

Bickerstaff brainstem encephalitis and Guillain-Barré syndrome overlap

Kombinace Bickerstaffovy kmenové encefalitis se syndromem Guillain-Barrého

Dear editors,

Bickerstaff brainstem encephalitis (BBE) is a rare acquired autoimmune inflammatory disorder of the nervous system related to Guillain-Barré syndrome (GBS) with ophthalmoplegia and ataxia. Patients present with hyperreflexia and altered consciousness. Miller Fisher syndrome (MFS) also associated with GBS is clinically characterized by ophthalmoplegia, ataxia and areflexia. Both MFS and BBE are observed after preceding infection and have some similarities with GBS. Since the discovery of GQ1b antibodies in early 1990, both disorders provided evidence that are of the same spectrum – GQ1b syndrome. MFS patients who developed limb weakness are overlapping with GBS or those who developed drowsiness are overlapping with BBE [1].

We report a patient, who developed bilateral ophthalmoplegia and ataxia, with weakness and numbness of upper extremities and later also weakness of lower extremities, accompanied by drowsiness with hyperreflexia. Based on clinical and EMG findings, high positivity of anti-GQ1b antibodies and prompt treatment response to intravenous immunoglobulins, we conclude that the final diagnosis of our patients is BBE overlapping with GBS.

On the January 27, 2022, pain on the right side of the face appeared in a 69-year-old woman who was diagnosed with recurrent maxillary sinusitis. The patient has been treated for mild bronchial asthma since childhood (salbutamol and ciclesonide were given for discomfort). She suffered from recurrent maxillary sinusitis on the right side and has undergone carpal tunnel surgery on the right side in 2016 with good outcomes. The patient was originally a civil servant and is now retired.

After 2 days, clarithromycin *per os* was administered on an outpatient basis. The pa-

tient was also examined in cardiology for subcompensation of arterial hypertension with a blood pressure of 149/104 mm Hg and was given amlodipine with a blood pressure adjustment. A neurological examination revealed gait uncertainty and a mild hypoaesthesia of the hands and feet and a brain CT revealed sinusitis maxillaris on the right side but otherwise was normal. On the following day, the patient presented with diplopia in the upward gaze and worsened stability gradually making walking impossible. Tingling and dullness of the hands and feet were also observed. A follow-up neurological examination and brain CT with angiography were performed and findings on the blood vessels were normal. On the next day (February 2), the patient was admitted to the department of neurology already displaying oculomotor disorder with diplopia, mild dysphagia and accentuation of trunk ataxia while sitting. Ischemic vascular etiology was considered, and acetylsalicylic acid (ASA) and a prophylactic dose of low molecular weight heparin (LMWH) were administered. On the next day, the patient displayed accentuated diplopia, dysphagia and backward and leftward thrusts while sitting. An MRI showed right sided maxillary sinusitis, but other findings were normal. On February 4, an EMG was performed indicating normal compound muscle action potential (CMAP) and sensory nerve action potential (SNAP) amplitudes which were otherwise within normal limits including repetitive stimulation. H-reflex was unobtainable (Tab. 1). The patient was no longer able to raise her head in a supine position and presented with worsened diplopia, preserved direct and consensual photoreactions as well as severely restricted vertical movements of the eyeballs, horizontal slow lateral movements of the eyeballs with a small range, and no convergence but with evident miosis. Reflexes C5–8 were low,

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J. Husárová-Cepková¹, E. Ehler^{1,2}, L. Ungermann³, I. Štětkařová⁴

¹ Department of Neurology, Pardubice Regional Hospital, Czech Republic

² Department of Neurology, Faculty of Health Studies, University of Pardubice, Czech Republic

³ Radiodiagnostic Department, Pardubice Regional Hospital, Czech Republic

⁴ Department of Neurology, Third Faculty of Medicine, Královské Vinohrady University Hospital, Prague, Czech Republic



Assoc. Prof. Edvard Ehler, MD, CSc., FEAN

Department of Neurology
Pardubice Regional Hospital
Kýjevská 44
530 03 Pardubice
Czech Republic
e-mail: eda.ehler@tiscali.cz

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upper extremities sustained for 10 s and impaired scapular fixation was observed with a right-hand grip of 44 and left-hand grip of 42 kPa. Lower extremities were sustained for 10 s and reflexes L2–S2 were brisk with a left-sided predominance, where extensor hypertonia and Babinski occurred. Hautant was observed in a seated position with a backward pull. The patient was somnolent with hypoesthesia on the left side of the

Tab. 1. Nerve conduction studies.

Nerve / Examination date	DML (ms)	A-CMAP (mV)	MCV (m/s)	F-wave lat (m/s)	A-SNAP (uV)	SCV (m/s)
I. median	4.30	8.4	53.3	28.6	7.4	39.0
4. 2. 2022	N (4.30)	(6.5)	(50.0)	(-29)	(-10)	(50.0)
24. 2. 2022	4.25	6.1	50.0	27.0	6.0	40.5
23. 3. 2022	4.60	8.6	51.2	31.1	11.3	40.0
I. ulnar	3.25	7.3	63.3	28.6	15.9	52.8
4. 2. 2022	(3.80)	(6.5)	(-50)	(-29)	(-12)	(-50)
24. 2. 2022	3.20	6.6	50.7	27.4	8.4	36.8
23. 3. 2022	2.95	8.4	46.2	25.2	6.8	35.9
I. peroneal	4.25	8.6	45.9	52.2	9.7	42.6
4. 2. 2022	(5.5)	(3.5)	(-45)	(-52)	(-9)	(-45)
24. 2. 2022	4.90	1.9	44.7	49.3	2.6	42.3
23. 3. 2022	4.15	4.5	48.2	55.3	3.8	40.6

A-CMAP – amplitude of compound muscle action potential; A-SNAP – amplitude of sensory nerve action potential; DML – distal motor latency; MCV – motor conduction velocity; SCV – sensory nerve conduction velocity

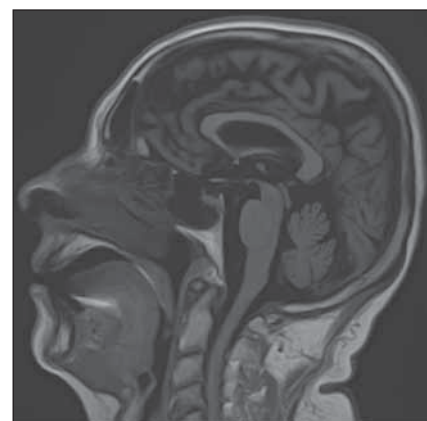


Fig. 1. Brain MRI. Sagittal T1 weighted image – normal structures of brainstem and corpus callosum.

Obr. 1. MR mozku. Sagitální T1 vážený snímek – normální struktury mozkové kmene a corpus callosum.

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face, acral to the hands and in the lower extremities from the mid-tibia distally. Findings were suggestive of an autoimmune etiology (BBE + GBS) and intravenous immunoglobulin G 2 g/kg was administered for 5 days. A lumbar puncture with cerebrospinal fluid examination was performed showing a nearly normal level of protein (0.56 g/L) and normal cytology.

During the following days, a progression of fatigue and decline were observed in the patient with further weakening of muscles. The patient became unable to hold out her arms and a gradual progression of respiratory insufficiency ensued. On day 6 of hospitalization, the patient was intubated with full artificial ventilation. On the 9th day of hospitalization, we received laboratory results with an anti-GQ1b level of 419% and no antibodies to other gangliosides were detected.

To improve mobility, weaning with intermittent connection to a humidifier began from February 20 and artificial ventilation was stopped on the following day. After two days, food intake per os began. On February 24, an EMG checkup examination was performed revealing lower CMAP amplitudes, a non-

elicitable H-reflex, normal blink reflex and a normal brainstem auditory evoked potential (BAEP). An MRI of the brain showed no evidence of a new lesion in the trunk, cerebellum, or corpus callosum (Fig. 1). On March 2, the patient was already alert, lifting and holding her head up and swallowing slowly but safely. The convergence of the eyeballs and the movement of the left eyeball upwards and sideways with the manifestation of diplopia was slightly restricted while walking was already possible with the help of two trekking sticks. A vibratory sensation was not present in the limbs but was adequate in the trunk with a measure of -6/8. A checkup EMG on March 23 showed no evidence of denervation syndrome in the lower extremities.

Bickerstaff brainstem encephalitis is a rare diagnosis, but with a clear diagnostic definition. Combination of BBE with a generalized form of GBS is a typical situation in most BBE patients. BBE is very rare in Europe and clinical characteristics are well defined. Combination of BBE and GBS is a presentation of the continuing spectrum of anti-GQ1b syndrome. These patients do not have a poor prognosis.