Case Report

Brachial plexus tuberculosis: A unique neurological variant of a common clinical disease

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ABSTRACT

Isolated primary tuberculous involvement of the brachial plexus has not been reported in the past. Here, we report the case of a 29-year-old male who developed neck pain and radiculopathy for 3 months. Weakness in the left shoulder-elbow (2/5 MRC grade) and wrist (4/5 MRC grade) was present with numbness to pain and temperature along the C4–C5 dermatome. Magnetic resonance imaging (MRI) revealed a T2 hyperintense heterogeneously contrast-enhancing lesion involving the upper trunk of the brachial plexus. Under suspicion of malignancy, surgical exploration was undertaken. Intraoperatively, pus was noticed with unhealthy granulation tissue extending along the left-sided C4–5 lamina and transverse process. The frozen section suggested granulomatous infection and histopathology confirmed tuberculosis (TB). After 18 months of antituberculous treatment, the patient gradually recovered complete strength in the left upper limb. Follow-up brachial plexus MRI showed near-complete resolution of the lesion. We describe a unique case of a young patient clinically suggestive of the rapid progressive lesion (mimicking malignancy) affecting brachial plexus, turning out as TB on histopathology.

Key words: Brachial plexus, Neuritis, Tuberculosis

Tuberculosis (TB) is a common condition affecting a variety of neural structures. TB proved to be the second most common infectious cause of death after coronavirus disease (COVID-19) in the year 2020. According to the Global TB report 2021, 43% of TB cases were reported from the Southeast Asia region and 26% were from India. Men represent 56% of the affected population, while women and children represent 33% and 11%, respectively. The burden of multidrug-resistant TB and rifampicin-resistant TB has remained stable at 3–4% (from 2015 to 2020) [1]. Developing countries have high endemicity for cranial and spinal tuberculous patients. The central nervous system is involved in approximately 10% of all TB cases [2]. Despite this, isolated involvement of the brachial plexus has not been described in the literature. Non-lesional lumbosacral plexopathy in a patient with pulmonary TB has been reported in the past [3].

CASE REPORT

We describe a patient with this chronic granulomatous infection of the upper trunk of the brachial plexus, which we believe is the first case of isolated brachial plexus TB to be reported in the literature to date.

A 29-year-old male patient presented with a complaint of neck pain radiating to the left upper limb for 3 months. The pain was localized to the posterior aspect of the left side of the neck, moderate in intensity, pricking type in nature, continuous throughout the day, not aggravated by any particular neck movements, not relieved by any specific neck posture/medications, and gradually began radiating to the left upper limb after 1 week of onset. Two weeks after the onset of pain, he developed rapidly progressive weakness involving the left upper limb. He noticed difficulty in lifting his hand above the shoulder while combing hair, wearing a sweater, lifting heavy objects above head level, and later difficulty in lifting a bucket of water and grasping objects with the left hand. There was no history of similar events in the past.

General examination revealed a conscious, alert, and oriented patient with a pulse rate of 74/min and blood pressure of 112/60 mmHg in the right arm. His body mass index was 20.76 kg/m². On clinical examination, bilateral cervical lymph nodes were enlarged, non-tender, and non-matted. The left-sided paravertebral tenderness was noticed adjacent to the C4 and C5 levels. No obvious spinal deformity could be identified. The patient had weakness in the left upper limb with a power of 2/5 (MRC grade) at the shoulder and elbow joint and 4/5 at the wrist.
joint. Sensory loss to pain and temperature was 50% along with 
C4, C5 dermatomes and biceps, triceps, and supinator reflexes 
were diminished.

Chest X-ray did not reveal any significant pulmonary findings. 
Cervical spine X-ray showed a paravertebral opacity near the 
C4–C5 lateral mass and transverse process region. Magnetic 
resonance imaging (Figs. 1 and 2) was suggestive of an ill-
defined heterogeneous contrast-enhancing lesion predominantly 
involving the upper trunk of the left-sided brachial plexus. 
A rapidly progressive clinical course was suggestive of malignant 
etiology and with the intent of obtaining a pathological diagnosis, 
extoration of the lesion was undertaken.

The left-sided linear paravertebral incision was given 
at the C4–C5 level. Pus was seen beneath the muscle layer at 
~3 cm depth with unhealthy granulation tissue extending up 
to the lamina/transverse process of C4–C5 on the left side. 
Decompression of lesion and evacuation of pus were carried 
out. Adequate decompression was achieved and no breach of the 
thecal sac or its contents was noticed. A frozen section of the tissue 
was suggestive of granulomatous infection. GeneXpert from the 
pus sample was positive. Histopathology revealed necrotumatus 
granulomatous inflammation with a central necrotic core 
surrounded by epithelioid macrophages, a few multinucleated 
giant cells, and lymphocytes. Hence, tuberculous etiology was 
confirmed and malignancy was ruled out.

Based on the frozen section report of granulomatous infection, 
antituberculous treatment was started with a standard regimen as 
per the Revised National TB Control Program guidelines [4,5]. 
Peroral (PO) medications isoniazid (5 mg/kg/day), rifampicin 
(10 mg/kg/day), pyrazinamide (25 mg/kg/day), ethambutol 
(15 mg/kg/day), and injectable streptomycin (15 mg/kg/day IM) 
were given for 2 months. This was followed by a continuation 
phase of isoniazid (5 mg/kg/day PO) and rifampicin 
(10 mg/kg/day PO) for 16 months. The patient showed gradual 
improvement in motor strength, subsidence of neck swelling, 
and radiological resolution of spinal compression. Since clinical 
recovery was maintained at a gradual rate as seen in follow-up 
outpatient department visits, complete antitubercular treatment 
(ATT) for 18 months was carried out. Periodic liver function tests 
were done to rule out the development of ATT-induced hepatitis. 
Pyridoxine supplementation (25 mg/day PO) was given during 
the entire treatment duration to prevent ATT-induced peripheral 
neuropathy. He completed 18 months course of ATT. MR brachial 
plexus at 4-year follow-up showed good resolution of neural 
compression and significant subsidence of the lesion (Fig. 3).

**DISCUSSION**

There are multiple mimickers of this clinicopathological entity 
which we considered in our differential diagnosis. There was 
a clinical possibility of a nerve sheath tumor (schwannoma); 
however, imaging characteristics suggested otherwise. 
Cervicothoracic junction TB, acute brachial plexus neuritis, 
Parsonage-Turner syndrome (PTS), Refsum disease, and 
hypertrophic polyneuritis of Dejerine are some of the likely 
differential diagnoses. Cervicothoracic junction TB may present 
with radiculopathy of the upper limbs [6]. The radiculopathy, 
lower motor neuron weakness in the distal upper limb muscles, and 
resultant deformity respond well to debridement and appropriate 
medical management. Myelopathy is an uncommon feature, in 
spite of extensive bony destruction. Adult and pediatric-onset 
cervical TB is differentiated by a frequent occurrence of paraplegia 
and larger abscesses in the former. Needle electromyography and 
nerve conduction studies usually show evidence of distal axonal 
degeneration. In the absence of radiological indicators, various 
differentials need to be considered.

Acute brachial plexus neuritis causes painful restriction of 
the upper limb movements. Involvement of the brachial plexus 
in various infective diseases has been described [7]. See et al. 
demonstrated a patient with *Streptococcus agalactiae* infection 
causing pyogenic shoulder joint arthritis and subsequent brachial 
plexus neuritis. They postulated an immune reaction to the 
streptococcus antigen as the probable etiological mechanism.
PTS is a similar condition with sudden onset pain and restriction of shoulder movements followed by atrophy of the upper limb muscles. Possible etiology includes trauma, recent immunization, surgery on the brachial plexus region, preceding viral infection, unusual strenuous physical activity, and the predisposing connective tissue or autoimmune disorders. Physiological recovery is seen in 70–90% of patients, while 10–20% have impaired exercise tolerance and persistent pain. The use of intravenous immunoglobulins as a therapeutic tool for PTS is under investigational stages.

Refsum disease and interstitial hypertrophic polyneuritis of Dejerine present a similar clinical picture. Joffroy in 1879 first elucidated neuralgia amyotrophy of the brachial plexus in a patient with sudden onset restriction of shoulder movement [8]. Mononeuritis multiplex with brachial plexus predilection is a condition characterized by sharp stabbing pain, weakness, and atrophy involving the shoulder. Taylor described this entity with a heredofamilial background [9]. The prognosis of neurological recovery in patients with brachial plexus neuritis is good with 75–80% recovery in 2 years and almost complete recovery in 3 years.

In our patient, GeneXpert of the pus sample was positive. It has been studied that GeneXpert can be 100% specific and 90–95% sensitive for diagnosing TB [10]. Histopathology is considered at least 85% specific and 93–94% sensitive as compared to TB-polymerase chain reaction [11]. TB can predispose to neuropathy through various mechanisms vis-à-vis spinal tuberculosis, meningeal TB, peripheral neuropathy due to immune-related mechanisms, and secondary to antitubercular therapy.

Pyridoxine deficiency is common in patients using combinations of pyrazinamide and isoniazid. This usually manifests with loss of proprioception and vibration sense. Hence, we supplement ATT with oral pyridoxine till the entire course is complete. Multidrug antitubercular chemotherapy is the mainstay of treatment. Debridement and surgical stabilization are required only in certain situations such as progressive neurological deficit and worsening mechanical construct stability. The standard ATT regime we follow comprises 2 months of isoniazid (H), rifampicin (R), pyrazinamide (Z), ethambutol (E), and streptomycin (S) followed by HR for 16 months. A Medical Research Council reported similar functional outcomes in patients with spinal TB undergoing chemotherapy, debridement, and/or fusion [12–14]. In our patient, debridement and evacuation of pus enabled early decompression of neural elements.

CONCLUSION

TB is a chronic granulomatous disease that can involve any system or component of the human body. Isolated involvement of the brachial plexus has not been reported in the medical literature. We describe this unique presentation of weakness in the upper limb secondary to tubercular involvement of the upper trunk of the brachial plexus. Debridement and evacuation of pus for relieving symptoms of compression of neurovascular structures must be timely done. A complete course of antitubercular chemotherapy is vital for a good neurological outcome.

REFERENCES

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Sreenivasan et al. Brachial plexus tuberculosis: A unique neurological variant of a common clinical disease


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