Autoimmune disorders of the neuromuscular junction

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The neuromuscular junction (NMJ) is a specialized synapse with a complex structural and functional organization. It is a target for a variety of immunological disorders and these diseases usually respond well to immunotherapies. The understanding of the immunological basis of myasthenia gravis, the most common neuromuscular junction disorder, has improved in recent years. Most patients have antibodies to the acetylcholine receptor (AChR), but around 10% have AChR antibodies that are only identified by novel methods, and up to 5% have muscle-specific kinase antibodies which define a different subgroup of myasthenia. The spectrum of antibodies and their pathophysiological aspects are being elucidated. Even though less common, Lambert Eaton myasthenic syndrome (LEMS) is important to recognize. The abnormality in LEMS is a presynaptic failure to release enough packets of ACh, caused by antibodies to the presynaptic voltage-gated calcium channels. More than half these patients have a small cell carcinoma of lung. Acquired neuromyotonia (NMT) is a condition associated with muscle hyperactivity. Clinical features include muscle stiffness, cramps, myokymia, pseudomyotonia and weakness. The immune mechanisms of acquired NMT relate to loss of voltage-gated potassium channel function. This review will focus on the important recent developments in the immune-mediated disorders of the NMJ.

Key words: Gravis, Lambert Eaton myasthenic syndrome, neuromyotonia, neuromuscular junction, autoimmunity, acetylcholine receptor, voltage-gated channel

The Neuromuscular Junction

The neuromuscular junction (NMJ) is a prototypic synapse although its structure is rather different from those of the central nervous system (CNS). The unmyelinated motor nerve terminals are separated from the postsynaptic membrane by a cleft that contains a basal lamina. This includes many proteins such as collagens, laminins, fibronectin and perlecan which help anchor some of the key elements involved in NMJ development and function; for instance acetylcholine esterase (AChE) is localized via ColQ, a collagen-like molecule, to the basal lamina. Agrin and neuregulins, secreted from the nerve terminal, are bound by the basal lamina and interact with their receptors, playing an important role in the location of postsynaptic membrane proteins, voltage-gated calcium channels and the dystroglycans. The postsynaptic membrane at the NMJ forms a series of deep folds. The acetylcholine receptors (AChRs) are found at the top one-third of these folds, whereas the voltage-gated sodium channels are anchored at the bottom of the folds. The development of the NMJ is a fascinating area of research and many of the proteins involved are relevant to disease (Figure 1 for a simple representation).

The nerve action potential opens voltage-gated calcium channels (VGCCs) that are located in the motor nerve terminal [Figure 1]. The resulting influx of calcium leads to the release of about 30 (in human muscle) individual packets of acetylcholine (ACh). Some of the ACh is hydrolysed by AChE but about 65% reaches the AChRs
on the postsynaptic membrane. Binding of two ACh to each AChR leads to the opening of the AChR-associated ion channel, influx of cations (mainly sodium) and generation of an endplate potential (EPP). The EPP in normal human muscles is around 20-30 mV. Miniature EPPs (MEPPs) are the result of the spontaneous release of single packets of ACh and their amplitudes are much smaller (around 1 mV) and generally reflect the density of functional AChRs.

The EPP rapidly depolarises the postsynaptic membrane and, when this reaches a critical firing threshold, the voltage-gated sodium channels open and an action potential is propagated along the muscle fiber leading to contraction. The extent to which the EPP exceeds that necessary to initiate the action potential is usually called the safety factor for neuromuscular transmission.[2] The EPP is short-lived because the AChRs close spontaneously, ACh dissociates and escapes by diffusion or is hydrolysed by AChE. The calcium channels also close spontaneously. Opening of voltage-gated potassium channels on the presynaptic membrane is important in restoring the membrane potential and limiting calcium channel opening.

The topic of this review is the chronic conditions caused by serum autoantibodies and which are treatable by immunotherapies. The NMJ is particularly vulnerable to circulating factors because it has no blood-brain barrier. This is well illustrated by envenomation by snakes, spiders, scorpions and other species, that results in NMJ paralysis or hyperexcitability.[3] Some of these neurotoxins are particularly valuable in studying the NMJ and for tagging the proteins for autoantibody assays, as will be described below.

**Myasthenia Gravis**

**Clinical features**

Myasthenia gravis (MG) usually presents in young adult or later adult life as muscle weakness and excessive fatigue during repetitive movements. It most often involves the extraocular muscles of the eye with double vision and ptosis at onset, but usually progresses to generalized weakness. Involvement of the facial and bulbar muscles is particularly disabling and respiratory involvement can be life-threatening. Cholinesterase inhibitors, by prolonging the action of ACh, tend to lead to clinical improvement.[4,5] The history of myasthenia research is fascinating[6] but cannot be reviewed here. The essential findings in the 1960s and 1970s were that the MEPP amplitudes are substantially reduced in muscle biopsies from MG patients[7] and that this is due to loss of the postsynaptic AChRs.[8] The latter finding depended on the discovery of bungarotoxin, a snake toxin that binds specifically to the AChRs.[9] At much the same time, purification of the AChR from electric organs of certain fish, and immunization of rabbits with the purified AChRs, led to the first experimental autoimmune myasthenia gravis (EAMG) model by Patrick and Lindstrom.[10] Lindstrom and his colleagues then detected antibodies to AChRs in the serum of MG patients, establishing the radioimmunoprecipitation assay that has changed little to this day.[11]

The levels of AChR antibodies vary widely between patients and there is only modest correlation with clinical severity. This is probably because the antibodies are highly heterogeneous in their characteristics and the epitopes to which they bind on the AChR.[12] However, during treatments such as plasma exchange the levels of AChR antibody correlated well with clinical severity within an individual.[13] This clinical experiment established the role of the antibodies in causing the disease. Equally important was the passive transfer of purified IgG from MG patients to mice that resulted in NMJ deficits with reduced AChRs.[14]

**Clinical Heterogeneity**

**Ocular myasthenia gravis**

It is unclear why the extraocular muscles are so often the first to be affected and may remain the only muscles involved. Indeed, they should be resistant to fatigue with high blood flow, mitochondria content and metabolic rate.[15] However, the motor unit sizes are small and the firing frequencies are high, and some of the muscle fibers have multiple NMJs in which the EPPs, rather than action potentials, are directly responsible for activating the contractile apparatus. This means that any reduction in the EPP could have a direct effect on the strength of muscle contraction.
There has been some controversy about the role of fetal rather than adult AChRs at extraocular muscle endplates,[16] but another factor that might make them more susceptible is the low expression of complement regulators;[17] this would make them more vulnerable to complement-mediated damage. Of possible relevance to this, in the Lambert Eaton myasthenic syndrome (LEMS, see below), in which complement-mediated damage is not thought to occur, ocular muscle weakness is uncommon.

**Generalized Acetylcholine Receptor**

Most patients in the Western world progress to generalized weakness, although in some Eastern countries ocular MG, particularly in young children, is very common.[18] Based on age at onset, HLA associations, AChR antibody positivity and thymic pathology the patients can be divided into early onset, late onset and thymoma-MG [Table 1].[19]

In the older patients, the Human leukocyte antigen (HLA) association is less marked and the thymus is essentially normal for age. Despite these important subgroups, the disease mechanisms appear to be very similar (see below). The late onset patients are being recognized much more frequently in the population and may well be underdiagnosed or misdiagnosed as stroke or motor neuron disease.[20]

**Neonatal Myasthenia Gravis and Arthrogryposis Multiplex Congenita**

A relatively small proportion of babies born to MG mothers have neonatal MG, due to the placental transfer of the maternal antibodies. In a small number of reported cases, the mothers give birth to babies with arthrogryposis multiplex congenita. This condition, which involves multiple fixed joint contractures and other abnormalities, can be caused by anything that reduces fetal movement in utero. When it is associated with maternal AChR antibodies it typically recurs in subsequent pregnancies unless the mother is adequately treated. Since some mothers have little, or even no, evidence of MG themselves, multiple affected offspring have been reported in some cases.[21] This condition appears to be caused by maternal antibodies that inhibit the function of the fetal form of the AChR paralyzing the developing fetus.[22] We established a maternal-to-fetal passive transfer model of this condition[23] and the possibility of maternal antibodies should be considered in other developmental conditions that recur in each consecutive pregnancy.

**The Cause of Myasthenia Gravis**

Despite many years of study, the cause(s) of MG are unknown. The antibodies are highly specific, generally high affinity for the AChR and variable in their characteristics. The binding sites on the AChR can be defined by competition with monoclonal antibodies raised against purified AChRs.[24,25] Many bind to a “main immunogenic region” (MIR) on the two AChR alpha subunits, but many antibodies bind to the other subunits, including the fetal-specific gamma subunit. Overall, the characteristics suggest that the human AChR is the immunogen and that the antibodies do not arise as the result of a cross-reaction with microbial antigens, as was proposed in the 1980s. However, it is possible that a low-affinity or cross-reactive antibody may lead subsequently to a higher affinity reaction.

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**Table 1: Subgroups of autoimmune neuromuscular junction diseases**

<table>
<thead>
<tr>
<th>AChR-MG</th>
<th>Age at onset</th>
<th>HLA association</th>
<th>Thymic pathology</th>
<th>Antibodies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early onset</td>
<td>&lt;40 years by definition</td>
<td>DR3 B8</td>
<td>Hyperplasia</td>
<td>AChR, AChR</td>
</tr>
<tr>
<td>Late onset</td>
<td>&gt;40 years by definition</td>
<td>DR2B7 but not very strong</td>
<td>Normal for age</td>
<td>Striated muscle antigens e.g. Titin, Ryanodine receptor</td>
</tr>
<tr>
<td>Thymoma</td>
<td>Variable</td>
<td>None known</td>
<td>Thymic tumor</td>
<td>AChR</td>
</tr>
<tr>
<td>Other subtypes</td>
<td></td>
<td>None known</td>
<td>Hyperplasia in some</td>
<td>Striated muscle antigens e.g. Titin, Ryanodine receptor</td>
</tr>
<tr>
<td>Low affinity AChR MG</td>
<td>Variable</td>
<td>None known</td>
<td>Hyperplasia in some</td>
<td>Low affinity antibodies detected against clustered AChR</td>
</tr>
<tr>
<td>Ocular MG</td>
<td>Variable</td>
<td>None known</td>
<td>Not known</td>
<td>AChR, AChR 50%</td>
</tr>
<tr>
<td>MuSK-MG</td>
<td>Variable</td>
<td>DR14DQ5</td>
<td>Normal for age</td>
<td>AChR, low affinity 15%</td>
</tr>
<tr>
<td>AChR/MuSK antibody negative MG (SNM)</td>
<td>Variable</td>
<td>None known</td>
<td>Not clear yet</td>
<td>MuSK</td>
</tr>
<tr>
<td>LEMS</td>
<td>Mostly 20-60</td>
<td>DR3 B8</td>
<td>None reported</td>
<td>Negative on all assays to date</td>
</tr>
<tr>
<td>LEMS with SCLC</td>
<td>40+</td>
<td>None known</td>
<td>None reported</td>
<td>VGCC</td>
</tr>
<tr>
<td>Neuromyotonia</td>
<td>Mostly 20-60</td>
<td>None known</td>
<td>May be thymoma</td>
<td>VGKC in 40%</td>
</tr>
</tbody>
</table>

Myasthenia gravis with acetylcholine receptor antibodies (AChR-MG), Myasthenia gravis with muscle specific kinase antibodies (MuSK-MG), Lambert Eaton myasthenic syndrome (LEMS), Small cell lung cancer (SCLC)
against the AChR. We now believe that lower-affinity antibodies do exist in some patients.

Mechanisms of Disease

The loss of AChRs at the NMJ is the underlying disease mechanism. This leads to reduced amplitude of the EPPs. The EPPs are often supra-threshold to begin with (and therefore induce an action potential in the muscle) but, during repetitive activity, they rapidly become sub-threshold leading to muscle weakness. The loss of AChRs results from several different processes. Relatively few antibodies directly block the function of the ACh-induced ion channel, but these antibodies can be important in some patients and some MG sera appear to have a transient inhibitory effect on AChR function in vitro. A more important mechanism is a reduction in AChRs due to the cross-linking of AChRs by divalent antibodies. This results in a temperature-dependent loss of AChR which can best be demonstrated in muscle cell lines in cultures, but has also been demonstrated at the mouse NMJ. Interestingly, there was an increase in synthesis rate of new AChRs in these experiments and in MG muscle biopsies, there is an increase in expression of the AChR.

Probably the most important mechanism of AChR loss is complement-mediated destruction of the postsynaptic membrane. The AChR antibodies are usually IgG1 or IgG which are both complement-activating subclasses. IgG and complement components C3 and C9, and of the membrane attack complex (MAC), are localized at the NMJ and AChR numbers appear to be inversely related to the amount of complement. There is also morphological damage to the endplates and simplification of the postsynaptic folds (Figure 1). This could be important as the voltage-gated sodium channels responsible for initiating the action potential are located at the bottom of the folds.

Several factors could modify the degree of AChR loss and thereby disease severity. Complement regulatory proteins are present which probably reduce the damage caused by complement-activating antibodies and as mentioned above there is increased AChR synthesis. In addition, the number of ACh packets released is increased at individual endplates. All of these factors are likely to be genetically regulated and will modify the severity of the disease and perhaps explain some of the variability between individuals, within an individual in different muscles and at different times.

The Thymus in Myasthenia Gravis

There are two main thymic pathologies in MG patients [Table 1]. Early-onset MG is associated with enlarged thymus glands with frequent lymphocytic infiltrates and germinal centers, similar to those found in lymph nodes. These glands, unusually, contain B cells and plasma cells as well as T cells. Moreover, in the thymic medulla, and sometimes surrounding or within the germinal centers, there are muscle-like myoid cells that express fetal AChR and other muscle proteins. It is known that the thymic cells can actively produce AChR antibody when cultured in vitro and removal of the thymus results in a moderate fall in AChR antibody although the levels usually plateau out after some months; nowadays most patients are also given immunosuppressive treatments. Further work on the thymic pathology has also shown that complement is activated on the abnormal epithelium, and on the myoid cells; it seems very likely that the AChRs expressed by these myoid cells initiate or maintain the immune response in early-onset MG. A recent report describes some of the factors that control AChR expression in these cells.

About 10% of MG patients have a thymoma, usually presenting in middle age and sometimes associated with other autoimmune disorders. Thymomas are epithelial tumors but often contain many T lymphocytes (and few B cells). They appear to be capable of exporting mature T cells but these cells may lack the normal regulatory mechanisms. The role of the thymoma is somewhat different from that of the thymus in early-onset MG. The AChR is not generally found in the thymoma itself, although there may be AChR subunits expressed individually in different epithelial cells. However, if AChR-sensitive T cells escape from the disordered thymic environment, they could get stimulated by the presence of AChRs on the myoid cells present in the adjacent “normal” thymus. Some of these possibilities are discussed by Vincent et al. Thymectomy generally does not lead to clinical improvement and most patients require immunosuppressive treatments.

The thymus in late-onset MG is usually considered to be atrophic or “normal for age”. However, there are few histological studies on this age group and thymectomy is seldom performed for MG patients over the age of 60 years, unless a thymoma is present.

Myasthenia Gravis without Acetylcholine Receptor Antibodies

Up to 20% of patients with generalized MG do not have detectable AChR antibodies by current laboratory methods, but their disease is also antibody mediated because they respond to plasma exchange, and their IgG can passively transfer disease to mice. A number of studies over many years have tried to define the nature of the antibodies in these patients.
Myasthenia Gravis with Muscle-specific Kinase Antibodies

A proportion of patients without AChR antibodies have instead antibodies to a protein called MuSK (muscle-specific kinase), which is restricted to the NMJ in adult muscle. MuSK is essential for the development of the NMJ but its role in adult muscle is less clear although it probably plays an important role in maintaining the structure of the postsynaptic membrane. MuSK-MG often presents with ocular involvement, but frequently evolves to include severe bulbar and facial weakness, sometimes with marked muscle atrophy in these muscles.

Antibodies to MuSK were demonstrated in around 70% of patients without AChR antibodies, and in none of the AChR antibody positive patients. Subsequently radioimmunoprecipitation assays found the antibodies in varying proportions of patients and the incidence seems to vary throughout the northern hemisphere (there are insufficient data from the southern hemisphere).

There are several interesting differences between MuSK-MG and typical AChR-MG patients. Firstly, the thymus in MuSK-MG is usually normal with very small and infrequent germinal centers and little complement activation. Secondly, the disease is associated with HLA DR14-DQ5 suggesting that the genetic susceptibility is very different from typical MG. Moreover, the MuSK antibodies are mainly IgG4 unlike the IgG1 and IgG3 of AChR antibodies.

We hypothesized that these patients had antibodies that bind to the AChR with low affinity in solution but would be able to bind to the AChRs when they were expressed at the NMJ where the density of the receptors is very high. To reproduce this high density, we expressed AChRs in a cell line and clustered the receptors with the clustering protein rapsyn. Curiously, there was little evidence of AChR loss either, and therefore the mechanisms by which MuSK-MG patients become weak are not clear.

There have not yet, however, been any detailed studies of neuromuscular transmission in the most affected muscles. The limb muscles are often relatively unaffected and limb muscle electrophysiology may be normal in MuSK-MG, while facial muscles are abnormal; this suggesting that the MuSK antibodies are particularly pathogenic in these muscles, as supported by magnetic resonance imaging (MRI) studies. Further in vitro and animal studies are required to determine the role of MuSK in adult muscle, and to look for differences between facial and limb NMJs.

Myasthenia without Muscle-specific Kinase or Acetylcholine Receptor Antibodies

There remain approximately 10% of MG patients who have neither MuSK nor AChR antibodies by the radioimmunoprecipitation methods used worldwide. However, we observed that these patients have clinical symptoms indistinguishable from those of typical AChR-MG patients, although they are often less severely affected. Moreover, their thymus glands show hyperplasia and germinal centers very similar to those of AChR-MG patients. Their sera frequently inhibited AChR function in vitro and seemed to increase desensitization of the AChR, but this effect appeared to be reversed by washing suggesting that perhaps these antibodies were low affinity rather than the high affinity antibodies detected in serum.

We hypothesized that these patients had antibodies that bound to the AChR, but were not able to bind to the AChRs when they were expressed at the NMJ where the density of the receptors is very high. To reproduce this high density, we expressed AChRs in a cell line and clustered the receptors with the clustering protein rapsyn. We were then able to detect the presence of antibodies in about 60% of the sera that were negative for binding to AChR in solution. It appears that this “cell-based” method can also be applied to measuring antibodies to MuSK, and should provide a more sensitive assay for all MG patients.

The Lambert Eaton Myasthenic Syndrome

Clinical features

The Lambert Eaton myasthenic syndrome is another autoimmune NMJ disorder, but it is also a prototype paraneoplastic neurological syndrome as about 50% of the patients have small cell lung cancer. The distribution of muscle weakness is different from MG as the trunk and legs are most often involved, and ocular involvement is uncommon. Autonomic symptoms
(dry mouth, constipation, impotence) are frequent. Typically, weakness improves during sustained effort as can be shown clinically or by electromyography. The compound muscle action potential is very small but increases during high-frequency stimulation or following a brief period of voluntary contraction. The reflexes are absent or depressed but may become stronger after voluntary contraction.

**Immunological basis**

In patients with small cell lung cancer (SCLC) it is highly likely that the immune response is primarily directed against the tumor (see below), but in the remaining patients, who have a chronic disease, the cause is unknown. An HLA association with HLA B8DR3 suggests an underlying genetic predisposition to autoimmunity and these patients often have other autoimmune disorders, such as thyroid disease or diabetes. These observations led to the first experimental studies implicating the immune system. It was shown that patients with LEMS responded clinically to plasma exchange, and daily injection of LEMS IgG antibodies into mice reproduced the principal neurophysiologic changes of LEMS.

**Mechanisms of disease**

The defect in LEMS is a presynaptic failure to release enough packets of ACh. The EPSPs are very small, as they are in MG, but the MEPPS are of normal amplitude. In these classic studies, repetitive stimulation of the nerve showed that the endplate potential increased during repetitive stimulation and also when extracellular calcium was raised. These results suggested a possible defect in the presynaptic voltage-gated calcium channels (VGCC).

In the 1980s Engel and colleagues used freeze fracture electron microscopy to study the motor nerve terminals in LEMS and in a passive transfer model of the disease (see below). They identified double parallel rows of intramembranous particles in the motor nerve terminal membrane which are thought to represent VGCCs. These channels were markedly reduced in number and their distribution was very abnormal.

Mice injected with LEMS IgG showed similar changes to those described in the patients. Moreover, IgG could be demonstrated on the motor nerve terminal, apparently at the position where VGCCs are thought to be concentrated. Other evidence implicated divergent binding of antibodies leading to reduction in the VGCCs, and complement did not seem to be involved.

It was some time before the presence of antibodies to the VGCCs was unequivocally demonstrated. VGCCs are a family of transmembrane proteins that can be distinguished by various drugs and neurotoxins. Conotoxin (ω-CmTx) is a snail toxin that binds to the P/Q-type VGCC and inhibits neuromuscular transmission. Antibodies in LEMS can be detected by radioimmunoprecipitation of VGCCs extracted from human or mammalian cerebellum that are labeled with 125-I-ω-Conotoxin (ω-CmTx) MVIIC. The antibodies are present in over 85% of patients and are highly specific for the disease, although they can be found in some patients with SCLC without LEMS. SCLCs express VGCCs on their surface, and LEMS IgG reduces the numbers of VGCC in these cells and in other cell lines expressing the calcium channels. Just as AChR synthesis is increased in MG muscle, so also is likely to be increased synthesis of P/Q-type VGCC or alternative channels in motor nerves in LEMS. P/Q-type VGCCs are also present at autonomic synapses explaining the frequent autonomic involvement in these patients.

**Acquired Neuromyotonia**

**Clinical features**

Acquired neuromyotonia (NMT) is a condition associated with muscle hyperactivity and probably under-diagnosed as it is not life-threatening. Clinical features include muscle stiffness, cramps, myokymia, pseudomyotonia and weakness. These are most common in the limbs and trunk but can be found in isolated muscle groups. The muscle hyperactivity continues during sleep and excessive sweating is a common complaint. The typical finding on electromyography is spontaneous motor unit discharges occurring in distinctive doublets, triplets, or longer runs with high intraburst frequency. Most of this activity is generated distally, perhaps at the NMJ itself, but in some cases more proximally. Cramp fasciculation syndrome may form part of the same clinical spectrum.

**Immunological basis**

As with LEMS, the first clinical observations pointed to an autoimmune basis. Although no HLA association has yet been demonstrated, other autoimmune diseases are commonly present, and about 20% of patients have a thymoma which can predate the symptoms or be identified after presentation of the neurological disease. Some patients also have MG. However, in contrast to MG and LEMS, some patients appear to have a monophasic illness that recovers spontaneously within a few months or years. There is circumstantial evidence that these forms of the disease occur following an infection or other immune response; one patient developed the disease after multiple wasp stings.

**Immune mechanisms**

Although neuromyotonia is a peripheral disease, the cerebrospinal fluid may contain oligoclonal bands, and since it can be associated with other autoimmune
diseases and thymoma, plasma exchange and passive transfer experiments were undertaken.\cite{82} Injection of IgG into mice resulted in a modest increase in the number of packets of ACh released, and IgG also increased neuronal activity in dorsal root ganglion cultures.\cite{83} Importantly, these findings were very similar to those in the presence of low concentrations of potassium channel blockers such as 4-aminopyridine. This suggested that loss of voltage-gated potassium channel (VGKC) function may underlie the electrophysiological findings in patients.

VGKCs are a large family of potassium channels, and the full spectrum of VGKC involved in neuromyotonia has not been explored. The antibodies are currently measured by radioimmunoprecipitation of VGKC Kv1.1, 1.2 and 1.6 that are conveniently identified by binding of the snake toxin, dendrotoxin.\cite{84} Kv1.1 and 1.2 are highly expressed in the peripheral nervous system, particularly in the juxtaparanodal region of the nodes of Ranvier\cite{85} and this is the most likely target for the antibodies, but they may also be expressed at the motor nerve terminal.

Antibodies to VGKCs can also be detected by immunostaining of xenopus oocytes or cell lines engineered to express the different Kv subtypes individually.\cite{84,86} The precise specificity of the antibodies is difficult to determine as the VGKCs are tetrampic proteins made up of different combinations of different Kv subunits. There are several reports that the antibodies, or IgG fractions, reduce potassium currents in cells transfected with the Kv1.2 subtype\cite{87} and that this does not require complement.\cite{88} It is likely that the antibodies act by reducing the VGKC numbers and that complement-mediated attack on the motor nerves does not occur.\cite{89}

**Central Nervous System Disease with Voltage-gated Potassium channel Antibodies**

Although the typical neuromyotonia patient does not have central symptoms, some have sensory nerve involvement and sleep problems or anxiety are not infrequent.\cite{90} Some patients have severe neuromyotonia with additional autonomic and marked central symptoms. This is usually called Morvan’s syndrome.\cite{90} In addition, patients with subacute onset of limbic encephalitis, with memory loss, seizures and disorientation\cite{90} or epilepsy with minimal cognitive involvement,\cite{91} are increasingly being recognized. It is still unclear as to which antibody specificity determines the clinical phenotype or whether other factors are involved.

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