

Case Report

Intramedullary dermoid cyst associated with Wallerian degeneration in a dog

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Abstract

Dermoid cysts are benign congenital lesions of ectodermal origin lined by stratified squamous epithelium and containing cutaneous adnexal structures, keratin debris, and sebaceous material. These lesions are commonly identified in the skin of dogs and cats, particularly along embryonic fusion lines.

25 Occurrence within the central nervous system is uncommon and intramedullary localization is
26 exceptionally rare. This report describes an intramedullary dermoid cyst associated with Wallerian
27 degeneration in an eight-month-old female Boxer dog presenting with a one-month history of paraplegia,
28 self-mutilation of the right pelvic limb paw, and loss of deep pain perception. Due to severe neurological
29 dysfunction, euthanasia was performed and the animal was submitted for pathological examination.
30 Gross examination revealed a 2 cm thin-walled, pearly white cystic lesion located within the L4–S3
31 spinal cord segments, containing friable brownish material with keratin filaments. Histopathological
32 examination demonstrated a cyst lined by well-differentiated stratified squamous keratinizing epithelium
33 with a prominent stratum granulosum and abundant orthokeratotic laminated keratin within the lumen.
34 Cutaneous adnexal structures, including sebaceous and apocrine glands, were observed within the cyst
35 wall. Adjacent spinal cord tissue exhibited severe compression, focal malacia, and Wallerian
36 degeneration characterized by digestion chambers and gutter cells. Immunohistochemically, the epithelial
37 lining showed strong cytokeratin immunostaining and absence of vimentin expression, whereas stromal
38 and adnexal components were vimentin-positive. The morphological diagnosis was an intramedullary
39 dermoid cyst associated with Wallerian degeneration. This report highlights the rare intramedullary
40 presentation of dermoid cysts in dogs and reinforces the importance of histopathological and
41 immunohistochemical evaluation for definitive diagnosis.

42

43 **Keywords:** intramedullary dermoid cyst, spinal cord, canine, Wallerian degeneration, neuropathology.

44

45 **Introduction**

46

47 Dermoid cysts are congenital developmental anomalies of ectodermal origin lined by stratified
48 squamous epithelium and containing keratin, sebaceous material, hair follicles, and other cutaneous

49 adnexal structures (5, 13, 15, 21). These lesions are commonly identified in the skin of dogs and cats,
50 particularly along embryonic fusion lines, and may vary from 1 to 5 cm in diameter (4, 13). The most
51 frequently affected anatomical sites include the dorsal cervical region, periorbital area, suprasternal
52 region, and head (4, 13). Dermoid cysts have been reported more commonly in large dog breeds such as
53 Saint Bernard and Boxer dogs, as well as in smaller breeds, including Dachshunds and Basset Hounds
54 (21). Cats may also be affected (15).

55 Dermoid cysts involving the central nervous system are uncommon and may occur intracranially
56 or, less frequently, within the spinal canal and spinal cord parenchyma (3, 5, 13, 20). These lesions are
57 considered congenital malformations associated with neural tube dysraphism, resulting from ectodermal
58 inclusion during neural tube closure in embryogenesis (10-13, 20). During this process, ectodermal
59 elements become entrapped within the developing neural tissue before complete meningeal formation,
60 giving rise to ectopic cutaneous structures within the neuraxis (10-13, 20). As these lesions progressively
61 accumulate keratinaceous debris and sebaceous material, they may act as space-occupying masses,
62 causing spinal cord compression and compressive myelopathy associated with neurological signs such
63 as paraplegia and sensory dysfunction (3).

64 Histologically, these lesions are classified according to the structures present within the cyst wall.
65 Epidermoid cysts are lined by stratified squamous epithelium without cutaneous adnexal structures,
66 whereas dermoid cysts contain skin appendages such as sebaceous glands, apocrine glands, and hair
67 follicles. Teratoid cysts additionally contain tissues derived from all three embryonic germ layers (1, 5).
68 Malignant transformation is considered rare but has occasionally been described, particularly as
69 squamous cell carcinoma (1, 5).

70 The differential diagnoses for dermoid cysts affecting the central nervous system include
71 encephalocele, meningocele, lipoma, hemangioma, hematoma, sebaceous cyst, pilonidal cyst, and
72 pericranial cysts (6, 8). Most spinal dermoid cysts reported in the literature are located in the intradural-

73 extramedullary compartment, whereas intramedullary localization is considered exceptionally rare (16).
74 Therefore, the objective of the present report is to describe the clinical, pathological, and
75 immunohistochemical findings of an intramedullary dermoid cyst associated with Wallerian
76 degeneration in a young Boxer dog.

77

78 **Case description**

79

80 An eight-month-old female Boxer dog was evaluated at the “Luiz Quintiliano de Oliveira”
81 Veterinary Hospital, School of Veterinary Medicine of Araçatuba (FMVA), UNESP, Araçatuba, São
82 Paulo State, Brazil. The animal had a one-month history of neurological signs, including self-mutilation
83 of the right pelvic limb paw, paraplegia, and loss of deep pain perception. Due to the severe neurological
84 dysfunction and poor prognosis, euthanasia was elected, and the animal was submitted for pathological
85 examination.

86 Tissue samples were collected during necropsy, fixed in 10% buffered formalin, routinely
87 processed for histopathology, embedded in paraffin, sectioned at 5 µm, and stained with hematoxylin
88 and eosin (H&E). Immunohistochemical analysis was subsequently performed to confirm the epithelial
89 origin of the lesion.

90 Gross examination revealed a thin-walled, pearly white cystic lesion located within the L4–S3
91 spinal cord segments, in the region of the lumbosacral plexus, measuring approximately 2 cm in diameter.
92 On cross-section, the cyst contained friable brownish material suggestive of keratin filaments (Fig. 1A),
93 and the adjacent spinal cord tissue was edematous and whitish.

94 Histologically, a cystic formation occupying more than 50% of the spinal cord parenchyma was
95 observed. The cyst was lined by well-differentiated stratified squamous keratinizing epithelium with a
96 prominent stratum granulosum. The lumen contained abundant orthokeratotic laminated keratin.

97 Cutaneous adnexal structures, including sebaceous and apocrine glands, were present within the cyst wall
98 and were associated with a moderate multifocal lymphoplasmacytic inflammatory infiltrate (Fig. 1B).
99 Adjacent adipose tissue was also observed. The lesion caused marked compression of the spinal cord
100 parenchyma, resulting in focal areas of necrosis of neurons, astrocytes, and oligodendrocytes with the
101 presence of Gitter cells and adjacent white matter Wallerian degeneration characterized by digestion
102 chambers and gitter cells. The morphological diagnosis was intramedullary dermoid cyst associated with
103 Wallerian degeneration.

104 Immunohistochemical analysis was performed using antibodies against cytokeratin (clone
105 AE1/AE3, monoclonal, mouse anti-human, Dako, dilution 1:250) and vimentin (clone V9, monoclonal,
106 mouse anti-vimentin, Dako, dilution 1:100). Antigen retrieval was performed in a pressure chamber
107 (Pascal, Dako). Endogenous peroxidase activity was blocked using methanol and 10% hydrogen
108 peroxide, followed by blocking of nonspecific reactions with Protein Block (Dako, code X0909). Primary
109 antibodies were incubated for 18h at 4°C. Immunoreactions were detected using the LSAB Kit (Dako,
110 code K0690-1) and visualized with 3,3-diaminobenzidine (DAB; Dako, code K3468-1). Sections were
111 counterstained with Harris hematoxylin.

112 Immunohistochemically, the epithelial lining of the cyst showed strong cytokeratin
113 immunolabeling, confirming its epithelial origin (Fig. 1C). The epithelial cells were negative for
114 vimentin, whereas the stromal tissue and cutaneous adnexal structures exhibited positive immunostaining
115 for this marker (Fig. 1D).

116

117 **Discussion**

118

119 Epidermoid and dermoid cysts involving the neuraxis are rare lesions reported in humans and
120 several animal species, including dogs, horses, rodents, and mice (1, 4, 6, 10-15, 18). In canine breeds,

121 Boxer dogs have been described as a predisposed breed for the development of dermoid cysts, including
122 lesions affecting the central nervous system (15).

123 Dermoid cysts are congenital lesions associated with neural tube dysraphism and originate from
124 abnormal ectodermal inclusion during neural tube closure in embryogenesis (2, 10-13, 20, 16). During
125 this developmental process, ectodermal elements become entrapped within the developing spinal canal
126 or spinal cord parenchyma before complete meningeal formation, resulting in ectopic cutaneous
127 structures within the neuraxis (10-13, 20). Although congenital, clinical signs may only become evident
128 later in life due to the gradual accumulation of keratinaceous debris and progressive compression of
129 adjacent neural tissue (2, 16).

130 Most spinal dermoid cysts described in the literature are located in the intradural-extramedullary
131 compartment, whereas intramedullary localization is considered exceptionally rare (16). The unusual
132 intramedullary location observed in the present case highlights the importance of including this lesion
133 among the differential diagnoses for progressive spinal cord compression in young dogs presenting
134 neurological dysfunction.

135 The dog described in the present report exhibited paraplegia, loss of deep pain perception, and self-
136 mutilation of the right pelvic limb associated with severe neurological dysfunction (21-23). The self-
137 mutilation behavior was likely associated with neuropathic sensory alterations, such as paresthesia or
138 dysesthesia, resulting from compression of the dorsal horns and/or lumbosacral nerve roots by the cystic
139 lesion. Similar manifestations have been reported in dogs with neuropathic pain secondary to spinal cord
140 and peripheral nerve injury (7, 23).

141 Gross findings in the present case were consistent with previous descriptions of spinal dermoid
142 cysts, characterized by a pearly white cystic lesion containing friable keratinaceous material (5, 12, 15,
143 20, 21). Histologically, dermoid cysts resemble cutaneous dermoid lesions and are characterized by cyst
144 walls lined by well-differentiated stratified squamous keratinizing epithelium supported by fibrous

145 stroma and associated with cutaneous adnexal structures, including sebaceous glands, apocrine glands,
146 and hair follicles (1, 5, 10, 12, 20). In the present case, the identification of cutaneous adnexal structures
147 within the cyst wall was essential for distinguishing the lesion from an epidermoid cyst.

148 The main histopathological differential diagnosis for dermoid cyst is epidermoid cyst, since both
149 lesions are lined by stratified squamous keratinizing epithelium and contain laminated keratin within the
150 lumen (9, 19). However, epidermoid cysts lack cutaneous adnexal structures, whereas dermoid cysts
151 contain skin appendages such as sebaceous glands, apocrine glands, and hair follicles within the cyst
152 wall (9, 19). The presence of these adnexal structures supported the diagnosis of intramedullary dermoid
153 cyst in the present case.

154 In addition to the cystic lesion, adjacent spinal cord tissue exhibited focal necrosis and Wallerian
155 degeneration characterized by digestion chambers and Gitter cells. These findings were likely secondary
156 to chronic compression of the spinal cord parenchyma by the expanding cystic lesion, resulting in
157 ischemia, hypoxia, and compressive myelopathy. The loss of deep pain perception observed clinically
158 further supported the severity of the spinal cord injury (22).

159 Immunohistochemically, the epithelial lining of the cyst showed strong cytokeratin
160 immunoreactivity and absence of vimentin expression, confirming its epithelial origin. In contrast,
161 stromal tissue and cutaneous adnexal structures exhibited positive vimentin immunostaining, consistent
162 with their mesenchymal origin. Similar immunohistochemical findings have been previously described
163 in dermoid cysts affecting the nervous system (10, 12, 20).

164 Although dermoid cysts are more frequently reported intracranially, particularly involving the
165 fourth ventricle and cerebellopontine angle (13, 20), spinal cord involvement is considerably less
166 common (10, 12, 21, 22). Therefore, the present report contributes to the limited veterinary literature
167 describing intramedullary dermoid cysts associated with severe neurological dysfunction and secondary
168 Wallerian degeneration in dogs.

169 Intramedullary dermoid cysts are exceptionally rare congenital lesions in dogs and should be
170 considered among the differential diagnoses for progressive spinal cord compression associated with
171 severe neurological dysfunction in young animals. In the present case, the presence of cutaneous adnexal
172 structures within the cyst wall was essential for differentiating the lesion from an epidermoid cyst.
173 Additionally, the associated Wallerian degeneration and neuropathic clinical manifestations highlighted
174 the severe secondary effects of chronic spinal cord compression. Immunohistochemical findings
175 supported the epithelial origin of the lesion and complemented the histopathological diagnosis.
176 Nevertheless, histopathological evaluation remains the gold standard for the definitive diagnosis and
177 differentiation of dermoid cysts from other intramedullary lesions.

178

179 **Data Availability**

180 All the original contributions presented in this study are included in the article/supplementary
181 material. Further inquiries can be directed to the corresponding author.

182

183 **Author Contributions**

184 **Mayara Caroline Rosolem:** Conceptualization, Data curation, Formal analysis, Investigation,
185 Writing – original draft, Writing – review & editing. **Pamela Rodrigues Reina Moreira:** Methodology,
186 Investigation, Formal analysis, Writing – review & editing. **Rosemeri de Oliveira Vasconcelos:**
187 Conceptualization, Resources, Supervision, Writing – review & editing. **Gisele Fabrino Machado:**
188 Investigation, Formal analysis, Resources, Writing – review & editing. **Maria Gisela Laranjeira:**
189 Investigation, Formal analysis, Resources, Writing – review & editing.

190

191 **Conflict of Interest**

192 The authors declare no competing interests.

193

194 **Generative AI Use Statement**

195 The authors did not use generative artificial intelligence tools or technologies in creating or editing
196 any part of this manuscript.

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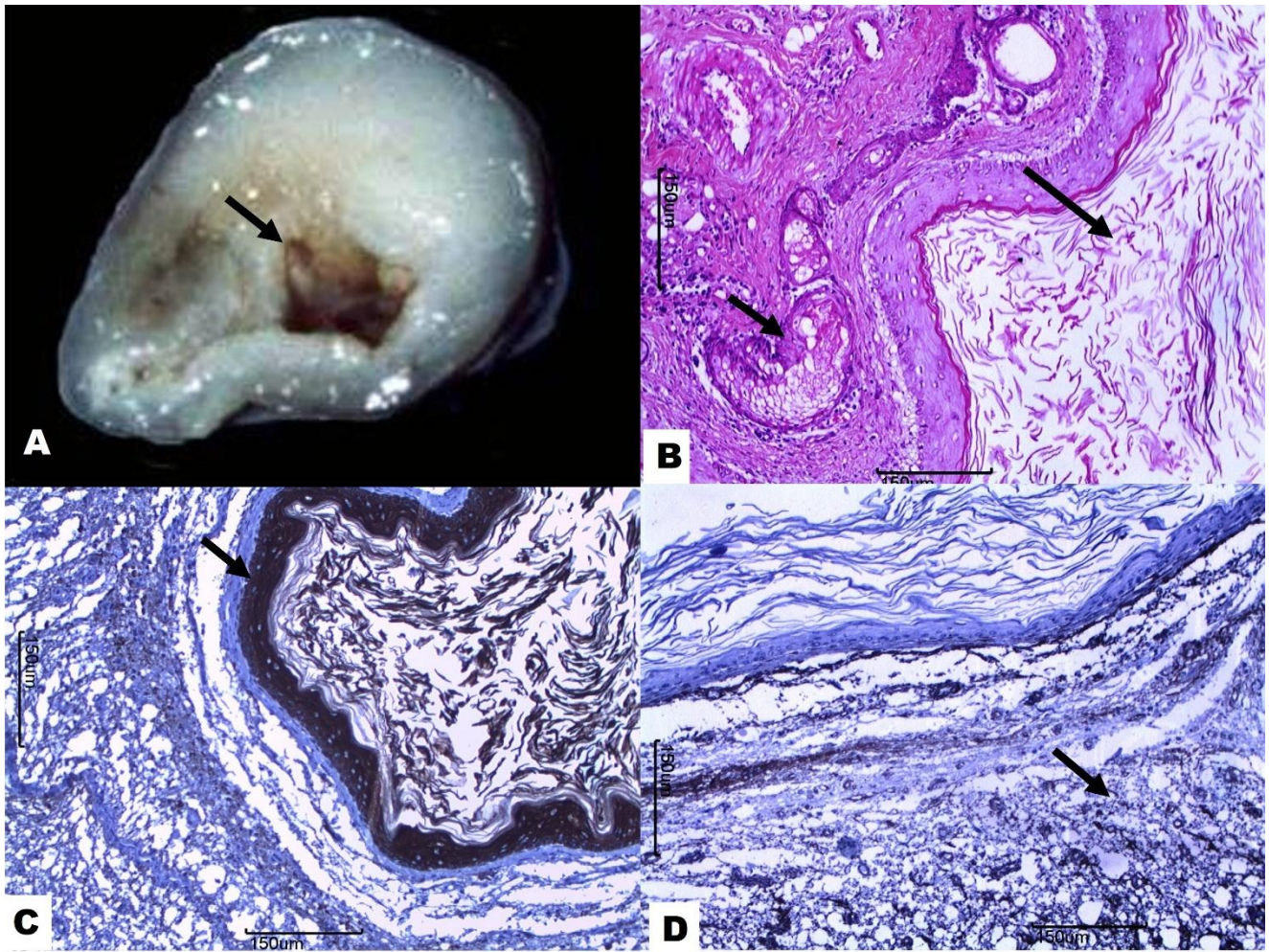
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253

254 **Figure 1.** Intramedullary dermoid cyst, dog. (A) Gross appearance of a tan swelling (arrow). On cross-
 255 section, a pearly white cyst measuring approximately 2 cm in diameter and containing friable brownish
 256 material suggestive of keratin filaments was observed. (B) Histopathological appearance showing
 257 cutaneous adnexal structures (short arrow) and well-differentiated stratified squamous keratinizing
 258 epithelium with a prominent stratum granulosum surrounding the nervous tissue (long arrow) (H&E,
 259 400×). (C) Immunohistochemistry showing positive cytokeratin immunostaining in the epithelial lining
 260 of the cyst. (D) Immunohistochemistry for vimentin showing negative staining in the epithelial cells and
 261 positive immunolabeling in the stromal tissue (arrow) and cutaneous adnexal structures (peroxidase
 262 polymer method, bar = 100 μm).