Case Report

CONGENITAL MESOBLASTIC NEPHROMA IN A YOUNG BEAGLE DOG

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Abstract: A 10-month-old male beagle dog had a congenital tumor in the right kidney. Histopathologically, the tumor was characterized by fingerlike extensions projecting into the adjacent normal tissue, and often entrapping the normal glomerulo-tubular structure. The tumor was composed of spindle cells arranged in an interlacing fascicular pattern with some collagen fibers. Ultrastructurally, the tumor cells had long projections on the surface, rarely in contact with the junctional structures among the neighboring cells, and formed thin discontinuous basement membranes at the cell ends. Immunohistochemically, the tumor cells showed positive reactions with vimentin and S-100 protein antisera, and some weak reactions with actin, desmin, or myoglobin, but no reactions with cytokeratin.

This present tumor should be diagnosed as a congenital mesoblastic nephroma (CMN) on the basis of characteristic features in HE staining, other immunohistochemical reactions, and electromicroscopic findings, as well as the young age of the dog. (J Toxicol Pathol 9: 101–105, 1996)

Key words: Dog, Renal tumor, Congenital mesoblastic nephroma, Nephroblastoma, Spontaneous tumor

Introduction

Though spontaneous renal neoplasms in the dog are uncommon, renal tumors such as adenoma1,2, carcinoma1,2, nephroblastoma3,4, interstitial cell tumor5, fibroma6, fibrosarcoma1,2, lipoma1, hemangiomma1,2, hamartoma7, lymphosarcoma8, and papilloma9, or carcinoma of transitional cells1,2 have been reported. Recently, the extremely rare diagnoses which are congenital mesoblastic nephroma8 (CMN) and renal neurofibroma9 (NF) have also been reported in beagle dogs. CMN is one kind of benign nephroblastoma of early infancy in dog8, but detailed ultrastructural and immunohistochemical studies are lacking. Our description is the first instance of a spontaneous congenital mesoblastic nephroma in young beagle dog, carried out with the aid of ultrastructural and immunohistochemical techniques.

Materials and Methods

The present neoplasm was found in a 10-month-old male beagle dog among control animals of a subacute toxicity study for safety assessment in Toxicology Research Laboratories of Fujisawa Pharmaceutical Co., Ltd. The dog was purchased when six months old from Ichiyanagi Farm (Shizuoka, Japan) and had been vaccinated for several pathogens. In our facilities, the animal was housed alone in a stainless steel cage in an environment of 23±2°C temperature, 55±5% relative humidity and 12:12 hr light-dark sequence, and was fed a commercial dog diet (TC-1, Maruha Pet Food, Tokyo, Japan) of 300 g daily with free access to water. The animal was routinely observed for clinical signs twice a day and weighted once a week during the acclimation and the study periods. Routine hematological and blood-chemical examinations were done twice during the
acclimation and once a week during the study. At the conclusion of the study, the animal was killed by bleeding from the cervical aorta under deep anesthesia. Detailed gross pathological examination and weight of many organs including bilateral kidneys were performed. The organs were fixed in 10% phosphate buffered formalin, embedded in paraffin, sectioned, and stained with hematoxylin–eosin (HE) for routine histopathologic examinations. Additionally, renal lesions suggestive of abnormality were sectioned and stained with periodic acid–Schiff (PAS), azan, silver impregnation, and alcian blue (pH 2.5). Immunohistochemical examination was conducted with an avidin–biotin–complex immunoperoxidase method (ABC-Elite, Vector Laboratories, CA, USA) using anti-cow S-100 protein (1:200 in dilution, Dako, Glostrup, Denmark), anti-swine vimentin (V9, 1:50, Dako), anti-human cytokeratin (MNF116, 1:100, Dako), anti-human desmin (Clone 33, 1:50, Sanbio, Uden, Netherlands), anti-human myoglobin (1:200, Dako), and anti-α-smooth actin antibody (HHF35, 1:200, ENZO, New York, USA). For routine electron microscopy, some examples of renal lesions were further fixed in a 2.5% glutaraldehyde solution and postfixed in a 1% solution of osmium tetroxide, then dehydrated in ethanol and embedded in Epon812. Ultrathin sections was stained with uranyl acetate and lead citrate.

Results

No abnormalities were detected in clinical observations, bodyweight changes, hematology or blood-chemistry during the acclimation or the study periods. In gross appearance, a pale and slightly firm focus, 10 × 10 × 12 mm in size, in the right kidney was found. The longitudinal cut-surface showed a white to milky white focus extending from the cortex to the medulla with an ambiguous border (Fig. 1). The weights of both kidneys were within normal range. The other organs and tissues had no abnormalities in detailed gross and histopathological examinations except for the above mentioned right kidney.

Histopathologically, the tumor had infiltrated the adjacent renal tissue but there was no compression or formation of capsule at the border (Fig. 2). The normal glomerulus and tubules with few atrophic or cystic appearances were often entrapped by the tumor,
Fig. 3. Normal glomerulotubular structure in the myxomatous area of the tumor. HE, ×203.

Fig. 4. The tumor is composed of long spindle-shaped cells arranging in an interlacing fascicular pattern. HE, ×203.

Fig. 5. Electron microscopical findings. (a) Tumor cells show spindle or rod-shaped nuclei and long cell processes, and produce a large amount of mature collagen fibers. ×9,600. (b) Arrows show infrequent thin discontinuous basement membrane. ×28,800.
but the migration of the tumor cells to the structure was not seen (Fig. 3). In the loosely organized and myxomatous area (Fig. 3), the stroma was positive for alcian blue staining. The tumor was composed of long or short spindle cells, mostly arranged in a compact interlacing facicular pattern or presenting a wavy appearance (Fig. 4). The tumor cells had spindle-shaped or oval basophilic nuclei with a few nucleoli, eosinophilic cytoplasm with indistinct cellular border and some intercellular collagen, which was weakly positive for silver impregnation and PAS, and which was blue with azan staining. In all fields of the tumor, mitotic figures were scarce, and necrotic or hemorrhagic changes were not observed. Immunohistochemistry gave positive staining for vimentin and S-100 protein in most tumor cells and weaker positive staining for actin, desmin, and myoglobin in some cells, but no reaction to cytokeratin.

Ultrastructurally, the tumor cells had spindle or rod-shaped nuclei (Fig. 5a) with occasional irregular nuclear membrane. A few mitochondria, long rough endoplasmic reticulum, free ribosome, microfilament, fatty droplets, and pinocytic vesicle were observed in the cytoplasm. The tumor cells had long projections on the surface, rarely contacted by junctional structures among neighboring cells and formed thin discontinuous basement membrane at the cells ends (Fig. 5b), but lamellar structures were not observed. While mature collagen fibers were prominent in the intracellular area of high cellularity (Fig. 5a), a myxomatous matrix was seen in the intracellular area of low cellularity.

Discussion

Spontaneous renal neoplasms in the dog are scarce. However, we have found reports on three general types of canine renal tumor: epithelial tumors such as adenoma\textsuperscript{1,2}, carcinoma\textsuperscript{1-3}, and nephroblastoma\textsuperscript{24}; mesenchymal tumors including interstitial cell tumor\textsuperscript{8}, fibroma\textsuperscript{8}, fibrosarcoma\textsuperscript{1,2}, and lipoma\textsuperscript{7}; and others such as hemangioma\textsuperscript{1,2}, hamartoma\textsuperscript{7}, and lymphosarcoma\textsuperscript{2}, in our search of the literature to date. However, only two papers on renal tumors in laboratory beagle dogs\textsuperscript{8,9} are available. Recently, a congenital mesoblastic nephroma (CMN)\textsuperscript{8} was reported for the first time in beagle dogs, as one kind of benign type of nephroblastoma of early infancy, but sufficient ultrastructural and immunohistochemical investigations were not performed. Histopathologically, our case possessed such characteristic features as showing infiltrative growth pattern into the normal adjacent renal tissue and the existence of normal glomerulo-tubular components in the tumor. Based on the microscopic similarities to the abovementioned canine CMN\textsuperscript{8}, the present lesion was diagnosed as a congenital mesoblastic nephroma. In our case, the tumor cells showed positive reaction to vimentin and S-100 protein in immunohistochemistry, and also showed elongated cell processes, poor junctional structure, poor basement membrane, and abundant collagen fibers by electromicroscopy. Therefore the tumor cells were thought to be derived from mesenchymal cells, and possibly from fibroblasts or schwann cells. Furthermore, the immunoreactivity of some of our tumor cells to actin, desmin, and myoglobin would be evidence of the myofibroblastic differentiation.

The present tumor in this case must be distinguished from such tumors as schwannoma, neurofibroma, fibroma, leiomyoma, and nephroblastoma (mesenchymal type). Our case was different from some cases of CMN in human\textsuperscript{10,11} and dogs\textsuperscript{8} in point of immunoreactivity to anti-S-100 protein as a marker of schwannoma and neurofibroma (NF)\textsuperscript{12}. We could distinguish between this case and schwannoma in the absence of compression, encapsulation, interdigitation with renal elements, immunohistochemical reactivity except for anti-S-100 protein, and lamellar structures. In this connection, most of the reported cases of human renal schwannoma\textsuperscript{13} were malignant and we found no literature about canine renal schwannoma in our investigation. Moreover, our case could be distinguished from NF, because this type of tumor generally shows compression of the adjacent tissue and some of the electromicroscopical properties of schwannoma, but no immunoreactivity with the markers of muscular tissue\textsuperscript{12}. Formerly, Zwicker et al.\textsuperscript{9} had reported a case of canine renal neurofibroma, regardless of the same characteristic features in HE staining as interdigitation with renal elements and no compression of adjacent nephrons in CMN and our case. As the article did not describe the electromicroscopical and immunohistochemical analysis except S-100 protein, more detailed evidence would be needed to make a definitive diagnosis of
neurofibroma, and we dare say this case of CMN would hold true for our foregoing discussion. In addition to the aforesaid description, our case would be distinguishable from leiomyoma and fibroma in point of absence of compression, encapsulation, remnants of normal structures, and reactivity with alcian blue5–8,14. Also, there is often a proliferation of blastomal cells and primitive glomerulo–tubular elements in the mesenchymal type of nephroblastoma5–8, our present case did not have these features. Therefore, we could differentiate it from the mesenchymal type.

Human CMN is said to be a particular type of nephroblastoma rarely having such hamartomatous properties as rhabdomyocytic differentiation, angionoid forms, hematopoietic foci, cartilaginous foci, or dysplastic nephron constituents10,15,16. Additionally, human CMN can be classified by morphology into three major groups: classical (fibromatous), cellular, and combined (mixed) type10,16. The cellular type tends to have higher cellularity with many mitotic figures. The cellular and mixed types seem to be more malignant than the classical type. In our case, in spite of the infiltration to the adjacent nephrons, we concluded that the tumor belonged to the benign classical type on the basis of the above three human categories, because malignant features such as cytological atypia, high mitotic rate, focal hemorrhage, and necrosis were lacking. Additionally, the pattern of infiltration suggested that the tumor grew more or less harmoniously during fetal life as the cases in human14.

The present report is the first description of a spontaneous congenital mesoblastic nephroma in a young beagle dog undertaken by the use of both of ultrastructural and immunohistochemical investigations. From now on, more canine cases of CMN will need to be studied immunohistochemically and electronmicroscopically in order to further clarify the pathognomonic features and pathema of CMN.

References