An orphan disease: IgG4-related spinal pachymeningitis: report of 2 cases

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IgG4-related disease is relatively new disease entity and a rare one, and our knowledge of this entity continues to evolve. It was first described in the pancreas and since then has been described in virtually every organ. Spinal involvement resulting in pachymeningitis is rare, and there are only 8 reported cases of the same to date, with the cervicothoracic spine being the most commonly affected region.

The authors describe 2 cases in which the patients presented with spinal compression resulting in myeloradiculopathy (Case 1) and radiculopathy (Case 2). Imaging of spine in both cases revealed an ill-defined contrast-enhancing lesion at the lumbar level. Preoperatively, a diagnosis of spinal tumor was made, but intraoperatively no spinal tumor was found. The diagnosis was established histopathologically.

The disease has no particular defining features clinically or radiologically and can mimic common spinal tumors. It is important to accurately diagnose this rare entity because of its multisystem involvement and progressive course. Strict treatment guidelines have yet to be formulated. Although histologically this disease can mimic other inflammatory conditions, the presence of storiform fibrosis and an increased number of IgG4-positive plasma cells can help in clarifying the diagnosis.

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KEY WORDS IgG4; pachymeningitis; spinal; intradural; spinal compression

THE newly recognized fibroinflammatory condition IgG4-related disease is characterized by tumefactive lesions, a dense lymphoplasmacytic infiltrate rich in IgG4-positive plasma cells, storiform fibrosis, and, often but not always, elevated serum IgG4 concentration.10 Because it is a relatively newly recognized entity, the picture of IgG4-related disease continues to evolve in the published reports. Even the nomenclature is still evolving, the latest to be adopted being IgG4-related disease.11 This disease was initially described in the pancreas. In the last decade or so as the disease had been diagnosed with increased frequency clinically, it has been described in virtually every organ system: the biliary tree, salivary glands, periorbital tissues, kidneys, lungs, lymph nodes, meninges, aorta, breast, prostate, thyroid, pericardium, and skin.3,4,6–8 The treatment guidelines are being continually modified.

We report 2 cases of spinal pachymeningitis due to IgG4-related disease. The presentation was like any other intradural spinal lesion. Although the intraoperative features were not typical of a spinal tumor, the final diagnosis was clinched on histopathology.

Case Reports
Case 1
A 50-year-old man presented with a 6-month history of progressive bilateral lower-limb weakness. He had wasting of lower limbs and difficulty in voiding for 3 months. Examination revealed significant atrophy and Grade 3/5 weakness in the lower limbs bilaterally.

Magnetic resonance imaging (Fig. 1) revealed an ill-defined intradural lesion at L1–2 that was isointense on T1-weighted imaging (T1WI), hypointense on T2-weighted

ABBREVIATIONS T1WI = T1-weighted imaging; T2WI = T2-weighted imaging.
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IgG4-related spinal pachymeningitis

Case 1

A 19-year-old man presented with 1-year history of backache and radiation of pain down to the left lower limb that had been present for 10 days. Examination revealed subtle weakness (Grade 4/5) bilaterally at the knee extensors with no sensory deficits. Results of complete blood count, electrolytes, and renal function tests were within normal limits. MRI showed an ill-defined intradural lesion at L2–3 (Fig. 3A–D). The lesion was isointense on T1WI and hypointense on T2WI and showed dense enhancement after gadolinium administration. The patient underwent L–2 laminectomy, and during surgery the dura was found to be diffusely thickened and hypertrophic, and it showed moderate vascularity. No definite tumor was seen underneath the dura. The margins between normal and abnormal dura were difficult to discern. Since no apparent lesion was seen, a mere biopsy of the thickened tissue was taken. Histopathological examination suggested IgG4-related pachymeningitis (Fig. 4 and Table 1). The patient was started on a course of oral steroids, was pain free at 6 months, and had no additional neurological deficits. Postoperative imaging showed near-complete resolution of the lesion (Fig. 3E–H).

Discussion

IgG4-related pachymeningitis is a rare disease. The predominant involvement in such cases is of cranial dura and spinal involvement is more rare. Only a handful of case reports are present in the literature. Because the disease is relatively new, the definition, diagnostic criteria, treatment protocols, and even nomenclature continue to evolve and are frequently revised/modified. The importance of establishing an early diagnosis lies in the fact that the disease mimics commonly encountered lesions in spinal locations and this can lead to a diagnostic pitfall. Another important factor in favor of an early diagnosis is the multisystem involvement in this disease, which is chronically progressive.

In the present study, both patients presented with features of neural compression at the level of involvement (i.e., myeloradiculopathy in Case 1 and radiculopathy in Case 2). MRI revealed an ill-defined lesion that was isointense on T1WI with intense contrast enhancement. The lesion was hypointense on T2WI. No particular distinguishing radiological features have been described for this disease to distinguish it from malignant lesions.

In both cases the patients were taken to surgery with a preoperative diagnosis of intradural neoplasia (e.g., meningioma/schwannoma/neurofibroma/malignant nerve sheath tumor).

Intraoperatively, no discernable neoplastic lesion was found, but the dura was thickened, hypertrophic, and diffusely involved. Defined borders between normal and abnormal areas were not found. In both cases the lesion was moderately vascular.

IgG4-related pachymeningitis is a histopathological diagnosis and is characterized by a plasma cell–rich inflammatory infiltrate, admixed with variable numbers of eosinophils, storiform fibrosis, and obliterator phlebitis. The plasma cells are predominantly IgG4 positive, and there is an increase in the IgG4/IgG–positive plasma cell ratio.

No consensus exists on the number of IgG4-positive plasma cells required for establishing a diagnosis of IgG4-related pachymeningitis. However, Lindstrom et al. proposed that > 10 IgG4-positive plasma cells/HPF were necessary for a diagnosis of IgG4-related pachymeningitis. Compared with other organs, obliterator phlebitis is a rare finding in meningeal IgG4-related disease. Important mimics from which this disease should be distinguished include the periphery of an abscess, autoimmune diseases, vasculitis, and any infectious pathology like tuberculosis.

Standard therapeutic regimens are still lacking for the treatment of IgG4-related disease. There are international consensus guidelines on how to manage it. The summary
includes following: 1) All patients with symptomatic, active IgG4-related disease require treatment. 2) Glucocorticoids are the first-line agent for achieving disease remission. However, the dosage and duration of treatment are still not standardized. 3) If there is relapse, either retreatment with steroids or other immunosuppressive agents may be required.

Immunosuppressive therapy in cases of relapse has been recommended based on an extrapolation from the treatment of other autoimmune conditions. These agents include azathioprine, mycophenolate mofetil, rituximab, and bortezomib.

Although most patients exhibit an initial response to glucocorticoids, the disease follows a progressive course. Clinical response is inversely proportional to the degree of fibrosis.

Spinal decompression provides relief by increasing space for neural elements. In addition, surgery provides adequate tissue for detailed histopathological examination.

**Conclusions**

We wish to state that IgG4-related disease is a rare differential diagnosis of leptomeningeal disease. No specific clinical or radiological hallmarks/diagnostic criteria have yet been defined to label the disease. Therefore, the diagnosis rests with histopathological examination, and the pathologist should have high index of suspicion. Although histologically this disease can mimic other inflammatory conditions, the presence of storiform fibrosis and an increased number of IgG4-positive plasma cells can help in confirming the diagnosis. Because the disease is relatively new, its characterization is evolving and newer additions are being made in the literature. Until the time definite

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<th>TABLE 1. Histopathological features of IgG4-related spinal pachymeningitis</th>
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* Positive cell count per ×40 field.
FIG. 3. Case 2. Preoperative (A–D) and postoperative (E–H) images. Sagittal T1-weighted image showing a hypodense lesion at L2–3 (A). Sagittal T1-weighted postcontrast image demonstrating intense contrast enhancement (B). Postcontrast axial T1-weighted image showing the ill-defined, contrast-enhancing lesion (C). Axial T2-weighted image demonstrating the hypodense lesion causing thecal sac compression (D). Sagittal T1-weighted image showing no lesion at the L2–3 level (E). Postoperative changes are seen in the region of the lamina and paraspinal muscles. Sagittal T1-weighted postcontrast image showing no contrast enhancement at the site of the resected lesion (F). Axial T1-weighted postcontrast image showing the absence of a lesion (G). Postoperative changes can be appreciated, for example, on this T2-weighted axial image in which no lesion is seen (H).

FIG. 4. Case 2. Photomicrograph showing dense fibrosis with obliterative phlebitis (A) and numerous plasma cell infiltrates (B). H & E, original magnification ×200. These plasma cells show positivity for CD138 (C), plasma cells showing positivity for IgG (D), and numerous IgG4 positive plasma cells (E). Immunoperoxidase, original magnification ×400. Figure is available in color online only.
diagnostic criteria are well established, it is extremely important to accurately diagnose this disease entity since the involvement is multiorgan and the course progressive.

References

Disclosures
The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions
Conception and design: Singla, Radotra. Analysis and interpretation of data: Radotra, Aggarwal, Kapoor. Drafting the article: Aggarwal, Kapoor. Critically revising the article: Singla, Radotra, Aggarwal. Administrative/technical/material support: Chatterjee.

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