Intracranial oligodendroglioma with optic nerve infiltration in a Labrador retriever

Intracraniaal oligodendroglioma met uitbreiding naar de nervus opticus bij een labrador retriever

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ABSTRACT

A seven-year-old neutered male Labrador retriever was presented with behavioral changes and reduced vision. Neurological examination revealed blindness of the left eye with a positive pupillary light reflex and a decreased mental status. Blood and cerebrospinal fluid analyses showed no abnormalities. MRI revealed a deviant area located in the thalamic, hypothalamic and caudate nuclear regions. Invasion of the left optic nerve was also observed. The dog was euthanized after 38 days of palliative treatment. Necropsy showed a non-encapsulated extruding white lardaceous mass, with a diameter of 2 cm, in the white matter, located in the ventral thalamic region near the chiasma opticum, which extended along the left optic nerve. Histologically, neoplastic cells had a “fried egg” appearance. Immunolabelling for glial fibrillary acidic protein demonstrated the presence of numerous reactive astrocytes. The tumor was diagnosed as a low grade (II) intracranial oligodendroglioma with infiltration of the optic nerve.

SAMENVATTING


INTRODUCTION

Neoplasia of the nervous system is a relatively frequent finding in dogs. There are reports of incidence rates varying from 0.145 to 3 % (Snyder et al., 2006). Over 70 % of the primary brain tumors occur in dogs over six years of age, while only 10 % occur in dogs of three years of age or younger (Koestner and Higgins, 2002). Overall, neoplasia of the nervous system has been reported in 1 to 3 % of the dogs in which an autopsy is performed (Braud et al., 2005). Breed and sex predilections have not been reported.

Primary central nervous system tumors may originate from neuroectodermal, ectodermal and/or mesodermal cells. These tumors rarely metastasize via cerebrospinal fluid or hematogenous route, but can sometimes invade the surrounding tissues. Secondary tumors may originate from a hematogenous metastasis of a different site of the body or from infiltrating neoplasms of the surrounding tissues (Braud et al., 2005).

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The common primary brain tumors in dogs are meningiomas, followed by astrocytomas and oligodendrogliomas. Undifferentiated sarcomas, choroid plexus tumors, medulloblastomas and ependymomas are less frequently observed (Snyder et al., 2006).

Oligodendrogliomas are neoplasms originating from oligodendrocytes, which play an important role in the maintenance of the axonal myelin sheaths. In dogs, this type of tumor has been reported to comprise up to 14% of the primary brain tumors (Snyder, 2006). Most often, these tumors occur in dogs of median age (Bullard et al., 1987). They are generally located in the white or gray matter of the cerebral hemispheres, and are usually well-demarcated, gelatinous and soft tumors, with multifocal hemorrhages (Koestner and Higgins, 2002).

Based on the most frequent localization of the tumor, which is at the cerebral hemispheres, patients are expected to show symptoms typically associated with supratentorial disorders, such as proprioceptive deficits, abnormal mental status or behavior, impaired vision and epileptic seizures (Kraft and Gavin, 1999; LeCouteur, 1999). Different symptoms may occur due to secondary changes in the brain or a different localization of the tumor (Braund et al., 2005).

Central nervous tumors have generally been classified by the World Health Organization (WHO) into four different grades. This classification includes a grading scheme that is a ‘malignancy scale’ ranging across a wide variety of neoplasms rather than a strict histological grading system whereby grade IV possesses the most malignant features. According to this system, oligodendrogliomas are divided into two groups: grade II (benign) and grade III (anaplastic, malignant) oligodendrogliomas (Louis et al., 2007). Oligodendrogliomas rarely metastasize and only a few reports describe multiple oligodendrogliomas in dogs (Koch et al., 2011).

The best way to visualize brain tumors is by using computed tomography (CT) or, even better, magnetic resonance imaging (MRI) (Kraft and Gavin, 1999). Unfortunately, medical imaging cannot differentiate oligodendrogliomas from other tumors (Young et al., 2010). Therefore, the definitive diagnosis is still based on histological examination (Braund et al., 2005).

CASE REPORT

Case history

A seven-year-old neutered male Labrador retriever was presented with three main complaints. During the last six months, behavioral changes and peruria (urinating randomly in the house) were noticed by the owner. The dog alternately gave no response or showed aggressive and hostile behavior as a reaction to the owners calls. On top of this, reduced vision had started to become obvious in the last four weeks as the dog had begun running into objects.

Clinical examination revealed a decreased mental status and a body temperature of 37.9 °C. On neurological examination, blindness of the left eye was observed as both the menace response and reaction to the cotton ball test were absent. The pupillary light reflex appeared to be present in both eyes. These results could have been explained by central blindness with a lesion in the right visual cortex, but to confirm this diagnosis further examination by the use of medical imaging was required.

Since abnormal behavior, altered mental status and vision loss, possibly due to central blindness, were the major symptoms, this dog was suspected of having a forebrain lesion (Wheeler, 1991). The peruria was most likely a secondary problem caused by the decreased mental status.

Hematology, serum biochemistry and analysis of the cerebrospinal fluid (CSF) did not reveal any abnormality.

On MRI, basic T1- and T2-weighted images and a fluid attenuated inversion recovery (FLAIR) sequence were taken. T1-weighted images, where fluids such as CSF color hypointensely, typically provide anatomical information, whereas T2-weighted images, where fluids color hyperintensely, give more information about the actual disease process. A FLAIR sequence is similar to a T2-weighted image, but nulls out “free fluid” so that CSF becomes hypointense instead of hyperintense, and oedema remains hyperintense. The T1-weighted sequence was repeated after the intravenous administration of Magnevist® (Bayer, gadopentetate dimeglumine) for contrast enhancement to detect lesions associated with abnormal vascularity or those thought to alter the blood-brain barrier.

The T1-weighted study showed a hypointense, ill-defined area located in the region of the thalamus, hypothalamus and the caudate nucleus. On the T2-weighted and FLAIR images, this intracranial lesion together with the left optic nerve colored hyperintensely, which revealed left optic nerve infiltration (Figure 1). There was no abnormal contrast enhancement visible on the T1-weighted sequence after the intravenous administration of Magnevist®. Altogether, MRI supported the diagnosis of an intracranial tumor with infiltration of the left optic nerve, and suspicion of a forebrain lesion was abolished.

Figure 1. MRI images. The T1-weighted transverse image (left) shows a hypointense ill defined area (black arrow) at the thalamus region. The T2-weighted image in a dorsal plane (right) shows a hyperintense aspect (black arrow) of the intracranial lesion and the left optic nerve (white arrow).
The patient was treated with Prednisolone 5MG® (Kela Laboratoria, Prednisolone, 1 mg/kg p.o. gradually decreased). Since the dog did not make any improvement after 38 days, he was euthanized and necropsied.

**Necropsy**

Necropsy showed a non-encapsulated extruding white fleshy mass with a diameter of 1.5 cm. This mass was located in the white matter of the ventral thalamic region near the chiasma opticum, and seemed to extend along the left optic nerve (Figure 2). No other brain lesions were detected. Besides congestion of the spleen and lungs, there were no other remarkable abnormalities.

**Histology**

Hematoxylin and eosin staining revealed a highly cellular intracranial mass of small uniform neoplastic cells with a small basophilic nucleus and a moderate amount of eosinophilic cytoplasm. In the left optic nerve, multifocal infiltration of neoplastic cells was detected. In the brain, the neoplastic cells had a slightly elongated nucleus in contrast to those in the left optic nerve, which had a round nucleus. Both in the brain and in the left optic nerve, the nuclei were surrounded by a perinuclear cytoplasmic vacuole, although the so called ‘fried eggs’ appearance was only obvious in the latter (Figure 3). The tumoral stroma consisted of fine branching capillaries.

**Immunohistochemistry**

Immunolabelling for the demonstration of astrocytes (glial fibrillary acidic protein, GFAP, Dako, 1:2000), neurons (neuron specific enolase, NSE, Dako, 1:800; neuron filament, NF, Dako, 1:1000; synaptophysin, Sy38, Dako, 1:400), endothelial cells (von Willebrand factor, Dako, 1:6400; CD31, Dako, 1:5), cell proliferation (Ki67, Dako, 1:20), T-lymphocytes (CD3, Dako, 1:400) and B-lymphocytes (CD20, Dako, 1:400) were performed. Staining was done using the Envision™ System (Dako).

The GFAP staining was clearly positive and showed that the tumor had a large population of non-neoplastic, reactive astrocytes (Figure 4).

Vascularization was demonstrated by the positive labelling of endothelial cells by vWF and CD31. The Ki67 stain showed a low mitotic activity within the tumor. Neoplastic cells did not react in NSE, NF synaptophysin CD3 and CD20 staining.

**DISCUSSION**

Snyder et al. (2006) studied 173 primary brain tumors in 172 dogs, of which 25 were identified as oligodendrogliomas. The most common presented signs in these animals were seizures (18 dogs), altered mental status (10 dogs) and loss of vision (5 dogs). Other symptoms included neck pain and vestibular symptoms. Although the symptoms seen in the patient of the
present case did not include epileptic seizures, there were clear signs of an altered mental status and loss of vision of the left eye. In the same study by Snyder et al. (2006), the results of the analysis of CSF fluid of seven dogs diagnosed with oligodendroglioma were reported. In five animals, the protein levels were elevated, and in three dogs the amount of white blood cells was increased. No abnormalities were detected in the cerebrospinal fluid of one dog, similar to the results of the dog in this case. At necropsy, most of the tumors were localized at the telencephalon (twenty dogs) (Snyder et al., 2006). The diencephalon (six dogs), mesencephalon (three dogs), myelencephalon (four dogs) and the olfactory region (five dogs) were less common targets. In half of the cases, the tumor had already stretched out to more than one of these regions. These findings suggest that the localization of the tumor in the thalamic region (which is localized in the diencephalon) in the present case is less common, but not exceptionally rare for this type of tumor.

To the authors’ knowledge, the infiltration of the optic nerve by an intracranial oligodendroglioma has not been reported in dogs yet. Naranjo et al. (2008) described a retinal oligodendroglioma with moderate invasion in the optic nerve. In human medicine, oligodendrogliomas originating from the optic nerve are rarely described (Offret et al., 1995; Lucarini et al., 1990). Although the tumor was morphologically classified as grade II, the tumor in the patient behaved rather malignant as neoplastic cells invaded the optic nerve. This probably concerned a slow process given the gradual worsening of the clinical symptoms.

Due to the positive pupillary light reflex, the blindness was suspected to be of central origin. However, no abnormalities were detected in the visual cortex on MRI or at necropsy.

This could be due to two reasons. Firstly, Koch et al. (2011) described a case report of a patient with multifocal oligodendroglioma without abnormalities at MRI or necropsy but with neoplastic cells present at histology. It is possible that the patient of the present case suffered from microscopical neoplastic infiltration of the visual cortex causing central blindness. Unfortunately, this could not be demonstrated. Secondly, it is also possible that the invading neoplastic cells did not destroy the optic nerve completely, resulting in a positive pupillary light reflex.

For the treatment of the patient, a palliative approach was chosen. Other options might have been radiation therapy or radiosurgery (Lester et al., 2001). The first method uses fractionation schemes to minimize the damage, caused by the ionizing radiation, to the surrounding tissues. Radiosurgery operates by directing highly focused beams of ionizing radiation with high accuracy to the neoplastic tissue via a stereotactic head frame. The radiation source rotates in an arc around the tumor, with several noncoplanar arcs. This method allows the use of a larger single treatment dose and thus a lower total dose. It therefore decreases the complications compared to classical radiation therapy. The availability of these treatments as well as the owner’s motivation may have influenced the choice of therapy.

In most cases, an oligodendroglioma has a typical appearance on histology, which is sufficient for the diagnosis. In some cases however, immunohistochemical staining may be necessary to distinguish an oligodendroglioma from other gliomas. In human medicine, the most frequently used primary antibodies for immunohistochemistry in tumors of the central nervous system are neuron-specific nuclear protein (a neuronally specific nuclear protein in vertebrates), synaptophysin (protein of the presynaptic vesicle), neuron-specific enolase (glycolytic isoenzyme in neurons and neuroendocrine cells), high molecular weight microtubule associated protein (required for initial neurite growth), neurofilament protein (intermediate filament found specifically in neurons), glial fibrillary acidic protein (intermediate filament protein expressed by astrocytes), vimentin (intermediate filament expressed by mesenchymal cells) and epithelial membrane antigen (heavily glycosylated transmembrane proteins expressed in several epithelial and non-epithelial cells) (Koperek et al., 2004). A study by Koperek et al. (2004) showed that markers for the detection of neuron-specific nuclear protein, vimentin and epithelial membrane antigen by immunohistochemistry can be used to differentiate a neurocytoma from a oligodendroglioma or ependymoma. Markers for vimentine, epithelial membrane antigen and GFAP can be used to differentiate between an ependymoma and oligodendroglioma. Unfortunately, these markers have not yet been evaluated for canine oligodendrogliomas. Furthermore, a specific antibody for the detection of oligodendrocytes in dogs is not yet available. Therefore, veterinary pathologists have to rely on the classical histological evaluation of tissue slides combined with immunohistochemistry. In the present case, histological examination revealed cells with a “fried egg” appearance, which was not always typical due to post-mortem autolysis and high numbers of astrocytes as demonstrated by immunohistochemistry. Careful examination revealed that these astrocytes were non-neoplastic and rather reactive, thus complicating the morphological diagnosis. It has been described that oligodendrogliomas may include reactive, proliferating but non-neoplastic astrocytes. When the amount of astrocytes exceeds 30% of the mass’ total cell population, these tumors are classified as oligoastrocytomas (Meuten, 2002).

CONCLUSION

The present case study describes a unique case of a low grade (II) intracranial oligodendroglioma in a dog with infiltration of the optic nerve. The more elongated nuclei in the neoplastic cells in the brain are a point of discussion and make differentiation between an oligodendroglioma and an oligoastrocytoma difficult. The GFAP-negativity of the neoplastic cells favors the first diagnosis. Nevertheless, this case points out the need of a specific immunohistochemical marker for oligodendrogliomas in small animal medicine.
REFERENCES


Uit het verleden

“De longen worden de vergaderplaats van tuberculeuze stof die zich in gezwellen verzamelt en eigenlijk de tuberkels uitmaakt, of die zich in longenweefsel verspreidende, eene tuberkuleuze oplooping uitmaakt. De gezwellen en verzamelingen zijn grys, geelachtig of doof wit; ze zijn van een vasten aerd, spekachtig of week; ze worden langzamerhand weeker, en veranderen in een dik kaesachtig vocht, dat een etter gelykt en schynt te zullen uitbreken. Zy veranderen ook in merlyk voorbereidsel bestaet; en dat dezelve eigen is aan de koeijen met eenen zwakken lichaemsbouw, fyne, gedweeë melk geven. Of de Letsels blijkt dat de ziekte nog niet duidelijk gedifferentieerd werd van echinococcus en fasciolasis.

Al bij al een vrij accurate weergave van deze ooit sterk verspreide plaag. Uit een hier weggelaten stukje beschrijving van de letsels blijkt dat de ziekte nog niet duidelijk gedifferentieerd werd van echinococcus en fasciolasis.

Luc Devriese