Introduction

The liver is the largest gland of the organism with both endocrine and exocrine functions. Endocrine functions of liver include the secretion of plasma proteins as albumin and $\alpha$- and $\beta$-globulins and cholinesterase. Bile excretion in the gastrointestinal (GI) tract constitutes the exocrine function of the liver (Frandson and Spurgeon, 1995). Moreover liver’s functions include clearance of drugs and toxins from the circulation, metabolism of carbohydrate and fat, synthesis of proteins as well as catabolism of immunoglobulin A and insulin, metabolism of several nitrogenous compounds, uptake and conjunction of bilirubin, free fatty acids and bile acids and storage of vitamins and minerals (Divers, 1998). The diversity and complexity of these functions are thus of utmost importance to the overall health of the animal.

Liver’s susceptibility to disease relies on its functions as a clearance organ for many toxins and drugs, and grazing animals like horses are particularly prone to liver disease (West, 1996). Hepatic disease in equines is frequently reversible because of the liver’s wide functional capacity that offers the possibility of regeneration.

The purpose of this review is to update the knowledge on hepatobiliary diseases and in particular on equine hyperlipaemia. Hepatobiliary disease’s aetiology, clinical signs, diagnosis and nutritional management are thus described in the first part of the review the second part being devoted to hyperlipaemia’s lipid metabolism, epidemiology, clinical signs, post-mortem observations and nutritional management. Diagnosis of hepatic disease is usually based on the assessment of the serum activities while hepatic biopsy is considered as the golden standard of diagnosis of hepatic function. Nutritional management is often very useful in management of hepatic diseases: diet should be low in protein (of good biological value) and high in non-structural carbohydrates except for chronic hepatic disease (slightly high protein). Equine hyperlipaemia’s mortality is around 70%. It consists of a disorder of lipid metabolism, characterized by increase in plasma triglycerides and deposition of fat on organs. From a nutritional point of view, hyperlipaemia in horses can be approached by maintaining positive energy balance, fighting dehydration and metabolic acidosis, and by the use of lipotropic factors.
clinical signs, post-mortem observations and nutritional management.

**Hepatobiliary disease**

The hepatocyte constitutes the functional cell of the liver. Along the hepatocyte lines, bile ducts are responsible for the transport of bile products into the duodenum (Frandson and Spurgeon, 1995). Liver diseases most commonly affect both the two structures but yet at different levels concerning its nature (Divers, 1998). The concept ‘liver disease’ includes several pathological conditions that affect liver’s functions and it can either consist of a temporary impaired functioning of the liver and/or it can progress to its failure (Lehrer, 2006). In equines, liver failure is rare. It becomes clinically apparent when more than around 70% of the liver function is lost (West, 1996; Divers, 1998; Durham et al., 2003a). The most common liver diseases in horses that lead to hepatic failure are Thelier’s disease, Tyzzer’s disease, pyrrolizidine alkaloid toxicity, ferrous fumarate toxicity, hepatic lipidosis, suppurative cholangitis, cholelithiasis and chronic active hepatitis (Kahn, 2006).

**Aetiology**

A number of different aetiological causes may be ascribed to the development of horse hepatic diseases. Toxic, infectious, non-infectious inflammatory, metabolic, obstructive and some unknown causes are recognized in the panorama of liver diseases in horses (Divers, 2005). Among toxic causes of hepatic disease, the most important is pyrrolizidine alkaloid hepatotoxicosis. It is induced by the major plant hepatoxins, the pyrrolizidine alkaloid (PA), which is present in Senecio spp., Amsinckia spp., Crotolaria spp., Cynoglossum officinale and others. Pyrrolizidine alkaloids are synthesized in the root of plants and then translocated to all other plant organs (Ober and Hartmann, 1999). Several case studies concerning Senecio spp. (Lessard et al., 1986; Small et al., 1993), Cynoglossum officinale (Knight et al., 1984) and Crotolaria spp. (Arzt and Mount, 1999) hepatotoxicosis in horses were reported in the last quarter of century. Pyrrolizidine alkaloid hepatotoxicosis is characterized by liver necrosis and fibrosis. According to a case study of Knight et al. (1984), the observed clinical signs of PA hepatotoxicosis were weight loss, icterus, photosensitization and hepatic encephalopathy, while Arzt and Mount (1999) observations included inappetence, emaciation, ataxia and icterus. Both levels of copper (Dewes and Lowe, 1985) and iron (Garrett et al., 1984) were found to increase in horse’s liver with Senecio poisoning being the cause in the first case, related to wood and shavings ingestion. In addition to the hepatic damage, the PA has carcinogenic, teratogenic and abortifacient properties, which are the consequences observed long after the ingestion of the plants (Knight, 1995).

Aliske clover (Trifolium hybridum) and Panicum grasses (Panicum coloratum) grazing during wet or humid weather have been observed to induce photosensitivity and hepatitis. Although unclear, the causes of this type of poisoning have been pointed out to be mycotoxins or plant metabolites because of its sporadic and weather-related occurrence (Knight, 1995). Aliske clover poisoning has been reviewed elsewhere (Nation, 1989, 1991) and literature on specific clinical cases dates back to the 20s and 30s (1928–33; Nation, 1989). Briefly, the pathological findings of aliske clover poisoning includes enlargement and discolouration of the liver with increased bile duct proliferation and perilobular fibrosis without inflammation (Nation, 1991). Literature on P. coloratum grasses hepatotoxicosis reveals deposition of crystalline substances in the bile ducts. The exact cause of liver damage in this case is rather controversial. The origin of these crystals is plant saponin derivatives. As a high-dose administration of isolated saponins to induce photosensibilization experimentally is necessary, the controversy of this condition relies on a possible synergistic action with mycotoxins (Cheeke, 1995). Details of hepatic toxicosis because of the consumption of P. coloratum in horses (Cornick et al., 1988) and sheep (Bridges et al., 1987) have been published in the late 80s. Recently Johnson et al. (2006) have reported a clinical case on hepatotoxicosis in 14 horses fed with fall panicum hay (Panicum dichotomiflorum) in USA. High levels of activity of aspartate aminotransferase, sorbitol dehydrogenase, gamma-glutamyl transferase and alkaline phosphatase were observed in this study in addition to necrosis of hepatocytes. Because of the absence of fibrosis was observed in the reported cases, the authors suggested that the withdrawal of hay was sufficient as a measure to allow recovery from acute exposure.

Indospicine (Knight, 1995) hepatotoxicosis constitutes another toxic cause of liver disease. Indospicine is a hepatotoxic amino acid that is present in various species of Indigofera (highly palatable to horses). As an antagonist of arginine, this amino acid inhibits protein synthesis. Diets containing high quantities of arginine, as cottonseed meal and peanut meal are known to protect horses against the effect of
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Indospicine. This type of poisoning leads ultimately to death from liver necrosis and nodular fibrosis and it affects similar animals that might consume the meat of poisoned horses (Knight, 1995).

Dietary iron and copper excesses can also be toxic to the liver: the frequent administration of iron and iron-containing preparations to improve horse's performance constitutes a risk for iron toxicity (Lewis, 1995b). Ferrous fumarate toxicosis was reported in the literature (Mullaney and Brown, 1988) to be one of the causes of hepatic failure (Kahn, 2006).

Oral administration of this compound to new-born Shetland foals provoked death by acute hepatic failure, which lesions were compared with the toxic hepatopathy syndrome reported in USA (Divers et al., 1983). These two previous studies reported a toxic hepatic failure in foals aged 2–5 days because of oral administration of (i) a product containing a microorganism (Divers et al., 1983) and (ii) a paste containing both Aspergillus sp and an iron compound (Acland et al., 1984). From their very first day of life, foals are particularly susceptible to iron toxicosis (toxic dose is 25 times greater in adult horses). Prior to death, animals to which iron was administered together with an inoculum, presented depression, diarrhoea, icterus, dehydration and coma. Association of excess in iron with deficiencies of selenium and/or vitamin E may boost this foals' susceptibility (Mullaney and Brown, 1988; Lewis, 1995b). Hepatic alterations because of excess of iron include liver deposition of iron and liver degeneration (Lewis, 1995b). According to Lewis (1995b), horses are quite resistant to copper excesses as, for some extent, they adapt their absorption of copper to ingestion levels. Lewis (1995b) reports a level of 2800 ppm of copper for a period of 2 months to provoke liver damage and 6 months to lead to death.

Finally mycotoxins can also be one of the toxic causes affecting liver functioning. Naturally occurring fungi in animal feeds produce mycotoxins in favourable environment conditions allowing its growth. Some of these mycotoxins can affect horse's liver (Pier et al., 1980). Aspergillus flavus in particular is a producer of some aflatoxins, the presence of which was reported in mouldy corn fed to horses prior to death. Aflatoxicosis in horses has been related with liver necrosis in this case report (Vesonder et al., 1991). The same observation (liver necrosis) was patent in a second case study in which one horse was fed with mouldy hay (McGavin and Knake, 1977). Bile duct hyperplasia and fibrous liver was reported in a third study concerning aflatoxicosis in horses (Angsubhakorn et al., 1981). Experimentally induced aflatoxicosis with aflatoxin B1 in weanling ponies (Bortell et al., 1983) has revealed hepatic necrosis and bile duct hyperplasia.

Infectious causes of hepatic disease include cholangiohepatitis and Tizzer's disease. Parasitism of horses by Fasciola hepatica or Parascaris equorum can also affect overall hepatic health. Cholangiohepatitis arises from a liver infection of Salmonella sp, E. coli, Pseudomonas sp or Actinobacillus equuli leading to inflammation of the bile ducts and adjacent liver. Other bacteria, such as Clostridium sp., Pasteurella sp., and Streptococcus sp. are less frequently recovered from horses suffering from cholangiohepatitis. On the other hand, Tizzer's disease is because of an infection of the liver by Clostridium piliforme. It is usually an infection that starts in the low intestinal tract and spreads via blood and lymph circulation (Kahn, 2006).

The most important parasites affecting the liver are P. equorum and F. hepatica. Larvae of strongyles also parasite the host's liver, while adult forms affect the large intestine. Strongyles parasitism [both large: Strongylus spp., Triodontophorus serratus, and small: Cyathostomum spp., Cylicostephanus spp., Poteriotostomum imperidentum (Mitfodlozze and Hutchinson, 1990)] is the most frequent parasitism observed in several coproscopical examination studies concerning horses (Mirck, 1978; Epe et al., 1993, 2004; Daugschies and Epe, 1995). A questionnaire study in Germany (Daugschies and Epe, 1995) including 3500 veterinarians reported a prevalence of strongyles in 42.3% of their patients. Coproscopical examination studies from Germany and the Netherlands have reported prevalances of 55.5% \(n = 9192\) (Epe et al., 1993), 37.4% \(n = 4399\) (Epe et al., 2004) and 57.3% \(n = 3791\) (Mirck, 1978).

The prevalence of P. equorum reported in the literature was of 12.0% from post-mortem intestinal collection \(n = 85\) (Kornaš et al., 2006). Results from coproscopical examination were much lower: 4.0% (Epe et al., 1993), 0.9% (Epe et al., 2004) and 6.1% (Mirck, 1978). A previous study on P. equorum administration to ponies has described both clinical evolution and lifecycle of these parasites (Srihakim and Swerczek, 1978). Parascaris equorum's lifecycle is characterized by transport of larvae to the liver by the bloodstream, migration to the lungs and development of larvae in the intestine because of caughing and swallowing of larvae by the animals. Besides caughing, clinical signs are very indistinct and include anorexia, rough coat and loss of weight. Blood parameters reveal mild anaemia, marked
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eosinophilia, and leukopenia. Post-mortem findings included haemorrhagic and necrosed liver (Srihakim and Swerczek, 1978). The prevalences of *F. hepatica* from coproscopical examination were of 0.2% (Epe et al., 1993), 0.04% (Epe et al., 2004) and 0.6% (Mirk, 1978). Several published studies of induced infection with *F. hepatica* have been reported (Nansen et al., 1975; Greck et al., 1977; Alves et al., 1988; Soulé et al., 1989). *Fasciola hepatica* eggs were hardly ever found in faeces in several studies (Greck et al., 1977; Alves et al., 1988; Soulé et al., 1989) revealing the inadequacy of this diagnosis tool in detecting infection of the liver by *F. hepatica* (liver fluke). According to Nansen et al. (1975), horses present a high resistance to liver fluke. Plasma enzymatic activity of glutamate dehydrogenase and gamma-glutamyl transferase was found to significantly increase 3 to 5 months after experimentally induced infection (Soulé et al., 1989).

Inflammatory causes of liver disease include chronic active hepatitis and liver neoplasia. Although chronic active hepatitis is often related to cholangiohepatitis, it can also arise from immune-mediated or toxic processes. Primary hepatic neoplasia is not commonly observed in horses. The bile ducts, hepatocytes or metastasis are the onset structures of carcinomas of the liver (Kahn, 2006).

The most common biliary duct obstructive cause of hepatic disease is choledolithiasis (Gerros et al., 1993). Cholelithiasis can occur because of ascariasis, biliary stasis, ascending biliary infection or variation of bile composition. According to the mentioned case report, the clinical signs observed were abdominal pain, icterus and pyrexia. Other obstructive causes include colon displacements and associated right hepatic lobe compression, papillary stricture, neoplasia, hepatic torsion and thrombosis of the portal vein. Recent case reports concerning hepatic neoplasia have been published (Patton et al., 2006; Schnabel et al., 2006; Loynachan et al., 2007). In Patton et al. (2006) case report, clinical signs of neoplasia in a thoroughbred mare included depression, head pressing and blindness. Further investigation observations following euthanasia included increased sorbitol dehydrogenase, gamma-glutamyl transpeptidase and alkaline phosphatase activities, increased total serum bilirubin (from which 95% was unconjugated) and serum activity of creatine kinase, increased plasma ammonia level, hyperglobulinemia and hyperglycemia. The liver presented neoplastic cell masses and a large thrombus of these cells was found in the portal vein.

Hepatic lipidosis and hyperammonemia are recognized metabolic aetiopathological causes of liver disease. Hepatic lipidosis will be described in detail further on this work. An impaired development of urease-producing bacteria in the intestine of the horse can be the cause of hyperammonemia, which is almost always associated with enteric disease, diarrhoea or colic (Kahn, 2006).

Thelier's disease and neonatal isoerythrolysis are some of not fully understood causes of hepatic disease. Thelier's disease or idiopathic acute hepatic disease is the most common cause of acute hepatitis.

Clinical signs

Some typical clinical signs associated with specific liver diseases have been described previously with the aetiopathological factors. Hepatic failure can lead to different clinical signs. The most common signs are weight loss, neurological troubles linked to ammonia excesses, icterus, hypoglycaemia, haemoglobinuria, while less frequently observed signs are photosensitivity, bleeding abnormalities, oedemas and ascites.

Weight loss can be one of the most common signs of liver failure (Cornick et al., 1988; McGorum et al., 1999; Kahn, 2006) in particular resulting from PA toxicosis. The decreased appetite and/or decreased metabolism at a hepatic level are factors that will direct liver disease clinical signs in this direction. In cases of neoplasia of the liver, both infection and circulating tumour-induced cytokines are the onset causes of weight loss.

Hepatoencephalopathy, neurological troubles associated with liver disease are also very often described (McGorum et al., 1999; Durham et al., 2003b) and it is related to the excess of ammonia in the blood stream. It consists mainly on depression, head pressing, apparent blindness, circling and other symptoms. The excess of ammonia in the blood stream is because of an insufficient conversion by the liver of ammonia in urea, arriving from the colon. This hyperammonemia will interfere with normal neurotransmission; will structurally change the blood brain barrier and will interfere with biochemical or electrophysiological pathways in the brain. Aromatic amino acids metabolism in the liver will also be affected leading to its increased concentration in the blood stream. As false neurotransmitters, aromatic amino acids in the blood will also have neurological consequences. Gut-synthesized gamma-aminobutyric acid’s increased concentration in the blood stream, associated with augmented aromatic amino acids and bile acids concentration, will potentially have an
Oedema, ascites and diarrhoea are not common.

Diagnosis

Diagnosis of hepatobiliary disease and determination of hepatic function can be achieved basically with three types of diagnosis tools: serum biochemical analysis, liver biopsy and ultrasound imaging. Despite all work that has been carried out concerning hepatic diseases in horses, only recently published works have focused seriously on the diagnostic value of the above-mentioned tools. These very interesting works of Durham et al. (2003a,b) have focused on ultrasound imaging, clinical and clinicopathological data and its adequacy to discriminate affected from non-affected horses.

Serum biochemical diagnosis is based on the assessment of the serum activities of gamma-glutamyl transferase and sorbitol dehydrogenase, bilirubin and bile acids (Divers, 1998). Although blood ammonia is commonly high when loss of hepatic function occurs, it is not so for liver failure so its use as a diagnosis tool is reserved. In plant poisoning, particularly PA hepatotoxicosis, diagnosis is often difficult because of late suggestion of poisoning. Often the cause of liver damage remains unknown besides the diagnosis tests mentioned and the bromsulphophthalain clearance test. In these cases, biopsy or necropsy is frequently the only reliable diagnostic tool (Knight, 1995). Durham et al. (2003c) have developed a scoring system for liver biopsies that has taken in consideration five pathological processes. Hepatic biopsy is useful for diagnosis and to plan for a specific therapy, but generally blood analysis is the most common diagnosis tool to detect liver disease. In several studies, biopsy is considered as the ‘gold standard’ technique for hepatic disease diagnosis (Spycher et al., 2001; Durham et al., 2003c).

The results obtained by Durham et al. (2003b) have highlighted that neither single nor grouped results of clinical data, serum biochemical analysis and ultrasonography were found to fully discriminate affected from non-affected horses when compared to liver biopsy. Hepatic encephalopathy, gamma-glutamyl transferase, hyperglobulinaemia, hypoalbuminaemia, high alkaline phosphatase activity, increased total bile acids and increased total bilirubin were the most valuable single positive results. Ultrasonography on the other hand has proven to be useful to detect particular abnormalities or add some
insights on the diagnosis together with the other mentioned tools.

Moreover, techniques using radiopharmaceuticals have also been already applied to analyse hepatic function: scintigraphy (Theodorakis et al., 1982) and clearance tests (Morandi et al., 2005). The necessity of both radiopharmaceuticals and adequate imaging machinery are the disadvantages that restrain scintigraphy’s daily life application. Despite the practical issues related to authorization of institutions to use radiopharmaceuticals, Morandi et al. (2005) studied the $^{99m}$Tc-mebrofenin clearance from the blood in seven horses, reporting the simplicity and reliability of the method. The results of this study have shown that with a clearance plasma-based method it is possible to access hepatic function.

**Nutritional management**

Among therapies, nutritional management is very often useful. Literature regarding nutritional requirements in horses affected with liver disease is scarce and focus mainly on hyperlipaemia (which will be described further on this review). Lewis (1995c) suggested five main points to have in consideration when feeding the liver diseased horse: (i) meet energy requirements; (ii) meet without exceeding dietary protein needs (low protein in case of hyperammonaemia); (iii) feed high ratio of branched-chain amino acids (BCAA)/aromatic amino acids; (iv) feed low fat and salt and (v) feed high-starch diets.

Diet must contain low levels of proteins (Hussein et al., 2004) of good biological value. High protein forages (e.g. alfalfa and fresh rich spring grass) are advised not to be fed to liver-diseased horses. On the contrary, first cut meadow hays and fall pasture grazing (avoiding long exposure to sun) can be considered optimal.

Increased number of daily meals could be also useful. Lewis (1995c) suggests feeding 0.5–0.6 kg/100 kg body weight BW/day divided in three to six daily meals. Ott (2001) suggests that higher levels of dietary protein induce higher urea-N serum concentration while blood ammonia remains constant. This observation was confirmed by Hussein et al. (2004), who added that after being fed, mares had statistically increasing serum ammonia concentrations. In the previous study, Hussein et al. (2004) have fed the mares twice daily. It is thus wise to consider that the higher number of meals can decrease the intensity of fermentation in the hindgut and consequently decrease ammonia peaks in the serum, which is a factor contributing for hyperammonaemia when considering liver-diseased horses.

As blood concentrations of BCAA may be decreased, and the aromatic amino acids (AAA) and ammonia are usually augmented in acute hepatobiliary disease, BCAA/tryptophan ratio in the diet must thus be kept high. Corn and molasses are often used. Table 1 summarizes some of BCAA/AAA ratio and protein content in several feedstuffs fed to horses. Horses with chronic hepatic disease, conversely, must be fed slightly high-protein diets. Non-structural carbohydrates are required in large amounts to meet the energy requirements and avoid mobilization of body glycogen, fat and protein.

As mentioned previously (aetiology), copper and iron excesses can affect liver function: liver damage was observed for horses consuming 2 800 ppm of copper (Lewis, 1995b) for 2 months and led to death in 6 months. Administration of supplements like iron-containing haematinics to horses to enhance performance must be precautious to avoid iron toxicity (Lewis, 1995b; Casteel, 2001). Pearson and Andreasen (2001) have studied the effect of iron oral administration on hepatic biopsies and serum analysis. Ferrous sulphate (50 mg/kg BW/day; 20% elemental iron) was orally administered to ponies during 28 weeks. By week 8, hepatic and serum iron concentration had increased when compared with control and baseline concentrations but these values have decreased by the twenty-eighth week. Hepatic biopsies did not present any lesions at hepatocyte’s level. This study suggests that a primary hepatopathy might be the cause for a secondary haemoriderin accumulation. The susceptibility to iron toxicosis is influenced also by selenium and vitamin E deficiencies. According to Lewis (1995b), despite the infrequent clinical selenium or vitamin E deficiencies,

<table>
<thead>
<tr>
<th>Feedstuffs</th>
<th>BCAA/AAA</th>
<th>Protein (% dry matter)</th>
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<tr>
<td>Soybean</td>
<td>1.8</td>
<td>50</td>
</tr>
<tr>
<td>Linseed</td>
<td>1.9–2.1</td>
<td>35–50</td>
</tr>
<tr>
<td>Rice</td>
<td>1.35</td>
<td>7–9</td>
</tr>
<tr>
<td>Wheat</td>
<td>1.5</td>
<td>11–14</td>
</tr>
<tr>
<td>Oats</td>
<td>1.65</td>
<td>10–13</td>
</tr>
<tr>
<td>Barley</td>
<td>1.65</td>
<td>13</td>
</tr>
<tr>
<td>Rye</td>
<td>1.85</td>
<td>14</td>
</tr>
<tr>
<td>Corn</td>
<td>2.15</td>
<td>8–10</td>
</tr>
<tr>
<td>Sorghum</td>
<td>2.3</td>
<td>12–13</td>
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</table>

BCAA, branched-chain amino acids; AAA, aromatic amino acids.
Horse diets are usually below the concentrations required (selenium requirement is of 0.1 ppm of diet dry matter against some cases of 0.05 ppm supply; vitamin E minimum requirement for maintenance being of 50 IU). In pigs, vitamin E bioavailability is positively related with fat content in the diet (Moreira and Mahan, 2002) but that is not the case regarding humans (Roodenburg et al., 2000) and horses (Kienzle et al., 2003).

Finally, fluid therapy is often very useful to improve perfusion, restore electrolytes and glucose levels and help toxin removal. Dextrose is used in hypoglycaemic horses (Divers, 1998). A successful treatment of a mare with encephalopathy and concurrent hypocalcaemia by continuous infusion of Ringer’s solution, calcium gluconate, dextrose, B-complex vitamins, sodium ampicillin and flunixin meglumine was previously reported by Scarratt et al. (1991).

Hyperlipaemia

Horses’ liver function is also severely affected as a result of equine hyperlipaemia, which is characterized, by increased plasma triglycerides and fatty infiltration of body organs. The incidence of this metabolic disorder has been reported in population surveys, varying between 3% and 5% while clinical studies of animals presented to veterinary hospitals have described incidences between 11% and 18% (Watson, 1998). Although of minor economical importance in the equine industry (Jeffcott and Field, 1985), mortality of affected animals is high, like in other species, ranging between 60% and 80% (Watson, 1998).

Lipid metabolism

Equine hyperlipaemia is a disorder of lipid metabolism, characterized by increase in plasma triglycerides. Because of hormone sensitive lipase activation in hyperlipaemic animals, an increase of plasma non-esterified fatty acids (NEFA) occurs, exceeding the oxidative, gluconeogenic and ketogenic pathways (Watson, 1998). Unlike other species, horses have a limited capacity for ketogenesis and do not develop pronounced ketonaemia associated with hyperlipaemia (Naylor et al., 1980). The synthesis of triglycerides in the liver is thus increased in the form of very low density lipoproteins (VLDL), which are secreted in the circulation increasing plasma triglycerides (Watson, 1998). Fatty infiltration of different organs may follow. Liver’s fatty infiltration is because of an increased VLDL synthesis that does not match the rate of triglycerides synthesis. Differences of lipid metabolism between hyperlipaemic and normal ponies have been described by Watson (1998) and are schematically shown in Fig. 1. Hyperlipaemic ponies have shown a much higher transportation of plasma NEFA to the liver (148.2 ± 8.4 mmol/h) when comparing to normal ponies (8.2 ± 5.2 mmol/h) as well as a higher VLDL-triglyceride (TG) synthesis (190 ± 3.9 mmol/h against 6.6±4.4 mmol/h). On the other hand, the VLDL-TG fractional catabolic rate is similar in both cases.

Epidemiology

Risk factors for hyperlipaemia can either precipitate or predispose to hyperlipaemia. Table 2 (source: Watson, 1998) summarizes the risk factors of hyperlipaemia which are followed by a general description. Although not being a condition observed strictly in ponies, a marked insensibility to insulin of these animals comparing to larger horses indicates a higher promptness for triglycerides mobilization and consequently hyperlipaemia (Jeffcott and Field, 1985). The presence of

Fig. 1 Lipid metabolism differences between healthy and hyperlipaemic ponies (Watson 1998).
Hyperlipaemia was observed both in ponies, donkeys and horses in a serum chemical profile study enrolling both clinically normal and sick, either with or without hyperlipaemia equids (Naylor et al., 1980). Although sex was thought to be involved as the predisposing factors, it seems that pregnancy and lactation are the factors that place females in a predisposing circumstances for hyperlipaemia. Moreover, a study with donkeys has evidenced the highest risk of females and body condition regardless of pregnancy status (Reid and Mohammed, 1996). Stated percentages of obese ponies that have developed hyperlipaemia vary between 60% and 90% (Watson, 1998) which indicates the tendency of obesity in ponies to be related with hyperlipaemia. Anorexia is very frequently observed immediately before the clinical signs onset. It can be because of underlying disease, insufficient supply of pasture.Supplementary feeding or even diet/environment modifications (Watson, 1998).

Stress is one of the major risk factors pointed out in literature (Jeffcott and Field, 1985; Watson, 1998). Transportation, change of environment, nutritional stress and pregnancy, lactation and disease-associated stress are among the stress situations that can represent a risk regarding hyperlipaemia. Its association with animal-focused predisposing factors have tendency to enhance the probability for hyperlipaemia occurrence.

Gastrointestinal tract diseases have been observed as a primary disease leading to hyperlipaemia. Among which, case records stated in literature (Moore et al., 1994) have observed associated septicaemia, colitis, parasitism, oesophageal obstruction, faecalith and gastric impaction. Other references include additional systemic diseases as pneumonia (Lewis, 1995a) and pancreatic pathology, renal insufficiency and laminitis (Watson et al., 1992). Although age is considered an important factor for development of hyperlipaemia, which trend tends to be higher for older animals, occurrence in foals from 2 days to 6 months has been reported (Moore et al., 1994).

### Clinical signs

Reported clinical signs of hyperlipaemia include dullness and depression, loss of appetite, production of scant faeces, obesity, weight loss, production of diarrhoeic faeces, increased pulse rate, pyrexia, increased respiratory rate, ventral oedema, jaundiced membranes, petechial haemorrhages, lipaemic serum (West, 1996) and failure to drink, progressive drowsiness, muscle fasciculations, colic, incoordination and coma (Lewis, 1995a). Plasma TG increase in horses with hyperlipaemia reaching values as high as 1500 mg/dl as well as plasma alkaline phosphatase activity (Lewis, 1995a). Often hyperlipaemia is associated with a primary disease. Gastrointestinal diseases are commonly referred to in this context (Moore et al., 1994; Watson, 1998) and recovery is thus often closely related to the success of treatment of the primary disease (Lewis, 1995a).

### Post-mortem observations

Post-mortem observations of affected animals comprise deposition of fat in the liver, kidneys, skeletal muscle, adrenal cortex and myocardium, vascular thrombosis in the lungs, kidneys and brain, renal nephrosis and proliferative glomerulonephritis, atrophy of the exocrine pancreas, lymphoid follicle depletion of the spleen, lymph nodes and mucosal ulceration of the GI tract as well as excessive fluidity of the GI contents (Jeffcott and Field, 1985).

### Nutritional management

Hyperlipaemia in horses, from a nutritional point of view can be approached by maintaining positive energy balance with low-fat diets, fighting dehydration and metabolic acidosis, and by the use of lipotropic factors. Fluid therapy includes the use of lactate or pyruvate; bicarbonate can be used with caution (Watson, 1998). High energy feedstuffs can be used for hyperlipaemic horses; sometimes nasogastric tube feeding is required. Dextrose is often used at quite high dosages (100 g/day for miniature horses) with galactose being used on alternate days to avoid/recover from metabolic acidosis. Nevertheless, it is important to monitor for insulin sensitivity and hyperglycaemia situations (Watson, 1998). Schmidt et al. (2001) have studied the effect of dietary fat on plasma metabolites in healthy ponies and they have concluded that fat feeding increases clearance of TG from the blood stream but it deteriorates glucose tolerance. To our knowledge, further work

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**Table 2 Risk factors for hyperlipaemia**

<table>
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<tr>
<th>Predispose</th>
<th>Precipitate</th>
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<tr>
<td>Breed</td>
<td>Anorexia</td>
</tr>
<tr>
<td>Sex (female)</td>
<td>Malnutrition</td>
</tr>
<tr>
<td>Obesity</td>
<td>Stress</td>
</tr>
<tr>
<td>Pregnancy/lactation</td>
<td>Disease</td>
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on this topic including hyperlipaemic animals has not been performed so far.

Enteral low fat and partially hydrolysed feeding preparation for humans (Vital HN, Ross laboratories, Columbus, Ohio, USA) has been successfully used in a miniature horse (Golenz et al., 1992). Fasting ponies have also been fed with a fat-free preparation consisting of glucose, casein, minerals and vitamins, which has also shown to successfully decrease the level of plasma triacylglycerol, although being much lower in fasting animals than in sick horses (Hallebeek and Beynen, 2001). Though supplementation in sick horses has not been studied, the authors state that, in association with conventional therapy, the potential of the diet to aid hyperlipaemia treatment is real (Hallebeek and Beynen, 2001). Complete feeds and increased daily meal numbers are also useful. Both Golenz et al. (1992) and Moore et al. (1994) propose an energetic level of approximately 1.2 to 2.0 maintenance level corresponding this value to a ‘stress factor’. Prevention of hyperlipaemia relies on ensuring an adequate intake of good quality feeds mostly in situations at risk and in predisposed animals.

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