RESEARCH ARTICLE

ONE CASE OF REFRACTORY STATUS EPILEPTICUS AND AUTOIMMUNE ENCEPHALITIS.

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Abstract

Purpose: Encephalitis is a severe inflammatory disorder of the brain with many possible causes and with difficult differentiate of diagnosis. Existing criteria for autoimmune encephalitis are too reliant on antibody testing and response to immunotherapy, which might delay the diagnosis. Several particular features raise the suspicion of an autoimmune cause in otherwise unexplained seizure disorders. One of the most important challenges is the elucidation of the causes of seizure disorders.

We presented one case of unexplained coma and refractory status epilepticus. Patient, 20 years old, man, was admitted in hospital with coma, stable hemodynamics, on mechanical ventilation. On MRI research was discovered existence of inflammatory infiltration. At primary stage, EEG discovered polymorphic dysrhythmia, bilateral activity, than plural peak sharp-slow wave paroxysms and status epilepticus. It was conducted combined treatment with anticonvulsive drugs, Immunoglobulin, pulse corticosteroid therapy with methylprednisolone and plasma exchange therapy.

Conclusion: An autoimmune cause was suspected based on frequent or medically intractable seizures and the presence of neural antibody, inflammatory changes indicated in spinal fluid and on MRI. Patients had abnormal findings on brain MRI, with extensive multifocal or diffuse cortical and subcortical involvement. EEG showed generalized periodic discharges and status epilepticus. Within investigation of IgG autoantibodies against proteins on the surfaces of neurons, we received difference result from two different laboratory, but refractory status epilepticus, radiological changes, EEG pattern and progress of illness has indicated autoimmune character of process.

Introduction:

Encephalitis is a severe inflammatory disorder of the brain with many possible causes and with difficult differentiate of diagnosis. Existing criteria for autoimmune encephalitis are too reliant on antibody testing and response to immunotherapy, which might delay the diagnosis.
Several particular features raise the suspicion of an autoimmune cause in otherwise unexplained seizure disorders.

One of the most important challenges is the elucidation of the causes of seizure disorders. IgG autoantibodies against proteins on the surfaces of neurons were identified as markers and pathogens in autoimmune encephalitides that are accompanied by repetitive focal seizures. The antibodies against surface antigens contribute directly to the disease processes. In particular, Anti-NMDA receptor encephalitis is a disease occurring when antibodies produced by the body’s own immune system attack NMDA receptors in the brain. In the absence of a reliable biomarker, it is difficult to gain systematic and general insights into these patients.

Because autoantibody test results and response to therapy are not available at disease onset, treatment is based on the initial diagnostic approach on neurological assessment and conventional tests that are accessible to most clinicians.

We presented one case of unexplained coma and refractory status epilepticus.

**Case report:**

Patient, 20 yars old. men, Caucasian was admitted in hospital at 06.07.2017. with coma, stable hemodynamics, on mechanical ventilation (patient was transferred from another clinic) CT of brain could not discovered any pathological changing. MRI within admission (07.07.2017) was without acute pathological intracerebral changing, EEG - discovered polymorphic dysrhythmia, bilateral activity (Fig. 1).

**Fig 1:** The patient was examined on viruses, parasites, bacteria.

<table>
<thead>
<tr>
<th>Borrelia burgdorferi</th>
<th>IgM-30u/ml, IgG-negative.</th>
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<tbody>
<tr>
<td>Mycobacterium tuberculosis</td>
<td>PCR in CSF -- negative.</td>
</tr>
<tr>
<td>Criptococcus Ag quantitative analysis</td>
<td>negative.</td>
</tr>
<tr>
<td>Brucella</td>
<td>IgM-10lu/ml, IgG - negative</td>
</tr>
<tr>
<td>Leptospira</td>
<td>IgM, IgG -negative</td>
</tr>
<tr>
<td>Blood test of gama interferon (ESAT-6, CFP-10, TB7.7)</td>
<td>negative</td>
</tr>
<tr>
<td>Measles IgG ab - 9600IU/L</td>
<td>N&gt;350, immunity confirmed</td>
</tr>
<tr>
<td>Measles-Virus – Ak. IgM negative</td>
<td>Normal range -negative</td>
</tr>
<tr>
<td>Tick borne encephalitis virus</td>
<td>(NORM 0.8-1.1)</td>
</tr>
<tr>
<td>0.7</td>
<td>Ig M</td>
</tr>
<tr>
<td>0.1</td>
<td>Ig G</td>
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Patient CSF and serum was examined on autoantibodies against proteins on the surfaces of neurons, in particular for diagnosing of NMDA receptor antibody encephalitis. CSF was sent to the lab at 7.07.17.

Treatment was started empirical, with antiviral (Acyclovir) and antibacterial drugs. After 6 day from the beginning of investigation, autoantibody test result was positive NMDAR AAB in CSF was 1:2, in serum 1:160. (serum and CSF was investigated in Heidelberg laboratory of Limbach)

After 4 day EEG revealed status epilepticus (Fig.2)
Fig.2: On repeated MRI research (1.07.17) was discovered existence of inflammatory infiltration (encephalitis) in right frontal and parietal lobes and around of hippocampus, intracranial arteriovenous hyperemia. (Fig.3).

Fig.3: For status epilepticus was initiated treatment with valproic acid (Depakin), Levetiracetam (Epixx) and carbamazepin. Regardless of this drugs EEG at 17.07 revealed plural Peak sharp-slow wave paroxysms (Fig.4)
After detection of NMDAR AAB in CSF, was started immunoglobulin therapy (0.4g/kg/day for 5 day). Treatment was lasted with peroral intake of Epixx and Depakin, intravenous infusion of levetiracetam (Keppra), Valproate sodium (Depakin) and intravenous infusion of propofol (4mg/kg/day), pulse corticosteroid therapy with methylprednizolone. Propofol infusion rate and maximum dosage was selected by considering to avoid of propofol infusion syndrome. We periodically were testing the concentration of anti-conventional drugs in the blood and the dosage was adjusted accordingly.

Additionally was used thiopental infusion and ketamin infusion with dosage 2.75mg/kg.h(3 day). But the convulsion on EEG was continued.

Epi pattern (27.07.17) had negative dynamics, increased amplitude of epileptic form activities (Fig. 5)

Plasma exchange (PE) was used as one of the treatment modalities in addition to immunosuppressive therapy, including corticosteroids, intravenous immunoglobulin (IVIG). Regardless of such combined and multilateral treatment, EEG (31.07.17) detected negative electrophysiological dynamics (Fig.6), MRI revealed negative radiologic changes (Fig.6)
Fig. 6: In large hemispheres, hyperintensiveness of subarachnoid spaces was reduced. However after contrasting, the focal point of contrast substance was much more manifested in parahippocampus and fragmentately in the area of the membranes (more left).

Fig. 7: For maximal captured and decreasion of brain bioactivity was used thiopental with high dose infusion. We achieved absolute decreasion of brain bioactivity and full resolvation of convulsion (8.08.17).
After 48 h. from cessation of thiopental infusion, we received normal EEG and later improvement of MRI parameters. Fig. 8

Fig. 8: Patient CSF and serum were sent to The Austrian Clinical Institute of Neurology.

a) This institute investigated serum and cerebrospinal fluid of our patient for presence of anti-neuronal antibodies.

b) In this laboratory screening for surface receptor antibodies (avidin-biotin-peroxidase technique; rat brain; IIFT Euroimmun) was negative for antibodies such as NMDAR, LGI1, CASPR2, AMPAR, GABA(B)R, and DPPX. A cell-based assay was negative for glycine-receptor antibodies (in-house; GlyR alpha1).

Institute has noticed that negative test result does not exclude autoimmune logically mediated syndromes of other origin.
Later, after 36 days from admission, the patient's state became stable, and the patient was weaned from artificial ventilation. The level of consciousness had gradually improved and finally, the patient was discharged from the hospital.

**Discussing:**

We presented drug-resistant status epilepticus that persisted following the continuous administration of intravenous anesthetics and anticonvulsant drugs for more than several days.

Acute onset, resistance to anti-seizure medication, evidence of CNS inflammation was clinical features suggestive of autoimmune epilepsy.

For treatment, we used first-line therapies—steroids, intravenous immune globulin, plasma exchange therapy, combinations of these interventions. Anticonvulsant drugs, valproate sodium, levetiracetam, midazolam, propofol infusion and some drugs (carbamazepin, depakinetix) intake from gastrointestinal route, was not effective.

Many researchers have reported that, during prolonged seizures, the number of activated GABA-A receptors on the postsynaptic membrane gradually decreases, whereas the number of inactive GABA-A receptors increases. These changes cause a significant reduction in the efficacy of antiepileptic drugs (AEDs). Recent studies have found that, during prolonged seizures, when the numbers and activities of GABA receptors gradually decrease, simultaneously, the numbers and activities of glutamatergic NMDA receptors increase. Thus, the commonly used first-line and second-line AEDs gradually fail, often causing RSE and thus providing the possibility of the use of ketamine to treat RSE. Ketamine is a noncompetitive NMDA receptor antagonist that might play a role in treating SE by blocking NMDA receptor-mediated glutamatergic transmission. But after ketamine infusion, its anticonvulsant effects have not been reached. For maximal capture and decrease of brain bioactivity, we used thiopental with high dose (400mg/h) infusion. We achieved absolute decrease of brain bioactivity and full resolution of convulsion.

**Conclusion:**

An autoimmune cause was suspected based on frequent or medically intractable seizures and the presence of neural antibody, inflammatory changes indicated in spinal fluid and on MRI. Patients had abnormal findings on brain MRI, with extensive multifocal or diffuse cortical and subcortical involvement. EEG showed generalized periodic discharges and status epilepticus. Within investigation of IgG autoantibodies against proteins on the surfaces of neurons, we received different result from two different laboratory, but refractory status epilepticus (Resistance to anti-seizure medication), radiological changes, EEG pattern and progress of illness has indicated autoimmune character of process.

**References:**

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