Diagnostic pitfalls in a young Romanian ranger with an acute psychotic episode

Abstract: The identification and distinction of the pathological conditions underlying acute psychosis are often challenging. We present the case of a 35-year-old ranger who had no history of acute or chronic infectious disease or any previous neuropsychiatric symptoms. He arrived at the Psychiatry Clinic and was admitted as an emergency case, displaying bizarre behavior, hallucinations, paranoid ideation, and delusional faults. These symptoms had first appeared 7 days earlier. An objective examination revealed abnormalities of behavior, anxiety, visual hallucinations, choreiform, and tic-like facial movements. After the administration of neuroleptic and antidepressant treatment, he showed an initial improvement, but on day 10 entered into a severe catatonic state with signs of meningeal irritation and was transferred to the intensive care unit. An electroencephalogram showed diffuse irritative changes, raising the possibility of encephalitis. Taking into consideration the overt occupational risk, Borrelia antibody tests were prescribed and highly positive immunoglobulin (Ig)M and IgG titers were obtained from serum, along with IgG and antibody index positivity in cerebrospinal fluid. In parallel, anti-N-methyl-D-aspartate receptor antibodies and a whole battery of other autoimmune encephalitis markers showed negative. A complex program of treatment was applied, including antibiotics, beginning with cefazidime and ciprofloxacin – for suspected aspiration bronchopneumonia – and thereafter with ceftiraxone. A gradual improvement was noticed and the treatment continued at the Infectious Disease Clinic. Finally, the patient was discharged with a doxycycline, antidepressant, and anxiolytic maintenance treatment. On his first and second control (days 44 and 122 from the disease onset), the patient was stable with no major complaints, Borrelia seropositivity was confirmed both for IgM and IgG while the cerebrospinal fluid also showed reactivity for IgG on immunoblot. On the basis of the putative occupational risk, acute psychotic episode, and the success of antibiotic therapy, we registered this case as a late neuroborreliosis with atypical appearance.

Keywords: Borrelia burgdorferi, neuroborreliosis, neuropsychiatric symptoms, encephalitis, anti-NMDAR

Introduction
Neuropsychiatric manifestations characterize the disease course of ~15% of Borrelia-infected individuals.1 The involvement of the central nervous system is rare, but encephalitis or encephalomyelitis may appear both in acute and late neuroborreliosis (LNB).2 Chronic or LNB as a tertiary disease stage is a well-known entity; however, the underlying pathomechanisms are the subject of intense scientific debate. Supporters of the pathogen persistence theory claim successful cultivation of Borrelia burgdorferi and/or isolation of its DNA from leptomeningeal infiltrates, subpial, or subependimal lesions.1,3 Regarding its histological forms, LNB appears either as an infiltrative or an atrophic meningoencephalitis.1 Overt psychiatric manifestations of encephalitis...
of LNB as acute psychosis, hallucinations, paranoia, or obsessive compulsive disorder are not typical, but have been reported in the literature. Sometimes, psychiatric symptoms may be the only evident disorder in neuroborreliosis. Muscular hallucinations with sudden onset may appear without hearing loss. Body schema disturbances, metamorphosia, and auditory hallucinations were described in a 7-year-old boy without any neurological signs.

According to the European Federation of Neurological Societies’ guidelines, in order to confirm neuroborreliosis, three criteria (neurological symptoms, cerebrospinal fluid [CSF] pleocytosis, and specific intrathecal antibodies) must be present, and to raise the suspicion of neuroborreliosis, two of these conditions should be present.

The 2013 European Concerted Action on Lyme Borreliosis diagnostic guidelines (www.eucalb.com) determine that the common CSF findings of LNB are intrathecal antibody production and presence of oligoclonal immunoglobulin (Ig)G bands, along with the impairment of the blood–brain barrier.

Here, we present the case of a young adult male patient who was admitted to the Psychiatric Clinic with symptoms of acute psychosis: paranoic delusions, hallucinations, and transient confusional states. He presented advanced symptoms of an encephalitis-like syndrome and catatonia and was sent to the intensive care unit (ICU), where anti-Borrelia serum and central nervous system-specific antibodies were determined. Antibiotic treatment was applied and the patient was transferred to the Infectious Disease Clinic, where his condition slowly began to improve.

Written consent was obtained from the patient to accept the diagnostic and therapeutic procedures, and processing of his medical data for scientific purposes. Approval was received from the Ethics Committee of the Mures Clinical County Hospital (no. 20305/09.12.2015).

Case report

A previously healthy 35-year-old male was referred to our emergency room following a 7-day history of disturbed behavior, persecutory type paranoid delusions, visual hallucinations, sleeplessness, agitation, and transient confusional states. The family and his employer reported that in this period he could not fulfill his daily tasks, either at his workplace or in the family, and he seemed on two or three occasions “disconnected” from reality for short periods of time. On examination, he presented bizarre behavior, tic-like orofacial movements, choreiform movements of the arms, bradypsychia, persecutory type paranoid delusional thoughts, suspiciousness, visual hallucinations, flat affect, psychomotor anxiety, insomnia, and social withdrawal with poor insight. Additionally, he showed reluctance to stay in hospital for further investigations and treatment.

Neuroleptic (4 mg risperidonom) and antidepressant (30 mg mirtazapinum) treatment was prescribed. The next day, he became logorhheic and 1,000 mg of sodium valproate was added to the treatment program. Routine laboratory workup for biochemical parameters and complete blood count along with the cranial computed tomography were normal.

On the fourth day of hospitalization, he presented slight muscle rigidity of the upper limbs (then thought to be caused by neuroleptics) for which 6 mg of trihexyphenidyl hydrochloride was added to his treatment.

After a gradual improvement, on the sixth day, the patient presented a short (a few minutes long) transitory confusional state (confusing the medical personnel with his parents).

On the ninth day, he was complaining of dizziness, postural imbalance, and increasing fatigue. A cranial native computed tomography scan showed mild cerebral edema and manitol infusion was administered. The same day, his condition worsened and he entered into a catatonic state. Shortly after, he presented an epileptic seizure. Neurological examination found no focal signs and no pathological reflexes. On the tenth day, the patient re-entered a state of catatonic stupor interrupted with short periods of catatonic agitation. His general condition worsened showing nuchal rigidity, mydriasis, bilateral Babinsky sign, vomiting, and after aspiration of gastric fluid, he was transferred to the ICU with a 92% oxygen saturation value. Laboratory testing showed a mild elevation of the white blood cell and neutrophil count (11.96×10^9/L and 9.68×10^9/L). Leukocytosis and neutrophilia increased further (15.97×10^9/L and 12.55×10^9/L), being accompanied by monocytosis (2.15×10^9/L). Neurological examination did not disclose any focal signs but electroencephalogram findings raised the suspicion of encephalitis (Figure 1A–C). The infectious disease specialist could not find any clinical evidence of ongoing infectious disease.

A blood test for Borrelia-specific antibodies was ordered taking into account that the patient was a ranger, although he denied he had had tick-bites or obvious erythema migrans previously. Borrelia antibodies were determined by an automated recombinant antigen mixture-based enzyme-linked immunosorbent assay approach applying MIKROGENs’ recomWell Borrelia IgG/IgM diagnostic kits (Neuried, Germany). Both serum IgM and IgG were positive (26.6 and 81.2 U, respectively). On the 13th day in the ICU, a lumbar puncture was performed. The CSF showed slightly elevated glucose (81 mg/dL), but normal protein level and
Figure 1 (A) First EEG pattern of the patient: atypical delta wave complexes on our patient’s EEG. Left postero-temporal focal lesion with polymorph delta waves (1 – shaded). (B) Second EEG pattern of the patient: alpha-basic rhythm with 8–9 cycles per second and with diffuse irritative changes, unmodified during eye opening, with asymmetric slow sharp wave discharges in the C-T region bilaterally (1 and 2 – shaded). (C) Normal EEG pattern for comparison: a basic activity with subalpha-theta waves, without pathological changes and asymmetry. (A and B) Registrations with time constant 0.10 s, high frequency filter 30.0 Hz, notch filter: yes, sensitivity: 5.0 μV/mm. (C) Registration with time constant 0.10 s, high pass filter 5.30 Hz, low pass filter 20.0 Hz, notch filter: yes.

Abbreviation: EEG, electroencephalogram.
LDH activity. CSF presented a few red blood cells, but no pleocytosis, with leucocytes being completely absent.

Paired serum and CSF samples confirmed Borrelia sensu lato serum IgM and IgG positivity (39.7 and 116.2 U) and, also, a positive IgG in CSF (24 U). The antibody index (AI) for IgG was calculated with MIKROGENs’ software and found to be positive: 2.83. Contrast computed tomography scan of the brain performed on the same day was normal.

A day later, immunoblot assay was performed to confirm Borrelia infection by applying Mikrogen’s RecomLine Borrelia IgG/IgM kit: this gave a clearly positive result for serum IgM (8 points, with Osp C reactivities for B. burgdorferi, Borrelia afzelii, Borrelia garinii, and Borrelia spielmanii) and IgG (25 points, with reactivities for p100, p58, p39, p41, variable major protein-like sequence (VlsE), and p18) but it was negative for CSF IgM and IgG (tests performed only with high-dilution, 1:20, due to the limited volume of CSF). A 16s rRNA PCR was performed on CSF at the Institute of Clinical Microbiology, Szeged, Hungary, and a negative result was obtained. After interdisciplinary consultations, the suspicion of autoimmune encephalitis was raised, and a comprehensive autoantibody test panel was performed for CSF at the Institute of Immunology, University of Pécs (Hungary) including anti-N-methyl-D-aspartate receptor (anti-NMDAR), anti-γ-aminobutyric acid B receptor (anti-GABA-BR), anti-α-amino-3-hydroxy-5-methyl-4 isoxazolepropionic acid receptor 1 (anti-AMPA1) and 2 (AMPA2), anti-leucin-rich glioma-inactivated 1 (anti-LGI1), and anti-contactin-associated protein-like 2. The autoantibody detection method was an indirect immunofluorescence assay (Euroimmun Gmbh, Lübeck, Germany), with antibody detection method was an indirect immunofluorescence assay. The repeated Borrelia enzyme-linked immunosorbent assay tests of paired serum and CSF samples confirmed serum IgM and IgG (tests performed only with high-dilution, 1:20, due to the limited volume of CSF). A 16s rRNA PCR was performed on CSF at the Institute of Immunology, University of Pécs (Hungary) including anti-N-methyl-D-aspartate receptor (anti-NMDAR), anti-γ-aminobutyric acid B receptor (anti-GABA-BR), anti-α-amino-3-hydroxy-5-methyl-4 isoxazolepropionic acid receptor 1 (anti-AMPA1) and 2 (AMPA2), anti-leucin-rich glioma-inactivated 1 (anti-LGI1), and anti-contactin-associated protein-like 2. The autoantibody detection method was an indirect immunofluorescence assay (Euroimmun Gmbh, Lübeck, Germany), with antibody detection method was an indirect immunofluorescence assay. The repeated Borrelia enzyme-linked immunosorbent assay tests of paired serum and CSF samples confirmed serum IgM and IgG positivity, along with CSF IgG positivity. This time, the CSF Borrelia IgM and IgG immunoblots could be repeated with low dilutions (1:2 for IgM and 1:4 for IgG) and were positive for IgG (p100, VlsE, B. afzelii p18, and lesser reactions for p58 and p39), but not in the case of IgM. The patient was monitored for 24 hours and discharged with recommendation for antidepressant and anxiolytic treatment. On the occasion of the second control visit on the 122nd day, there was no change in his serological status according to the Western blot analysis. The patient showed a good general condition, coherent thinking and speaking, with the only complaint of right arm paresthesia, but without any other pathological findings on clinical examination. In summary, the outcome of the disease was very favorable, the patient being able even to drive his car. He was discharged with low-dose alprazolam and mirtazapine maintenance treatment.
Diagnostic pitfalls in a ranger with acute psychosis

**Discussion**

The intrathecal presence of Borrelia-specific antibodies is the immunological hallmark of neuroborreliosis. The diagnosis of acute neuroborreliosis is partially clinical, being supported by the following CSF findings: elevated leukocyte count, blood–brain barrier dysfunction, elevated IgM levels, and AI for IgM and IgG. In a study performed on an LNB cohort, AI for IgM and IgG was elevated in ~70% of the patients, being more prominent in the subgroups presenting polyradiculoneuritis and isolated facial palsy than in the meningitis subgroup.13

Very few cases with an organic psychotic syndrome and lacking peripheral neurologic signs of LNB are mentioned in literature.4,5 In a similar case, a 31-year-old woman with acute psychosis and high-titer Borrelia seropositivity, but no history of tick-bite, reacted promptly to a combined ceftriaxone and doxycycline treatment.6 Another patient, with an initially untreated meningopolyradiculitis, evolved into an acute schizophrenia-like psychosis, and recovered after 1 week of ceftriaxone treatment.11 However, overt or putative LNB is sometimes difficult to differentiate from primary psychiatric disorders, such as depression or schizophrenia-like psychosis.14

Chronic or late-phase neuroborreliosis is characterized by the presence of IgG type antibodies and a positive CSF/serum IgG AI. The risk of negative serological and intrathecal findings is low at the time of initial clinical presentation. Kaiser15 reported a 49% prevalence of serum, 35% of CSF IgM type, 81% of serum, and 90% of CSF IgG type antibodies in blot and enzyme immunoassay results from patients with acute or chronic neuroborreliosis, applying antigens from three different Borrelia strains: *B. burgdorferi*, *B. afzelii*, and *B. garinii*.

According to Ljostad et al,16 the diagnostic sensitivity of AI is 100% when symptoms have a duration of at least 6 weeks, but it is not optimal within the first 6 weeks of progression. In patients with a negative pretreatment, AI generally remains with a negative finding at follow-up.

It should be noted that our patient presented a serum antibody profile specific for different Borrelia strains (p39, p100 – *B. afzelii*, p58 – *B. garinii*, ViSE, p18-mixtures of strains). He also possessed an increased IgG AI and, at the second CSF testing, Borrelia antibody-specific Western blot proved the presence of anti-p100, anti-p18, and anti-ViSE IgG type antibodies. In the absence of pleocytosis and
typical peripheral neurological signs – but with the presence of psychosis, cognitive disorders, and a positive IgG AI – the suspicion of neuroborreliosis is real, but according to a recently proposed diagnostic algorithm, it indicates a previous central nervous system infection, not an active disease stage. However, in our case, the clinically successful antibiotic therapy is supportive for an ongoing, active Lyme disease with a putative neurological involvement.

In Europe, the most common neuroborreliosis-causing species is *B. garinii*. Clinical diagnosis is more specific in the case of *B. garinii* infection, but difficult and often missed when *B. afzelii* is the causative agent. In Europe, associated inflammatory response is more commonly met: meningitis, meningoradiculitis (Bannwarth syndrome), peripheral neuropathy, and (more rarely) encephalitis and encephalomyelitis being described. Our patient possessed IgG type antibodies against *B. afzelii* (p100, p18, p39) and *B. garinii* (p58) and common epitopes (VisE) both in serum and CSF, and also IgM type antibodies against Osp C of *B. burgdorferi*, *B. afzelii*, *B. garinii*, and *B. spielmanii* in serum: a condition which raises the possibility of repeated infections with different *Borrelia* species. The probability of reinfection is the subject of intensive debate, but in our case, where the professional exposure was real, it has to be taken into consideration.

In a previous study, tests applying recombinant *Borrelia* antigens were shown to have the best sensitivity and specificity. The frequency of antibody specificities were described as follows: VisE (100%), Osp C (79%), p18 (52%), and p58 (50%). In this study, Osp C positive samples were also characteristically reactive for two or more *Borrelia* species.

Taking into consideration the suggested clinical picture, anti-NMDAR encephalitis had to be excluded as an important condition for differential diagnosis.

Anti-NMDAR antibodies were also observed in demyelinating encephalomyelitis and neuromyelitis optica disorders. Anti-NMDAR encephalitis is often misclassified as catatonic schizophrenia, but neurometabolic investigations may elucidate the hypo-glutaminergic status. Availability of NMDAR testing in different Eastern-European countries and medical centers is highly variable. Our patient presented negative anti-NMDAR serum and CSF tests, and also an absence of other neuromune autoantibodies (CASP2, GABA-2R, AMPA1, AMPA2, and LGI1).

CSF analysis in anti-NMDAR encephalitis presents variable findings. Lymphocytic pleocytosis is characteristic for 80%, with total protein increase in ~33% and oligoclonal bands in 25% of the cases. These conditions were lacking in our case. In the absence of prodromal signs, a typical clinical picture, CSF pleocytosis, and body temperature changes, infectious encephalitis was not considered as an underlying pathology.

Acute encephalopathy with catatonic manifestation of chronic neuroborreliosis has very rarely been documented before. Autoimmune reactions based on molecular mimicry and *Borrelia* glycolipid induction of the proinflammatory cytokines interleukin (IL)-6, IL-8, and IL-12 as well as chemokines, such as chemokine (C-X-C motif) ligand 12 and 13 (CXCL12 and CXCL13), may be responsible for these scenarios.

Prolonged, up to 26–52-week antibiotic therapy, was proposed to improve cognitive functions, fatigue, and myalgia in patients with neuroborreliosis. However, in this case, antibiotic therapy of conventional duration applied in conformity with the European Concerted Action on Lyme Borreliosis recommendations proved to be efficient.

**Conclusion**

When occupational risk factors emerge in patients with acute neuropsychiatric disease, the possibility of Lyme neuroborreliosis should be considered even in the absence of a tick-bite history and lack of previous erythema migrans. Taking into consideration the limited diagnostic value of serology and molecular biology, and the limited availability of CSF for repeated analysis, a differential diagnosis approach in the suspicion of neuroborreliosis sometimes has to rely on elimination strategies. Analyzing the possibilities in our patient – based on the clinical picture and laboratory findings – we interpret the case as either an LNB with atypical neuropsychiatric appearance or LNB associated with a neuropsychiatric disease onset. It should be noted that this classification is supported by the significant and sustainable clinical improvement obtained by conventional antibiotic therapy.

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**Author contributions**

EEN analyzed and interpreted the laboratory data and drafted the article. AR provided a substantial proportion of clinical data and contributed to the study concept. EU, GT, and TB
contributed to data acquisition, data analysis, and study concept. EH and AG performed the critical revision of the manuscript. IK-Z elaborated the concept of the case presentation and provided clinical data. All authors contributed toward data acquisition, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

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