Subacute cognitive impairment, hyponatremia and mesial temporal lobe lesions
A typical presentation of voltage-gated potassium channel (VGKC) antibody-associated limbic encephalitis

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For many years, several reports have called attention for non-viral limbic encephalitis (LE), particularly with paraneoplastic etiology. However, an increasing number of data have demonstrated that some of those disorders are immune-mediated with antibodies to voltage-gated potassium channels (VGKC)\(^1\).

Herein, we present a case of VGKC antibody-associated LE and discuss the main phenotypic and imaging features of this unusual disease.

CASE
A 50-year-old man presented to our hospital with three months history of progressive personality change, sleep disturbance, memory impairment, visual hallucinations and seizures. Neurological examination showed disorientation, cognitive impairment (Mini Mental State Examination: 13) and memory dysfunction. Blood tests showed hyponatremia - 130 mmol/L (135-145 mmol/L). Thyroid function, antibodies and serologic tests were normal. Cerebrospinal fluid showed high level of proteins (558 mg/dL) and mild lymphocytic pleocytosis. Brain magnetic resonance imaging (MRI) disclosed bilateral T2 and FLAIR hyperintensity of mesial temporal lobes (Figure). Cancer investigation resulted negative. VGKC serum level was higher than 650 pmol/L (normal range less than 450 pmol/L). These findings confirmed the diagnosis LE with VGKC antibodies. Methylprednisolone was started, with partial improvement. Three weeks later, immunoglobulin 0.4 g/Kg daily for five consecutive days was initiated. One week after immunoglob-
ulin therapy, the patient presented a clear improvement and achieved a MMSE score of 26.

**DISCUSSION**

Recent data have described LE caused by antibodies to VGKC, a specific cell-membrane antigen\(^1,2\). Other antibodies related to LE include NMDA, AMPAR and GABA\(^3,4\). Although LE is frequently related to auto-immune etiology, a paraneoplastic syndrome is found in 30% of the cases\(^3\). When hyperhidrosis, autonomic dysfunction, psychiatric symptoms and neuromyotonic discharges are present, Morvan syndrome must be suspected\(^5\).

Although often unrecognized, the neurological spectrum of VGKC antibody-associated LE includes seizures and neuropsychiatric symptoms, ranging from alterations in memory, behavior, and cognition\(^1\). Neuroimaging shows bilateral hyperintense sign in mesial temporal lobe\(^1\). Imaging differential diagnosis includes: herpetic and paraneoplastic encephalitis, status epilepticus, other immunomediated encephalitis (e.g., anti-NMDA, anti-VGKC) and Alzheimer disease due to the S170F mutation in the presenilin 1 (PS-1) gene\(^2\). Hypo-natremia occurred in 60% of patients and might be attributed to the expression of leucine-rich, glioma-inactivated 1 (LGI1) in the hypothalamus\(^2\).

Recently, Lai et al. described that LGI1 is the autoantigen associated with LE previously attributed to VGKC antibodies. Several patients with the typical syndrome presented negative VGKC and positive LGI1 antibodies\(^2\). The majority of the patients favorably respond to immunotherapy. Administration of human immunoglobulin or plasma exchange is the first line treatment. Methylprednisolone can also be effective. Several patients required chronic therapy on lower dose steroids, with or without azathioprine\(^5\).

**REFERENCES**


**DECLÍNIO COGNITIVO SUBAGUDO, HIPONA TREMIA E LESÕES TEMPORAIS MESIAIS: UMA APRESENTAÇÃO TÍPICA DE ENCEFALITE LÍMBICA ASSOCIADA A ANTICORPOS CONTRA CANAIS DE POTÁSSIO VOLTAGEM DEPENDENTES (VGKC)**

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