PHARYNGEAL-CERVICAL-BRACHIAL VARIANT: AN UNUSUAL PRESENTATION OF GUILLAIN-BARRÉ SYNDROME

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ABSTRACT
Pharyngeal-cervical-brachial (PCB) variant is a rare presentation of Guillain–Barré syndrome (GBS), which is symbolized by oroparhyngeal, cervical and upper limb weakness. It is often wrongly diagnosed as brainstem stroke, myasthenia gravis or botulism. The presence of anti-GT1a and anti-GQ1b ganglioside IgG antibodies and axonal electrophysiological pattern are the hallmark of PCB variant. The treatment is similar to any other variant of GBS. We present here, report of a 20-year-old male, who presented with findings suggestive of PCB variant of GBS but had absent anti-GT1a and anti-GQ1b antibodies and a normal cerebrospinal fluid. He made a remarkable recovery after administration of intravenous immunoglobulins.

KEY WORDS: Guillain–Barré syndrome (MeSH); Pharyngeal-cervical-brachial variant (Non-MeSH); anti-ganglioside antibodies (Non-MeSH); Immunoglobulin G (MeSH); Cerebrospinal Fluid (MeSH); Gangliosides (MeSH); Immunoglobulins, Intravenous (MeSH); Electrophysiology (MeSH).

INTRODUCTION
Guillain–Barré syndrome (GBS) is the most common cause of flaccid paralysis of acute onset worldwide. The pharyngeal-cervical-brachial (PCB) presentation of GBS, is a rare variant that has an incidence rate of 0.07-0.25/100,000. It is described by weakness of facial muscles and upper limbs, dysarthria and loss of muscle stretch reflexes. The triggers for GBS e.g. infection, surgery and inflammation are also shared by its variants, particularly the PCB variant. The electro-diagnostic (EDX) evaluation of the PCB variant generally demonstrates an axonal pattern of polyneuropathy rather than demyelination. As this presentation of GBS is quite rare, the physicians often wrongly diagnose such patients as suffering from a brainstem stroke, botulism or myasthenia gravis. Anti-GT1a and anti-GQ1b antibodies provide supportive evidence for the diagnosis of PCB variant though the former are positive in only 50% of the patients. The early disease may also lack the characteristic cerebrospinal fluid (CSF) or EDX abnormalities. We present here, report of a 20-year-old male, who presented with findings suggestive of the PCB variant of GBS but had absent anti-GT1a and anti-GQ1b antibodies and a normal CSF. He made a remarkable recovery after administration of intravenous immunoglobulins.

CASE SUMMARY
A 20-year-old male presented to us with complaints of fever and productive cough for one month and inability to swallow, diplopia and nasal twang in voice for the last two days. Fever was insidious in onset, documented up to 100°F, and intermittent in character with no diurnal variation. He was unable to swallow both liquid and solid diet and had episodes of nasal regurgitation of food particles. The diplopia was evident on lateral gaze. He also had complaints of progressive weakness involving upper limbs more than lower limbs and difficulty in combing hairs, brushing teeth and writing. There was no history of loose stools.

On examination, he had a blood pressure of 150/100 mmHg with a heart rate of 106 beats/min. He was maintaining oxygen saturation at air without oxygen support. The neurological examination revealed bilateral 6th, 9th, 10th, 11th, and 12th nerves' palsies. The muscle power was 3/5 in upper limbs and 4/5 in lower limbs according to the Medical Research Council Scale. The muscle stretch reflexes were grade 0 in upper limbs and 1+ in lower limbs with absent Babinski reflex. The sensations were intact. He had an ataxic gait; however, the cerebellar signs were absent. Based on history and clinical examination, a provisional diagnosis of PCB variant of GBS was made while keeping brain stem stroke, botulism and myasthenia gravis as differentials.

The laboratory evaluation revealed hemoglobin of 17g/dl and hematocrit of 45%. The CSF evaluation revealed a normal study. Arterial blood gases and the pulmonary function tests were also normal. Rest of the baseline investigations were within normal limits. The magnetic resonance imaging (MRI) of the brain, cervical spine and chest x-rays did not show any abnormality. The EDX evaluation, performed on the third day of admission revealed a normal study. His antiganglioside antibodies profile was negative.
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Based on progressive and persistent weakness, absent muscle stretch reflexes and normal MRI brain and CSF, the patient was treated as a case of PCB variant of GBS. He was managed in the intensive care unit of the hospital. The feeding was started through a nasogastric tube. He received 125 g of IV IG over five days. His symptoms started improving on 6th day of the start of treatment and the nasogastric tube was removed. An EDX assessment was recalled that suggested acute motor axonal neuropathy. The second CSF evaluation revealed a high protein count (albuminocytological dissociation). His follow-up examination after one week showed persistent inability to abduct eyes (Figure-1) while rest of the weakness was fully resolved. He was further advised wide range of motion exercises of the eyeballs. Written informed consent was obtained from the patient for publication of his personal information.

DISCUSSION

The first description of the PCB variant of GBS dates to 1986 when Ropper identified it as an atypical presentation of GBS in three patients who exhibited rapidly progressive weakness of the muscles of oral cavity, pharynx, neck and shoulder. The legs were spared and there were no sensory symptoms.

The above-mentioned clinical features are still the hallmark of the PCB variant. History of a recent infection strongly supports the diagnosis. Investigations that validate the disease include albuminocytological dissociation of CSF, antibodies against GT1a or GQ1b and neuropathy on EDX studies. Albuminocytological dissociation of CSF may possibly be absent in 50% of the patients during first week of the illness. Anti-GT1a antibodies are also nonexistent in half of the patients. The EDX studies in PCB reveal an axonal pattern of the polyneuropathy rather than demyelination. MRI brain and cervical spine are normal in the PCB variant.

The management guidelines for the PCB variant are analogous to other variants of GBS. The do’s and don’ts of IVIG or plasma exchange therapy are similar. In early disease, death may occur due to sepsis, pulmonary embolism or cardiac arrest so all patients should remain admitted in the hospital until no clinical progression of the morbidity is established. Monitoring of bulbar function and respiratory effort is obligatory and directs the corresponding use of nasogastric tube and/or ventilator support. Ventilatory support may be required in patients with inefficient cough, atelectasis, hypoxemia, hypercapnia or vital capacity < 15 ml/kg body weight.

There has been a report of the PCB variant of GBS from Pakistan. He was a 25-year-old male soldier with similar symptoms, normal CSF and axonal neuropathy on EDX evaluation. He remained on mechanical ventilation for 170 days. He received plasmapheresis and had a sub-total recovery after seven months. Our case had a milder course and a rapid recovery probably because of good initial motor power, early treatment and no requirement for the ventilatory support.

In conclusion, recognition of the PCB variant and other atypical cases of GBS is important, as it permits early, effective and lifesaving treatment in patients with a potential of highly morbid or fatal outcome.

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AUTHORS’ CONTRIBUTIONS

Following authors have made substantial contributions to the manuscript as under:

MF & FA: Identified & managed the case, manuscript writing, final approval of the version to be published
SBA: Diagnosis of the case, manuscript writing, final approval of the version to be published

Authors agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

CONFLICT OF INTEREST

Authors declared no conflict of interest

GRANT SUPPORT AND FINANCIAL DISCLOSURE

NIL

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