The term ‘limbic encephalitis’ (LE) describes the condition when limbic areas of the brain are inflamed (swollen) and consequently not functioning properly. The main regions of the limbic system include the hippocampus and amygdala, hypothalamus, cingulate gyrus, limbic cortex). The limbic areas of the brain control many functions including memory, learning, and emotions such as aggression. In addition, some of these limbic areas are susceptible to seizures, which are a common feature of limbic encephalitis.

**SYMPTOMS OF LIMBIC ENCEPHALITIS**

The symptoms of LE include memory loss, seizures, confusion, disturbances of sleep and psychological problems such as altered personality or behavior.

**CAUSES OF LIMBIC ENCEPHALITIS**

Most forms of LE fall into two main categories:

1. Infectious encephalitis – caused by direct invasion of the limbic area of the brain by a bug, usually a virus.
2. Autoimmune encephalitis – caused by the person’s own immune system reacting against parts of the limbic system.

1. **Infectious causes**

Many infections of the brain can potentially cause inflammation of the limbic areas. A number of viruses, such as the herpes simplex virus (HSV) seem to preferentially target this area. Some people may therefore be given the diagnosis of LE whilst others are given the diagnosis herpes simplex encephalitis for the same condition. A clearer way for people would be to say that the person has ‘herpes simplex virus encephalitis affecting mainly the limbic areas of the brain’, but this is rather long-winded.
2. Autoimmune causes

A major role of our immune system is to recognize and eliminate infection. But sometimes parts of the immune system called ‘antibodies’ may instead react with proteins of our own body to cause autoimmune diseases. When this reaction is against proteins of the limbic areas of the brain, this is called ‘autoimmune limbic encephalitis’.

There are broadly two forms of autoimmune limbic encephalitis: paraneoplastic limbic encephalitis (PLE) and non-paraneoplastic limbic encephalitis (NPLE).

a) Paraneoplastic limbic encephalitis (PLE)

Sometimes when the immune system starts to react with the limbic areas, this happens because the person has a tumor in their body which activates the immune system. This activated immune system can, in turn, attack the brain. Doctors call this paraneoplastic limbic encephalitis as the tumor (neoplasm) affects the brain from a distance, via the immune system.

In many cases, PLE can be diagnosed by testing for one of paraneoplastic autoantibodies in the patient’s blood and spinal fluid. Most individuals with PLE have a cancer of the lung, thymus gland, ovary, breast or testes. More rarely, other cancers can initiate the condition. The outcome is very dependent on the underlying tumor and the precise condition, often classified by the antibody. In some cases, the condition may improve or at least stabilize if the cancer is detected and treated effectively. However, unfortunately, in many cases treatment does not improve the patient’s neurological symptoms, probably because the immune system has irreversibly damaged the brain cells and the tumor cannot be controlled successfully.

b) Non-paraneoplastic limbic encephalitis (NPLE)

NPLE has only been clearly recognized in the last few years when doctors began to identify patients with symptoms like those with PLE but who did not have any of the marker paraneoplastic antibodies in their blood and never developed a tumor. NPLE is far more common than PLE.

It is becoming increasingly clear that NPLE is caused, at least in part, by specific antibodies in the patient’s blood that target the patient’s brain, particularly the hippocampus and other limbic areas. Limbic encephalitis is the main manifestation of several of the autoimmune encephalitis syndromes that may occur without a cancer association. Many of these patients improve if they are treated with drugs that suppress the immune system and reduce the levels of the antibodies. These drugs include steroids, intravenous immunoglobulins and plasma exchange.

Types of antibodies

Several specific brain protein targets for these antibodies have been discovered over the last years and this variety may explain why people have different symptoms. The main established antibodies and their associated features are described below:
The Most Common Syndromes

NMDAR-antibodies

Another antibody that can cause PLE is the NMDAR antibody. This disease may be associated with a growth such as a cancer in around 30% of cases. This antibody usually causes encephalitis involving several brain regions, but it can sometimes cause a pure LE (Please see the anti-NMDAR encephalitis factsheet for further details).

It should be noted that the brain imaging and the routine lumbar puncture results may be normal in autoimmune limbic encephalitis.

LGII/CASPR2 (previously termed voltage-gated potassium channel complex antibodies (VGKC))

These two forms of Autoimmune encephalitis are associated with LGII (leucine-rich-glioma inactivated 1) and CASPR2 (contactin-associated protein 2) antibodies. This AE may also present with rapid eye movement sleep behavior disorder, hypothermia, startle syndrome, ataxia and intestinal pseudo-obstruction. A recently described and highly distinctive feature is the presence of faciobrachial dystonic seizures (FBDS), characterized by brief and frequent dystonic paroxysms typically involving the face and ipsilateral arm; these patients consistently have antibodies against LGII. Around 40% of cases have a normal MRI, and the Cerebral Spinal Fluid (CSF), tends to show the absence of pleocytosis or intrathecal synthesis.

Less Common Syndromes

Patients with an encephalopathy and prominent insomnia, neuromyotonia and dysautonomia are often termed Morvan's syndrome. These patients show high levels of CASPR2-antibodies, often with lower levels of LGI1-antibodies. Morvan's patients have a high risk of an underlying tumor which is usually a thymoma: most patients with a tumor have CASPR2 antibodies.

AMPAR and GABA_{B/A}R antibodies

Antibodies against two other receptors in the brain, AMPA and GABA_{B/A}R, are less common causes of autoimmune limbic encephalitis. AMPAR antibody has been identified in women with relapsing AE. GABA_{B}R has a typical presentation apart from early and prominent epileptic seizures. Although most of these patients have an underlying tumor, this is a form of Paraneoplastic Limbic Encephalitis that can often respond to treatment relatively well.

GLycineR antibody

Patients with GlycineR may present with progressive encephalomyelitis with rigidity and myoclonus (PERM), a rare condition showing limb and axial rigidity, muscle spasms, brainstem signs and hyperekplexia.

Treatments of autoimmune limbic encephalitis
The diagnosis of autoimmune encephalitis is particularly important because the disease is potentially treatable with medicines that dampen down the immune system. These medications are called immunosuppressive and include steroids, immunoglobulins (a blood product given into the vein in a drip) and plasma exchange (when some of a person’s blood is taken out from a vein, and the plasma part of the blood which contains antibodies is separated and replaced with new plasma and then put back into the vein in a drip). All these drugs have known side-effects, but their benefits are generally felt to outweigh possible side-effects in these conditions.

**Future challenges in autoimmune encephalitis**

As these autoimmune diseases have only been recently described, there is still much to be done to raise awareness among the medical community. Future research aims to understand the biological mechanisms by which these antibodies affect the excitability of the brain that causes these diseases. Researchers also hope to discover further antibodies which may allow other autoimmune encephalitis forms to be diagnosed. In addition, ongoing research is trying to understand how to best target the cells which produce antibodies and tailor therapies in patients with autoimmune encephalitis.