Hypertrophic osteopathy associated with systemic granulomatous disease in a horse

Osteopatia hipertrófica associada à doença granulomatosa sistêmica em um equino

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Abstract

A 4-year-old Criollo stallion was presented at the equine clinic of veterinary hospital of the Federal University of Santa Maria, RS, with a 30-day history of progressive weight loss, anemia and swelling of the forelimbs and face. Physical examination revealed that the swelling was firm and had a bone-like consistency, also radiographs showed extensive periosteal proliferation on the forelimb long bones that suggested hypertrophic osteopathy (Marie’s disease). Physical examinations identified no respiratory findings. However, during ultrasound examination, superficial lung disease was identified. The animal was treated with antibiotics and nonsteroidal anti-inflammatory drugs for 12 days. Due to a complete lack of response to this treatment, the horse was euthanized. At necropsy several granulomatous lesions were identified in the thorax, abdomen and testicular tunics. Bony proliferation was evident on many bones of the appendicular skeleton and face. Based on these findings the diagnosis of hypertrophic osteopathy associated with sarcoidosis was established. It is important to perform a thorough clinical examination and include hypertrophic osteopathy in the differential diagnosis of diseases that are accompanied by swelling of the face and limbs as edema from various causes, fibrous osteodystrophy, for example.

Key words: Pulmonary. Periosteal. Sarcoidosis. Pathology.

Resumo

Um garanhão Crioulo, 4 anos de idade, foi atendido na clínica de equinos do hospital veterinário da Universidade Federal de Santa Maria, RS, com histórico de emagrecimento progressivo há 30 dias, anemia e aumento de volume dos membros torácicos e da face. O exame físico revelou que o
aumento de volume era firme e apresentava consistência óssea. Radiografias também mostraram extensa proliferação periosteal nos ossos longos dos membros anteriores, o que sugeriu a osteopatia hipertrófica (doença de Marie). Os exames físicos não identificaram alteração respiratória. No entanto, durante o exame ultrassonográfico, foi identificada doença pulmonar superficial. O animal foi tratado com antibióticos e anti-inflamatórios não esteróides durante 12 dias. Devido a uma completa falta de resposta ao tratamento, o cavalo foi sacrificado. Durante a necropsia, várias lesões granulomatosas foram identificadas no tórax, abdômen e túnica testicular. Proliferação óssea era evidente em vários ossos do esqueleto apendicular e da face. Com base nestes resultados o diagnóstico de osteopatia hipertrófica associada à sarcoidose foi estabelecido. É importante realizar um exame clínico completo e incluir a osteopatia hipertrófica no diagnóstico diferencial de enfermidades que vêm acompanhadas de aumento de volume da face e dos membros como edema por diversas causas e osteodistrofia fibrosa, por exemplo.


**Introduction**

Hypertrophic osteopathy (HO) is a symmetric bilateral progressive proliferation of subperiosteal bone substance and fibrous connective tissue of the appendicular and axial skeleton, and facial bones. The clinical suspicion of HO can be confirmed by radiographic identification of periosteal new bone formation involving the diaphyses and metaphyses of the appendicular and facial bones (KATZ, 2015). It is a relatively rare disease in horses. It occurs more frequently in man and dogs, usually associated with lung tumors (KAWCAK; BAXTER, 2011).

In horses, this syndrome has been associated with various diseases, especially thoracic injuries, such as lung abscesses, lung neoplasms, granulomatous pneumonia, rib fractures with pleural adhesions and pericarditis (CHAFFIN et al., 1990; MAIR et al., 1996; MAIR; TUCKER, 2004; BAYLESS et al., 2014). HO has also been described associated to gastric squamous cell carcinoma (SCHLEINING; VOSS, 2004).

Systemic granulomatous disease (SGD) is a rare condition that affects horses, characterized by non-caseous granulomatous inflammation of skin and/or multiple organs. Clinical signs vary depending on the organic system involved, but progressive weight loss, inappetence and fever are clinical findings frequently observed in horses with SGD (REES, 2004).

HO secondary to SGD was previously described in one horse in a retrospective study of 24 horses with HO. However, the horse recovered after prolonged treatment with corticosteroids, and there is no further information regarding the distribution of granulomatous lesions (MAIR et al., 1996). The present study describes a case of HO associated with systemic granulomatous disease in a horse.

**Case Report**

A 4-year-old Criollo stallion was referred with a one-month history of progressive weight loss accompanied by edema of the forelimbs and face (Figures 1A and 1B). Clinical examination, except for the high body temperature (39.5 ° C), showed no changes (heart rate: 42; respiratory rate: 12; normal colored mucous). Palpation of the lower limbs showed hard swelling, pointing to the radiological evaluation of the limbs. Radiographs taken from both third metacarpal bones showed severe periosteal proliferation suggesting HO. Laboratory tests showed anemia (Ht 27% / reference range: 32-53%), leukocytosis (16,500 μL⁻¹ / reference range: 5400-14,500 μL⁻¹) with neutrophilia (73%), hyperproteínaemia (8.8 g dL⁻¹ / reference range: 5.8-8.7 g dL⁻¹), hyperfibrinogenemia (800mg dL⁻¹ / reference range: 100-400mg dL⁻¹) and hypoalbuminemia (1.3 g dL⁻¹ / reference range: 2.6-3.7 g dL⁻¹). During thoracic
ultrasonography bilateral presence of “comet tail” artifacts (suggestive of pleural lesion), most evident in the ventral portion of the lungs, was identified. Analysis of peritoneal fluid showed an increase in nucleated cells (18,800 μL⁻¹), predominantly neutrophils (96%) and 3.4 g dL⁻¹ total protein. Treatment was started with sulfamethoxazole and trimethoprim (20mg kg⁻¹ b.i.d.), ketoprofen (2.2mg kg⁻¹ b.i.d.) and omeprazole (4mg kg⁻¹ s.i.d.) for six days. No clinical improvement occurred and the association trimethoprim sulfa was replaced by ampicillin sodium (10mg kg⁻¹ t.i.d.) for six more days. On the 12th day, laboratory tests showed progressive anemia (Ht 21%), hyperfibrinogenemia (800mg dL⁻¹) and hypoalbuminemia (1.18g dL⁻¹). Due to a complete lack of response to treatment and a further deteriorating condition, euthanasia was recommended. Euthanasia was performed by licocaine intrathecal injection (20 ml), with the horse under general anesthesia (ketamine 2.2mg kg⁻¹ and xylazine 1.1mg kg⁻¹), according to the rules of the Federal Council of Veterinary Medicine.

Figure 1. A: Forelimbs swelling. B: Evident face swelling. C: Numerous granulomatous nodules in the lung pleura, with predominantly cranial-ventral distribution. D: Numerous granulomatous nodules in smallcolon (serous membrane).

At necropsy, numerous small scabs on the skin of the limbs, ventral abdomen and scrotum were observed. Multiple white, firm, coalescing nodules with 0.5 to 1 cm in diameter were distributed on the pleura and in the lung parenchyma, predominantly in the cranio-ventral portions, compressing the underlying lung parenchyma (Figure 1C). Similar nodules were observed in
the epicardium, myocardium. Multiple nodules were also observed in the mesenteric, hilar and tracheobronchial lymph nodes and on mesenteric, gastric, intestinal serous membrane (Figure 1D), spleen, diaphragma, liver and bladder wall, and infiltrating the testicular tunics. Fragments of all these organs, brain and bones were fixed in formalin, processed and stained with hematoxylin and eosin, Ziehl-Neelsen, Grocott, Giemsa, Masson’s trichrome and by reaction with periodic acid-Schiff (PAS), by methods of routine. Nodules were cultured both aerobically and anaerobically on 5% sheep blood Agar. Fragments nodules were also cultured on Lowenstein-Jensen and Stonebrink Leslie solid culture medium, subsequently incubated at 37ºC for up to 60 days (WAYNE; KUBICA, 1986).

After maceration of bones, it was observed periosteal new bone formation in the tarsal bones, metatarsus, tibia, carpus, metacarpus, radius and ulna, phalanges of pelvic and thoracic limbs, maxillary bone, mandible and nasal bones (Figures 2A and 2B). There were no articular changes.

**Figure 2.** Macro and microscopic lesions of hypertrophic osteopathy. A: Osteoproliferative lesions in maxilar bone. B: Osteoproliferative lesions in a third metacarpal bone. The cortex throughout the length of the bone is covered by an irregular layer of fragile and porous bone spiculae. C: Non-caseating granuloma focally extensive on the lung. The center contains mainly Langhans giant cells (arrow) and numerous epithelioid macrophages, surrounded by lymphocytes and plasma cells. (Hematoxylin and eosin, 10x objective). D: Sub macroscopic image of a transversal section of radius. There is a continuous limit (arrow) between the cortical bone and the newly formed bone. (Hematoxylin and eosin, 4x objective).
Histological evaluation of the lungs showed numerous coalescing nonencapsulated foci of nodular non-caseous granulomatous inflammation. Foci were composed of numerous epithelioid macrophages and multinucleated giant cells, mostly Langhans type. Surrounding these areas plasma cells and lymphocytes (Figure 2C) were observed. In areas adjacent to the granulomatous inflammation a marked proliferation of fibrous connective tissue made the pleura markedly thicker. Similar granulomas to those described in the lungs were also seen in the epicardium and myocardium, mesentery, mesenteric, tracheobronchial and hilar lymph nodes, liver, gastric serosa, small intestine, small colon, bladder and testicular tunics. There were no granulomas on the mucosal sections. Aggregates of epithelioid macrophages and giant cells were occasionally observed in the bone marrow. On the skin sections, variably circumscribed foci of lymphocytes, plasma cells, epithelioid macrophages and occasional giant cells were predominantly located in the superficial dermis. Histologically, the skeletal lesions consisted of periosteal bone trabeculae of irregular size and thickness that were arranged perpendicularly to the original bone cortex (Figure 2D). Masson’s trichrome staining showed marked difference between cortical bone and new bone formation, as evidenced by the blue color of woven bone in recently proliferated bone tissue.

Histochemical evaluation (Ziehl-Neelsen, Grocott, Giemsa, Masson’s trichrome and periodic acid-Schiff) of affected tissues did not detect the presence of infectious agents. There was no bacterial growth in aerobic and anaerobic cultures.

These findings corroborate with those observed by Mair et al. (1996), who described 24 cases of HO in horses. Of these, 9 had granulomatous inflammatory lesions at necropsy, and 14 had thoracic injuries, although extrathoracic lesions coexisted in some cases. Tomlinson et al. (2011) described a case of equine HO associated with nodular pulmonary fibrosis. In this case, the animal also had severe swelling of limbs, with progressive weight loss during six months with normal appetite and intermittent fever.

The HO pathophysiology is not well defined. The mechanism appears to involve an initial increase in blood flow to the limbs with consequent fluid retention, followed by proliferation of vascular connective tissue and periosteum and subsequent bone deposition. However, the initial events leading to increased blood flow are not known (Tomlinson et al., 2011). Numerous theories have been proposed, including a hormonal and neurogenic cause. The hormonal disease refers to an increase in the estrogen level, which can be present in some cases. The neurogenic cause relies on the fact that vagotomy in humans can lead to regression of symptoms (Kawcak; Baxter, 2011). Chaffin et al. (1990) reported HO regression after treatment of an intrathoracic abscess in a horse. Enright et al. (2011) described HO regression in two animals treated with nonsteroidal anti-inflammatory drugs (NSAIDs) and corticosteroids. The horses had HO secondary to splenomegaly and inflammatory airway disease, respectively. In the present study, however, the injuries that led to hypertrophic osteopathy could only be identified during the necropsy and were characteristic of SGD.

SGD is also called equine sarcoidosis because of similarities with the systemic non-caseous granulomatous disease, sarcoidosis, which affects humans. However, it is preferable not to use this term to avoid confusion with the equine sarcoid. Like human sarcoidosis, SGD has no defined etiology. It is suggested that the disease results from an exaggerated immune response, against infectious agents or allergens that induce antigenic stimulus (Rees, 2004). In horses, no agent was directly associated with SGD. All research using histochemical, immunohistochemical, microbiological culture and molecular tests were negative for etiologic agents (Rees, 2004). The intake of vetch (Vicia villosa) was first incriminated as a cause of SGD in two horses, but later cases have been reported in horses that grazed in uncultivated
areas without the plant (SPIEGEL et al., 2006). The horse of this report never grazed in a field planted with vetch.

The definitive diagnosis of SGD is made by exclusion of other agents and requires histopathological evaluation and immunohistochemistry. The main differential diagnoses include bacterial and fungal infections of the skin, poisoning, and neoplasms (REES, 2004; SPIEGEL et al., 2006). Where granulomatous disease appears to be limited to the skin the prognosis is good, as horses respond to treatment with corticosteroids. However, when there is systemic involvement, as in this case, the prognosis is poor. Horses with SGD usually are euthanized a few weeks or months after the onset of clinical signs.

It is important to perform a thorough clinical examination and include hypertrophic osteopathy in the differential diagnosis of diseases that are accompanied by swelling of the face and limbs as edema from various causes or fibrous osteodystrophy, for example.

References


