Rosai-Dorfman Disease of the Spine Masquerading as Meningioma

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Rosai-Dorfman disease (RDD) is histiocytic proliferative disorder of idiopathic etiology and the characteristic of the disease includes sinus histiocytosis and massive lymphadenopathy.¹ The central nervous system (CNS) is involved in less than 5% of the cases (75% brain and 25% spinal cord).² Isolated involvement of the spine in RDD (without involvement of the lymph nodes) is rare with only few reported cases in literature.³⁻⁷ On imaging isolated RDD of the spine can mimic meningiomas, making its preoperative diagnosis difficult.³,⁴ A 65-year-old woman presented with the history of progressive onset of spastic paraparesis. There was no history of bladder or bowel involvement. On examination, the patient had grade 3 power in all muscle groups of the lower limbs; her sensory level was at D10 level. She had reduced sensations to touch and pin prick as well as decreased proprioception below this level. Axial T1-weighted (T1W) magnetic resonance imaging (MRI) of the brain showed an extradural isointense lesion compressing the cord and pushing it towards the left side. The lesion was becoming hypointense on T2-weighted (T2W) axial images, and it was enhancing after contrast administration (►Fig. 1). The patient underwent D7–D10 laminectomy. Intraoperatively the lesion was pale yellow in color, was relatively avascular, and had both intra- and extradural components. Histopathologic examination of the lesion showed features suggestive of extranodal RDD. Microscopic examination demonstrated mixed inflammatory background consisting of lymphocytes, plasma cells, and polymorphonuclear leukocytes. Singly scattered and clusters of pale staining histiocytes demonstrating emperipolesis are seen. Foci of fibrosis are noted (►Fig. 2). The lesional histiocytes that showed emperipolesis express S100 protein and CD30 did not reveal any R-S cells. Special stains for microorganisms were negative. Postoperatively the patient gradually improved in her paraparesis and sensory symptoms. On discharge, she had grade 4/5 power in all muscle groups of the lower limbs. She is doing well at 1-year follow-up.

The exact etiology of RDD is not known and it is considered a benign idiopathic histiocytosis. Infectious agents (i.e., human herpes virus 6 [HHV-6] and parvovirus B19) or an abnormal immunologic response have been suggested as causative factors.⁴,⁸ Although there is no age preponderance for the CNS involvement in RDD, the disease usually involves male patients between 20 and 40 years of age.¹,² The systemic RDD most commonly involves the orbit, skin, respiratory tract, bones, kidneys, heart, and head and neck region, and is characterized by massive and painless cervical lymphadenopathy with variable symptoms of fever, leukocytosis, raised erythrocyte sedimentation rate (ESR), and weight loss.⁵,⁹,¹⁰ Involvement of the spine is uncommon, and usually there is absence of lymphadenopathy.¹¹

It is difficult to make a diagnosis of isolated RDD on imaging as MRI features of this disease are almost similar to other common lesions of the spinal cord (i.e., meningiomas).⁴ The signal intensity of these lesions on MRI ranges from iso- to hypointensity on T1W images to iso- to hyperintensity on T2W images, and after the contrast administration, the lesions show an intense and uniform enhancement (including dural tail).¹²,¹³ Because of this, the definitive diagnoses are possible only after the histopathology (if necessary immunohistochemistry), which will be the characteristic pattern of the disease.⁴,¹² On histopathology, RDD is characterized by emperipolesis “phagocytosis of intact lymphocytes by macrophages” and

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presence of large histiocytes immunoreactive for S100 protein and CD68. These cells are negative for CD1 distinguishing from Langerhans' cell histiocytosis. The mainstay of treatment is surgical decompression, and the main objective of the management of spinal RDD is to relieve the compression over the neural structures. In cases of recurrent or progressive cases of RDD, several therapies (e.g., radiotherapy and combination of chemotherapy with corticosteroids, 6-mercaptopurine, and methotrexate) with variable success have been reported. The natural history of RDD ranges from complete and spontaneous remission, periodic exacerbations and remissions, and persistence of the disease, and, in some cases, it may be associated with fatal outcome due to systemic complications.

Fig. 1  MRI T1W (A), T2W (B), and axial images showing extradural hypointense lesion compressing the cord and pushing it toward left side. MRI dorsal spine postcontrast sagittal image (C) showing intensely enhancing lesion with dural tail.

Fig. 2 (A) H and E stain 10X magnification—microscopic examination demonstrating that mixed inflammatory background consists of lymphocytes, plasma cells, and polymorphonuclear leukocytes. Singly scattered and clusters of pale-staining histiocytes seen and (B) H and E stain 40X magnification—microscopic examination demonstrating that mixed inflammatory background consists of lymphocytes, plasma cells, and polymorphonuclear leukocytes. Singly scattered and clusters of pale-staining histiocytes demonstrating that emperipolesis are seen. Foci of fibrosis are noted.
Isolated involvement of the spine in RDD is uncommon, and it needs to be considered in the differential diagnosis of patients presenting with spinal cord compression. Early surgical intervention, resection of the lesion, and relief in the mass effect help increase chances of recovery of neurologic functions.

References
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