NOTE

Branchial osteogenetic neoplasm in barbel *Barbus barbus plebejus*

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ABSTRACT: A branchial osteogenetic neoplasm affecting a barbel *Barbus barbus plebejus* (Valenciennes, 1829) is described. The osteoblasts' pleomorphism, the lack of a well-developed and complete separation, the presence of eccentric, terminal proliferative edges infiltrating the lining tissues and the abundant tumour matrix suggest a histopathological diagnosis of a 'productive osteoblastic osteosarcoma.' The occurrence of eosinophilic granule cells (EGCs) scattered among neoplastic tissue is discussed in relation to the neoplastic growth and the inflammatory reaction, with reference to recent discoveries in mammalian mast cell biology.

KEY WORDS: Barbel, *Barbus barbus plebejus*, Gills, Neoplasm, Osteoblastic osteosarcoma, Eosinophilic granule cells (EGCs)

An emaciated barbel *Barbus barbus plebejus* (Valenciennes, 1829) measuring approximately 40 cm in length was caught in the Adige River (North Italy). The fish exhibited a partial opening of the left branchial operculum and when the branchial chamber was dissected, an irregularly shaped, almond-sized tissue mass of reddish-grey color was found on the fourth gill arch. Necropsy performed on the entire fish indicated no gross pathology apart from serious cachexia. The affected gill arch and other tissue samples were preserved in 10% buffered formalin for later laboratory examination.

The tissue mass appeared to originate at the rostral edge of the gill arch; however, the osteo-chondroid axis of gill arch was not part of the tumour. Greyish-reddish ulcerations of the superficial pharyngeal mucous-epithelium were evident and a trabecular mass of bony appearance could be detected. Subsequent dissection of the tumour proved difficult because an osteo-chondroid consistency made the tumour resistant to cutting, prompting us to decalcify the tissue mass with an EDTA-based calcium chelating agent. Following decalcification, the tumour was sectioned with the sectioning plane oriented perpendicular to the gill arch. Tissue sections were processed for light microscopy and stained using the following techniques: Haematoxylin & Eosin, Alcian-Pas (McManus & Mowry 1960), Gomori Trichromic Stain (Mazzi 1977), Von Kossa (Pearse 1985).

Histologically, 3 tissue areas could be identified:

(1) The core of the mass appeared to be composed primarily of trabecular, lamellar bone forming a reticule which lined loose, well-vascularised fibro-connnective tissue and displayed proliferative, osteogenetic edges (Fig. 1).

(2) The proliferative edges of the osseous core merged gradually into a region of inflammatory fibrohistiocytic tissue. A shape progression from osteoblasts to fibroblastic-like cells was observed. Abundance of eosinophilic granule cells (EGCs) could also be seen in that tissue.

(3) The osseous, fibro-connective mass was superficially lined by the pharyngeal mucous-epithelium, displaying erosions, ulceration and mucous-epithelial cysts in the underlying connective tissue (propria-submucosa) (Fig. 2). Several EGCs were evident in the epithelium and in the propria-submucosa (Fig. 3). Histologically, the trabecular osseous tissue had a lamellar appearance and was lined by pleomorphic osteoblasts. Laterally, the cells appeared flattened and fibroblast-shaped; at the proliferative edges, the cells were arranged in a palisade formation with the major axis oriented perpendicular to the osteoid spicule (Figs. 4 & 5). Osteoblasts characteristically displayed basophile cytoplasm and contained large nuclei that were rich in euchromatin and had prominent nucleoli,
Fig. 1. Barbel, fourth gill arch, tumorous mass. A trabecular bone reticule, defining regions of loose, well-vascularised fibro-connective tissue (arrowheads), is apparent. Proliferative edges are evident (arrows), eliciting a fibro-histiocytic reaction (on the right). Haematoxylin & Eosin. Decalcified tissue. x40

Fig. 2. Barbel, fourth gill arch. Pharyngeal mucous-epithelium. A mucous-epithelial cyst embedded in the propria-submucosa is recognisable (arrowheads) beneath the pharyngeal epithelium. Mucous cell clusters are also evident (arrow). Haematoxylin & Eosin. Decalcified tissue. x400
Fig. 3. Barbel, fourth gill arch. Pharyngeal mucosa. The intense inflammatory reaction of the pharyngeal propria-submucosa lining the tumourous mass is apparent. Several blood vessels (arrowheads) and EGCs (arrows) scattered within the tissue are detectable. Haematoxylin & Eosin. Decalcified tissue. ×400

Fig. 4. Barbel, fourth gill arch. Close view of the proliferative edges. Large osteoblasts in palisade formation lining a newly formed bone spicula are visible (large arrows). The portion of osteoid immediately in contact with the osteoblasts appears clearer than the rest, suggesting a maturative progression from osteoid tissue to true osseous tissue (arrowheads). An osteocyte-like cell is apparent embedded within the bone matrix (small arrow). Haematoxylin & Eosin. Decalcified tissue. ×400
suggestive of a high biosynthetic rate. At the proliferative edges, osteoblasts appeared more voluminous and displayed a foamy, clearer, paranuclear portion of the cytoplasm (Fig. 5). Moreover, the portion of osteoid located nearest to the osteoblasts appeared less stain-reactive than the rest, suggesting a maturative progression from osteoid tissue to true osseous tissue (Fig. 4). Loose, well-vascularised, fibro-connective tissue was scattered among the trabeculae and lined by fibroblast-like osteoblasts.

At proliferative edges of the tumour, osteoblasts merged into a region of inflammatory fibro-histiocytic tissue embedded in a loose fibro-connective matrix that was rich in lymphocytes, histiocytes and spindle-shaped fibroblast-like cells. Haemorrhages, EGCs and karyorrhectic debris were also detected, suggesting a strong inflammatory and immunological reaction by the fish.

Because the tumour originated from the fish's gill arch, these structures were examined histologically to compare normal and pathological aspects of the tissue. The branchiospines of a normal gill arch had a lamellar axis of spongy bone. Very small, elongate fibroblast-like osteoblasts lined the bone axis and small, sparse osteocytes were embedded in the bone matrix. Normal branchiospines lacked both the palisade-arranged osteoblasts and the proliferative edges found in the previously described tumour bone. The bone lacunae of normal branchiospines were filled with loose, well-vascularised, fibro-connective tissue and EGCs were rare (less than 1 cell per 400× microscopic field) at the branchiospine base.

The pharyngeal mucous epithelium lining of the tumour mass displayed focal hyperplasias, erosions and ulceration involving the underlying propria-submucosa. Spongiosis and lymphocytic infiltration were also detected; moreover, EGCs were scattered among the mucous epithelium and the propria-submucosa (up to 15 cells per 400× microscopic field; Fig. 3). Mucous epithelium cysts were embedded in the propria-submucosa and extended into the inflammatory tissue described above (Fig. 2).

Our histopathological observations are consistent with those of an osteogenetic neoplasm (Pool 1990, Palmer 1992, Woodard 1996). The histopathology described here, however, differs from previous descriptions of osteogenetic tumours. In fact, the majority of osteogenetic tumours described in fishes are osteomas, of which only a few—mainly they are osteophytes or reactive osseous hyperplasias—can be
considered true neoplasms (cf. Wellings 1969, cf. Mawdesley-Thomas 1975, Hayes & Ferguson 1989). Osteoma itself has been considered by some pathologists to be hamartomas (Aegerter & Kirkpatrick 1968). According to Pool (1990), these tumours cannot be differentiated from exostosis by microscopic examination. In veterinary medicine, classification, diagnosis and prognosis of osteogenetic neoplasms are accomplished by correlating histopathological patterns with clinical and radiological findings (Pool 1990). The lack of suitable radiological and clinical findings, as well as the lack of a codified histopathological classification for osteogenetic fish neoplasms may explain the problematic and controversial diagnostic findings in fish.

Among vertebrates, teleosts are unique in having 2 major bone types: cellular or osteocytic bone and acellular or anosteocytic bone (Weiss & Watabe 1979, Ellis et al. 1989). Cellular bone possesses osteocytes embedded within the bone matrix is found in 'ancestral' teleosts (e.g. Clupeidae, Salmonidae, Cyprinidae) and shares morpho-functional features with the bone tissues of other vertebrates (Weiss & Watabe 1979, Ellis et al. 1989, Ferguson 1989). By contrast, acellular bone lacks osteocytes, is found only in 'advanced' teleosts (e.g. Percidae and Centrarchidae) and possesses unique morpho-functional features (Moss 1962, 1965, Weiss & Watabe 1979).

Reviewing the available diagnostic features concerning osteogenetic tumours in domestic animals (Pool 1990, Palmer 1992, Woodard 1996), and based solely on our histopathological examination, we tentatively propose a diagnosis of 'productive osteoblastic osteosarcoma' following the classification reported by Pool (1990). The terms 'productive' and 'osteoblastic' are meant to indicate that the osteosarcoma produces an abundant tumour matrix and is composed of anaplastic osteoblasts. We are led to this diagnosis by (1) the occurrence of osteoblast pleomorphism (i.e. cell morphology ranging from clearly delineated osteoblasts to spindle-shaped, fibroblast-like cells); (2) the lack of a well-developed, complete periosteal demarcation; (3) the presence of eccentric, terminal proliferative edges infiltrating the lining tissues; and (4) the histological evidence that a strong inflammatory, and possibly immunological, reaction occurred.

Concerning the inflammatory response and EGCs, the role of the latter in normal and pathological fish tissues is still a matter of debate, although recent experimental surveys suggest a homology with mammalian mastocytes (Reite & Evensen 1994, Reite 1997). Mammalian mast cells are involved in acute inflammation, in cellular growth modulation, and in leukocyte differentiation and activation (Aloe & Levi-Montalcini 1977, Norrby 1983, Wodnar-Filipowicz et al. 1989, Katayama et al. 1992, Leon et al. 1994, Paus et al. 1994, Galli & Costa 1995, Galli 1997). The ability of mammal mast cells to synthesise and release Tumour Necrosis Factor (TNF) is particularly interesting (Gordon & Galli 1991). In humans TNF is known to modulate a variety of physiological and pathological events such as: inflammation (acute phase reaction, endothelial effects, fibroblast effects, leukocyte effects), delayed hypersensitivity (endothelial effects), cell proliferation and tissue regeneration (mitogenic for hepatocytes and other cells), apoptosis (TNF receptor - TNFR1 - activation), and food intake control (tumour cachexia, obesity; cf. Cotran et al. 1999). Unfortunately, apart from the involvement of fish EGCs/mast cells in the inflammatory reaction (Secombes 1996, Reite 1997, 1998), there is no corresponding information about their role in fish tissues. Hence, the abundance of EGCs within the tumour mass, particularly at the proliferative edges and the inflammatory interface, must be tentatively interpreted in light of mammalian mast cell function and the apparent homology shared by fish EGCs and the latter. Similarly, the abundance of EGCs in the pharyngeal mucous epithelium and propria-submucosa can presumably be related to the epithelial modifications (mainly hyperplasia) and to epithelial cyst formation observed by means of extrapolation from mammalian mast cell action. Regarding cytochines in fish, although TNF has been reported in rainbow trout macrophages by Zelikoff et al. (1990) and in the serum of virus-infected fish by Ahne (1993), their detection and evaluation of function has just begun. Thus, the existence of TNF is rarely determined by identification at cDNA level, rather TNF is detected by the cross reaction with antibodies against mammalian cytochines and by the biological cross reactivity with mammalian cytochines (cf. Manning & Nakanishi 1996).

Although osteogenetic tumours are well known and described in fish oncology, they have not been reported to affect gills (cf. Wellings 1969, cf. Mawdesley-Thomas 1975). Considering the exposure gills have to the environment, and the probability that they will encounter carcinogens, it is surprising that the incidence of neoplasm in fish gills is so low compared other tissues and organs (cf. Wellings 1969, cf. Mawdesley-Thomas 1975, cf. Roberts 1989). To the best of our knowledge, and in view of the available literature on fish tumours (Wellings 1969, Mawdesley-Thomas 1975), this is the first description of a osteogenetic neoplasm affecting barbel gills.

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