Autoimmune Encephalitis—Antibody Targets and Their Potential Pathogenicity in Immunotherapy-responsive Syndromes

Katarzyna D Bera, DPhil,1 Angela Vincent, FRCPath, FRS2 and Sarosh R Irani, DPhil, MRCP (Neurol)3

1. Medical Student; 2. Professor; 3. Clinical Fellow, Nuffield Department of Clinical Neurosciences, University of Oxford, UK

Abstract
Autoimmune encephalitis (AIE) associated with neural autoantibodies is increasingly recognized as a cause of subacute onset amnesia, confusion, and seizures. In the past decade, several key antibody targets have been identified in AIE. These include the N-methyl D-aspartate (NMDA) receptors, voltage-gated potassium channel complexes—in particular leucine-rich glioma inactivated 1 (LG1) and glutamic acid decarboxylase (GAD). There is accumulating clinical and laboratory evidence that antibodies targeting the extracellular domains of cell-surface molecules are directly pathogenic. Each antibody target associates with a spectrum of clinical features and relative response to immunotherapies. These immunotherapies have been shown to improve short- and long-term clinical outcomes in affected patients. AIE is an important differential diagnosis to consider in patients presenting with symptoms of encephalitis as early diagnosis can lead to successful treatment.

Keywords
Autoimmune encephalitis, N-methyl D-aspartate receptor, VGKC complex, LG1, CASPR2, immunotherapy, autoantibody

Disclosure: Katarzyna D Bera, DPhil, has no conflicts of interest to declare. Angela Vincent, FRCPath, FRS, and the Department of Clinical Neurology in Oxford receive royalties and payments for antibody assays and Angela Vincent is the named inventor on patent application WO/2010/046716 entitled 'Neurological Autoimmune Disorders'. The patent has been licensed to Euroimmun AG for the development of assays for LG1 and other VGKC-complex antibodies. Sarosh R Irani, DPhil, MRCP (Neurol), is a co-inventor and may also receive future royalties.

Received: February 23, 2013 Accepted: March 18, 2013 Citation: US Neurology, 2013;9(1):55–60 DOI: 10.17925/USN.2013.09.01.55

Correspondence: Sarosh R Irani, DPhil, MRCP (Neurol), Level 6, West Wing, John Radcliffe Hospital, Oxford, OX3 9DS, UK. E: sarosh.irani@ndcn.ox.ac.uk

Autoantibody-mediated diseases of the central nervous system (CNS) are a rapidly expanding field within contemporary neuroimmunology. Autoimmune encephalitis (AIE) is a treatable cause of subacute-onset memory loss and confusion. In some patients, the evidence for inflammation is limited and the term autoimmune encephalopathy may be preferred. The associated spectrum of autoantibodies against specific molecules in the CNS is widening, and serologic testing can help reach a definitive diagnosis. Although each individual autoimmune encephalitis syndrome has some distinctive demographics and associated clinical features, an autoimmune cause for recent-onset amnesia, confusion, and seizures should be considered in men and women, adults and children alike. Importantly, AIE is often a treatable neurologic condition and hence its consideration and recognition is of great importance.

Autoantibodies can be divided into those that target the extracellular domains of membrane molecules and those against intracellular targets. While the former are believed to be pathogenic since they can access their targets on the membrane of live cells, the latter are likely to be markers of immune-mediated disease but unlikely to themselves be pathogenic.

Initially, AIE was considered a paraneoplastic phenomenon, typically associated with a number of tumors and many of the antibodies (Abs) targeted intracellular antigens (such as Hu, CV2/CRMP5 and Ma2). Patients with these antibodies and associated tumors are unlikely to respond to immunotherapy and have in general a poorer prognosis.1,2

More recently described antibodies associated with AIE have been directed against cell surface molecules and ion channels in particular. Ion channels are involved in crucial processes within the nervous system including direct signal transmission, synaptic plasticity, and neuronal network modulation. Any interruption of ion channel function can therefore be expected to impact on the performance and output of the CNS. By contrast to those with paraneoplastic encephalitis, patients with antibodies against surface antigens often have no underlying tumor and respond well to immunotherapy.3,4

A Glimpse at the Neuromuscular Junction
To understand how antibodies can cause disease, the neuromuscular junction (NMJ), and pathophysiology of myasthenia gravis (MG) provides a useful paradigm. Eighty percent of patients with MG have serum antibodies directed at nicotinic acetylcholine receptors (nAChRs) located at the post-synaptic aspect of the NMJ. In individual patients, the reduction of antibody levels is accompanied by clinical improvement. The study of nAChR Abs has revealed a variety of different mechanisms of pathogenicity. Most Abs cross-link adjacent nAChRs and lead to internalization and subsequent destruction of the receptor-Ab complex.1 Equally, many Abs lead to activation of the lytic cascade of
This illustration of antibody-mediated disease in the peripheral nervous system illuminates some of the ways by which Abs can cause disease at a simple synapse and might offer some insight as to the potential complexity associated with studying their actions on central neuronal networks.

### Autoimmune Encephalitides—An Evolving Spectrum of Antibody-associated Diseases

The subacute onset of memory loss, confusion, and seizures are characteristic features of AIE associated with neural antibodies. Viral encephalitis and metabolic conditions, such as herpes simplex encephalitis and Wernicke’s encephalopathy, respectively, can present with similar features. The clinical features can help make the diagnosis of an autoimmune process and then guide the rational identification of an associated Ab (see Table 1). Therefore, autoimmune encephalitides may be subdivided according to the antigen recognized by patient Abs.

As reviewed by Vincent and colleagues, the most common and well researched associated Abs are directed against the voltage-gated potassium channel-complex (VGKC-complex), N-methyl D-aspartate receptor (NMDAR) and glutamic acid decarboxylase (GAD). Small case series have also described patients with Abs against glycine receptors, amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPA-R), and gamma-aminobutyric B receptors (GABA-BR).

### NMDAR-antibody Encephalitis

NMDAR-antibody encephalitis was first described as a paraneoplastic phenomenon in young females with an ovarian teratoma. The Ab target was then confirmed as the extracellular domain of the NR1 subunit of the NMDA receptor. In the following years, the clinical spectrum associated with NMDAR-Abs has widened significantly to include adult and pediatric, male and female, paraneoplastic and non-paraneoplastic patients. NMDAR-Ab encephalitis is now the most common AIE in younger patients and is a differential diagnosis to be considered in any patient presenting with subacute alterations of cognition, behavior and amnesia.

Other consistently reported features of NMDAR-antibody encephalitis include seizures, dysautonomia, central hypoventilation, and sleep pattern alterations. However, the movement disorder is often the most distinctive single feature of the condition. This involves stereotyped, complex, prolonged oro-facial, and limb movements which are often refractory to drugs usually used to treat chorea and dystonia. Most patients develop these varied symptoms within the first 4 weeks of their illness, often in a characteristic pattern.

The majority of NMDAR-Abs are of the IgG1 subclass, however recently other NMDAR-Ab subclasses have also been reported. NMDAR IgM Abs were found in a patient with NMDAR-Ab encephalitis with features of bipolar disorder and the clinical improvement mirrored a fall in NMDAR IgM. In addition, 7/24 patients with a subacute decline of cognitive function were shown to have NMDAR IgA but not IgG and again, clinical improvement correlated with the serum levels of NMDAR IgA.

### Abs to the Voltage-gated Potassium Channel-complex

For over a decade, VGKC-complex Abs have been associated with clinically different syndromes: neuromyotonia (NMT), Morvan’s syndrome, and...
Patients with VGKC-complex Ab LE are more often male, older than 50 years of age and many respond well to immunotherapy. The majority of patients do not have tumors. The clinical features of VGKC-complex-Abs include subacute onset of amnesia, disorientation, neuropsychiatric changes, and seizures. Serum hyponatremia and signs of hippocampal inflammation on MRI are useful paraclinical pointers towards the diagnosis. Seizures are typically of medial temporal lobe semiology but it was recently reported that faciobrachial dystonic seizures (FBdS) are a frequent accompaniment. FBdS are brief (few seconds in duration), frequent (median 50 per day) events which typically affect the arm and face and precede VGKC-complex Ab LE in 75% of cases.24–26

NMT is a peripheral nerve hyperexcitability condition that manifests with symptoms such as muscle cramps, stiffness, and fasciculations and shows an underlying thymoma in about 20% of cases. Morvan’s syndrome describes the combination of NMT with CNS features. The latter are often with more psychiatric symptoms than in cases with LE and tumors (such as thymomas or small cell lung cancer) are found in about 40% of cases.27,28

The diagnosis of VGKC-complex Abs relies on Ab-mediated precipitation of mammalian brain membranes that are radioactively labeled with alpha-dendrotoxin. The nature of the assay means that Abs against components closely associated with the VGKCs themselves will co-immunoprecipitate the radioactive-dendrotoxin-VGKC. Further methods have to be used to identify the individual antigenic targets associated within the complex, and so far, three have been described: LGI1 (leucine-rich glioma inactivated 1), CASPR2 (contactin associated protein 2), and Contactin-2.29,30 LGI1 was shown to be the target of the Abs found in patients with VGKC-complex Ab LE.27,29 and also in cases with FBdS.30 By contrast, CASPR2-specific Abs are more commonly found in Morvan’s syndrome and NMT.31

**Patients with Glutamic Acid Decarboxylase Abs**

Patients with GAD Abs may present with a subacute onset of temporal lobe epilepsy (TLE) and cognitive impairment. Abs to the intracellular protein GAD are also present in other diseases such as stiff-person syndrome. Patients with GAD-Ab LE are often younger adult women (median age at onset = 47 years), mostly without a tumor. The disease has a progressive course, without phases of remission.22,23

Although GAD is an intracellular antigen and Abs against intracellular targets are believed less likely to be pathogenic per se, the typical changes seen on MRI in GAD LE patients are indicative of an inflammatory process within the temporal lobe. It is possible that as yet unidentified Abs against cell surface structures co-exist in the serum and are the actual pathogenic molecules driving the process. The patients’ response to oral immunotherapy is variable, but there are some successful reports of possible improvement following plasma exchange or intravenous immunoglobulin correlated with a reduction in Abs.34–36

**Other Antigenic Targets**

Other proteins involved in physiologic CNS synaptic processes have also been shown to be antigens in AIE. A small case series in 2009 identified 10 patients with LE and Abs against the GluR1 and GluR2 subunits of AMPAR.37 The patients were predominantly female (9/10) with an average age of 60 and the majority were also diagnosed with tumors (7/10). One year later, 15 LE patients were described with Abs against the GABA-R.38 These patients were older, 62 years on average, mainly male (8/15) and 7/15 had lung tumors (5/7 were small cell lung cancer). In a further study of LE patients, 10/70 were shown to have GABA-R-Abs, again the average age of the patients was 60 and 8/10 had small cell lung cancer.39 Antibodies against mGluR5 and DPPX have more recently been discovered.

**Pediatric Autoimmune Encephalitis**

Pediatric patients are an interesting emerging subgroup with AIE. Insults affecting the CNS may have more impact on the long-term outcome especially in very young patients where brain maturation and development is still a delicate ongoing process. Alternatively, children may have an intrinsically higher potential for plasticity. The two Abs that have been most commonly associated with pediatric AIE are NMDAR and VGKC-complex Abs.

In one study, NMDAR-Abs were reported in 32 pediatric patients and ovarian teratomas were present in one-third of the female patients under 18 and only one-tenth of those younger than 14 years.40 Many of the clinical features were similar to the adult cases. Subtle differences include a higher rate of prodromal infectious symptoms and the early presence of a movement disorder. The fact that pediatric cases are less likely to have tumors is relevant since a recent study has suggested the need for early aggressive treatment in non-paraneoplastic pediatric NMDAR-Ab encephalitis.41

The presentation of pediatric patients with LE and VGKC-complex-Abs appears to show some differences from the adult patient group. There are different rates of psychiatric/psychotic features, seizures, and infrequent reports of hyponatremia.42–44 Most of the reported pediatric cases are also female, whereas VGKC-complex-Abs are more common in male adult patients. Interestingly, many pediatric cases were VGKC-complex-Ab positive, without Abs against the known antigenic targets (LG1, CASPR2, and contactin-2).44 42% of autoantibody positive (NMDAR or VGKC-complex) patients had a full recovery in a recently published case-series.45 A further case-series reported only 1/4 with complete recovery in pediatric VGKC-complex autoantibody patients who presented with status epilepticus.46

**Treatments for Patients with Autoimmune Encephalitis**

Although no systematic studies of treatment regimens or RCTs for AIE have been undertaken, the available case series show a few patterns and overall, prompt immunotherapy and tumor management, when relevant, is believed to be key to improving patient outcomes.9,10,42,43,46

AIE can be subdivided into diseases with different antigens and these often show slightly different responses to treatment. It is therefore key to diagnose AIE promptly and exclude other causes for the patient’s presentation. The published case studies for all AIE associated with antibodies targeting extracellular domains of membrane proteins suggest that immunotherapies can help expedite recovery. As to which immunotherapy regimen is best to follow, the verdict is still out, although recently studies have approached the question of patient management in NMDAR and VGKC-complex Ab encephalitis. As VGKC-complex Abs are often associated with a monophasic illness, therapy is usually required for shorter periods and is less aggressive than in NMDAR-Ab encephalitis.
Autoimmune Encephalitis

An observational cohort study summarized clinical findings in nearly 600 NMDA receptor (NMDAR) patients treated at several centers. The authors suggested that 53% of cases showed a good response to first-line immunotherapy (steroids, IVIG, PEX, alone or combined) within 1 month. Among patients who did not respond to this treatment, the study concluded that long-term outcome was better if rituximab and/or cyclophosphamide were also administered. In 2004, VGKC-complex antibody (VGKC) was shown to respond to immunosuppressive treatments using steroids, IVIG or PEX with a fall in Abs levels and marked neuropsychologic improvement. The only published small prospective study in the field of VGKC-complex antibody (VGKC) showed favorable outcome in all nine VGKC patients where the protocol consisted of plasma exchange (PEX, 50 ml/kg), intravenous immunoglobulin (IVIG, 2 g/kg) and methylprednisolone (1 g x3), followed by maintenance oral prednisolone (1 mg/kg/day). Efficacy was also seen using monthly intravenous methylprednisolone pulses (500–1,000 mg/day on 3–5 consecutive days) which successfully improved seizure control (9 of 9 patients) and reduced VGKC-complex Abs levels in 6/9 patients. However, the same regimen did not show a change in GAD levels (6/9 patients) or reduction in their seizure frequencies (0/7 patients). Amongst the different AEs, most research into the pathogenicity of the Abs has been published in NMDAR-Ab encephalitis. Many of the mechanisms previously established in myasthenia gravis appear to be relevant to Abs-associated with CNS pathologies (see Figure 1). Application of NMDAR-Abs can cause internalization of the NMDARs in vitro and in vivo after infusion of patient IgG into the hippocampus of rats. Internalization can also be caused by NM2A IgM or IgA Abs. In addition, NMDAR-Abs can show other potentially pathogenic actions—they may alter the trafficking and distribution of NR2A and 2B subunits around glutamatergic synapses (in some via an interaction between NMDARs and ephrin receptors). This effect appears to be more rapid than the internalization of receptors, since the first changes to the composition at the glutamatergic synapse can be seen as early as 2 hours after Ab application. The effects of NMDAR-Abs also acutely interfere with the synapse’s ability to change in response to long-term potentiation protocols, limiting the ability of neuronal networks to remodel in response to activity.

NMDAR-IgG Abs are mainly of the IgG1 subclass and they were shown to bind and activate the complement system in vitro, both in HEK cells expressing NMDARs and in neuronal cultures from the hippocampus. However, a role for NMDAR-Abs in the activation of complement has not been demonstrated in the limited numbers of biopsies or post mortem reports.

Figure 1: Overview of the Antibody Targets in NMDAR, VGKC-complex, and GAD-antibody-associated Encephalitis and their Synaptic Localizations

The mechanisms associated with the antibodies (Abs) in each of these three types of autoimmune encephalitis (AIE) are listed below. N-methyl D-aspartate receptor (NMDAR)-Abs (left) Ab-mediated NMDAR internalization and changes to the ratio of tetramers containing NR2A or NR2B at the synapse. Microgliosis and lymphocytic infiltration have been implicated in the pathogenesis. Voltage-gated potassium channel (VGKC)-complex Abs (center): several antigens are tightly associated with the VGKC complex and can be targeted by the Abs, leading to neuronal loss, axonal injury, and complement activation. The commonest target is leucine-rich, glioma inactivated 1 (LGI1). Both neuronal loss and axonal injury have been detected in hippocampi from patients with glutamic acid decarboxylase (GAD) Abs (right). The pathogenesis is less clear but in some cases, antibodies target alternative cell surface proteins.)
Effects of NMDAR-ABS on more complex neuronal networks and CNS physiology have also been demonstrated. Intrahippocampal infusions of patient IgG showed that NMDAR-ABS increase extracellular concentrations of glutamate and alter glutamatergic pathways and metabolism. Not all of the symptoms in NMDAR-ab encephalitis are restricted to hippocampal functions. Indeed, NMDAR-AB effects on neocortical networks were shown to be involved in corticomotor excitability after infusion of patient IgG into the prefrontal cortex.

However, in vivo features which reproduce clinical features seen in affected patients are awaited. Indeed, many of the above studies used models that reduce the complexity of the hippocampal and neuronal networks and model the mechanism in vitro. Post mortem and biopsy studies of patients may offer more insights into the human pathophysiology or, alternatively, may only capture an end-stage disease. A detailed analysis of the cortical biopsy and autopsy pathology seen in NMDAR, VGKC, and GAD LE was recently published. The specimens showed infiltration with cytotoxic T-cells and lymphocytes, however this was much less commonly seen in cortex of NMDAR-ab encephalitis patients.

Previous post mortem studies also showed prominent microgliosis in brain tissue from NMDAR-Ab positive patients. VGKC-complex and GAD-antibody encephalitis both showed a marked degree of neuronal loss within the hippocampus and evidence of axonal injury. These findings were not, however, observed in the tissue obtained from patients with NMDAR-Ab encephalitis patients. Therefore, this is likely to be in line with a mechanism by which antibodies primarily affect synaptic processing and act to modulate neuronal network activity in NMDAR-ab encephalitis. Evidence of activation of the complement system was only present in VGKC-complex Ab positive cases, highlighting this pathway as a possible explanation for loss of neurons within the hippocampus.

There is ongoing doubt whether Abs against intracellular targets can be truly pathogenic. However the MRI findings, pathology analyses and a recent report showing a decrease in cortical GABA concentrations in patients with GAD-ABS indicate the presence of a functionally significant inflammatory process. Whether the GAD-ABS are causing the processes or a simply a marker that coincides with unknown other Abs is currently difficult to assess.

More Questions to Answer

In 1957, Witebsky proposed criteria to determine if a disease is autoimmune in nature, based on Koch’s postulates describing the relationship between pathogens and disease. The criteria required an autoantibody-mediated autoimmune response, recognition of the target antigen, an autoimmune response which was inducible in animal models and animals immunized against the identified antigen developing a similar disease.

As reviewed above, few of these aspects have been shown for AIE to date. A recent case study by Hansen and colleagues however shows that intrathecal synthesis of NMDAR-ABS does not necessarily correlate with symptoms in patients. More research is needed to explain how Abs can be pathogenic, access the CNS, and how subsequently may lead to changes in the neuronal networks and cause the typical pattern of disease in patients. In particular, the recently described FBDS with its window of treatment opportunity might give further insight into the mechanisms involved in the disease progression. It will be very interesting to understand the localization of these early events and whether their treatment can halt the frequently observed progression to the amnesia and confusion which characterize AIE.

It is rather difficult to model complex AID without an active immunization model, since each passive transfer relies on the fact that the animals’ immune system would respond to the human antibodies in the same way as the patients. However, this is not always the case: passive transfer animal models of neuromyelitis optica (NMO) require the presence of human complement to fully show the pathogenic potential of NMO Abs.

We are only just beginning to understand the complex mechanisms of disease caused by Abs within the CNS. Further research into early phases of disease and the Ab-mediated pathology might lead to novel targeted approaches which could complement the existing array of immunotherapy. In addition, formal prospective studies are needed to determine ideal treatment regimens for this potentially reversible cause of encephalitis.
Autoimmune Encephalitis

49. Manto M, Dalmau J, Didebold A, et al., Differentiation of cortical motor responses is increased by IgGs of patients with NMDA receptor antibodies, J Neuro, 2011;268:27–33.
50. Stagg CI, Leng B, Blett VS, et al., Autoantibodies to glutamic acid decarboxylase in patients with epilepsy are associated with low cortical GABA levels, Epilepsy, 2013;51:1918–901.