CASE REPORT

Coexistence of occipital dysplasia and occipital hypoplasia/syringomyelia in the cavalier King Charles spaniel

Concurrent occurrence of occipital dysplasia and occipital hypoplasia in two dogs is described in this report. Occipital hypoplasia results in reduced volume of the caudal fossa, leading to overcrowding of the neural structures and, in severe cases, development of syringomyelia. In occipital dysplasia, there is a failure of complete ossification of the supraoccipital bone. When the two conditions occur concurrently, it is possible that syringomyelia may develop more slowly, resulting in presentation with clinical signs in middle to old age. This has implications for screening tests for early detection of syringomyelia, with a view to using the dog for breeding purposes, as dogs with an apparently mild phenotype for occipital hypoplasia/syringomyelia may actually have a more severe genotype.

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INTRODUCTION

The foramen magnum is a ring of bone formed from four occipital bone centres: the supraoccipital bone dorsally, the basilar part ventrally and the exoccipitals that bear the occipital condyles laterally (Evans 1993). In addition to providing an exit for the spinal cord, the foramen magnum allows cerebrospinal fluid (CSF) to shunt rostrally and caudally between the head and the spine. This rapid efflux and influx compensates for brain expansion and contraction during the cardiac cycle (Oldfield and others 2001). Obstruction to CSF movement can result in development of syringomyelia, a condition whereby fluid-containing cavities develop within the spinal cord. The most common cause of syringomyelia in veterinary medicines is occipital bone hypoplasia (Rusbridge and others 2000), which is inherited in the cavalier King Charles spaniel (CKCS) and may be seen in other toy breeds (Rusbridge and Knowler 2003, 2004).

It is hypothesised that the bony defect of the supraoccipital bone is shortened, reducing the volume of the caudal fossa. The cerebellar vermis is often pushed through the foramen magnum and the medulla is deviated dorsally. Syringomyelia may vary in severity, but for many affected animals, it is a debilitating neurological disease, with clinical signs such as dysesthesia, cervical/occipital pain, paresis, ataxia and scoliosis. One of the most common signs is a tendency to scratch at the shoulder or the neck area. This has implications for screening tests for early detection of syringomyelia, with a view to using the dog for breeding purposes, as dogs with an apparently mild phenotype for occipital hypoplasia/syringomyelia may actually have a more severe genotype.
with any impairment of function. In contrast, Parker and Park (1974) did find neurological deficits in some of the dogs they studied (miniature and toy poodles, Yorkshire terriers, Lhasa apsos, Chihuahuas, beagle, Pomeranian, shih tzu and Maltese terriers); however, it was not established whether these were related to occipital dysplasia. Some of the dogs had concurrent hydrocephalus. Syringomyelia was not reported as a finding in any of the cases that received a postmortem examination and in no cases was there any apparent permeant protrusion of the cerebellum into the spinal canal. However, not all the dogs had a full postmortem examination of the spinal cord for syringomyelia. In particular, three dogs with a large dorsal notch and a long-standing history of ataxia did not have postmortem examination.

In the current report, two dogs with concurrent occipital dysplasia and occipital hypoplasia with secondary syringomyelia are described. The dogs were related to each other and to other dogs with occipital hypoplasia and secondary syringomyelia but without occipital dysplasia.

**CASE HISTORIES**

**Dog V**

A 10-year-old, male CKCS was presented for examination as the owner was concerned about the possibility of occipital hypoplasia/syringomyelia. In a previous study (Rusbridge and Knowler 2003), this popular stud dog had been identified as being an important ancestor in an extended family of CKCS with syringomyelia occurring secondarily to occipital bone hypoplasia and it was the sire and grandsire of several cases confirmed by magnetic resonance imaging (MRI). The only reported irregularity was that the dog had a mild tendency to scratch at its right mid-cervical area. This had been noticed since it was approximately 18 months old. There were no other neurological deficits at this time.

Ten months after the initial examination, the owner reported that the dog was becoming more sensitive around its right ear, and over the following three months, the dog developed tetraparesis more severe on the right, with more severe pain. The dog deteriorated to the stage that the owner elected for it to be euthanased. Gross postmortem findings are shown in Fig 1A and B. There was a large defect in the supraoccipital bone which was originally covered with a tough membrane that was confluent with the...
atlantooccipital membrane. In a normal neck position, the cerebellum extended into the foramen magnum. There were other gross lesions that could provide an explanation for the neurological signs and histopathological examination confirmed syringomyelia (Fig 1C).

A simplified familial relationship between dog V and other dogs with occipital hypoplasia/syringomyelia is illustrated in Fig 2. For the majority of dogs, it is not known whether or not there is concurrent occipital dysplasia as this cannot be readily appreciated on MRI. However, there were two descendants where the occipital bone was inspected intraoperatively. Dog T (grandson) and dog O (great grandson) had onset of the signs of syringomyelia at 14 and 28 months, respectively. Both had severe signs of pain and neither could be exercised as a consequence. Both were confirmed by MRI and subsequently had a suboccipital cranietomy with atlas cranial laminectomy and durotomy to relieve the obstruction at the foramen magnum. In both dogs, there was no dorsal notch to the foramen magnum, that is, no occipital dysplasia.

**Dog H**

Dog H was an eight-year-old, male CKCS, with a five-year history of brief episodes of unexplained pain. These episodes had become more frequent over the past six months. There was an approximate nine-month history of a tendency to scratch at the shoulders (both sides), with a six-month history of pelvic limb ataxia and a three-week history of thoracic limb weakness. A scoliosis had also been noticed. There had been a partial but not sustained response to 4 mg methylprednisolone (Medrone; Pfizer) daily and 200 mg gabapentin (Neurontin; Pfizer) twice daily. Neurological examination revealed a bilateral thoracic limb weakness more severe on the right. There was carpal hyperextension of this limb, with atrophy of the shoulder muscles and a tendency to stumble. Proprioceptive responses were delayed in the pelvic limbs. A tendency to scratch at the right shoulder was noted.

MRI of the brain and vertebral column revealed a small caudal fossa, overcrowding of the foramen magnum and syringomyelia from the level of C1 to L3/L4 (Fig 3). The width of the syrinx was variable, and at its maximum at the level of C2, and was two-thirds of the diameter of the spinal cord. Cervical scoliosis was confirmed. Other than the hypoplasia, no other irregularity of the occipital bone was identified. The caudal fossa had been imaged in both sagittal and axial orientations.

Due to the rapidly progressive clinical signs and inadequate response to corticosteroids and gabapentin, the owner elected for surgical management. A standard approach to the caudal fossa was made. When the supraoccipital bone was exposed, it was apparent that it was dysplastic. There was an arch-shaped dorsal widening of the foramen magnum. The bony defect was filled by a thick membrane. Above the bony defect, there was a band of normal bone (3 mm wide) above which was another hole 5×3 mm. The bony defect was widened, and the thick connective tissue membrane over the defect and the thickened atlantooccipital membrane were removed. The surgery was then continued with a cranial C1 laminectomy which was extended to approximately 5 mm below the tip of the vermis (about one-third of the length of the atlas arch).

Finally, a durotomy was made from just below the tip of the vermis to the level of the foramen magnum. This allowed further decompression of the cerebellum and allowed the surgeon to ensure that the vermis had been adequately exposed and to facilitate removal of arachnoid adhesions, but there were none. The resulting triangular defect was patched with biocompatible collagen matrix (Vet BioSILt; Cook/Global Veterinary Products). Closure was routine. Three weeks after surgery, the owner reported a marked improvement in demeanour, exercise tolerance, strength and coordination. The scratching behaviour had reduced. The familial relationship of dog H to dog V is illustrated in Fig 2.

**DISCUSSION**

The occipital bone forms from fusion of the mesenchyme of at least three occipital somites (Marin-Padilla 1991). The mesenchyme forms cartilage which in turn undergoes the process of endochondral ossification to form bone. In addition, there is a membranous tissue caudal to the cartilaginous supraoccipital bone plate which undergoes intramembranous ossification and ultimately fuses to the

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_Discussion about the role of the occipital bone and its relevance to the observed cases._

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**FIG 2.** Simplified diagram of one familial relationship between dogs V, H, T and O. The pivotal ancestral dogs had been identified in previous studies (Rusbridge and Knowler 2003, 2004). D and F Pivotal ancestral dogs, H and V Dogs with occipital dysplasia, T and O Dogs without occipital dysplasia.
cartilaginous part (Matsumura and others 1994). It is proposed that occipital dysplasia occurs when the ventromedial portion of the developing supraoccipital bone fails to ossify (Watson and others 1989). In contrast, it is proposed that occipital hypoplasia occurs because of an early paraxial mesodermal insufficiency (Marin-Padilla 1991).

Occipital dysplasia appears not to cause a functional problem because the overall shape and size of the caudal fossa are unchanged. In contrast, occipital hypoplasia results in a reduced volume of the caudal fossa which in turn can lead to the development of syringomyelika. Occipital dysplasia is common in dogs with a rounded skull shape (Watson and others 1989), and occipital hypoplasia is common in the CKCS; therefore, it is not surprising that the two conditions should occur in the same dog. What is unusual about dogs V and H is that the progression of the signs of syringomyelia was initially very slow and neither dog displayed severe signs until middle to old age. The majority of dogs with syringomyelia occurring secondarily to occipital hypoplasia present with severe compromise before seven years of age (Rusbridge and others 2004). It is possible that the membrane covering the supraccoxial defect allows for a dynamic expansion and less severe obstruction of CSF movement through the foramen magnum. As a consequence, it is possible that syringomyelia could develop more slowly, resulting in later onset signs. In other words, dogs with occipital hypoplasia and occipital dysplasia potentially may have a milder phenotype than those with occipital hypoplasia alone.

If this is the case, then there are implications for breeding. If the dog is a breeding male, then the mild or subclinical signs may not be recognised by the owner, especially when the dog is young. If the dog becomes a popular stud dog, then the potential for occipital hypoplasia but not necessary occipital dysplasia may be disseminated widely in the breed. Dog V sired more than 50 litters and has hundreds of descendants across the world. Many breeders arrange for their potential breeding stock to have a brain and/or upper cervical MRI, with the aim of selecting those without occipital hypoplasia/syringomyelia for at least one-half of a mating. If the onset of syringomyelia has been delayed by occipital dysplasia, then a dog may be erroneously thought to have a milder phenotype and used for breeding purposes. This is especially important if the screening is carried out before the age of 18 months. Occipital dysplasia is difficult to identify on MRI; however, the foramen magnum can be radiographed in a manner described by Parker and Park (1974), with the dog in dorsal recumbency, the nose flexed at 25 to 40°, and the x-ray beam centred on the frontal sinus. Computed tomography is also likely to be useful. However, before recommendations are made for screening, it should be established what is acceptable and unacceptable – that is, what degree of caudal fossa volume reduction leads to syringomyelia.

**Conclusions**

Occipital dysplasia may be seen in conjunction with occipital hypoplasia, possibly resulting in less obstruction of the foramen magnum and later/slower onset of syringomyelia. However, the affected dogs may still pass on a tendency for a more severe phenotype to their descendents. The presence of occipital dysplasia in conjunction with occipital hypoplasia should be taken into account in any future studies on imaging, CSF flow or genotyping, and further work is needed to establish whether occipital dysplasia does affect the pathogenesis of syringomyelia.

**Acknowledgements**

The authors are grateful to Alexander de Lahunta for reading and commenting on this text and to Dr Andrew Torrance of TDDS Laboratories and Dr Malcolm Silkstone of Abbey Veterinary Services for preparing and interpreting the histology of dog V.

**References**


Watson, A. G., De Lahunta, A. & Evans, H. E. (1989) Occipital bone hypoplasia (Chiari type I malformation) in cavalier King Charles spaniels. Veterinary Record 185 (Pt 2), 295-300


