Nutritional aspects of Haemorrhagic Syndrome ('Hitra Disease') in farmed Atlantic salmon Salmo salar

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ABSTRACT: Haemorrhagic Syndrome (HS) or 'Hitra Disease' currently constitutes the most important disease problem in Norwegian salmonid farming. Histopathological and chemical findings strongly support the hypothesis of HS being a multifactorial disease complex in which nutritional and/or metabolic disorders are important components. The resemblance to vitamin E/selenium deficiency syndromes in other domesticated animals is discussed.

INTRODUCTION

Since 1977, Haemorrhagic Syndrome (HS) or 'Hitra Disease' has been the most important and widespread disease in farmed Atlantic salmon Salmo salar L. in Norway (Poppe et al. 1985b).

HS has been considered to be a disease complex characterized by high winter mortality, anaemia and haemorrhages (Egidius et al. 1981, 1984, Poppe et al. 1985a). So far, this disease has only been reported to occur in Norway, although an outbreak of a disease tentatively defined as 'Cold water vibriosis', but very similar to HS, has recently been reported from the Shetland Islands (Bruno et al. 1985).

It has been suggested that HS arises because of a basic disorder of non-infectious origin (Fjølstad & Heyeraas 1985, Poppe et al. 1985b). In order to acquire a broader basis for a nutritional theory, histological and chemical investigations were carried out in different groups of salmon. In this paper the results of histological and chemical investigations in farmed (healthy and diseased) and wild salmon are reported.

MATERIALS AND METHODS

Histological examinations. Materials examined originated from 80 farmed Atlantic salmon from 20 different sea-water fish farms along the Norwegian coast. Wild salmon (*n* = 10) were caught in bag-nets off the Trøndelag coast.

Samples were collected from moribund, diseased, and apparently healthy fish at different seasons during the period 1982 to 1985. The majority of the farmed fish were juveniles, 0.5 to 2.0 kg, and had been in seawater cages for 5 to 12 mo. The wild fish were from 3 to 6 kg and had spent 1 yr in the sea. Shortly after the fish had been killed by a light blow to the head, bled and autopsied, samples of brain, gills, pseudobranch, heart, liver, spleen, head, kidney, stomach with pyloric caeca and pancreatic tissue, anterior and posterior gut, skeletal musculature and skin, were taken and fixed in 10% neutral, buffered formalin.

The samples were prepared according to standardized histological techniques and stained with haematoxylin & eosin (HE). Some selected samples were also stained with Martius scarlet blue (MSB), Giemsa, and elastin/van Giessen (EVG). Unstained liver sections were studied by fluorescent microscopy using a Zeiss standard fluorescent microscope (excitation 350 to 410 nm, emission 445 nm) according to Roald et al. (1981).

Chemical analyses. Liver samples were collected from farmed and wild Atlantic salmon during the period 1984–1985. Farmed salmon material (*n* = 666) was taken from fish forwarded to the National Veterinary Institute for health control or disease investigation. Wild salmon (*n* = 54) were caught in bag-nets off the Trøndelag coast in Mid-Norway, in traps in a river in the Oslo area, and on long-lines in the fishing grounds around the Faroe Islands.

Samples were stored at −20°C until all analyses
were performed. The liver samples were analysed for copper (Cu), zinc (Zn), iron (Fe) and selenium (Se). Selected samples (see 'Results') were analysed for mercury (Hg) and cadmium (Cd). All samples were digested in a mixture of nitric and perchloric acid. Cu, Zn, Fe and Cd were determined by flame atomic absorption spectrometry, Hg by the cold vapour technique (Haugen et al. 1985), and Se by a fluorimetric method (Ihnat 1974, Norheim & Nymoen 1981), or a hydride generator method (Norheim & Haugen 1985). Results are expressed as µg g⁻¹ wet weight. Statistical analyses were carried out according to Nissen (1982).

Fig. 1. *Salmo salar*. Macroscopic appearance of farmed salmon with HS. Note pale organs and multiple haemorrhages in the liver.

Fig. 2. *Salmo salar*. Body musculature of salmon with HS. Transverse section, HE, 400x. Oedema is seen between the fibrils and there is central migration of sarcolemmal nuclei.
RESULTS

About 75% of the farmed salmon material was characterized as ‘diseased’. Similar pathological changes were seen in all the diseased fish examined, and no differences were observed between different size-classes or stocks.

Diseased farmed salmon were macroscopically characterized by abdominal reddening and haemorrhages, especially at the base of the fins and vent area. Most internal organs (gills, heart, liver, spleen) were pale and the abdominal and pericardial cavity was filled with moderate to large amounts of serohaemorrhagic fluid. Petechiae on the peritoneum, swim bladder and perivisceral fat tissue were found in most cases, while subcapsular petechiae in the liver were also occasionally present (Fig. 1). The kidney appeared swollen in most cases. The stomach was usually filled with mucoid fluid. A catarrhal, or sometimes haemorrhagic enteritis was common. These results are in accordance with those described in earlier studies on HS (Egidius et al. 1981, 1984, Poppe et al. 1983, 1985b).

The pathological changes described above were not present in healthy fish and wild salmon. However, a moderate hyperaemia in the vent area could be seen in some cases in healthy farmed fish, as well as petechiae on the anterior part of the peritoneum. The liver was pale or mottled in some farmed salmon which were otherwise apparently healthy. In wild salmon fatty infiltration of the liver was occasionally seen.

The most severe histopathological changes in diseased fish were found in the skeletal and heart musculature. In other organs congestion and oedema were frequently found, as well as focal haemorrhages in some cases. Moderate hepatocellular fatty infiltration and focal liver degeneration were typical findings. Congestion and subcapsular (superficial) haemorrhages were also frequently found, although the liver tissue was otherwise without visible pathological changes. No ceroid-like material was observed.

Severe signs of circulatory failure with congestion, peritubular oedema and degeneration could be found in the kidney as well as haemorrhages in the haematopoietic tissue. Focal haemorrhages and clusters of bacteria were frequently found in the spleen. Congestion, oedema, focal haemorrhages and bacteria were frequently seen in the subcutaneous connective tissue. Similar changes together with desquamation of the epithelium were found in the digestive tract.

In the skeletal musculature the most characteristic feature was a mottled appearance together with a jumbled mixture of normal and dystrophic muscle fibrils. The intermuscular spaces were wider than normal and often filled with a proteinaceous fluid. Although the sarcolemmal membrane was intact, the contents showed varying degrees of degeneration (Fig. 2). Most typical was diffuse swelling and loss of striations and homogenization of the muscular tissue. The most severe lesions were located close to the intermyotomal fascial tissue. Central migration of sarcolemmal
nuclei was found in some cross sections. ‘Piling’ of sarcolemmal nuclei could be seen in some cases, probably indicative of regenerative processes. The muscular lesions described above could also be observed in farmed fish otherwise characterized as ‘healthy’, especially in winter.

In fish characterized as obviously diseased, i.e. in later or advanced stages of the disease, the lesions were of the same type as those already described, though more pronounced. Exudation was more evident with discoid and/or granular degeneration (Fig. 3 & 4). Haemorrhages could occasionally be seen near the intermuscular septae. Bacteria could be seen as clusters or single cells in the degenerated muscle fibres and in the exudates, especially in Giemsa stained preparations. Vacuolation of the degenerated musculature was frequently seen.

In the myocardium longitudinal and cross-striations were usually diffuse or absent. In the borderline between firm and spongy musculature, small focal haemorrhages and clusters of bacteria could frequently be seen, especially in advanced stages of the disease. The epicardium was infiltrated with mononuclear cells and was moderately to severely thickened. No pathological changes were observed in the heart or skeletal musculature of wild salmon.

The results of the analyses for Cu, Zn, Se and Fe are summarized in Table 1. Selenium levels were on aver-

Table 1. Salmo salar. Mean concentrations and ranges of copper, zinc, iron and selenium (µg g⁻¹ wet weight) in liver from wild salmon and 3 groups of farmed salmon n: number of samples

<table>
<thead>
<tr>
<th>Group</th>
<th>Copper µg Cu g⁻¹</th>
<th>Zinc µg Zn g⁻¹</th>
<th>Iron µg Fe g⁻¹</th>
<th>Selenium µg Se g⁻¹</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>n</td>
<td>n</td>
<td>n</td>
<td>n</td>
</tr>
<tr>
<td>Wild salmon</td>
<td>37</td>
<td>37</td>
<td>37</td>
<td>54</td>
</tr>
<tr>
<td></td>
<td>184 ± 78 (29–390)</td>
<td>31 ± 5 (19–40)</td>
<td>148 ± 89 (48–440)</td>
<td>14 ± 6.3 (2.4–35)</td>
</tr>
<tr>
<td>Healthy farmed salmon</td>
<td>233</td>
<td>233</td>
<td>196</td>
<td>260</td>
</tr>
<tr>
<td></td>
<td>75 ± 45 (9–230)</td>
<td>33 ± 11 (14–95)</td>
<td>136 ± 89 (24–820)</td>
<td>2.4 ± 0.9 (0.6–6.0)</td>
</tr>
<tr>
<td>Farmed salmon with HS</td>
<td>85</td>
<td>85</td>
<td>80</td>
<td>181</td>
</tr>
<tr>
<td></td>
<td>41 ± 26 (5–140)</td>
<td>26 ± 11 (12–76)</td>
<td>70 ± 45 (2–340)</td>
<td>1.8 ± 0.6 (0.8–4.7)</td>
</tr>
<tr>
<td>Farmed salmon with diseases other than HS</td>
<td>225</td>
<td>225</td>
<td>102</td>
<td>108</td>
</tr>
<tr>
<td></td>
<td>83 ± 74 (5–630)</td>
<td>30 ± 9 (5–65)</td>
<td>118 ± 65 (25–320)</td>
<td>2.2 ± 1.0 (0.6–5.4)</td>
</tr>
</tbody>
</table>

Fig. 4. Salmo salar. Detail from Fig. 3. HE, 400x

Notes: Figures in the same column without a common superscript are statistically different from each other (P < 0.01, t-test)
Fig. 5. *Salmo salar*. Correlation between hepatic levels of copper and selenium in wild (△) \((n = 36)\) and farmed salmon (○) \((n = 320)\). Average of 10 samples of farmed salmon. The 2 straight lines represent the range covered by 90% of samples in the 2 groups, respectively.

Average of 6 to 7 times higher in liver of wild salmon as compared to farmed salmon. Levels of Cu were also significantly higher in wild salmon, while those of Fe and Zn did not differ between farmed salmon as a whole, and wild salmon. Among farmed salmon, the group suffering from HS had slightly lower hepatic levels of all 4 elements than the other groups. There was a strong positive correlation \((r = 0.70)\) between liver levels of Cu and Se in farmed salmon and a fairly good correlation \((r = 0.56)\) in wild salmon. The regression coefficients, however, differed considerably, being 0.31 and 0.06, respectively (Fig. 5). No further correlations were found between the elements. Mercury levels were almost the same in farmed and wild salmon, being on average 0.02 to 0.04 μg g⁻¹ and 0.03 to 0.05 μg g⁻¹, respectively. Cadmium, on the other hand, differed considerably, with levels below the detection limit of 0.1 μg g⁻¹ in farmed salmon and 0.17 to 0.46 μg g⁻¹ in wild salmon.

**DISCUSSION**

The predominant necropsy findings in HS are anaemia, congestion, oedema, serohaemorrhagic fluid in the body cavities and haemorrhages. The histopathological findings in this investigation were characterized by marked congestion of internal organs, exudative haemorrhagic diathesis and muscular dystrophy and degeneration. Although no macroscopic lesions were seen in the musculature, principal lesions in fish with HS were degeneration and exudative diathesis both in skeletal and cardiac musculature. Such findings have not been associated with any of the classical infectious diseases of fish, but they are well recognized in nutritional and/or metabolic diseases.

Exudative and haemorrhagic diathesis, as well as anaemia, are typical findings in HS (Poppe et al. 1983, Waagebø et al. 1985). In addition, increased activity of serum aspartate aminotransferase (ASAT) and in some cases alanine aminotransferase (ALAT) have been recorded in fish with HS (Braekkan pers. comm. 1985).

Similar changes are typical findings in selenium/vitamin E deficiency syndromes in domestic animals (Blood et al. 1983). These diseases have been well documented both pathologically and aetiologically (Obel 1953, Grant 1961, Nañstad 1970, Nordstoga 1983). Typical for these diseases are congestion, oedema, exudative and haemorrhagic diathesis, anaemia, microangiopathy, muscular and myocardial dystrophy and/or degeneration ('mulberry heart disease'), focal liver degeneration ('hepatosis dietetica'), steatitis ('yellow fat') and catarhal enteritis. On the basis of findings in a series of experiments in pigs, Nañstad (1970) concluded that vitamin E deficiency is manifested by a wide variety of organ changes that are related disorders forming different parts of a syndrome. Extensive haemorrhages in the subcutaneous tissue and internal organs of rapidly growing male mink kits with vitamin E/selenium deficiency have been described by Nordstoga (1983). In this case the haemorrhages dominated the picture and camouflaged
the muscular dystrophy which also was present. The actual way in which such disorders manifest themselves is probably dependent upon a wide range of factors, such as diet, the composition and condition of fatty acids and their interaction with other nutrients, growth rate, animal species, stress and other environmental circumstances.

As regards fish, muscular degeneration, anaemia and exudative diathesis have been demonstrated in experimental selenium and vitamin E deficiency in Atlantic salmon (Poston et al. 1976). Muscular degeneration has also been demonstrated in rainbow trout Salmo gairdneri Richardson (Roberts et al. 1979), and in turbot Scophthalmus maximus L., following selenium/vitamin E depletion (Pearse et al. 1984). Recently, Fjølstad & Heycraas (1985) described muscular and myocardial degeneration in Atlantic salmon with HS.

Oxidized fats and oils or easily oxidizable polyunsaturated fatty acids are important triggers in the selenium/vitamin E deficiency syndrome in domestic animals (McMurray & Rice 1982). The feed given to farmed salmon is comprised to a large extent of unsaturated fat of marine origin. Under practical conditions raw materials for feed are often stored for prolonged periods in freezers before use. This may result in considerable rancidity of the fatty fraction of the feed (Aasgaard & Bergsrønning 1985). Oxidized oils have been shown to induce increased erythrocyte fragility and anaemia in rainbow trout (Moccia et al. 1984). Oxidized oils have also induced anaemia in farmed salmon (Rørstad 1985).

The hypothesis that HS is basically a nutritional or metabolic disorder is also supported by the finding in the present study of muscular dystrophy in apparently healthy fish from farms with HS. Bacteria are seldom detected or isolated from such fish, although easily demonstrated in later stages of the disease. The slow-growing, cold-water Vibrio spp. isolated from salmon with HS (Egidius et al. 1981, 1984, Holm 1985, Poppe et al. 1985b) may therefore be secondary invaders in a host already weakened and stressed by nutritional or metabolic disorders in addition to thermal and other physical stress factors (Snieszko 1974). Decreased immune response has been found in rainbow trout in association with vitamin E deficiency (Blazer & Wolke 1984), while depressed antibacterial activity in salmon serum has been demonstrated by feeding rancid oils (Rørstad 1985).

The considerable difference registered between hepatic selenium levels in wild and farmed salmon is remarkable, although not unequivocally indicative of selenium deficiency in farmed fish. Critical and optimal selenium levels in fast-growing farmed salmon have not yet been ascertained. Marine fish, e.g. cape-mallotus villosus, constitutes the greater part of the feed for farmed salmon. This type of raw material is considered to be a good selenium source, and one should therefore not expect the occurrence of selenium deficiency. Requirements for selenium and its availability from different sources under practical fish farming conditions are not known. Nor is it possible to interpret the difference in hepatic selenium levels in HS-diseased salmon, compared to other groups, as an indication of selenium deficiency of causal significance. The lower selenium levels, as well as the lower levels of Cu, Zn and Fe, may very well be a result of reduced feed intake, reduced transport or storage capacity or enhanced clearance. Certain heavy metals such as mercury (Hg) may induce selenium accumulation in marine mammals (Koeman et al. 1975, Kari & Kauranen 1978), a molar Hg:Se ratio close to 1 being often found. Accumulation of these 2 elements thus occurs in parallel, although the greater part of selenium is used in the detoxification of mercury. The interaction between selenium and mercury in fish seems to be more complex and is far from being completely understood (Froslie et al. 1985). There was a fairly strong positive correlation between copper and selenium in farmed salmon in the present investigation. Such a relation between these elements in salmon has also recently been reported by Julshamn et al. (1985). The weaker correlation between Se and Cu in wild salmon and the different regression coefficients make it difficult to explain the higher hepatic selenium levels in wild salmon on the basis of a linear relation to copper. There may well be, however, a non-linear relation between the 2 elements when wild and farmed salmon are considered together.

The levels of other heavy metals found are probably of only minor importance in relation to the selenium levels.

Vitamin E and other antioxidants are generally added to commercial fish feeds in Norway, although no data are available regarding actual vitamin E levels in fish feed which is used.

As regards serum vitamin E levels in fish, data, albeit limited, indicate that levels are higher in wild salmon than in farmed salmon (Poppe et al. 1985a). The significance of this finding, the critical levels of serum vitamin E in salmon and the importance of vitamin E per se in the etiology of HS are, however, difficult to evaluate. It is interesting to note that HS seldom causes any significant losses in rainbow trout raised on the same farms and fed the same diet as salmon which develop the disease (Hästein pers. comm.). This observation may indicate interspecies differences in the utilization of essential nutrients and/or response to nutritional disorders (Moccia et al. 1984).
Selection for rapid growth in farmed salmon has resulted in an increase in the growth rate of about 3 % yr⁻¹ (Gjedrem pers. comm.). This genetic improvement, together with the expanded use of intensive practices in salmon farming in general and higher stocking densities in particular, probably means that the provision of a well-balanced diet is of ever-increasing significance and that requirements for essential nutrients may perhaps have increased.

**CONCLUSION**

As a whole, the pathological changes described in this investigation are indicative of a metabolic or nutritional disorder. It is, however, not possible to pin-point a single factor or factors that induce HS in farmed salmon. Nevertheless, it may be suspected from the present findings that the syndrome arises because of a basic disorder of non-infectious origin, probably of nutritional or metabolic nature, and that the primary task for future research is to clarify the etiology of the syndrome so that improved prophylactic measures can be introduced.

**LITERATURE CITED**


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