Smadotth H, Thorarensen H: Effects of reduced salinities on growth, feed conversion efficiency and osmoregulatory status in the spittle wolfish. Fish Biol 2001;59:416–426.
33. Adeyemo OK, Agbede SA, Olaniy AO, Shoaga OA: The haematological response of C. gariepinus to changes in acclimation temperature. Afr J Bioc Res 2003;6:105–108. | Article
34. Zeitoun IH, Ulrey DE, Tack PI: Effects of water salinity and dietary protein levels on total serum protein and haematocrit of rainbow trout (Salmo gairdneri) fingerlings. J Fish Res Bd Canada 1974;31:1133–1134. | Article
35. Jawad LA, Al-Mukhtar HK, Ahmed HK: The relationship between haematocrit and some biological parameters of the Indian shad, Tenualosa ilisha (Family Clupidae). Anim Biodivers Conserv 2004;27:478–483. | Article
36. Rambhaskar B, Srinivasa Rao K: Comparative haematology of ten species of marine fish from Visakhapatnam Coast. Fish Biol 1986;30:59–66. | Article
37. Xiaoyun Z, Mingyun L, Khalid A, Weinmin W: Comparison of haematology and serum biochemistry of cultured and wild Dojo loach Misgurnus anguillicaudatus. Fish Physiol Biochem 2009;35:435–441. | Article
38. Brown JA, Moore WM, Quabius ES: Physiological effects of saline waters on zanders. Fish Biol 2001;59(15):1544–1555. | Article
39. Coz-Rakovac R, Strunjak-Perovic I, Hacmanjek M, Topic PN, Lipez Z, Sostaric B: Blood chemistry and histological properties of wild and cultured sea bass (Dicentrarchus labrax) in the North Adriatic Sea. Vet Res Commun 2005;29:677–687. | Article | Pubmed
41. Kavadias S, Castritsi-Catharios J, Dessypris A: Annual cycles of growth rate, feeding rate, food conversion, plasma glucose and plasma lipids in the population of European sea bass (Dicentrarchus labrax) farmed in floating marine cages. J Appl Ichthyol 2004;19:29–34. | Article

Citation:
Francesco F, Satheeshkumar P, Senthil Kumar D, Caterina F and Giuseppe P: A Comparative study of hematological and blood chemistry of Indian and Italian Grey Mullet (Mugil cephalus Linneaus 1758). HOAJ Biology 2012, 1:5. http://dx.doi.org/10.7243/2050-0874-1-5